Working Resources List on Dementia Care Management and Intellectual Disability

Preparing Community Agencies for Adults Affected by Dementia - "PCAD" Project

v.26k

PCAD Project
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To find resource or content of interest, search by keyword (e.g., assessment, carer, decline, etc.) in PDF reader.
Abstract: Education and training is vital in improving age-related care. Findings suggest that improved training of five electronic databases identified 11 articles that reported on the role of process of mapping existing evidence of dementia education and training preferred reporting items for systematic reviews and meta-analyses to guide the additional training needs. A systematic scoping review was completed using examine the impact dementia education has on caregivers' confidence to be a factor in meeting the age-related needs of people. This paper aims to provide the age-related care. Findings suggest that improved training provision is needed to support early diagnosis and increase caregivers' confidence in meeting the physical and psychological needs of older adults with intellectual disability.

Adams, D., Oliver, C., Kalsy, S., Peters, S., Broquard, M., Basra, T., Konstandinidi, E., & McQuillan, S.

Behavioural characteristics associated with dementia assessment referrals in adults with Down syndrome.


Abstract: Behavioral changes associated with dementia in Down syndrome are well documented, yet little is known about the effect of such behaviors on carers and referral. By comparing the behavioral and cognitive profiles of individuals referred for a dementia assessment with those of individuals not referred, some insight can be gained into behavioral characteristics that initiate referral for specialist support or interventions. Forty-six adults with Down syndrome were divided into two groups dependant upon method of entry into the study; post-referral to a specialist service for older adults with intellectual disabilities and Down syndrome for a dementia assessment (n = 17) or after receiving information sent out to day centers and residential homes (n = 29). These groups were compared on established measures of dementia alongside two informant measures of behavior. Those referred for a dementia assessment evidenced scores indicative of cognitive decline on both informant and direct Neuropsychological Assessments and showed more behavioral excesses, but not deficits, and lower socialization and coping skills than those in the comparison group. Carers of those referred for a dementia assessment reported a greater impact of behavioral excesses on staff than on the individual showing the behavior in contrast to the comparison group. The behavioral differences between those referred and the comparison group suggest that two factors are involved in the instigation of a referral for a dementia assessment: the nature of the behavioral presentation (excesses rather than deficits) and the effect of that behavioral change upon the care staff.

Adam, E., Sleeman, K.E., Brearley, S., Hunt, K., & Tuffrey-Wijne, I.

The palliative care needs of adults with intellectual disabilities and their access to palliative care services: A systematic review.


Abstract: There is evidence that people with intellectual disabilities experience healthcare inequalities, including access to specialist palliative care, but to date, there has not been a systematic review of empirical evidence. To identify the palliative care needs of adults with intellectual disabilities and the barriers and facilitators they face in accessing palliative care. Systematic review using a narrative synthesis approach (International prospective register of systematic reviews (PROSPERO) registration number: CRD42019138974). Five databases were searched in June 2019 (MEDLINE, Embase, PsycINFO, the Cochrane library and CINAHL) along with hand searches and a search of the grey literature. All study designs were included. A total of 52 studies were identified, all of which were conducted in high-income countries, the majority in the United Kingdom (n = 28). From a total of 2,970 participants across all studies, only 1% were people with intellectual disabilities and 1.3% were family members; the majority (97%) were health/social care professionals. Identified needs included physical needs, psychosocial and spiritual needs, and information and...
communication needs. Barriers and facilitators were associated with education (e.g. staff knowledge, training and experience), communication (e.g. staff skill in assessing and addressing needs of people with communication difficulties), collaboration (e.g. importance of sustained multidisciplinary approach) and health and social care delivery (e.g. staffing levels, funding and management support). This review highlights the specific problems in providing equitable palliative care for adults with intellectual disabilities, but there is a lack of research into strategies to improve practice. This should be prioritised using methods that include people with intellectual disabilities and families.


Individual cognitive stimulation therapy (iCST) for people with intellectual disability and dementia: a feasibility randomised controlled trial

*Aging & Mental Health, 2020, 26(4), 698-708. DOI: 10.1080/13607863.2020.1869180*

Abstract: Authors examined the feasibility, acceptability and fidelity of individual Cognitive Stimulation Therapy (iCST) in people with intellectual disability (ID) and dementia. We aimed to recruit forty dyads (carer and individual with dementia and ID) who were randomised to iCST or a waiting list control group. Both groups received treatment as usual. Family and paid carers delivered the manualised intervention (40 sessions over 20 weeks). Recruitment and retention of participants, intervention adherence, fidelity and acceptability were assessed. Outcome measures of cognition, adaptive functioning, quality of life (QoL) and carer outcomes were collected at baseline, midpoint (11 weeks) and at 21 weeks. Qualitative interviews were conducted with six carers about their experience of iCST. Forty dyads were recruited over 10 months from 12 National Health Service trusts. One dyad dropped out and 97.5% completed the midpoint and end-point assessments respectively. Assessment of fidelity indicated that the correct session structure was not followed: 70% completed at least 20 sessions and there was a high level of satisfaction with iCST. QoL was significantly higher in the iCST arm at 21 weeks (adjusted mean difference: 3.11; 95% CI: 0.64 to 5.58). There were no differences in the other outcome measures. The intervention was feasible and acceptable. A full-scale trial is warranted but some modifications are needed, including improved training and supervision for carers to improve fidelity.

**Ahlström, G., Aaxon, M., Sandberg, M., & Flygare Wallén, E.**

Health care utilisation among older people with Down syndrome compared to specific medical guidelines for health surveillance: a Swedish national register study


Abstract: Specific medical guidelines for health surveillance exist for people with Down syndrome (DS) since 25 years but knowledge of adherence to the guidelines is lacking. The guidelines were developed to avoid unnecessary suffering from preventable conditions. The aims of the study were to investigate 1) planned health care visits in relation to the co-morbidities described in specific medical guidelines as a measure of adherence, 2) unplanned health care visits as a measure of potentially unmet health care needs and 3) gender differences in health care utilization among older people with DS. This register-based study includes people with DS (n = 472) from a Swedish national cohort of people with intellectual disability (n = 7936), aged 55 years or more, and with at least one support according to the disability law, in 2012. Data on inpatient and outpatient specialist health care utilization were collected from the National Patient Register for 2002-2012. A total of 3584 inpatient and outpatient specialist health care visits were recorded during the 11 years, of which 54.6% (n = 2103) were planned, 44.0% (n = 1695) unplanned and 1.4% (n = 56) lacked information. More than half of the visits, 67.0% (n = 2352) were outpatient health care visits thus inpatient 33% (n = 1272). Most planned visits (29.4%, n = 618) were to an ophthalmology clinic, and most unplanned visits to an internal medicine clinic (36.6%, n = 621). The most common cause for planned visits was cataract, found at least once for 32.8% in this cohort, followed by arthritis (8.9%), epilepsy (8.9%) and dementia (6.6%). Pneumonia, pain, fractures and epilepsy each accounted for at least one unplanned visit for approximately one-fourth of the population (27.1, 26.9, 26.3 and 19.7% respectively). Men and women had similar numbers of unplanned visits. However, women were more likely to have visits for epilepsy or fractures, and men more likely for pneumonia. Increased awareness of existing specific medical guidelines for people with DS is vital for preventive measures. The relatively few planned health care visits according to the medical guidelines together with a high number of unplanned visits caused by conditions which potentially can be prevented suggest a need of improved adherence to medical guidelines.

**Aldred, M.J., Martini, A.C., Patterson, D., Hendrix, J., & Granholm, A.C.**

Aging with Down syndrome: where are we now and where are we going?

*Journal of Clinical Medicine, 2021 Oct 13, 10(20), 4687. doi: 10.3390/jcm10204687*

Abstract: Down syndrome (DS) is a form of accelerated aging, and people with DS are highly prone to aging-related conditions that include vascular and neurological disorders. Due to the overexpression of several genes on Chromosome 21, for example genes encoding amyloid precursor protein (APP), superoxide dismutase (SOD), and some of the interferon receptors, those with DS exhibit significant accumulation of amyloid, phospho-tau, oxidative stress, neuronal loss, and neuroinflammation in the brain as they age. In this review, we will summarize the major strides in this research field that have been made in the last few decades, as well as discuss where we are now, and which research areas are considered essential for the field in the future. Authors examine the scientific history of DS bridging these milestones in research to current efforts in the field. We extrapolate on comorbidities associated with this phenotype and highlight clinical networks in the USA and Europe pursuing clinical research, concluding with funding efforts and recent recommendations to the NIH regarding DS research.

**Alftberg, Å., Johansson, M., & Ahlström, G.**

Ambivalence among staff regarding ageing with intellectual disabilities: Experiences and reflections.


Abstract: This study explores the experiences and reflections of staff in intellectual disability (ID) services concerning ageing with ID. Qualitative interviews were conducted with 24 staff members in group homes and daily activity centres. The findings showed that the staff were uncertain about the signs of ageing in people with intellectual disabilities; they compared the life conditions of these people with conditions in older people without intellectual disabilities. Their emphasis on an active lifestyle was very strong. The staff members also mentioned uncertainty about how to facilitate assistive devices and whether ‘ageing in place’ was the best solution. The overall theme was manifested as ambivalence where notions of older people with intellectual disabilities seemed incompatible with notions of old age in general and could be explained by the theoretical concept of age coding. The findings of this study indicate the need to provide education about ageing to staff working in ID services.


Individual cognitive stimulation therapy (iCST) for people with intellectual disability and dementia: a feasibility randomised controlled trial


Abstract: To examine the feasibility, acceptability and fidelity of individual Cognitive Stimulation Therapy (iCST) in people with intellectual disability (ID) and dementia. We aimed to recruit forty dyads (carer and individual with dementia and ID) who were randomised to iCST or a waiting list control group. Both groups received treatment as usual. Family and paid carers delivered the manualised intervention (40 sessions over 20 weeks). Recruitment and retention of participants, intervention adherence, fidelity and acceptability were assessed. Outcome measures of cognition, adaptive functioning, quality of life (QoL) and carer outcomes were collected at baseline, midpoint (11 weeks) and at 21 weeks. Qualitative interviews were conducted with six carers about their experience of iCST. Forty dyads were recruited over 10 months from 12 National Health Service trusts. One dyad dropped out and 97.5% completed the midpoint and end-point assessments respectively. Assessment of fidelity indicated that the correct session structure was not followed; 70% completed at least 20 sessions and there was a high level of satisfaction with iCST. QoL was significantly higher in the iCST arm at 21 weeks (adjusted mean difference: 3.11; 95% CI: 0.64 to 5.58). There were no differences in the other outcome measures. The intervention was feasible and acceptable. A full-scale trial is warranted but some modifications are needed, including improved training.
and supervision for carers to improve fidelity

Altuna, M., Giménez, S., & Fortea, J.
Epilepsy in Down syndrome: A highly prevalent comorbidity
Abstract: Individuals with Down syndrome (DS) have an increased risk for epilepsy during the whole lifespan, but especially after age 40 years. The increase in the number of individuals with DS living into late middle age due to improved health care is resulting in an increase in epilepsy prevalence in this population. However, these epileptic seizures are probably underdiagnosed and inadequately treated. This late onset epilepsy is linked to the development of symptomatic Alzheimer's disease (AD), which is the main comorbidity in adults with DS with a cumulative incidence of more than 90% of adults by the seventh decade. More than 50% of patients with DS and AD dementia will most likely develop epilepsy, which in this context has a specific clinical presentation in the form of generalized myoclonic epilepsy. This epilepsy, named late onset myoclonic epilepsy (LOMEDS) affects the quality of life, might be associated with worse cognitive and functional outcomes in patients with AD dementia and has an impact on mortality. This review summarizes the current knowledge about the clinical and electrophysiological characteristics, diagnosis and treatment of epileptic seizures in the DS population, with a special emphasis on LOMEDS. Raised awareness and a better understanding of epilepsy in DS from families, caregivers and clinicians could enable earlier diagnoses and better treatments for individuals with DS.

Alzheimer's Association
Guidelines for dignity: Goals of specialized Alzheimer/dementia care in residential settings
47 pp.
Abstract: Standards for care and structure of care settings housing persons affected by Alzheimer's disease. Includes sections on philosophy, pre-admission activities, admission, care planning and implementation, adapting to changes in condition, staffing and training, physical environment and "success indicators."

Alzheimer's Australia
Down syndrome and Alzheimer's disease
12 pp.
[Place of publication not provided] (no date)
Abstract: Informational booklet on dementia and people with Down syndrome jointly issued by Alzheimer's Australia, Down Syndrome Victoria, and Centre for Developmental Disability Health Victoria. Contains three main sections: (1) About Alzheimer's disease and Down syndrome, (2) Diagnosis, and (3) Support, as well as a section on local resources.

Alzheimer's Disease International
Planning and design guide for community based day care centres
21 pp.
Abstract: An illustrated 21-page booklet highlighting main design issues and suggestions for organizing an effective environment for adults with dementia - with applications for residential environment.

Alzheimer's Disease Society
Safe as houses -- Living alone with dementia (A resource booklet to aid risk management)
30 pp.
Abstract: A 30-page booklet designed for the carer who is concerned about an older person with early to mid-stage dementia who may be living on their own. The booklet examines risks that the older adult may encounter and suggests how they could be minimized. The intent of the booklet is to aid the older person remain functional at home, with as minimal risk, for as long as possible. Covers personal care, finances, wandering, security, medication, utilities, and household safety. Whilst information is generic, resource information is geared toward the UK.

Alzheimer's Society
Learning disabilities and dementia
https://www.alzheimers.org.uk/about-dementia/types-dementia/learning-disabilities
Abstract: Web-based information produced in the UK on the topic of intellectual disabilities and dementia. Contains background information, as well as diagnosis, identification of symptoms and support and care services.

Alzheimer's Society
Supporting a person with dementia who also has a learning disability
Abstract: Informational webpage on how to approach helping a relative or friend with an intellectual disability who's diagnosed with dementia.

Anderson, M., Oak, K., Goodey, R., Dodd, K., & Shankar, R.
Do the severity of Intellectual Disability and/or the presence of neurodevelopmental disorders influence the onset of dementia in people with Down syndrome?
Abstract: Having a diagnosis of Down syndrome (DS) is associated with intellectual disability (ID), pervasive developmental disorders and Alzheimer's dementia (AD). The association between these conditions has not been well evaluated. This paper looks to examine the current evidence pertaining to the relationship between dementia in people with DS and severity of ID and the presence of pervasive developmental disorders. A scoping review using PRISMA guidance was undertaken. Medline, Cochrane database, NHS evidence, Trial registers and Open Grey were searched in December 2018 and an updated search was completed in July 2020. Three search strategies were used to retrieve articles relating to DS, dementia, pervasive developmental disorders (including autism, autism spectrum disorder (ASD) and attention-deficit hyperactivity disorder (ADHD)) and severity of ID. Studies were included if they met the pre-defined inclusion criteria of investigating an association between autism/ASD, ADHD, or severity of ID and the development of dementia in people with DS. Studies were excluded if they did not include primary data, if the population included non-Down causes of ID, or if no specific outcome measure related to comorbid autism/ASD, ADHD, or severity of ID and dementia in people with a diagnosis of DS were reported. There were no exclusions related to study design. Papers were assessed for quality using the Mixed Methods Appraisal Tool (MMAT). The search identified 15 papers, publishing results from 12 studies, relating to severity of ID, DS and dementia. No papers were identified relating to pervasive developmental disorders, DS and dementia. There is limited evidence on how severity of ID impacts on the presentation, diagnosis, management or prognosis of dementia in people with DS. However, no evidence was found on comorbid pervasive developmental disorders, DS, and dementia. This paper has identified multiple areas for future research. There is an urgent need for longitudinal studies into the presentation, development and progression of dementia in people with DS ensuring the severity of ID and comorbid pervasive developmental conditions are captured regularly to understand their influence on the dementia etiology and outcome.

Gait dyspraxia as a clinical marker of cognitive decline in Down syndrome: A review of theory and proposed mechanisms
Epub 2016 Feb 27.
Abstract: Down syndrome (DS) is the most common genetic cause of intellectual disability in children. With aging, DS is associated with an increased risk for Alzheimer's disease (AD). The development of AD neuropathology in individuals with DS can result in further disturbances in cognition and behavior and may significantly exacerbate caregiver burden. Early detection may allow for appropriate preparation by caregivers. Recent literature suggests that declines in gait may serve as an early marker of AD-related cognitive disorders; however,
this relationship has not been examined in individuals with DS. The theory regarding gait dyspraxia and cognitive decline in the general population is reviewed, and potential applications to the population with individuals with DS are highlighted. Challenges and benefits in the line of inquiry are discussed. In particular, it appears that gait declines in aging individuals with DS may be associated with known declines in frontoparietal gray matter, development of AD-related pathology, and white matter losses in tracts critical to motor control. These changes are also potentially related to the cognitive and functional changes often observed during the same chronological period as gait declines in adults with DS. Gait declines may be an early marker of cognitive change, related to the development of underlying AD-related pathology, in individuals with DS. Future investigations in this area may provide insight into the clinical changes associated with development of AD pathology in both the population with DS and the general population, enhancing efforts for optimal patient and caregiver support and propelling investigations regarding safety/quality of life interventions and disease-modifying interventions.

Andrews, E.J., Martini, A.C., & Head, E. Exploring the role of sex differences in Alzheimer's disease pathogenesis in Down syndrome. Front Neurosci. 2022 Aug 12;16:954999. doi: 10.3389/fnins.2022.954999. Abstract: Women are disproportionately affected by Alzheimer's disease (AD), yet little is known about sex-specific effects on the development of AD in the Down syndrome (DS) population. DS is caused by a full or partial triplication of chromosome 21, which harbors the amyloid precursor protein (APP) gene, among others. The majority of people with DS in their early- to mid-40s will accumulate sufficient amyloid-beta (Ab) in their brains along with neurofibrillary tangles (NFT) for a neuropathological diagnosis of AD, and the triplication of the APP gene is regarded as the main cause. Studies addressing sex differences with age and impact on dementia in people with DS are inconsistent. However, women with DS experience earlier age of onset of menopause, marked by a drop in estrogen, than women without DS. This review focuses on key sex differences observed with age and AD in people with DS and a discussion of possible underlying mechanisms that could be driving or protecting from AD development in DS. Understanding how biological sex influences the brain will lead to development of dedicated therapeutics and interventions to improve the quality of life for people with DS and AD.


Antonangeli, J.M. The Alzheimer project: formulating a model of care for persons with Alzheimer’s disease and mental retardation American Journal of Alzheimer’s Disease, 1995, 10(4), 13-16. Abstract: Article speaks to a pilot project conducted in Massachusetts to increase staffing, education and Alzheimer case management supports. Special supports were designed and offered to a number of adults with Down syndrome affected by dementia, including specialize assessments, team care planning meetings, home adaptations and behavior loss supports.

Arvio, M., & Bjelocigro-Laasko, N. Screening of dementia indicating signs in adults with intellectual disabilities Journal of Applied Research in Intellectual Disabilities, 2021 (May 01), https://doi.org/10.1111/jar.12888 Abstract: In intellectual disability, the cognitive delay is observed during developmental age, whereas in dementia, cognitive decline occurs during post-developmental period. So far, the risk of dementia in people with intellectual disability, excluding those with Down syndrome, is poorly known.

We screened dementia signs in a study group of 230 adults (34–80 years of age) with the help of the British Present Psychiatric State—Learning Disabilities assessment. Of the study members, 42% showed two or more signs. The overall frequency of symptoms did not differ between age groups. The number of individuals with a genetic syndrome or disease manifesting with a shortened lifespan was greater in the younger age groups when compared to the older age groups. People with an intellectual disability represent numerous rare syndromes with comorbidities. It seems that dementia signs may affect any age groups of adults with intellectual disability.

Arvio, M., & Bjelocigro-Laasko, N. Down syndrome - Onset age of dementia. Journal of Alzheimer’s Disease & Parkinsonism, 2017, 7(3), 1000329. doi: 10.4172/2161-0460 Abstract: Alzheimer’s disease is the most common cause of death in people who have Down syndrome (DS). This prospective, population-based, 15-year follow-up study aimed to define the onset age of dementia. At baseline 98 adults were screened for the first time by using the Present Psychiatric State-Learning Disabilities assessment; 19 also participated a second parallel follow up survey. These screenings were repeated twice more during the study. The number of study subjects during the 15 year follow-up period decreased. Another study found that adaptive skills in people with DS start to deteriorate at the age of 35 and our current findings indicate that also the first signs of dementia appear at the same age. The indicative signs for dementia increased rapidly after the age of 35 and appeared most frequently as reduced self-care skills, loss of energy, impaired understanding and forgetfulness. Regular follow-up of people who have Down syndrome from the age of 30 onward enables appropriate interventions to delay the progression of dementia. Early recognition of dementia is of utmost importance, because any deterioration in daily performance due to some treatable but undiagnosed condition may result in more restrictive residential care. Once memory changes have been observed, they may not reflect the onset of AD (depression, sleep disorders, B12-vitamin deficiency, hypothryoidism, urinary tract infections or polypharmacy) may be the underlying causes behind the first dementia signs. When periodic assessments are completed at regular intervals by a carer (who is familiar with the person for a long time) explains why the first dementia signs were recognized at clearly younger age in our study than in other reported surveys (i.e., 35 versus 48-56 years).

Aschenbrenner, A.J., Baksh, R.A., Benejam, B., Beresford-Webb, J.A., Coppus, A., Fortea, J., Handen, B.L., Hartley, S., Head, E., Jaeger, J., Levin, J., Loos;i. S.V., Rebillat, A-S., Sacco, S., Schmitt, F.A., Thurlow, K.E., Zaman, S., Hassenstab, J., & Strydöm, A. Markers of early changes in cognition across cohorts of adults with Down syndrome at risk of Alzheimer’s disease Alzheimer’s and Dementia (Amsterdam, Netherlands), 2021,13(1), e12184. doi: 10.1002/dad2.12184 Abstract: Down syndrome (DS), a genetic variant of early onset Alzheimer’s disease (AD), lacks a suitable outcome measure for prevention trials targeting pre-dementia stages. We used cognitive test data collected in several longitudinal aging studies internationally from 312 participants with DS without dementia to identify composites that were sensitive to change over time. We then conducted additional analyses to provide support for the utility of the composites. The composites were presented to an expert panel to determine the most optimal cognitive battery based on predetermined criteria. There were common cognitive domains across site composites, which were sensitive to early decline. The final composite consisted of memory, language/executive functioning, selective attention, orientation, and praxis tests. We have identified a composite that is sensitive to early decline and thus may have utility as an outcome measure in trials to prevent or delay symptoms of AD in DS.

Auty, E., & Scior, K. Psychologists’ clinical practices in assessing dementia in individuals with Down syndrome Journal of Policy and Practice in Intellectual Disabilities, 2008, 5(4), 259-268. DOI:10.1111/j.1741-1130.2008.00187.x Abstract: There are now ample guidelines for the assessment and diagnosis of possible dementia in individuals with intellectual disabilities (ID) and Down syndrome. However, little is known about their implementation in clinical practice. This study set out to examine the clinical practice of one key professional group, namely clinical psychologists. A national survey of clinical
psychologistsin ID services in the United Kingdom was undertaken. Detailed descriptions of clinical practice were obtained from 64 psychologists. Responses were further explored in focus groups. The results suggest marked variability in practice, assessment methods, and explanations clinicians give to service users and carers. Clinicians described struggling with the ethics and practicalities of how to present dementia assessments to individuals with ID and highlighted a need for more research, debate, and guidance. They also noted numerous shortfalls in service provision once aging individuals with ID show signs of dementia. Further research and clearer consensus guidelines are needed to improve assessment and diagnosis for this group.

**Axmon, A., Björne, P., Nylander, L., & Ahlstrom, G.**

Psychiatric diagnoses in older people with intellectual disability in comparison with the general population: a register study


Abstract: To describe the occurrence of psychiatric diagnoses in a specialist care setting in older people with intellectual disability (ID) in relation to those found in the same age group in the general population. A cohort of people with ID (n = 7936), aged 55 years or more in 2012, was identified, as was an age- and sex-matched cohort from the general population (n = 7936). Information regarding psychiatric diagnoses during 2002–2012 was collected from the National Patient Register, which contains records on all inpatient care episodes and outpatient specialist visits in Sweden. The mean age at the start of data collection (i.e. January 1st, 2002) was 53 years (range 44–85 years). Seventeen per cent (n = 1382) of the people in the ID cohort had at least one psychiatric diagnosis recorded during the study period. The corresponding number in the general population cohort was 10% (n = 817), which translates to an odds ratio (OR) of 1.84. The diagnoses recorded for the largest number of people in the ID cohort were ‘other’ (i.e. not included in any of the diagnostic groups) psychiatric diagnoses (10% of the cohort had at least one such diagnosis recorded) and affective disorders (7%). In the general population cohort, the most common diagnoses were affective disorders (4%) and alcohol/substance-abuse-related disorders (4%). An increased odds of having at least one diagnosis was found for all investigated diagnoses except for alcohol/substance-abuse-related disorders (OR = 0.56). The highest odds for the ID cohort was found for diagnosis of psychotic disorder (OR = 10.4) followed by attention deficit/hyperactive disorder (OR = 3.81), dementia (OR = 2.71), personality disorder (OR = 2.67), affective disorder (OR = 1.74) and anxiety disorder (OR = 1.36). People with ID also had an increased odds of psychiatric diagnoses not included in any of these groups (OR = 8.02). The percentage of people with ID who had at least one diagnosis recorded during the study period decreased from more than 30% among those aged 55–59 years in 2012 (i.e. born 1953–1957) to approximately 20% among those aged 75+ years in 2012 (i.e. born in or before 1937). Older people with ID seem to be more likely to have psychiatric diagnoses in inpatient or outpatient specialist care than their peers in the general population. If this is an effect of different disorder prevalence, diagnostic difficulties or differences in health care availability remains unknown. More research is needed to understand the diagnostic and treatment challenges of psychiatric disorders in this vulnerable group.

**Axmon, A., Karlsson, B., & Ahlstrom, G.**

Health care utilisation among older persons with intellectual disability and dementia: a registry study


Abstract: Both persons with intellectual disability (ID) and persons with dementia have high disease burdens, and consequently also high health care needs. As life expectancy increases for persons with ID, the group of persons with the dual diagnosis of ID and dementia will become larger. Through national registries, we identified 7936 persons who had received support directed to persons with ID during 2012, and an age- and gender-matched sample from the general population. A national registry was also used to collect information on health care utilisation (excluding primary care) for the period 2002-2012. Health care utilization was measured as presence and number of planned and unplanned in-patient and out-patient visits, as well as length of stay. In comparison with persons with ID but without dementia, persons with ID and dementia were more likely to have at least one planned out-patient visit (odds ratio [OR] 8.07), unplanned out-patient visit (OR 2.41), planned in-patient visit (OR 2.76) or unplanned in-patient visit (OR 4.19). However, among those with at least one of each respective outcome, the average number of visits did not differ between those with and without dementia. Persons with ID and dementia were less likely to have at least one planned out-patient visit than persons with dementia in the general population sample (OR 0.40), but more likely to have at least one unplanned in-patient visit (OR 1.90). No statistically significant differences were found for having at least one unplanned out-patient or planned in-patient visit. Nevertheless, among those with at least one unplanned out-patient visit, the number of visits was higher in the general population sample. Persons with ID and dementia are less likely to receive planned health care than persons with dementia in the general population. They have, however, higher levels of unplanned health care utilization. This may be an indication that the current support system is not sufficient to meet the challenges of increased longevity among persons with ID.

**Aylward, E., Burt, D., Thorpe, L., Laï, & Dalton, A.J.**

Diagnosis of dementia in individuals with intellectual disability


Abstract: The foremost impediment to progress in the understanding and treatment of dementia in adults with intellectual disability is the lack of standardized criteria and diagnostic procedures. Standardized criteria for the diagnosis of dementia in individuals with intellectual disability are proposed, and their application is discussed. In addition, procedures for determining whether or not criteria are met in individual cases are outlined. It is the intention of the authors, who were participants of an International Colloquium on Alzheimer Disease and Mental Retardation, that these criteria be appropriate for use by both clinicians and researchers. Their use will improve communication among clinicians and researchers, and will allow researchers to test hypotheses concerning discrepancies in findings among research groups (e.g. dementia prevalence ranges and age of onset). (Report of the Task Force for Development of Criteria for the Diagnosis of Dementia in Individuals with [intellectual disability].)


Multiple morbidity across the lifespan in people with Down syndrome or intellectual disabilities: a population-based cohort study using electronic health records.


Abstract: The Down syndrome phenotype is well established, but our understanding of its morbidity patterns is limited. We comprehensively estimated the risk of multiple morbidity across the lifespan in people with Down syndrome compared with the general population and controls with other forms of intellectual disability. In this matched population-based cohort-study design, we used electronic health-record data from the UK Clinical Practice Research Datalink (CPRD) from Jan 1, 1990, to June 29, 2020. We aimed to explore the pattern of morbidities throughout the lifespan of people with Down syndrome compared with people with other intellectual disabilities and the general population, to identify syndrome-specific health conditions and their age-related incidence. We estimated incidence rates per 1000 person-years and incidence rate ratios (IRRs) for 32 common morbidities. Hierarchical clustering was used to identify groups of associated conditions using prevalence data. Between Jan 1, 1990, and June 29, 2020, a total of 10 204 people with Down syndrome, 38 814 controls, and 69 150 people with intellectual disabilities were included. Compared with controls, people with Down syndrome had increased risk of dementia (IRR 94.7, 95% CI 69.9–128.4), hypothyroidism (IRR 10.6, 9.6–11.8), epilepsy (IRR 9.7, 8.5–10.9), and haematological malignancy (IRR 4.7, 3.4–6.3), whereas asthma (IRR 0.88, 0.79–0.98), cancer (solid tumour IRR 0.75, 0.62–0.89), ischaemic heart disease (IRR 0.65, 0.51–0.85), and particularly hypertension (IRR 0.26, 0.22–0.32) were less frequent in people with Down syndrome than in controls. Compared to people with intellectual disabilities, risk of dementia (IRR 16.60, 14.23–19.37), hypothyroidism (IRR 7.22, 6.62–7.88), obstructive sleep apnoea (IRR 4.45, 3.72–5.31), and haematological malignancy (IRR 3.44, 2.58–4.59) were higher in people with Down syndrome, with reduced rates for a third of conditions, including new onset of dental inflammation (IRR 0.88, 0.78–0.99), asthma (IRR 0.82, 0.73–0.91), cancer (solid tumour IRR 0.78, 0.65–0.93), sleep disorder (IRR 0.74, 0.68–0.80), hypercholesterolaemia (IRR 0.88, 0.79–0.99), and type 2 diabetes (IRR 0.89, 0.80–0.99).
Cambridge Cognitive Examination (CAMCOG), the neuropsychological assessment from the CAMDEX schedule. Fifty-six were assessed again 6 years later (Time 2). Based on the CAMDEX informant interview, nine of the 74 at Time 1, and 11 of the 56 at Time 2, were found to meet clinical criteria for AD. Forty-one scored above floor on the CAMCOG at Time 1 and were included in the analysis of cognitive decline. Concurrent validity was established by comparing diagnosis at Time 2 with independent evidence of objective decline on cognitive tasks since Time 1. Predictive validity was established by examining how accurately diagnosis at Time 1 predicted both cognitive decline and future diagnosis. Inter-rater reliability was determined by comparing the level of agreement between two raters. CAMDEX-based diagnosis of AD was shown to be consistent with objectively observed cognitive decline (good concurrent validity) and to be a good predictor of future diagnosis. Although numbers are small, some support is also provided for the accuracy with which diagnosis predicts cognitive decline. Inter-rater reliability was good with Kappa > 0.8 for 91% of items and > 0.6 for all items. The use of the modified CAMDEX informant interview enables the structured collection of diagnostic information, so that a valid and a reliable diagnosis of dementia can be made in those with pre-existing ID, using established diagnostic criteria.


Abstract: Recent research suggests that preclinical Alzheimer's disease (AD) in people with Down syndrome (DS) is characterized by changes in personality/behavior and executive dysfunction that are more prominent than deterioration in episodic memory. This study examines the relationship between executive dysfunction and the clinical and preclinical features of AD in DS. To determine the specificity of this relationship, performance on executive function (EF) measures is contrasted with performance on memory measures. One hundred and three people with DS (mean age 49 years, range 36-72) with mild to moderate learning disabilities (LD) took part. Dementia diagnosis was based on the CAMDEX informant interview conducted with each participant's main carer. Reported changes in personality/behavior and memory were recorded. Participants completed six EF and six memory measures (two of which also had a strong executive component) and the BPVS (as a measure of general intellectual ability). First, performance was compared between those with and without established dementia of Alzheimer's type (DAT), controlling for age and LD severity using ANCOVA. Next, the degree to which informant-reported changes predicted cognitive test performance was examined within the non-DAT group using multiple regression analyses. The DAT group (N=25) showed a consistent pattern of impaired performance relative to the non-DAT group (N=78), across all measures. Within the non-DAT group, number of informant-reported personality/behavior changes was a strong predictor of performance on two EF and two 'executive memory' tests (but not on episodic memory tests). Informant-reported memory changes, however, were associated with impaired performance on a delayed recall task only. These findings provide further evidence for a specific impairment in frontal-lobe functioning in the preclinical stages of AD in DS.


Abstract: Down syndrome is the most common genetic cause of learning difficulties, and individuals with this condition represent the largest group of people with dementia under the age of 50 years. Genetic drivers result in a high frequency of Alzheimer's pathology in these individuals, evident from neuroimaging, biomarker, and neuropathological findings, and a high incidence of cognitive decline and dementia. However, cognitive assessment is challenging, and diagnostic methods have not been fully validated for use in these patients; hence, early diagnosis remains difficult. Evidence regarding the benefits of cholinesterase inhibitors and other therapeutic options to treat or delay progressive cognitive decline or dementia is very scarce. Despite close similarities with late-onset Alzheimer's disease, individuals with Down syndrome respond differently to treatment, and a targeted approach to drug development is thus necessary. Genetic and preclinical studies offer opportunities for treatment development, and potential therapies have been identified using these
approaches.


Abstract: In this article, the diagnosis of Alzheimer’s disease and its progressive behavioral impact on persons with Down syndrome is discussed. Several implications and suggestions for care and service provision for adults with Down syndrome are presented, including that Alzheimer’s disease in an adult with Down syndrome has an impact on the carer, adjusting communication strategies to correspond to the stage of dementia, aiding families to seek assistance from social agencies, stressing the remaining abilities and skills, aiding families and carers to develop realistic methods of providing care, and adapting the persons care and environment to help them cope with losses stemming from dementia. The authors also suggest proactive strategies for anticipating decline among adults with Down syndrome associated with dementia.


Abstract: The recent National Institute of Health (NIH) INCLUDE (Investigation of Co-occurring conditions across the Lifespan to Understand Down syndrome) initiative has bolstered capacity for the current increase in clinical trials involving individuals with Down syndrome (DS). This new NIH funding mechanism offers new opportunities to expand and develop novel approaches in engaging and effectively enrolling a broader representation of clinical trials participants addressing current medical issues faced by individuals with DS. To address this opportunity, the NIH assembled leading clinicians, scientists, and representatives of advocacy groups to review existing methods and to identify those areas where new approaches are needed to engage and prepare DS populations for participation in clinical trial research. This paper summarizes the results of the Clinical Trial Readiness Working Group that was part of the INCLUDE Project Workshop: Planning a Virtual Down Syndrome Cohort Across the Lifespan Workshop held virtually September 23 and 24, 2019.


Abstract: As the life expectancy of people with Down syndrome (DS) has markedly increased over the past decades, older adults with DS may be experiencing a higher incidence of aging conditions. In addition to longevity, the amyloid precursor protein gene located on chromosome 21 places individuals with DS at a high risk for developing Alzheimer disease. Yet, few studies have determined prevalence of dementia and comorbidities among older people with DS. To determine the prevalence of dementia and aging-related comorbidities in older adult individuals with DS, Cross-sectional analysis of 2015 California Medicare claims data. We examined 1 year of cross-sectional Medicare claims data that included 100% of Californian Medicare beneficiaries enrolled in both Medicare Part A and B in 2015. Of these 3 001 977 Californiaen Medicare beneficiaries 45 years or older, 878 individuals were identified as having a diagnosis of DS. Data were analyzed between April 2017 and February 2018. The frequency of DS dementia was assessed across different age categories. The number and frequency of 27 comorbidities were compared among individuals with DS and people with dementia and by age and sex groups. A total of 353 DS individuals (40%) were identified as having dementia diagnoses (mean, 59.7 years; 173 women [49%]) and 526 without dementia diagnoses (mean, 55.9 years; 250 women [48%]). The frequency of DS dementia among those 65 years or older rose to 49%. The mean number of comorbidities per individual increased with age in general. Comorbid conditions were more numerous among those with dementia compared with those with DS without dementia (mean, 3.4 vs 2.5, respectively), especially among those younger than 65 years. In particular, 4 treatable conditions, hypothyroidism, epilepsy, anemia, and weight loss, were much more frequent in DS dementia. Older Medicare beneficiaries in California with DS, especially those with dementia, have a high level of multimorbidity including several treatable conditions. While DS follow-up has long been confined to the pediatric sphere, we found that longevity in individuals with DS will necessitate complex adult and geriatric care. More evidenced-based and standardized follow-up could support better long-term comorbidity management and dementia care among aging adults with DS.


Abstract: Alzheimer disease (AD) is the leading cause of death in individuals with Down syndrome (DS). Previous studies have suggested that the APOE ε4 allele plays a role in the risk and age at onset of dementia in DS; however, data on in vivo biomarkers remain scarce. To investigate the association of the APOE ε4 allele with clinical and multimodal biomarkers of AD in adults with DS, his dual-center cohort study recruited adults with DS in Barcelona, Spain, and in Cambridge, UK, between June 1, 2009, and February 28, 2020. Included individuals had been genotyped for APOE and had at least clinical or AD biomarker measurement; 2 individuals were excluded because of the absence of trisomy 21. Participants were either APOE ε4 allele carriers or noncarriers. Participants underwent a neurological and neuropsychological assessment. A subset of participants had biomarker measurements: Aβ1-42, Aβ1-40, phosphorylated tau 181 (pTau181) and neurofilament light chain (NFL) in cerebrospinal fluid (CSF), pTau181, and NFL in plasma; amyloid positron emission tomography (PET); fluorine 18-labeled-fluorodeoxyglucose PET; and/or magnetic resonance imaging. Age at symptom onset was compared between APOE ε4 allele carriers and noncarriers, and within-group local regression models were used to compare the association of biomarkers with age. Voxelwise analyses were performed to assess topographical differences in gray matter metabolism and volume. Of the 464 adults with DS included in the study, 97 (20.9%) were APOE ε4 allele carriers and 367 (79.1%) were noncarriers. No differences between the 2 groups were found by age (median [interquartile range], 45.9 [36.4-50.2] years vs 43.7 [34.9-50.2] years; P = .56) or sex (51 male carriers [52.6%] vs 199 male noncarriers [54.2%]). APOE ε4 allele carriers compared with noncarriers presented with AD symptoms at a younger age (mean [SD] age, 50.7 [4.4] years vs 52.7 [5.8] years; P = .02) and showed earlier cognitive decline. Locally estimated scatterplot smoothing curves further showed between-group differences in biomarker trajectories with age as reflected by nonoverlapping CIs. Specifically, carriers showed lower levels of the CSF Aβ1-42 to Aβ1-40 ratio until age 40 years, earlier increases in amyloid PET and plasma pTau181, and earlier loss of cortical metabolism and hippocampal volume. No differences were found in NFL biomarkers or CSF total tau and pTau181. Voxelwise analyses showed lower metabolism in subcortical and parieto-occipital structures and lower medial temporal volume in APOE ε4 allele carriers. In this study, the APOE ε4 allele was associated with earlier clinical and biomarker changes of AD in DS. These results provide insights into the mechanisms by which APOE increases the risk of AD, emphasizing the importance of APOE genotype for future clinical trials in DS.


Abstract: We aimed to define prodromal Alzheimer’s disease (AD) and AD dementia using normative neuropsychological data in a large population-based cohort of adults with Down syndrome (DS). We employed a cross-sectional study. DS participants were classified into asymptomatic, prodromal AD and AD dementia, based on neuropsychologist’s judgment blinded to neuropsychological data (Cambridge Cognitive Examination for Older Adults with Down’s syndrome [CAMCOG-DS] and modified Cued Recall Test [mCRT]). We compared the
cutoffs derived from the normative data in young adults with DS to those from receiver-operating characteristic curve (ROC) analysis. Diagnostic performance of the CAMCOG-DS and modified Cued Recall Test (mCRT) in subjects with mild and moderate levels of intellectual disability (ID) was high, both for diagnosing prodromal AD and AD dementia (area under the curve [AUC] 0.73-0.83 and 0.90-1, respectively). The cutoffs derived from the normative data were similar to those derived from the ROC analyses.  

Diagnosing prodromal AD and AD dementia in DS with mild and moderate ID using population norms for neuropsychological tests is possible with high diagnostic accuracy.

Abstract: Diagnosis of prodromal Alzheimer's disease and Alzheimer's disease dementia in people with Down syndrome is a major challenge. The Cambridge Examination for Mental Disorders of Older People with Down's Syndrome and Others with Intellectual Disabilities (CAMDEX-DS) has been validated for diagnosing prodromal Alzheimer's disease and Alzheimer's disease dementia, but the diagnostic process lacks guidance. To derive CAMDEX-DS informant interview threshold scores to enable accurate diagnosis of prodromal Alzheimer's disease and Alzheimer's disease dementia in adults with Down syndrome. Psychiatrists classified participants with Down syndrome into no dementia, prodromal Alzheimer's disease and Alzheimer's disease dementia groups. Receiver operating characteristic analyses assessed the diagnostic accuracy of CAMDEX-DS informant interview-derived scores. Spearman partial correlations investigated associations between CAMDEX-DS scores, regional Aβ binding (positron emission tomography) and cortical thickness (magnetic resonance imaging). Diagnostic performance of CAMDEX-DS total scores were high for Alzheimer's disease dementia (area under the curve (AUC), 0.998; 95% CI 0.953-0.999) and prodromal Alzheimer’s disease’s (AUC = 0.954; 95% CI 0.887-0.982) when compared with healthy adults with Down syndrome. When compared with those with mental health conditions but no Alzheimer's disease, CAMDEX-DS Section B scores, denoting memory and orientation ability, accurately diagnosed Alzheimer's disease dementia (AUC = 0.998; 95% CI 0.892-0.984), but were unable to diagnose prodromal Alzheimer's disease. CAMDEX-DS total scores exhibited moderate correlations with cortical Aβ (r = 0.4 to 0.6, p = 0.05) and thickness (r = -0.4 to -0.05, p < 0.05) in specific regions. CAMDEX-DS total score accurately diagnoses Alzheimer's disease dementia and prodromal Alzheimer's disease in healthy adults with Down syndrome.

Bevens, S., Dawes, S., Kenschole, A., & Gaussen, K.  
Abstract: Despite the longstanding use of music therapy with people with intellectual disabilities and the growing evidence base for using music therapy as a tool to aid behavioural and psychological symptoms of dementia in the general population, there is little work published which details the use of music therapy groups for people with intellectual disabilities who have a diagnosis of dementia. The purpose of this paper is to report a qualitative evaluation of staff views of a music therapy group for people with intellectual disabilities and dementia. Carers of service users attending the group were interviewed either individually or through a focus group in order to ascertain their views about the music therapy group. The interview transcripts were then analysed using thematic analysis. Two core themes and eight sub themes emerged from the data. These themes show that the group was felt to be pleasurable and enjoyable for the service users and that some tangible benefits of attending the group were observed by staff members. Notwithstanding the positive feedback, the results also suggested that more work is needed to inform carers of the goals and purpose of such groups. Further psycho-education for carers is suggested as a strategy to support future groups to run successfully. There is little published research into the use of music therapy for people with intellectual disabilities who also have dementia. The current paper provides a starting point for future work in the area and further recommendations for future practice and research are considered.

Bigby, C., & Beadle-Brown, J.  
Abstract: Building on cultural dimensions of underperforming group homes this study analyzes culture in better performing services. In depth qualitative case studies were conducted in 3 better group homes using participant observation and interviews. The culture in these homes, reflected in patterns of staff practice and talk, as well as artefacts differed from that found in underperforming services. Formal power holders were undisputed leaders, their values aligned with those of other staff and the organization, responsibility for practice quality was shared enabling teamwork, staff perceived their purpose as "making the life each person wants it to be," working practices were person centered, and new ideas and outsiders were embraced. The culture was characterized as coherent, respectful, "enabling for residents, and "motivating" for staff. Though it is unclear whether good group homes have a similar culture to better ones the insights from this study provide knowledge to guide service development and evaluation.

Bigby, C., Bowers, B., & Webber, R.  
Planning and decision making about the future care of older group home residents and transition to residential aged care Journal of Intellectual Disability Research, 2011 Aug, 55(8), 777-789. doi: 10.1111/j.1365-2788.2010.01297.x  
Abstract: Planning for future care after the death of parental caregivers and adapting disability support systems to achieve the best possible quality of life for people with intellectual disability as they age have been important issues for more than two decades. This study examined perceptions held by family members, group home staff, and organizational managers about the future of older residents and the decisions made that a move to residential aged care was necessary. Grounded Dimensional Analysis was used to guide data collection and analysis by an interdisciplinary research team. Three sets of interviews over a period of 18 months were conducted with a family member, house supervisor and the program manager for each of seventeen older group home residents in Victoria, Australia. For the eight people for whom it was decided a move was necessary and the six who eventually moved focussed questions were asked about the decision-making process. While plans for lifelong accommodation in a group home proved unfounded, key person succession plans were effective. However, decisions move to a residential aged care facility was necessary were made in haste and seen as a fait accompli to involved family members. Although family members take seriously their mandate to oversee well-being of their older relative, they have little knowledge about their rights or avenues to safeguard untimely or inappropriate decisions being made by professionals.

Bigby, C., Knox, M., Beadle-Brown, J., Clement, T., & Mansell, J.  
Abstract: Culture recurs as an important but under-investigated variable associated with resident outcomes in supported accommodation for people with intellectual disability. This study aimed to conceptualize the potential impact of culture on quality of life outcomes for residents in 5 small group homes. Five categories of culture were developed: misalignment of power-holder values with organizations espoused values, otherness, doing for not with, staff centered, and resistance. Differences from institutional culture are discussed, and the potential of the findings as a starting point to consider culture in high performing group homes and develop a quantitative measure of culture.

Bishop, K.M., Hogan, M., Janicki, M.P., Keller, S.M., Lucchino, R., Mughal,

Abstract: Increasing numbers of adults with intellectual disabilities (ID) are living into old age. Though this indicates the positive effects of improved health care and quality of life, the end result is that more adults with ID are and will be experiencing age-related health problems and also exhibiting symptoms of cognitive impairment and decline, some attributable to dementia. Early symptoms of dementia can be subtle and in adults with ID are often masked by their lifelong cognitive impairment, combined with the benign effects of aging. A challenge for caregivers is to recognize and communicate symptoms, as well as find appropriate practitioners familiar with the medical issues presented by aging adults with lifelong disabilities. Noting changes in behavior and function and raising suspicions with a healthcare practitioner, during routine or ad hoc visits, can help focus the examination and potentially validate that the decline is the result of the onset or progression of dementia. It can also help in ruling out reversible conditions that may have similar presentation of symptoms typical for Alzheimer's disease and related dementias. To enable caregivers, whether family members or staff, to prepare for and advocate during health visits, the National Task Group on Intellectual Disabilities and Dementia Practices has developed guidelines and recommendations for dementia-related health advocacy preparation and assistance that can be undertaken by provider and advocacy organizations.


Abstract: Increased life expectancy generates greater ethical and legal dilemmas in the treatment of people with Down syndrome. Assumptions that younger cohorts of people with DS will experience healthier lives when compared to previous generations may not be realized as specific health issues associated with DS are genetically encoded and thus contemporary generations may face the same adverse health issues. With respect to dementia, authors note that by age 60 years, dementia involving memory loss, cognitive decline, and changes in adaptive behavior may be present in at least 56% of adults with DS and that some the neuropathological features of Alzheimer disease may be evident as early as age 40.


Abstract: None provided.

Bowers, B., Webber, R., & Bigby, C. Aging and health related changes of people with intellectual disabilities living in group homes in Australia Journal of Policy and Practice in Intellectual Disabilities, 2009, 6(2), 98. [SINGLE PAGE] 

Abstract: Group homes for people with ID are based on social models, emphasizing inclusion, engagement in community, and quality of life. As age related changes occur, group home staff members are faced with decisions about how to respond, how to support people experiencing health problems, and whether or for how long people can remain in the group homes. This study explored how group home staff members respond to aging and age related health conditions in group home residents and to identify factors that put people at risk of premature or inappropriate relocation. Using a longitudinal design in order to observe, over time, the onset of health problems, the initial responses of housing staff to health, the development of health conditions, the consequences of their initial responses, and the outcomes for both staff and residents were considered. In-depth interviews were conducted—at three 6-month intervals with 18 clusters of the housing manager, family member, the person with the disability, and in some cases, healthcare providers. A total of 91 interviews were completed, transcribed, and analyzed and in keeping with the theory-generating approach, early interviews were open and exploratory, evolving over time to facilitate comparative analysis across groups, strategies, conditions, and care issues. Staff and family members agreed that aging and the development of associated health conditions was increasingly becoming an issue for them. Significantly, there was wide variation among housing staff in terms of philosophy of care, with some believing that people should be supported to remain at the group homes for as long as possible. This, however, required the acquisition of new resources, a range of organizational changes to support staff and residents, changes to staffing patterns and levels, and a change in recruiting as a strategy to alter skill mix of house workers. Authors concluded that problems identified by most housing staff included: (a) inability of residents to retire despite age and health status; (b) risk of premature moves to reside care; (c) difficulties in identifying and managing behavioral issues; and (d) disruptive behavior. Early symptoms of dementia can be subtle and in adults with ID are often masked by their lifelong cognitive impairment, combined with the benign effects of aging. A challenge for caregivers is to recognize and communicate symptoms, as well as find appropriate practitioners familiar with the medical issues presented by aging adults with lifelong disabilities. Noting changes in behavior and function and raising suspicions with a healthcare practitioner, during routine or ad hoc visits, can help focus the examination and potentially validate that the decline is the result of the onset or progression of dementia. It can also help in ruling out reversible conditions that may have similar presentation of symptoms typical for Alzheimer's disease and related dementias. To enable caregivers, whether family members or staff, to prepare for and advocate during health visits, the National Task Group on Intellectual Disabilities and Dementia Practices has developed guidelines and recommendations for dementia-related health advocacy preparation and assistance that can be undertaken by provider and advocacy organizations.


Abstract: 20 chapter general text on adapting homes and living environments for persons with dementia; applicable to home and other residential situations for clinical, social, and ethical implications of changing life expectancy in Down syndrome Psychiatric comorbidity in older adults with intellectual disability Psychiatric Danubina, 2017, 29(Suppl 3), 590-593. PMID: 28953835 

Abstract: The majority of adults with an intellectual disability live with family carers, many of whom are ageing and have support needs of their own. Planning for the future thus becomes the key to preventing a crisis situation when family care is no longer viable because of death or ill health. Existing knowledge and practice are largely based upon the perspective of professionals and carers. This study explored the views, aspirations and concerns of adults with an intellectual disability, about living at home and planning for the future. Findings show that participants were very aware of the need for alternative housing or support in the future and had clear preferences about their future options. However, they also showed considerable concern for their family carers and this often impacted on their willingness to plan for the future or to move to alternative housing. Their demonstrable awareness of the inevitable death or ill health of family carers, and willingness to engage with the implications, emphasize the importance of involving adults with intellectual disability in planning for their future, as well as providing them with bereavement support.
adults with intellectual disabilities and dementia. Chapter sections include Aging and Alzheimer’s disease, Sensory environment (light and aging vision, lighting, impact of color, patterns and texture, acoustical changes, and wayfinding guidelines), Special care settings (creating a home feeling, designing spaces, therapeutic gardens and outdoor spaces), Implementing effective interior design (furniture and fabrics, floor-covering, wall and ceiling finishes, windows and window treatments), and The Design process. Contains a directory of resources and a glossary of terms.

Brodat, H., Seher,K., & Gibson, L.
Dementia time to death: A systematic literature review on survival time and years of life lost in people with dementia.
International Psychogeriatrics, 2012, 24(7),1034-1045. Published online on 13 February 2012. doi: 10.1017/S1041610211002924.
Abstract: Life expectancy with dementia directly influences rates of prevalence and service needs and is a common question posed by families and patients. As well as years of survival, it is useful to consider years of life lost after a diagnosis of dementia. Authors systematically reviewed the literature on mortality and survival with dementia which were compared to estimated life expectancies in the general population. Both were then compared by age (under 65 years vs. 65+ years), gender, dementia type, severity, and two epochs (prior to and after introduction of cholinesterase inhibitors in 1997). Survival after a diagnosis of dementia varies considerably and depends on numerous factors and their complex interaction. Relative loss of life expectancy decreases with age at diagnosis across varying sex, dementia subtypes (except for frontotemporal dementia and dementia with Lewy bodies), and severity stages. Numerous study deficiencies precluded a meta-analysis of survival in dementia. Authors concluded that estimates of years of life lost through dementia may be helpful for patients and their families.

Burt, D.B., & Aylward, E.H.
Abstract: A working battery of tests for the diagnosis of dementia, which is applicable to most adults with intellectual disability, is proposed by an international Working Group. The battery, reflecting contemporary research and practice findings, includes scales for informant report of functioning and tests for direct assessment. The Working Group recommends the international use of the battery both as part of ongoing and new longitudinal research, and in clinical practice. The widespread use of a common battery will enhance communication and collaborative opportunities among researchers and clinicians at various sites, and will help to standardize diagnostic protocols and research findings. The collaborative evaluation of such a battery will address one of the greatest challenges in the field, that of differentiating change associated with ageing from that associated with dementia.

Burt, D.B., & Aylward, E.
Assessment methods of diagnosis of dementia
In M.P. Janicki & A.J. Dalton (Eds.), Dementia, Aging, and Intellectual Disabilities. pp. 141-156
Abstract: Standardized diagnostic criteria and procedures are proposed to further progress in the understanding and treatment of dementia in adults with intellectual disabilities. This book chapter is a revised summary of previous reports prepared by participants of an international working group, which was conducted under the auspices of the International Association on Intellectual Disability and the American Association on Mental Retardation. Similarities in diagnostic issues between adults with intellectual disability and those in the general population are discussed, followed by a summary of issues unique to adults with intellectual disability. A brief overview of the application of CD-10 diagnostic criteria to adults with intellectual disability is presented, including a description of procedures for determining whether criteria are met in individual cases. Finally, clinical and research recommendations are made.

Burt, D.B., Loveland, K.A., & Lewis, K.R.
Abstract: The relation between dementia and depression in 61 adults with Down syndrome or 43 adults with mental retardation due to other causes was examined. Age-matched participants, ranging in age from 20 to 60 years, received a neuropsychological battery to assess declines in functioning and caregiver report measures to assess adaptive behavior and depression. Eight adults with Down syndrome had both depression and declines in functioning. No adults with mental retardation due to other causes had declines. Greater severity of depression was related to lower MA, poorer memory, and lower adaptive functioning in adults with Down syndrome only. Results suggest that dementia and depression are associated in Down syndrome but not in mental retardation due to other causes.

Abstract: Diagnosis of dementia in adults with intellectual disabilities requires documentation of clinically significant declines in memory and other cognitive skills, as well as changes in everyday and emotional functioning. To improve diagnostic accuracy in adults with Down syndrome, the authors examined conditions often associated with dementia, as well as tests useful for documentation of decline. Specific aims were to identify psychiatric disorders or medical conditions that increased the odds of a dementia diagnosis; to evaluate the sensitivity and specificity of widely used dementia scales; and to determine which tests, used singly or in combination, most accurately supported the presence of dementia. Participants were 78 adults with Down syndrome. Two methods based on a large test battery and one method based on clinical judgment were used to diagnose dementia. It was found that combinations of tests lead to increased levels of diagnostic sensitivity compared with single tests. When taken in combination with other investigations, our results suggest that assessment for psychiatric disorders, delayed memory decline, adaptive behavior decline, and the presence of seizures would be useful for the diagnosis of dementia and that dementia scales would provide additional useful information. The authors conclude that combinations of tests and scales will be most useful for diagnosing dementia in adults with intellectual disabilities. The authors suggest that further research is needed to promote rapid progress, with studies that focus on common diagnostic methodology, identification of screening instruments, and amounts of decline indicative of dementia.

Abstract: Accurate detection of dementia in adults with intellectual disabilities is important for clinical care, program planning, and clinical research. This paper reports on a study that examined two major diagnostic methods that varied in the following ways: (1) the extent to which they relied on clinical judgment; (2) the statistical method used to detect declines; and (3) the sensitivity to declines in functioning. Two methods based on testing were compared with one based on clinical judgment. Data were drawn from annual sequential assessments of 168 adults with intellectual disabilities (78 with Down syndrome and 90 with other etiologies). Agreement between testing and clinical judgment methods was 72-75% depending on testing method used. Clinical judgment produced a higher rate of dementia diagnosis for adults with Down syndrome compared with testing methods, suggesting a possible bias. The authors found that diagnostic criteria were useful both for identifying dementia and for describing its characteristics. Our results suggest that clinical judgment could result in a higher number of adults with Down syndrome diagnosed with dementia than methods based on test batteries. Common results across research studies indicate that combinations of sources of information (interviews/direct testing) would be most useful for dementia diagnosis. Future collaboration across research sites is needed to promote rapid progress in this important area, with emphasis on differential diagnosis.
Bush, A., & Beal, N.
Risk factors for dementia in people with Down syndrome: Issues in assessment and diagnosis
Abstract: It has been clearly established that there is an increased incidence of early onset dementia of the Alzheimer type (DAT) in people who have Down syndrome. There are variations in the age of onset of the clinical signs of DAT, which may be accounted for by different risk factors. In this review we examined the evidence that different biological and psychological factors may influence the risk for DAT. Limitations in design of early studies, the need for consistent diagnostic criteria for DAT in individuals with Down syndrome, and the lack of adequate psychometric tools to detect cognitive change are highlighted. Implications for research and clinical practice are considered in order to assess potential risk factors.

Cairns, D., Kerr, D., Chapman, A.
Difference realities: a training guide for people with Down’s syndrome and Alzheimer’s disease
pp. 54
University of Stirling (Dementia Services Development Centre), Stirling, Scotland FK9 4LA
A working guide for staff who are working with people with intellectual disabilities affected by Alzheimer’s disease. Topical sections cover the definitions of dementia and deal with diagnostic suggestions, as well as dealing with communication, helping maintenance of skills, dealing with challenging behaviors, structuring activities, and overall management of dementia. Written in an easy style, this guide is a very useful addition to any materials given to staff to help them understand and related to people affected by dementia.

Cairns, V., Lamb, I., & Smith, E.
https://doi.org/10.1111/j.1468-3156.2010.00851.x
Abstract: The high prevalence of dementia in individuals with Down’s syndrome has led intellectual disability services in the Hyndburn and Ribble Valley (HRV) area of England to develop a screening service to address this need. The authors offer reflections upon this process by its members after the first 12 months of operation. A multidisciplinary team, comprising professionals from intellectual disability psychology, intellectual disability speech and language therapy, intellectual disability community nursing and older adults psychiatry, has developed, and begun to implement, screening care pathways. The service conducts routine screening assessments, provides intervention for individuals where concerns arise and delivers training to carers. At the point of writing, 27 service users have received screening assessments and six have been identified as at moderate–high risk of developing dementia. Reflection and feedback has highlighted issues for consideration throughout the service development process, and an evaluation of the training provided by the service has found this to be effective in increasing carers understanding about dementia and intellectual disabilities.

Carfi, A., Antocioccio, M., Brandi, V., Cipriani, C., Fiore, F., Mascia, D., Vetrano, D.L., Onder, G.
Down syndrome in adulthood: A disease for geriatricians European Geriatric Medicine, 2014, 5(Suppl 1), S49.
Abstract: Authors evaluated 89 adults with Down syndrome at a clinic in Rome, Italy, using a range of physiological and neurological methods, including nutritional and sensory assessments. The S’s mean age was 42 years (range 18 to 72); 51% were females. Authors found behavioral disorders (53%), mood disorders (43%), seizures (22%), osteoporosis (40%), hypothyroidism (53%), diastolic dysfunction (80%), OSAS (90%), and hearing impairment (82%). Authors noted severe cognitive impairments in 67%, BMI greater than 25 in 66%, and low scores on physical performance measures (50%). Authors conclude that the pattern of diseases and conditions noted resemble those of other older adults and recommend that a mandatory geriatric evaluation be undertaken in older adults with Down.

Carling-Jenkins, R., Torr, J., Iacono, T., & Bigby, C.
Abstract: Australian research addressing the experiences of families of adults with Down syndrome and Alzheimer’s disease in seeking diagnosis and gaining support is limited. The aim of this study was to gain a greater understanding of these processes by exploring the experiences of families and carers in supporting people with Down syndrome and Alzheimer’s disease who had lived most or all of their lives with family. Three detailed case studies were created from multiple data sources, and then analyzed thematically. Families of adults with Down syndrome experienced stress and confusion as they negotiated a service system poorly equipped to meet their needs and professionals more focused on longstanding disability than the recent diagnosis of Alzheimer’s disease. Such overshadowing led to mismanagement by services. Authors conclude that this research advances understandings of the support needs of people with Down syndrome and Alzheimer’s disease and their families and exposes gaps in the service system.

Carmeli E, Ariav C, Bar-Yossef T, Levy R, Imam B.
Abstract: Adults with Down syndrome (DS) are often physically inactive, which may accelerate the onset of disease and aging symptoms. Eight older persons with DS (aged 54-61), and 10 younger persons with DS (aged 26-35) living in a residential care center were examined. Eighteen age- and gender-matched individuals without DS served as control groups. Sensory-motor tasks and Posture Scale Analyzer (PSA) were used to examine coordination and standing stability. The isokinetic muscle strength test was used for muscle strength investigation. The functional performance, coordination, and leg muscle strength of older adults with DS were more impaired than both the younger DS and the control groups. The older DS group showed lower sway rate and more symmetrical weight-bearing distribution during quiet standing than both the younger DS and the control groups. Our observations may have significant implications for understanding movement dysfunction in older adults with DS.

Carmeli, E., Ariav, C., Bar-Yossef, T., & Levy, R.
Abstract: Persons with Down syndrome (DS) are comparatively physically inactive, which could accelerate the onset of disease, resulting in symptoms associated with aging that are detrimental to health. The aim was to evaluate movement abilities across the life span in persons with DS. Eleven persons with DS (>50 years, mean age 58 years), and 10 younger persons with DS (<49 years, mean age 28 years) who resided in a residential living center were included in the study. Age- and gender-matched people without DS (n=22) served as control group. Five sensory-motor tasks that involved the integration of hand movements with visual information were used, as well as the posture scale analyzer system to examine postural stability. Results showed that the older persons with DS had more medical problems than the young persons with and without DS. The hand coordination and postural stability of the older adults with DS were more impaired in comparison with the young group and both control groups. It is postulated that their poor motor function and slower responses might be explained by a less active lifestyle, that could accelerate the onset of disease, resulting in symptoms associated with aging that are detrimental to health. Our observations could have significant implications for understanding the mechanisms underlying movement dysfunction in older adults with DS and might offer new approaches for possible prevention.

Carr, J.
Abstract: Longitudinal studies show that in general the population IQ declines...
with age: early and rapidly in the case of performance IQ, later and more slowly in the case of verbal IQ. These populations have not apparently included people with intellectual disabilities (ID). A literature search identified 11 studies, some cross-sectional and others longitudinal, which provided data on a variety of verbal and performance tests, over periods from 3 to 19 years, on older people with ID. Following statistical advice the results from the different tests were converted into the equivalent of, for verbal scores, British Picture Vocabulary Scale, and for performance scores, Leiter International Performance Scale, raw scores. Percentage change between earlier and later scores was then calculated. With one exception the studies considered tend to show overall ability declining relatively more, and performance ability declining relatively less, than has been shown to occur in the general population. Potential confounding factors, such as population attrition, cohort effects, etc., are thought not to have affected these results. The pattern of change with age in verbal and performance ability appears different in people with ID from that seen in the general population. Some possible reasons for this difference are discussed.

Carr, J., & Collins, S. 
Abstract: A population sample of people with Down syndrome has been studied from infancy and has now been followed up again at age 47 years. Intelligence and language skills were tested and daily living skills assessed. Memory/cognitive deterioration was examined using two test instruments. Scores on verbal tests of intelligence changed little. Those on a non-verbal test, on self-help skills and on both memory tests showed some decline, even when the scores of those already suffering from dementia were discounted. At age 47, scores on most tests of even the majority of the cohort (i.e. those not definitely diagnosed with dementia) showed some decline. While this includes the scores of people who may subsequently develop dementia, it may also reflect the normal ageing process in this population.

Castro, P., Zaman, S., & Holland, A. 
Abstract: People with Down's syndrome (DS) are at high risk for developing Alzheimer's disease (AD) at a relatively young age. This increased risk is not observed in people with intellectual disabilities for reasons other than DS and for this reason it is unlikely to be due to non-specific effects of having a neurodevelopmental disorder but, instead, a direct consequence of the genetics of DS (trisomy 21). Given the location of the amyloid precursor protein (APP) gene on chromosome 21, the amyloid cascade hypothesis is the dominant theory accounting for this risk, with other genetic and environmental factors modifying the age of onset and the course of the disease. Several potential therapies targeting the amyloid pathway and aiming to modify the course of AD are currently being investigated, which may also be useful for treating AD in DS. However, given that the neuropathology associated with AD starts many years before dementia manifests, any preventative treatment must start well before the onset of symptoms. To enable trials of such interventions, plasma, CSF, brain, and retinal biomarkers are being studied as proxy early diagnostic and outcome measures for AD. In this systematic review, we consider the prospects for the development of potential preventative treatments of AD in the DS population and their evaluation.

Centre for Developmental Disability Health Victoria 
Dementia and Intellectual Disability – A guide to supporting people with intellectual disabilities through their journey with dementia: Online Learning for Disability Support Workers http://www.cddh.monash.org/online-learning/ 
Abstract: These are on-line learning modules for disability staff supporting people who were at risk of developing, or had already been identified as having, dementia. There are 46 minute modules in the series addressing key questions you may have when supporting someone with dementia. They cover helpful information related to dementia and ID. Module 1: Understanding dementia and intellectual disability; Module 2: Taking action – The role of the support worker in assessment; Module 3: Supporting someone with intellectual disability and dementia; and Module 4: Supporting people through environment and activity. There are also a series self-taking test questions.

Chao, S.F., McCallion, P., & Nickle, T. 
Abstract: Burnout has been considered important to understanding the well-being of workers in the intellectual disabilities (ID) field and the quality of services delivered to clients/consumers. However, little research has examined the psychometric properties and applicability to staff in ID services of one of the most widely used burnout measurements – the Human Services Survey version of the Maslach Burnout Inventory (MBI-HSS). Data were gathered using a mailed questionnaire comprising the MBI-HSS and demographic information. The sample consisted of 435 staff delivering direct care and working in out-of-home community placements for persons with ID in New York state. The factorial structure of the scale was examined using confirmatory and exploratory factor analysis. Internal consistency estimates of reliability of the MBI-HSS were determined using Cronbach’s alpha. Confirmatory factor analysis supported the MBI-HSS as an acceptable measure to evaluate burnout in ID services staff. However, the reliability statistics obtained for the Depersonalization (DP) sub-scale was much lower than what has been reported in studies with other staff populations. An exploratory factor analysis suggested that a four-factor solution, dividing the DP sub-scale into two factors, provided a somewhat better fit for the sample. The use of the MBI-HSS as an instrument for measuring burnout among ID workers has its attraction but also its limitations. In particular, the DP sub-scale should be used with caution because there appear to be wording issues for staff in ID settings that may lead to inconsistent responses.

Chapman, M., Lacey, H., & Jervis, N. 
Abstract: Dementia prevalence rates are higher amongst people with intellectual (learning) disabilities than the general population. People with Down's syndrome are at even greater risk of developing dementia and of developing dementia at an earlier age. This study, conducted as part of a wider service evaluation, explored community learning disability team perspectives on screening, pathways, training, information and supports developed to improve services for people with learning disabilities and dementia. A focus group was held with health and social care professionals working in community learning disability services. Thematic analysis was used to analyze the data. The dementia screening, pathways and processes had become embedded in practice, leading to a common framework, an efficient, multidisciplinary, proactive approach, earlier detection and diagnosis of dementia and identification of other health needs and issues. This avoided crisis situations supporting people to remain at home longer. Training and information were felt to improve care quality and reduce caregiver anxiety. People with intellectual disabilities and caregivers were involved to varying extents. External influences impacting on support included the availability, appropriateness, cost and effectiveness of different models of service provision. Service developments have been made as a result of the findings which suggest that dementia pathways and supports improve service provision and outcomes for people with intellectual disabilities. It is important to develop the evidence base on the effectiveness of different service models for people with intellectual disabilities and dementia.

Chaput, J.L. 
Housing people with Alzheimer disease as a result of Down syndrome: a quality of life comparison between group homes and special care units in long term care facilities. Master’s thesis, Department of City Planning, University of Manitoba (1998) 
Abstract: Report of study to determine which form of housing, group homes or special care units (SCUs), provided an enhanced quality of life for individuals with Down syndrome (DS) and Alzheimer disease (AD). Ten long term care (LTC) facilities with SCUs for people with AD in the Winnipeg, Canada area and
Chaput, J.L., Udell, L.


Abstract: The purpose of the study was to determine which form of housing, i.e., group homes or special care units (SCUs), provided a better quality of life for individuals with Alzheimer disease (AD) as a result of Down syndrome (DS). The study also provided Winserv Inc. (a non-profit housing organization that houses people with mental disabilities) with important information. Using the study results, Winserv Inc. was able to determine that their group homes were suitable to maintain individuals with DS and AD and that their group homes were more cost-effective than SCUs in terms of caregiving. Twenty caregivers from both group homes and SCUs were selected to participate in this study. Ten long term care (LTC) facilities with SCUs for people with AD were selected in the Winnipeg area and ten group homes for people with Down syndrome and AD were chosen in Winnipeg and across Canada. The results indicated that the group homes seemed to provide the best quality of life for people with AD as a result of Down syndrome because they provided a home-like environment and they operated according to a therapeutic philosophy of care. In addition, costs for caregiving seemed to be more economical in group homes than in SCUs because group homes utilized lower staff wages and medical costs.

Chaput, J.L.

Adults with Down syndrome and Alzheimer’s disease: Comparisons of services received in group homes and in special care units.*Journal of Gerontological Social Work*, 2002, 38, 197-211 [https://doi.org/10.1080/01634370108233521]

Abstract: An increasing number of people with Down syndrome are at risk of dementia from Alzheimer’s disease. Many reside in residential group homes. When they are affected by dementia, the challenge to agencies providing group homes is how to best provide continued housing and provide effective dementia-related care management. In the general population, long term care is typically provided in nursing facilities, often in special care units (SCUs). This study evaluated select factors found in group homes and SCUs to determine which is able to provide a better quality of life for people with Down syndrome affected by dementia. Interviews, using quality of life indicators, were conducted at 20 sites, equally selected from group homes and SCUs, on the basis of their experience with people with dementia. Results indicate that group homes can provide conditions associated with better quality of life and, additionally, operate with lower staffing costs due to the non-utilization of medical staff.

Chicoine, B., Rivelli, A., Fitzpatrick, V., Chicoine, L., Jia, G., Rzetksy, A.


Abstract: Given the current life expectancy and number of individuals living with Down syndrome (DS), it is important to learn common occurrences of disease conditions to maintain the developmental lifespan. This study analyzed data from a large cohort of individuals with DS in an effort to better understand these disease conditions, inform future screening practices, tailor medical care guidelines, and improve utilization of health care resources. This retrospective, descriptive study incorporated up to 28 years of data, compiled from 6078 individuals with DS and 30,326 controls matched on age and sex. Data were abstracted from electronic medical records within a large Midwestern health system. In general, individuals with DS experienced higher prevalence of testicular cancer, leukemias, moyamoya disease, mental health conditions, bronchitis and pneumonia, gastrointestinal conditions, thyroid disorder, neurological conditions, atlantoaxial subluxation, osteoporosis, dysphagia, diseases of the eyes/adnexa and of the ears/mastoid process, and sleep apnea, relative to matched controls. Individuals with DS experienced lower prevalence of solid tumors, heart disease conditions, sexually transmitted diseases, HIV, influenza, sinusitis, urinary tract infections, and diabetes. Similar rates of prevalence were seen for lymphomas, skin melanomas, stroke, acute myocardial infarction, hepatitis, cellulitis, and osteoarthritis. While it is challenging to draw a widespread conclusion about comorbidities in individuals with Down syndrome, it is safe to conclude that care for individuals with DS should not automatically mirror screening, prevention, or treatment guidelines for the general U.S. population. Rather, care for those with DS should reflect the unique needs and common comorbidities of this population.

Choi, P., Motl, R.W., & Agiovlasitis, S.


Abstract: Adults with intellectual disability (ID) have a higher rate of fall events than the general population. Consequently, interventions for reducing fall events and improving health are highly required for individuals with ID. One essential step towards effectively delivering fall prevention interventions among adults with ID involves evaluating their feasibility. This study examined the feasibility of a home-based exercise intervention, supplemented with behavioural change strategies, among individuals with ID living in residential settings. This study provided an 8-week intervention, consisting of a workshop for support workers and sessions for participants with ID, focusing on behavioural reward/s, education regarding fall prevention/exercise and exercise training. One week prior to and 1 week following such an intervention, such participants underwent multiple measurements for (1) physical performance, (2) self-efficacy for activity and (4) social support. Participants having ID (n = 33), support workers (n = 11) and one administrator participated in this study. There were no adverse events during the intervention, and the mean adherence rate was 70.8 ± 19.5%. Two participants with ID dropped out of the programme due to a lack of interest. The participants with ID significantly improved individual physical performance, self-efficacy for activity, fall efficacy and support from friends and support workers. Fall prevention interventions for adults with ID living in group-homes were highly promising for eventual large-scale implementation within such communities.

Cipriani, G., Danti, S., Carlesi, C., & DiFiorino, M.


Abstract: People with Down syndrome (DS) enjoy a longer life expectancy now than they ever have before and are therefore at greater risk of developing conditions associated with aging, including dementia. Authors undertook at review to explore the phenomenon of dementia in DS. Medline and Google Scholar searches were conducted for relevant articles, chapters, and books published until 2017. Search terms included Alzheimer’s disease, cognitive impairment, dementia, DS, and trisomy 21. Publications found through this indexed search were reviewed for further references. Authors concluded that virtually, all subject aged 35 to 40 show key neuropathologic changes characteristic of Alzheimer’s disease, but only a part of them show clinical signs of dementia, usually around the age of 50 years. Early signs of dementia in people with DS may be different from those experienced by the general population. Failure to recognize this can delay diagnosis and subsequent interventions.

Cipriani, G., Picchi, L., Dolciotti, C., & Bonuccelli, U.

Abstract: People with Down's syndrome (DS) have an increased risk of developing Alzheimer's disease (AD) during middle age. Both disorders can present with a decline in cognitive skills and behavioral symptoms. Therefore, dementia, particularly in its early stages, can be difficult to diagnose in this population. We conducted a search of electronic databases for literature on the relationship between AD and DS. The key words used were: "Down syndrome", "Alzheimer's disease", "dementia", and "mental retardation". AD onset has been reported as early as age 30 in individuals with DS, and there is a dramatic increase in prevalence rates in older age groups. This trend reflects increased survival of persons with DS probably as a result of advances in medical treatment and improved living conditions. Even with careful clinical assessment, it can be very difficult to identify early symptoms of dementia when it is superimposed on a background of intellectual disability. The reasons include the wide intra-individual variability in cognitive functioning and difficulties involved in establishing baseline levels of the premorbid condition. Many frontal lobe-related symptoms usually associated with later stages of dementia in the general population are commonly seen in the early stage of the dementia that develops in adults with DS. After onset, the clinical symptoms of dementia progress rapidly in all subjects with DS. Research suggests that the presentation of dementia in people with DS may differ from that typical of AD in the later stages. A number of factors have been identified, and data from these researches rather than memory. DS can be best understood as a complex syndrome of genetic origin that has protean neurobiological consequences and numerous clinical characteristics. [Note: article text in Italian]


Abstract: The number of people with intellectual disability living into old age and developing dementia continues to increase. Dementia presents a wide range of challenges for staff due to progressive deterioration. This article presents the findings from a narrative literature review of professional carers’ experiences of caring for individuals with intellectual disability and dementia. Seven electronic databases were searched using Boolean operators and truncation to identify relevant literature. Search results were combined and narrowed to articles relevant to staff working with individuals with intellectual disability and dementia, and 14 articles met the criteria for review. Themes outlined in the review include staff knowledge of dementia, staff training in dementia, caregiving, challenging behavior, pain management, mealtime support and coping strategies. Overall carers must review and adjust their care delivery and support to people with intellectual disability and dementia, not only in terms of identifying and responding to their health needs but also through collaborative team working within and across services.


Abstract: The authors endeavored to explore nurses’ experiences of caring for older people with intellectual disability and dementia. Ageing and dementia prevalence is increasing along with the life expectancy of people with intellectual disability. As a population group, people with intellectual disability have a high prevalence of dementia, which is higher within the subpopulation of Down syndrome. People with intellectual disability live in residential care, community or residential settings, and nurses are required to adapt their practices to meet the changed needs of the individual. A qualitative Husserlian descriptive phenomenological methodology was undertaken by the researchers so as to be able to become absorbed in the quintessence of meaning and explore nurses’ experience of working with older people with intellectual disability and dementia. Ethical approval was obtained, and data were collected using semistructured interviews (n = 11). Interviews were transcribed and analyzed using Colaizzi's framework for data analysis. The authors extracted three key themes were identified: ‘knowledge of dementia’, ‘person-centred care’ and ‘transitions within the service’. The study highlights the need for proactive planning, life story books of the patient, and funding to support client and staff. The authors concluded that overall, the study highlights the importance of knowing the person, supporting the individual and recognizing presenting behaviors as part of the control of the individual. The article presents the experiences of nurses caring for the older person with intellectual disability and dementia. Transitions are often very difficult for both the person and their peers, and they experience benefit from the efforts of a multidisciplinary team facilitating a person-centered approach.


Abstract: People with an intellectual or developmental disability (IDD) experience complex age-related issues, including dementia-related disorders, at higher rates and earlier ages than the general population. Increased support needs of this subpopulation can strain caregivers and existing community supports. Patterns of resource awareness and utilization and unmet needs are not well understood for this subpopulation. A collaboration of the Massachusetts Council on Aging (MCOA), the Massachusetts Department of Developmental Services (DDS), and the Center for Developmental Disability Evaluation and Research (ODDER) at University of Massachusetts Medical School conducted a needs assessment with caregivers of people with intellectual and developmental disabilities to assess awareness and utilization of community-based resources, and unmet needs including for caregiver supports. Home visits, including an environmental assessment, and interviews with caregivers of 95 adults with dementia-related diagnoses were conducted. Interviews asked about changes in the adult's condition since diagnosis including a needs assessment about the dementia-related knowledge and training, care confidence levels, perceived barriers and/or concerns to care provision and resource use. About half of the caregivers reported significant changes in the person’s skills, function, and memory since diagnosis, as well as worsening of the person’s gait, continence, and swallowing. 78% of caregivers reported feeling confident providing care currently and 66% were confident about providing future care. 100% of the respondents found outreach provided by a nurse practitioner helpful. Caregiver concerns included lack of suitable day programming, future planning resources, and caregiver burnout/stress. Most caregivers were currently aware of some local resources but with scattered use. Environmental assessments indicated most homes, while accessible, warranted additional lighting and clutter removal. Caregivers requested additional training in addressing the behavioral and mental health needs. Findings suggest a need for increased collaboration across the intellectual and developmental disability and aging/human services systems, and additional training and resource navigation guides for caregivers.


Abstract: Widespread inquiry identified 378 adults with Down’s syndrome resident in Leicestershire, England. The immediate carer of 351 of these (92.8%) was interviewed for the purpose of establishing a past history of seizures, including the age at which the seizures began. The immediate carer was also invited to provide information to enable the completion of an Adaptive Behaviour Scale (A.B.S.) rating. Individuals with a history of seizures were divided into two groups on the basis of whether or not seizures commenced prior to or after age 35 years. Two control groups of individuals with Down’s syndrome, but without a history of seizures were selected. Adaptive Behaviour Scale scores for those in whom seizures commenced at a younger age were similar to those who had no recorded history of seizures. However, in those in whom seizures began in later life, scores on all domains of the A.B.S. were significantly reduced compared to both young epileptic patients and their...
controls. Adaptive Behaviour Scale scores for the older control group held an intermediate position, suggesting that late-onset epilepsy may be a late manifestation of a dementing process. A clinical diagnosis of dementia recorded in the case records was significantly associated with the presence of late-onset epilepsy. This is supportive of the hypothesis that late-onset epilepsy in individuals with Down’s syndrome is associated with Alzheimer’s disease.

Coppus, A.M.W

People with intellectual disability: What do we know about adulthood and life expectancy?


Abstract: Increases in the life expectancy of people with Intellectual Disability have followed similar trends to those found in the general population. With the exception of people with severe and multiple disabilities or Down syndrome, the life expectancy of this group now closely approximates with that of the general population. Middle and old age, which until 30 years ago were not recognized in this population, are now important parts of the life course of these individuals. Older adults with Intellectual Disabilities form a small, but significant and growing proportion of older people in the community. How these persons grow older and how symptoms and complications of the underlying cause of the Intellectual Disability will influence their life expectancy is of the utmost importance.


Dementia and mortality in persons with Down's syndrome.


Abstract: Numerous studies have documented that persons with Down’s syndrome (DS) are at an increased risk of Alzheimer's disease (AD). However, at present it is still not clear whether or not all persons with DS will develop dementia as they reach old age. We studied 506 people with DS, aged 45 years and above. A standardized assessment of cognitive, functional and physical status was repeated annually. If deterioration occurred, the patients were examined and the differential diagnosis of dementia was made according to the revised Dutch consensus protocol and according to the ICD-10 Symptom Checklist for Mental Disorders. We compared our findings with those reported in the literature. The overall prevalence of dementia was 16.8%. Up to the age of 60, the prevalence of dementia doubled with each 5-year interval. Up to the age of 49, the prevalence is 8.9%, from 50 to 54, it is 17.7%, and from 55 to 59, it is 32.1%. In the age category of 60 and above, there is a small decrease in prevalence of dementia to 25.6%. The lack of increase after the age of 60 may be explained by the increased mortality among elderly demented DS patients (44.4%) in comparison with non-demented patients (10.7%) who we observed during a 3.3-year follow-up. There was no decrease in incidence of dementia in the age group of 60 and above. Our findings are very similar to those published in the literature. Patients with dementia were more frequently treated with antiepileptic, antipsychotic and antidepressant drugs. The history of depression was strongly associated with dementia. The authors concluded that their study is one of the largest population-based studies to date. We found that despite the exponential increase in prevalence with age, the prevalence of dementia in the oldest persons with DS was not higher than 25.6%.


Early age at menopause is associated with increased risk of dementia and mortality in women with Down syndrome.


Abstract: In a prospective longitudinal cohort study of dementia and mortality in persons with Down syndrome aged 45 years and older, 85 postmenopausal women were followed for a mean follow-up time of 4.3 years (range 0.0 to 7.4 years). The effect of age at menopause on age at diagnosis of dementia and survival was estimated using correlation analysis and Cox Proportional Hazard Model. We found a significant correlation between age at menopause and age at diagnosis of dementia (rho=0.52; p<0.001), and between age at menopause and age at death (rho=0.49; p=0.001). Early age at menopause is associated with a 1.8 fold increased risk of dementia: Hazard Ratio (HR): 1.82 (95%Confidence Interval (CI): 1.31-2.52) and with risk of death: HR: 2.05 (95%CI: 1.33-3.16). Our study suggests that age at menopause in women with Down syndrome is a determinant of age at onset of dementia and mortality.

Cooper, S.A.

High prevalence of dementia among people with learning disabilities not attributable to Down's syndrome.


Abstract: For many years, it has been known that dementia can occur in people with learning disabilities, but there have been few research studies. Studies that do quote rates for dementia show these to be high, but this important fact has received remarkably little attention. Comprehensive psychiatric and medical assessments were undertaken on the whole population (ascertained as far as is possible) of people with learning disabilities aged 65 years and over living in Leicestershire, UK (N=134), in order to ascertain rates of DCR defined dementia, and associated factors. Dementia was diagnosed in 21.6%, against an expected prevalence of 5.7%, for a group with this age structure. The rate of dementia increased in successive age cohorts: 16.6% aged 65-74 years; 23.5% aged 65-84 years; and 70% aged 85-94 years. People with dementia tended to be older, female, with more poorly controlled epilepsy, a larger number of
additional physical disorders, less likely to be smokers and had lower adaptive behavior scores than did the elderly people without dementia. They were more likely to live in health service accommodation. Dementia occurs at a much higher rate among elderly people with learning disabilities than it does among the general population; this is independent of the association between dementia and Down’s syndrome. Whether this relates etiologically to genetics, lack of brain ‘reserve’ or history of brain damage is yet to be determined.

Cooper, S.-A., & Prasher, V.

Abstract: Dementia commonly occurs in elderly people with intellectual disability, especially those with Down’s syndrome. The non-cognitive symptoms of dementia can be of greater significance to individuals and carers than the cognitive changes caused by this condition. It is not known whether there are differences between people with Down’s syndrome and those with intellectual disability of other causes with regard to the prevalence of such symptoms. The present study was undertaken to draw a comparison between a group with Down’s syndrome and dementia (n= 19), and a group with intellectual disability of other causes and dementia (n= 26). Maladaptive behaviours and psychiatric symptomatology were assessed in both groups. The group with Down’s syndrome had a higher prevalence of low mood, restlessness/excessive overactivity, disturbed sleep, being excessively uncooperative and auditory hallucinations. Aggression occurred with greater frequency in those subjects with intellectual disability of other causes. These findings are of epidemiological importance in terms of service planning and understanding psychiatric presentation.

Cooper, S.-A., & van der Speck
Epidemiology of mental ill health in adults with intellectual disabilities. *Current Opinion in Psychiatry,* 2009, Sep, 22(5), 431-436. doi:10.1097/YCO.0b013e328323e2a1e

Abstract: Adults with intellectual disabilities experience higher rates of mental ill health than the general population. Despite this, the epidemiological knowledge base remains limited. The purpose of this article is to review mental health epidemiological studies relevant to adults with intellectual disabilities, published since January 2008. Several studies have aimed to build the epidemiological evidence base, particularly with regards to problem behaviours, which appear to be remitting-relapsing conditions rather than necessarily being chronic. Most of such work confirms prevalence and incidence rates, and conducts exploratory analyses to determine factors independently related to mental ill health. Down syndrome protects against problem behaviours and mental ill health (except dementia that occurs at a higher rate), whereas epilepsy does not appear to affect risk for mental ill health. Dementia is four times more common in older persons with intellectual disabilities without Down syndrome than in the general population. Persons with borderline intellectual disabilities also experience higher rates of mental ill health than the general population, but receive fewer treatments.


Abstract: The Patient Protection and Affordable Care Act added a new Medicare benefit, the Annual Wellness Visit (AWV), effective January 1, 2011. The AWV requires an assessment to detect cognitive impairment. The Centers for Medicare and Medicaid Services (CMS) elected not to recommend a specific assessment tool because there is no single, universally accepted screen that satisfies all needs in the detection of cognitive impairment. To provide primary care physicians with guidance on cognitive assessment during the AWV, and when referral or further testing is needed, the Alzheimer’s Association convened a group of experts to develop recommendations. The resulting Alzheimer’s Association Medicare Annual Wellness Visit Algorithm for Assessment of Cognition includes review of patient Health Risk Assessment (HRA) information, patient observation, unstructured queries during the AWV, and use of structured cognitive assessment tools for both patients and informants. Widespread implementation of this algorithm could be the first step in reducing the prevalence of missed or delayed dementia diagnosis, thus allowing for better healthcare management and more favorable outcomes for affected patients and their families and caregivers.

Cosgrave, M.P., Tyrell, J., McCarron, M., Gill, M., & Lawlor, B.A.

Abstract: In a cross-sectional study of aggression, and adaptive and maladaptive behavior in 128 subjects with Down’s syndrome (DS), 29 of whom had dementia, the current authors found that the presence of dementia was not predictive of aggression or maladaptive behavior. However, the level of adaptive behavior was shown to be lower in subjects with dementia, and in those with lower levels of cognitive functioning, as measured on a rating instrument, the Test for Severe Impairment. Although the presence of aggressive behaviors is not higher in subjects with dementia and DS on cross-sectional review, it remains to be seen whether aggression will increase in individual cases with the onset or progression of dementia. The decline in adaptive behavior shown in the present study confirms the findings of previous studies and indicates a direction for service development for persons with the dual diagnosis of dementia and DS.

Cosgrove, M.P., Tyrell, J., McCarron, M., Gill, M., & Lawlor, B.A.

Abstract: Menstrual status and the age of menopause were investigated in 143 Irish females with Down’s syndrome (DS). The average age of menopause in 42 subjects (44.7 years) was younger than in the general population. The age at onset of dementia correlated with the age of menopause. This finding may be a manifestation of accelerated ageing in DS or point to oestrogen deficiency being an independent risk factor for the development of Alzheimer’s.

Courtenay, K., Jokinen, N.S., & Strydom, A.

Abstract: Authors conducted a systematic review of the available Dutch, English, and German language literature for the period 1997–2008 on the current knowledge on social-psychological and pharmacological caregiving with respect to older adults with intellectual disabilities (ID) affected by dementia. Authors note that caregiving occurs on a personal level between the person and their caretaker and organizational and interorganizational supports have an impact on the quality of care provided. However, the lack of robust evidence to meet the needs of adults with ID affected by dementia means that service organizations often have to extrapolate from the evidence base of dementia care practices in the general population. The review showed that concerns over staff burden, behavioral interventions, and staff training, and applications of models of care were emerging, but were not systematically studied. Authors noted that pharmacological agents and nonpharmacological, psychosocial techniques were being used to assist carers manage behavior, but the evidence base of both nonpharmacological and pharmacological interventions that can help people with ID and dementia and their carers is insufficient because of the absence of systematic and robust studies. The authors note a need for an international research agenda that begins to address gaps in knowledge. With more adults projected to be affected by dementia, a robust evidence-based body of literature on dementia care in people with ID can help with planning for and providing quality dementia-capable services.

Cox, S.

Abstract: Publication details some 10 case studies of housing options and accommodations for persons affected by dementia (and applicable to adults with dementia).
neuropsychologic deficits in this brain region. Neuropsychologic testing parietal lobe has reductions in metabolic function before the onset of positron emission tomography in Alzheimer's disease have shown that the neurochemical data suggest that a cholinergic deficiency must be present for Alzheimer's disease are at increased risk of developing Alzheimer's disease, similarities. Genetic studies indicate that near relatives of patients with Down's syndrome and those with Alzheimer's disease reveal striking Annuals of Internal Medicine, Cutler, N.R., Heston, L.L., Davies, P., Haxby, J.V., & Schapiro, M.B.

Cognitive changes in memory precede those in praxis in aging persons with Down syndrome. Cognitive decline due to aging among persons with Down's syndrome. The aim was to determine whether or not there was a specific sequence of cognitive changes over a 3-year period for the test measures. When compared with a young group (17-39years of age at the start), an old group of persons with Down syndrome (40-58years of age at the start) showed small but statistically significant changes over time suggestive of "pre-clinical signs" of dementia. When the data were sorted into 4 subgroups on the basis of age, a more detailed analysis revealed that the subgroup that was 50 years of age and older at the start showed changes in scores which were of a magnitude more clearly indicative of early dementia on the test measures. Deterioration in learning/ memory functions began at a mean age of 54.2 years, followed later by deterioration in movement-related functions (praxis) at a mean age of 56.9 years. Deterioration in scores on an informant-based behavior rating scale (MOSES) occurred at an intermediate age of 55.0 years. The results provide preliminary support for the hypothesis that persons with Down syndrome who are 50 years of age and older may develop a specific sequence of functional changes during the early stage of dementia. They also illustrate ways in which small sample norms can be effectively used to increase the practical usefulness of tests intended to evaluate dementia in persons with intellectual disabilities.

Das J.P., Davis B., Alexander J., Rauno K.P. & Naglieri J.A.


Abstract: This study examined decline in cognitive functions in individuals with Down syndrome (DS) over the age of 40 in comparison to participants of the same age and comparable mental handicap without Down syndrome (NonDS). Both DS (n = 32) and NonDS (n = 31) samples were divided into "younger" (40-49years) and "older" (50-62) groups. Cognitive processes were examined by tests of general intellectual functioning (Dementia Rating Scale, Peabody Picture Vocabulary Test-Revised, and the Matrix Analogies Test-Expanded form), as well as planning, attention, simultaneous, and successive processing tests taken from Das-Naglieri Cognitive Assessment System. The older individuals with Down syndrome performed more poorly than those in the other three groups. The differences were particularly evident in tasks requiring cognitive skills than do younger patients. Brain metabolic function is excessively reduced in the demented adults with Down's syndrome.

Dalton, A.J., Fodor, B.L., Patti, P.J., Tsiouris, J.A., & Mehta, P.D.

Abstract: This report describes the results of five studies aimed at evaluating the usefulness, reliability, and validity of the Multidimensional Observation Scale for Elderly Subjects (MOSES) in the assessment of change in ageing persons with intellectual disability. Three hundred and thirty-six individuals with an average age of 49.8 years, including an equal number of men and women, were participants in one or more of the five studies. There were 220 participants with Down syndrome, 81 persons without Down syndrome with intellectual disability, and 35 persons from the general ageing population who were clinically diagnosed with Alzheimer's disease using NINCD/ADRDA criteria. Persons with Down syndrome 40 years of age and older could be distinguished from their diagnosed peers with Down syndrome by statistically significant poorer scores on the MOSES, with those 50 years of age and older showing the worst scores. Comparisons of adults with intellectual disability diagnosed with dementia of the Alzheimer type (DAT), using DSM-IV criteria, with or without Down syndrome, as well as a comparison of persons with clinically diagnosed depression, provided evidence that subtests of the MOSES were sensitive to DAT but less so to depression. These clinical groups also showed significantly poorer scores on the MOSES when compared with those of the normative sample. It was concluded that the MOSES is a behavioural observation scale that can provide useful information in clinical settings as well as in research.

Croy, N., Adams, M., Shorten, N., & Langdon, P. E.


Abstract: This study investigated whether a personalized life story book and rummage box enhanced well-being and led to changes in behavior for people with Down syndrome (DS) who have dementia. A randomized single case series design was used with five participants who had DS and a diagnosis of dementia. Participants were invited to take part in three conditions at random (i) life story book, (ii) rummage box and (iii) no-intervention condition. The two reminiscence conditions were significantly associated with enhanced well-being as compared to the no-intervention condition. However, for one participant, the life story book was associated with significantly higher well-being, while for another participant, the rummage box was associated with significantly higher well-being, suggesting some participants may prefer one method over another. Personalized life story books and rummage boxes are associated with higher levels of well-being for people with DS and dementia.

Cutler, C.E., Kramer, J., & Mutchler, J.E.


Abstract: Family care provision is the norm for adults with intellectual and developmental disabilities (I/DD), even as they and their support networks grow older. As families age together, the role of primary carer frequently transitions from the parent to a sibling, as aging parents die or become too frail to provide continued support. This paper explores the transition in care from the perspective of a sibling who has replaced parents as the primary carer for an individual aging with I/DD. Data are drawn from semi-structured, in-depth interviews with a sample of adults over age 40, living in the United States, and caring for a sibling with I/DD(n = 15). Data were analyzed using a constant comparative qualitative approach. Results reveal themes impacting the adjustment to the role of primary carer, the extent to which aging transformed the content of care needs, the importance of planning, and the availability of supplementary support. First, we found that the aging process permeated the caregiving role, requiring ongoing modification of that role as a result of the aging of the adult with I/DD and their family support system. A second key theme is that planning shapes the adjustment to the carer role. A third key theme is that the support systems surrounding the sibling dyad contribute to successful adjustment to care providing. Findings from this study underscore the need to develop long-term services and supports as well as educational resources that accommodate this population of carers as they age together with their sibling with I/DD.

Crook, N., Adams, M., Shorten, N., & Langdon, P. E.


Abstract: This study investigated whether a personalized life story book and rummage box enhanced well-being and led to changes in behavior for people with Down syndrome (DS) who have dementia. A randomized single case series design was used with five participants who had DS and a diagnosis of dementia. Participants were invited to take part in three conditions at random (i) life story book, (ii) rummage box and (iii) no-intervention condition. The two reminiscence conditions were significantly associated with enhanced well-being as compared to the no-intervention condition. However, for one participant, the life story book was associated with significantly higher well-being, while for another participant, the rummage box was associated with significantly higher well-being, suggesting some participants may prefer one method over another. Personalized life story books and rummage boxes are associated with higher levels of well-being for people with DS and dementia.
planning and attention. Authors note the possibility of using these tests as indicators of the early signs of Alzheimer's disease.

Davis, D.R.
A parent’s perspective
In M.P. Janicki & A.J. Dalton (Eds.), Dementia, aging, and intellectual disabilities. pp. 42-50
Abstract: Book chapter that provides an account of the experiences of a family with an adult son with Down syndrome who eventually succumbs to dementia of the Alzheimer’s type. Includes a discussion of the difficult early years of the son’s life and the challenges the family faced as he aged. It also examines the family’s problems in recognizing that their son was experiencing the onset of dementia and his gradual decline until his death at age 46.

Day, K., Carreon, D., & Stump, C.
The therapeutic design of environments for people with dementia: A review of the empirical research
Gerontologist, 2000, 40, 397-416
Abstract: Design of the physical environment is increasingly recognized as an important aid in caring for people with dementia. This article reviews the empirical research on design and dementia, including research concerning facility planning (relocation, respite and day care, special care units, group size), research on environmental attributes (noninstitutional character, sensory stimulation, lighting, safety), studies concerning building organization (orientation, outdoor space), and research on specific rooms and activity spaces (bathrooms, toilet rooms, dining rooms, kitchens, and resident rooms). The analysis reveals major themes in research and characterizes strengths and shortcomings in methodology, theoretical conceptualization, and application of findings.

Davies, M., McGilade, A., & Bickerstaff, D.
A needs assessment of people in the Eastern Health and Social Services Board (Northern Ireland) with intellectual disability and dementia
Abstract: Article details a study undertaken by the Eastern Health and Social Services Board (Northern Ireland) which aimed to identify the number of people with intellectual disability within this area who were diagnosed with or were thought to have dementia. The objectives of the study were to collate demographic details and to profile the needs of this group. Key workers were asked to provide this information and were invited to comment on gaps in existing service provision and on future needs. A number of findings emerged: diagnostic services were patchy; people with dementia were living in a range of residential settings; carers wished to care for their clients for as long as practically possible, but they required extra resources and training to do so; and some individuals with an intellectual disability were excluded from elderly services. A report was compiled incorporating 12 recommendations.

Davys, D., & Haigh, C.
Older parents of people who have a learning disability: Perceptions of future accommodation needs
doi:10.1111/j.1468-3156.2007.00447.x
Abstract: The aim of this qualitative study was to provide an insight into the perceptions of older parents of adults with intellectual disability on the future accommodation needs of their adult children. Semi-structured interviews were used to seek parental awareness of residential options available, concerns in relation to future accommodation and the preferred accommodation options for their offspring. Four couples who shared the family home with an adult who has an intellectual disability took part in the study and data were analyzed using a step-by-step form of content analysis as described by Bumard [Nurse Educ Today 11 (1991), 461]. Emergent themes from transcripts were then organized into main categories. The results of this study suggest that older parents are dissatisfied with both statutory and private services, that they have concerns for their other children, and their own aging. Being a parent to a person who has an intellectual disability is seen to be a difficult task, were worried about what the adult’s brothers and sisters may have to do in the future, but also wanted to provide support at home for as long a possible. Of the parents who participated in this study, three couples wanted to maintain their adult child at home for as long as possible and the parents who were actively seeking accommodation outside the home expected to be involved in all aspects of their daughter’s care for the long-term future.

de Franca Bram, J.M., Talib, L.L., Giroud Joaquim, H.P., Carvalho, C.L., Gattaz, W.G., & Forlenza, O.V.
Alzheimer’s disease-related biomarkers in aging adults with Down syndrome: systematic review
Current Psychiatry Research and Reviews, 2019, 15(1), 49-57. DOI: 10.2174/157340051566619021215285
Abstract: Down syndrome (DS) is associated with a high prevalence of cognitive impairment and dementia in middle age and older adults. Given the presence of common neuropathological findings and similar pathogenic mechanisms, dementia in DS is regarded as a form of genetically determined, early-onset AD. The clinical characterization of cognitive decline in persons with DS is a difficult task, due to the presence intellectual disability and pre-existing cognitive impairment. Subtle changes that occur at early stages of the dementing process may not be perceived clinically, given that most cognitive screening tests are not sensitive enough to detect them. Therefore, biological markers will provide support to the diagnosis of DS-related cognitive impairment and dementia, particularly at early stages of this process. To perform a systematic review of the literature on AD-related biomarkers in DS. We searched PubMed, Web of Science and Cochrane Library for scientific papers published between 2008 and 2018 using as primary mesh terms ‘Down’, ‘Alzheimer’, ‘biomarker’. 79 studies were retrieved, and 39 were considered eligible for inclusion in the systematic review: 14 post-mortem studies, 10 neuroimaging, 4 addressing cerebrospinal fluid biomarkers, and 11 on peripheral markers. There is consistent growth in the number of publication in this field over the past years. Studies in DS-related dementia tend to incorporate many of the diagnostic technologies that have been more extensively studied and validated in AD. In many instances, the study of CNS and peripheral biomarkers reinforces the presence of AD pathology in DS.

A new cognitive evaluation battery for Down syndrome and its relevance for clinical trials
Abstract: The recent prospect of pharmaceutical interventions for cognitive impairment of Down syndrome (DS) has boosted a number of clinical trials in this population. However, running the trials has raised some methodological challenges and questioned the prevailing methodology used to evaluate cognitive functioning of DS individuals. This is usually achieved by comparing DS individuals to matched healthy controls of the same mental age. We propose a new tool, the TESDAD Battery that uses comparison with age-matched typically developed adults. This is an advantageous method for probing the clinical efficacy of DS therapies, allowing the interpretation and prediction of functional outcomes in clinical trials. In our DS population the TESDAD battery permitted a quantitative assessment of cognitive defects, which indicated language dysfunction and deficits in executive function, as the most important contributors tother cognitive and adaptive behavior outcomes as predictors of functional change in DS. Concretely, auditory comprehension and functional academics showed the highest potential as end-point measures of therapeutic intervention for clinical trials: the former as a cognitive key target for therapeutic intervention, and the latter as a primary functional outcome measure of clinical efficacy. Our results also emphasize the need to explore the modulating effects of IQ, sex, and age on cognitive enhancing treatments. Noticeably, women performed significantly better than men of the same age and IQ in most cognitive tests, with the most consistent differences occurring in memory and executive functioning and negative trends rarely emerged on quality of life linked to the effect of age after adjusting for IQ and sex. In sum, the TESDAD battery is a useful neurocognitive tool for probing the clinical efficacy of experimental therapies in interventional studies in the DS population suggesting that age-matched controls are advantageous for determining normalization of DS.

De Vreese, L.P., Gomiero, T., Uberti M., De Bastiani, E., Weger, E.,
Mantesso, U., & Marangoni, A.

Functional abilities and cognitive decline in adult and aging intellectual disabilities. Psychometric validation of an Italian version of the Alzheimer's Functional Assessment Tool (AFAST-I), designed for informant-based assessment of the degree of impairment and of assistance required in seven basic daily activities in adult/elderly people with intellectual disabilities (ID) and (suspected) dementia, and a pilot analysis of its clinical significance with traditional statistical procedures and with an artificial neural network. AFAST-I was administered to the professional caregivers of 61 adults/seniors with ID with a mean age (± SD) of 53.4 (± 7.7) years (36% with Down syndrome).

Internal consistency (Cronbach’s a coefficient), inter/intra-rater reliabilities (intra-class coefficients, ICC) and concurrent, convergent and discriminant validity (Pearson’s r coefficients) were computed. Clinical significance was probed by analysing the relationships among AFAST-I scores and the Sum of Cognitive Scores (SCS) and the Sum of Social Scores (SOS) of the Dementia Questionnaire for Persons with Intellectual Disabilities (DMR-I) after standardization of their raw scores in equivalent scores (ES). An adaptive artificial system (AutoContractive Maps, AutoCM) was applied to all the variables recorded in the study sample, aimed at uncovering which variable occupies a central position and supports the entire network made up of the remaining variables interconnected among themselves with different weights. AFAST-I shows a high level of internal homogeneity with a Cronbach’s a coefficient of 0.92. Inter-rater and intra-rater reliabilities were also excellent with ICC correlations of 0.96 and 0.93, respectively. The results of the analyses of the different AFAST-I validities all go in the expected direction: concurrent validity (r=0.87 with ADL; convergent validity (r=0.63 with SCS; r=0.61 with SOS); discriminant validity (r=0.21 with the frequency of occurrence of dementia-related Behavioral Excesses of the Assessment for Adults with Developmental Disabilities (AAD-I)). In our sample age and gender do not correlate with the scale and comparing the distribution of the AFAST-I and DMR-SCS and DMR-SOS expressed as ES, it appears that memory disorders and temporal and spatial disorientation (SCS) precede the loss of functional abilities, whereas changes in social behaviour (SOS) are less specific in detecting cognitive deterioration sufficient to provoke functional disability and vice versa. The results of AutoCM analysis reveal that the hub (core) of the entire network is represented by the functional domain ‘personal/oral hygiene’ in the entire study sample and ‘use of toilet’ in a subgroup of subjects who obtained an ES equal to 0 at DMR-SCS. These results confirm the reliability and validity of AFAST-I and emphasise the complexity of the relationship among functional status, cognitive functioning and behaviour also in adults/seniors with ID.

De Vreese, L.P., Mantesso, U., De Bastiani, E., Marangoni, A.C., Weiner, M.F. & Gomiero, T.

Psychometric evaluation of the Italian version of the AADS questionnaire: A caregiver-rated tool for the assessment of behavioral deficits and excesses in people with intellectual disabilities and dementia. Clinical validity was assessed by comparing AADS-I scores obtained by subjects with and without dementia using the Mann-Whitney U test. A total of 40 adults/older people with ID at five ID-specific centers in the province of Trento and Cremona participated in the study. Findings show optimal levels of internal consistency (α = 0.80) and confirm the factors identified in the Spanish validation study (symptoms of discomfort, positive social interaction and depression). The scale has high inter-rater reliability (ICC = 0.95) and good test-retest reliability (ICC = 0.89). The total AQUALID score correlates significantly with the AADS sub-scores for behavioral excesses, but does not differ between individuals with and without dementia, though two out of the three identified factor scores are significantly higher in the dementia subgroup. The authors conclude that the Italian version of the AQUALID is a reliable and valid instrument for estimating quality of life in aging adults with ID and dementia.


Measuring quality of life in intellectually disabled persons with dementia using the Italian version of the Quality of Life in Late-Stage Dementia (QUALID) scale. Journal of Alzheimer's Disease and Parkinsonism, 2012, 2:104e.

De Vreese, L.P., U., De Bastiani, E., Weger, E., Marangoni, A.C. & Gomiero, T.

A psychometric validation of the Italian version of the Alzheimer's Functional Assessment Tool scale (AFAST-I), designed for informant-based assessment of the degree of impairment and of assistance required in seven basic daily activities in adult/elderly people with intellectual disabilities (ID) and (suspected) dementia, and a pilot analysis of its clinical significance with traditional statistical procedures and with an artificial neural network. AFAST-I was administered to the professional caregivers of 61 adults/seniors with ID with a mean age (± SD) of 53.4 (± 7.7) years (36% with Down syndrome).

Internal consistency (Cronbach’s a coefficient), inter/intra-rater reliabilities (intra-class coefficients, ICC) and concurrent, convergent and discriminant validity (Pearson’s r coefficients) were computed. Clinical significance was probed by analysing the relationships among AFAST-I scores and the Sum of Cognitive Scores (SCS) and the Sum of Social Scores (SOS) of the Dementia Questionnaire for Persons with Intellectual Disabilities (DMR-I) after standardization of their raw scores in equivalent scores (ES). An adaptive artificial system (AutoContractive Maps, AutoCM) was applied to all the variables recorded in the study sample, aimed at uncovering which variable occupies a central position and supports the entire network made up of the remaining variables interconnected among themselves with different weights. AFAST-I shows a high level of internal homogeneity with a Cronbach’s a coefficient of 0.92. Inter-rater and intra-rater reliabilities were also excellent with ICC correlations of 0.96 and 0.93, respectively. The results of the analyses of the different AFAST-I validities all go in the expected direction: concurrent validity (r=0.87 with ADL; convergent validity (r=0.63 with SCS; r=0.61 with SOS); discriminant validity (r=0.21 with the frequency of occurrence of dementia-related Behavioral Excesses of the Assessment for Adults with Developmental Disabilities (AAD-I)). In our sample age and gender do not correlate with the scale and comparing the distribution of the AFAST-I and DMR-SCS and DMR-SOS expressed as ES, it appears that memory disorders and temporal and spatial disorientation (SCS) precede the loss of functional abilities, whereas changes in social behaviour (SOS) are less specific in detecting cognitive deterioration sufficient to provoke functional disability and vice versa. The results of AutoCM analysis reveal that the hub (core) of the entire network is represented by the functional domain ‘personal/oral hygiene’ in the entire study sample and ‘use of toilet’ in a subgroup of subjects who obtained an ES equal to 0 at DMR-SCS. These results confirm the reliability and validity of AFAST-I and emphasise the complexity of the relationship among functional status, cognitive functioning and behaviour also in adults/seniors with ID.


Psychometric evaluation of the Italian version of the AADS questionnaire: A caregiver-rated tool for the assessment of behavioral deficits and excesses in people with intellectual disabilities and dementia. Clinical validity was assessed by comparing AADS-I scores obtained by subjects with and without dementia using the Mann-Whitney U test. A total of 40 adults/older people with ID at five ID-specific centers in the province of Trento and Cremona participated in the study. Findings show optimal levels of internal consistency (α = 0.80) and confirm the factors identified in the Spanish validation study (symptoms of discomfort, positive social interaction and depression). The scale has high inter-rater reliability (ICC = 0.95) and good test-retest reliability (ICC = 0.89). The total AQUALID score correlates significantly with the AADS sub-scores for behavioral excesses, but does not differ between individuals with and without dementia, though two out of the three identified factor scores are significantly higher in the dementia subgroup. The authors conclude that the Italian version of the AQUALID is a reliable and valid instrument for estimating quality of life in aging adults with ID and dementia.


Psychometric evaluation of the Italian version of the AADS questionnaire: A caregiver-rated tool for the assessment of behavioral deficits and excesses in people with intellectual disabilities and dementia. Clinical validity was assessed by comparing AADS-I scores obtained by subjects with and without dementia using the Mann-Whitney U test. A total of 40 adults/older people with ID at five ID-specific centers in the province of Trento and Cremona participated in the study. Findings show optimal levels of internal consistency (α = 0.80) and confirm the factors identified in the Spanish validation study (symptoms of discomfort, positive social interaction and depression). The scale has high inter-rater reliability (ICC = 0.95) and good test-retest reliability (ICC = 0.89). The total AQUALID score correlates significantly with the AADS sub-scores for behavioral excesses, but does not differ between individuals with and without dementia, though two out of the three identified factor scores are significantly higher in the dementia subgroup. The authors conclude that the Italian version of the AQUALID is a reliable and valid instrument for estimating quality of life in aging adults with ID and dementia.

Abstract: Dementia is common among adults with Down’s syndrome (DS); yet the diagnosis of dementia, particularly in its early stage, can be difficult in this population. One possible reason for this may be the different clinical manifestation of dementia among people with intellectual disabilities. The aim of this study was to map out the carers’ perspective of symptoms of dementia among adults with DS in order to inform the development of an informant-rated screening questionnaire. Unconstrained information from carers of people with DS and dementia regarding the symptoms, particularly the early symptoms of dementia, was gathered using a qualitative methodology. Carers of 24 adults with DS and dementia were interviewed. The interviews were recorded and fully transcribed. The transcripts were then analyzed using qualitative software.

There appeared to be many similarities in the clinical presentation of dementia in adults with DS and the non-intellectually disabled general population. Like in the non-intellectually disabled general population, forgetfulness especially, impairment of recent memory combined with a relatively intact distant memory and confusion were common, and presented early in dementia among adults with DS. However, many ‘frontal lobe’-related symptoms that are usually manifested later in the process of dementia among the general population were common at an early stage of dementia among adults with DS. A general slowness including slowness in activities and speech, other language problems, loss of interest in activities, social withdrawal, balance problems, sleep problems, loss of pre-existing skills along with the emergence of emotional and behavioral problems were common among adults with DS in our study. This study highlighted the similarities in the clinical presentation of dementia among the general population in adults with DS with a particular emphasis on the earlier appearance of symptoms associated with the frontal lobe dysfunction among adults with DS.


Abstract: With the increasing life expectancy in the last five decades, people with intellectual disabilities (ID) are exposed to the risk of age-related neurodegenerative disorders including dementia. The prevalence of dementia is increased in people with ID compared with that in the general population. People with Down syndrome (DS) are at even a higher risk of developing Alzheimer’s dementia (AD) compared with non-ID people with ID. However, there are difficulties in making an early and accurate diagnosis of dementia in individuals with ID and the screening instruments such as the Mini Mental Status Examination that are used in the general population often show floor effects when used for individuals with ID because of their pre-existing cognitive impairment, the level of which varies depending on the severity of ID. Both informant-rated and direct neuropsychological tests have been used for the case detection of dementia for individuals with ID. However, direct neuropsychological tests cannot be used for those who have severe ID and their validity could still be questionable in a number of cases of mild to moderate ID. There are many similarities and some differences in the clinical manifestation of dementia in individuals with ID and the non-ID general population. Impaired recent memory and confusion in the context of relatively intact distant memory are likely to be the early symptoms in individuals with ID who have a mild ID, whereas loss of skills and change in behavior are likely to be the early features for those with more severe ID. Many symptoms, including features of ‘frontal lobe dysfunction’ that tend to appear late in dementia in the general population, may appear early in individuals with ID and DS. Ideally, individuals with ID should be screened for signs of dementia from age 30 onwards. A multi-disciplinary approach should be taken for diagnosis of dementia in individuals with ID using a combination of informant-rated scales and neuropsychological tests in a longitudinal fashion over time. Important differential diagnoses include hypothyroidism, depression, and sensory impairment. Assessment should include physical, psychological, and social aspects including appropriate examinations and investigations. The evidence-base for the pharmacological management of dementia in ID and DS is poor, which does not allow to draw any definitive conclusion about their efficacy in this population. Therefore, the same pharmacological regime that is used in the non-ID population along with non-pharmacological interventions should be considered for people with ID.

Dekker, A.D., Coppus, A.M.W., Strydom, A., Vermeiren, Y., Grefelman, S., Elevedj, J., Beugelsdijk C., ... De Deyn, P.P.


Abstract: Core to the dementia are behavioral and psychological symptoms of dementia (BPSD), an umbrella term for various behavioral changes associated with the presence of dementia, such as apathy, anxiety, disinhibition, irritability, and agitation. The prevalence of BPSD in DS is high and increases with increased severity of dementia. The impact of BPSD on the individual, their caregivers and society is considerable. The authors sought to develop an adapted evaluation scale for comprehensive assessment of BPSD in DS, taking pre-existing behavior and limitations associated with intellectual disability into account. A multidisciplinary collaboration of major DS expertise centers in Europe has been established to develop the novel BPSD-DS evaluation scale. Development consisted of a series of phases, starting with a thorough evaluation of the rather limited knowledge on BPSD in DS and extensive expert feedback rounds to identify relevant behavioral items. A structured, informal interview has been adopted, since people with DS often have difficulty to verbalize their emotions and feelings. Using defined scoring definitions for frequency, severity and caregiver burden at two periods of time, the scales enables identification of specific changes over time, i.e. disentangling behavioral alterations from characteristic behavior that has always been typical for an individual. Multiple optimization rounds and a pilot study yielded the final version that is tested until summer 2016 in a multi-center European validation phase including approx. 250 DS individuals with and without dementia. The BPSD-DS evaluation scale exclusively focuses on behaviour, including 83 items in twelve clusters: anxiety & nervousness, sleep disturbances, irritability, obstinacy, agitation & stereotypical behaviour, aggression, apathy & aspontaneity, depression, delusions, hallucinations, disinhibition & sexual behaviour, and eating and drinking behavior. Scale development, the final version, as well as the first results from the validation phase will be presented. Current clinical experiences and preliminary results for interrater reliability (> 90) are promising. BPSD have been largely neglected in DS. The novel BPSD-DS evaluation scale is the first comprehensive tool for its systematic assessment in the DS population, and serves a three-fold purpose in the future: (1) to monitor behavioural changes in daily care, (2) to identify early BPSD in DS in longitudinal research studies, and (3) to assess behavioural outcome measures.
in clinical trials for dementia in DS.


Abstract: People with Down syndrome (DS) are prone to develop Alzheimer's disease (AD). Behavioral and psychological symptoms of dementia (BPSD) are core features, but have not been comprehensively evaluated in DS. In a European multidisciplinary study, the novel Behavioral and Psychological Symptoms of Dementia in Down Syndrome (BPSD-DS) scale was developed to identify frequency and severity of behavioral changes taking account of life-long characteristic behavior. 83 behavioral items in 12 clinically defined sections were evaluated. The central aim was to identify items that change in relation to the dementia status, and thus may differentiate between diagnostic groups. Structured interviews were conducted with informants of persons with DS without dementia (DS, n = 149), with questionable dementia (DS+Q, n = 65), and with diagnosed dementia (DS+AD, n = 67). First exploratory data suggest promising interrater, test-retest, and internal consistency reliability measures. Concerning item relevance, group comparisons revealed pronounced increases in frequency and severity in items of anxiety, sleep disturbances, agitation & stereotypical behavior, aggression, apathy, depressive symptoms, and eating/drinking behavior. The proportion of individuals presenting an increase was highest in DS+AD, intermediate in DS+Q, and lowest in DS. Interestingly, among DS+Q individuals, a substantial proportion already presented increased anxiety, sleep disturbances, apathy, and depressive symptoms, suggesting that these changes occur early in the course of AD. Future efforts should optimize the scale based on current results and clinical experiences, and further study applicability, reliability, and validity. Future application of the scale in daily care may aid caregivers to understand changes, and contribute to timely interventions and adaptation of caregiving.

Dekker, A.D., Strydom, A., Coppus, A.M.W., Nizetic, D., Vermeiren., Y., Naude, P.J.W., Van Dam, D., Potier, M-C., Fortea, J., De Deyn, P.P.
Behavioral and psychological symptoms of dementia in Down syndrome: Early indicators of clinical Alzheimer's disease? *Cortex*. 2015, 75, 36-61. doi: 10.1016/j.cortex.2015.07.032. Epub 2015 Aug 13. Abstract: Behavioral and psychological symptoms of dementia (BPSD) are a core symptom of dementia and are associated with earlier institutionalization and accelerated cognitive decline for adults with Down syndrome (DS) and increased caregiver burden. Despite the extremely high risk for DS individuals to develop dementia due to Alzheimer's disease (AD), BPSD have not been comprehensively assessed in the DS population. Due to the great variety of DS cohorts, diagnostic methodologies, sub-optimal scales, covariates and outcome measures, it is questionable whether BPSD have always been accurately assessed. However, accurate recognition of BPSD may increase awareness and understanding of these behavioral aberrations, thus enabling adaptive caregiving and, importantly, allowing for therapeutic interventions. Particular BPSD can be observed (long) before the clinical dementia diagnosis and could therefore serve as early indicators of those at risk, and provide a new, non-invasive way to monitor, or at least give an indication of, the complex progression to dementia in DS. This review found that various BPSD appear to be altered in demented DS individuals, but study results have not always been consistent. From childhood to adulthood, externalizing behavior likely decreases and internalizing behavior increases. Frontal lobe symptoms have been suggested as early signs of AD in DS. Disinhibition and apathy, as well as executive dysfunction, seem to be omnipresent in the prodromal phase, whereas reports are inconsistent for depression. Regarding activity disturbances, studies indicated decreasing hyperactivity levels towards adulthood. Excessive activity in demented DS individuals should be a fairly easy observable sign, however, general slowness has been reported and apathy itself might cause reduced activity. Agitation appears to be more prevalent in demented than in non-demented DS individuals, but reports on aggression are inconsistent, though aggression seems to be reduced in the overall DS population. Sleep disturbances are markedly present in both demented and non-demented DS individuals. Although sleep disorders may not yet differentiate between those with and without AD, they are important to consider as such sleep disorders may aggravate cognitive decline and BPSD.


Abstract: Differentiating dementia from baseline level of functioning is difficult among people with severe/profound intellectual (and multiple) disabilities. Moreover, studies on observable dementia symptoms are scarce. This study examined (a) the relevance of dementia diagnosis, (b) observable symptoms and (c) training/information needs. Four explorative focus groups were held with health professionals and family members who have experience with people with severe/profound intellectual (and multiple) disabilities (≥40 years) and decline/dementia. Thematic analysis showed that participants wanted to know about a dementia diagnosis for a better understanding and to be able to make informed choices (question 1). Using a categorization matrix, cognitive and behavioral changes were shown to be most prominent (question 2). Participants indicated that they needed enhanced training, more knowledge development and translation, and supportive organizational choices/policies (question 3). Timely identifying/diagnosing dementia allows for a timely response to changing needs. This requires a better understanding of symptoms.


Abstract: [Corrections: In Figure 8, p. 1519, the second graph (item 10.3) reads: “10.3 making inappropriate comments.” Making inappropriate comments is the description belonging to item 10.2 of the BPSD-DS II. Instead, the second graph (item 10.3) should read: “10.3 loss of decorum.”]


Abstract: People with Down syndrome (DS) are at high risk to develop Alzheimer's disease dementia (AD). Behavioral and psychological symptoms of dementia (BPSD) are common and may also serve as early signals for dementia. However, comprehensive evaluation scales for BPSD, adapted to DS, are lacking. Therefore, we previously developed the BPSD-DS scale to identify behavioral changes between the last six months and pre-existing life-long characteristic behavior. To optimize and further study the scale (discriminative ability and reliability) in a large representative DS study population. Optimization was based on item relevance and clinical experiences obtained in the initial study. Using the shortened and refined BPSD-DS II, informant interviews were conducted to evaluate 524 individuals with DS grouped according to dementia status: no dementia (DS, N = 292), questionable dementia (DS + Q, N = 119), and clinically diagnosed dementia (DS + AD, N = 113). Comparing item change scores between groups revealed prominent changes in frequency and severity for anxious, sleep-related, irritable, restless/stereotypic, apathetic, depressive, and eating/drinking behavior. For most items, the proportion of individuals displaying an increased frequency was highest in DS + AD, intermediate in DS +
Q, and lowest in DS. For various items within sections about anxiety, sleep-related, irritable, apathetic, and depressive behaviors, the proportion of individuals showing an increased frequency was already substantial in DS + Q, suggesting that these changes may serve as early signals of AD in DS. Reliability data were promising. The optimized scale yields largely similar results as obtained with the initial version. Systematically evaluating BPSD in DS may increase understanding of changes among caregivers and (timely) adaptation of care/treatment.

Denney, H., Allen AP, McClinchey E, Buttery N, Garcia-Dominguez L, Chansler R, Corr C, Dunne P, Kennelly S, Daly L, McCallon P, McCarron M. A scoping review of post-diagnostic dementia supports for people with intellectual disability and dementia. *Aging & Mental Health*, 2022 Oct 11:1-10. doi: 10.1080/13607863.2022.2130171. Epub ahead of print. PMID: 36218056. Abstract: People with intellectual disability, particularly people with Down syndrome, are at an increased risk for early-onset dementia, in comparison to people without an intellectual disability. The aim of this review was to scope the current landscape of post-diagnostic dementia supports for people with intellectual disability.Method: A systematic search of five electronic databases (CINAHL, Medline, PsycArticles, PsycInfo and Web of Science) was conducted for this scoping review. Results were screened independently by two reviewers, with a third reviewer for arbitration where necessary. Forty-two studies met the inclusion criteria, and relevant information was extracted. The articles included focused on the experiences of people with intellectual disability and dementia, as well as the role of carers, family members and staff. Key themes included ageing in place, environmental supports for people with intellectual disability and dementia, dementia-specific interventions and therapies, as well as the feasibility of these interventions. Besides the studies that focussed on these themes, other studies focused on staff training and family supports. This review highlights the importance of implementing timely and appropriate post-diagnostic supports for people living with intellectual disability and dementia. More controlled trials are required on post-diagnostic dementia supports for people with intellectual disability.

Devenny, D.A., Krinsky-McHale, S.J., Sersen, G., & Silverman, W.P. Sequence of cognitive decline in dementia in adults with Down's syndrome. *Journal of Intellectual Disability Research* 2000, 44(6), 654-665. doi:10.1046/j.1365-2788.2000.00305.x Abstract: Because of lifelong intellectual deficits, it is difficult to determine the earliest signs and characteristics of age-associated decline and dementia among adults with Down syndrome. In a longitudinal study in which all participants were healthy at the time of their entry into the study, the present authors compared the amount of decline on the subtests of the WISC-R to determine the sequence of cognitive decline associated with varying stages of dementia. Twenty-two individuals with varying degrees of cognitive decline were compared to 44 adults with DS who have remained healthy. All participants functioned in the mild or moderate range of intellectual disability at initial testing. On each subtest of the WISC-R, the amount of change experienced by the healthy participants over the study period was compared to the amount of change found for each of the groups with decline. Out of the individuals who showed declines, 10 adults with DS were classified as having 'questionable' decline based on the presence of memory impairment, and five and seven adults with DS were classified as in the 'early stage' and 'middle stage' of DAT, respectively, based on the presence of memory impairment, score on the Dementia Scale for Down Syndrome and a physician's diagnosis. It was found that participants who were identified as 'questionable', in addition to the memory loss that determined their classification, also showed significant declines on the Block Design and Coding subtests. The five adults in the early stage of dementia showed declines on these subtests, and in addition, on the Object Assembly, Picture Completion, Arithmetic and Comprehension subtests. The seven adults in the middle stage of dementia showed declines on these subtests, plus declines on Information, Vocabulary and Digit Span subtests. The Picture Arrangement and Similarities subtests were not useful in distinguishing between the groups because of baseline floor effects for a substantial proportion of participants. The present longitudinal study showed a sequence of cognitive decline associated with DAT, beginning with a possible 'pre-clinical' stage, and progressing through the early and middle stages. This approach begins to define the sequence of declining cognitive capacities that contributes to the observed functional deterioration caused by Alzheimer's disease and that is likely to reflect the involvement of cortical areas as the disease progresses.

Dick, M.B., Doran, E., & Phalen, M., & Lott,I. Cognitive profiles on the Severe Impairment Battery are similar in Alzheimer disease and Down syndrome with dementia. *Alzheimer Disease and Associated Disorders*, 2015, Dec 22 [Epub ahead of print] Abstract: Previous research has revealed similarities in the neuropathology, clinical presentation, and risk factors between persons with Alzheimer disease from the general population (GP-AD) and those with Down syndrome (DS-AD). Less is known, however, about the extent of similarities and differences in the cognitive profiles of these 2 populations. Fifty-one moderate to severely demented GP-AD and 59 DS-AD individuals participated in this study which compared the cognitive profiles of these 2 populations on the Severe Impairment Battery (SIB), controlling for sex as well as level of functional ability using a modified version of the Bristol Activities of Daily Living Scale. Overall, the neuropsychological profiles of the higher-functioning individuals within the DS-AD and advanced GP-AD groups, as represented by mean difference scores on the SIB as a whole and across the 9 separate cognitive domains, were very similar to one another after adjusting for sex and functional impairment. To our knowledge, this is the first study to directly compare the cognitive profiles of these 2 populations on the SIB. Findings suggest that the underlying dementia in GP-AD and DS-AD may have corresponding and parallel effects on cognition.

Dillane, I., & Dood, O. Nursing people with intellectual disability and dementia experiencing pain: An integrative review. *Journal of Clinical Nursing*, 2019, 10.1111/jocn.14834, 28(13-14), 2472-2485 Abstract: To explore the current evidence of nurses caring for people with intellectual disability and dementia who experience pain. People with intellectual disability are aging and are experiencing age-related health conditions including dementia and conditions associated with pain, but at an earlier age. Addressing the needs of people with intellectual disability who develop dementia is a new challenge for nurses. The authors undertook an integrative literature review and a systematic search of databases: CINAHL, MEDLINE, PsycINFO, Cochrane, EMBASE, Academic Search Complete, Scopus and Web of Science between 27 October 2017-7 November 2017. Hand searching and review of secondary references were also undertaken. Quality appraisal (Crove Critical Appraisal Tool), thematic data analysis (Braun and Clarke, Qualitative Research in Psychology, 3, 2006, 77) and reporting using the PRISMA guidelines. Seven papers met the inclusion criteria, and three themes emerged from this review: nurses knowledge of ageing, dementia and pain; recognising pain in people with intellectual disability and dementia; and the role of nurse education. People with intellectual disability and dementia have difficulty communicating their pain experience compounded by pre-existing communication difficulties. A pain experience can present similar to behavioral and psychological symptoms of dementia, and diagnostic overshadowing often occurs whereby a pain need is misinterpreted as behavioral and psychological symptoms resulting in inappropriate treatment. Nurses need greater knowledge about the presence of pain and potential causes in people with intellectual disability and dementia, and education can be effective in addressing this knowledge deficit. Pain assessment tools for people with intellectual disability and dementia need to include behavioural elements, and baseline assessments are required to identify changes in pain presentation. Nurses need to recognize and respond to pain based on the evidence in order to deliver quality care.
planning which causes high stress levels for caregivers. Policy makers and researchers working in this field need to take into consideration the needs of older caregivers when making future plans for adults with disabilities.

Dodd, K.
Abstract: The purpose of this paper is to consider the implications for people with Down’s syndrome and their families of identifying those people who are at risk of developing dementia from the research study “Estimating the number of people with Down’s syndrome in Scotland and the cohort at elevated risk of early onset dementia”. The commentary is based on a review of the associated literature. Estimating the numbers is important but has serious implications for people who have an elevated risk and their families. Preparation and ongoing support and planning are vital to ensure that quality of life is maintained as dementia is identified and progresses.

Dodd, K.
Supporting people with Down’s syndrome and dementia Tizard Learning Review. 2003, 4(4), 14-18
Abstract: Brief review of literature and concepts dealing with the prevalence of dementia among people with Down syndrome in England, ethical issues in assessment and diagnosis, the value of early diagnosis, and an explication of service options and management strategies. Review concludes with a prognosis for services in the future.

Abstract: Post diagnostic support (PDS) has varied definitions within mainstream dementia services and different health and social care organizations, encompassing a range of supports that are offered to adults once diagnosed with dementia until death. An international summit on intellectual disability and dementia held in Glasgow, Scotland in 2016 identified how PDS applies to adults with an intellectual disability and dementia. The Summit proposed a model that encompassed seven focal areas: post-diagnostic counseling; psychological and medical surveillance; periodic reviews and adjustments to the dementia care plan; early identification of behavior and psychological symptoms; reviews of care practices and supports for advanced dementia and end of life; supports to carers/ support staff; and evaluation of quality of life. It also explored current practices in providing PDS in intellectual disability services. The Summit concluded that although there is limited research evidence for pharmacological or non-pharmacological interventions for people with intellectual disability and dementia, viable resources and guidelines describe practical approaches drawn from clinical practice. Post diagnostic support is essential, and the model components in place for the general population, and proposed here for use within the intellectual disability field, need to be individualized and adapted to the person’s needs as dementia progresses. Recommendations for future research include examining the prevalence and nature of behavioral and psychological symptoms (BPSD) in adults with an intellectual disability who develop dementia, the effectiveness of different non-pharmacological interventions, the interaction between pharmacological and non-pharmacological interventions, and the utility of different models of support.

Donaldson S.
Abstract: Author assessed how staff ratings of challenging behavior for people with Down syndrome and dementia affected the self-reported well-being of care staff. Data were collected from 60 care staff in 5 day centers in a large city in England. The data were collected by use of a questionnaire. There was no significant difference between those who cared for individuals with Down syndrome and dementia and those caring for service users with other non-specified learning disabilities without dementia, regarding their self-reported well-being. Self-reported well-being did correlate with staff rating of challenging behavior in both those who cared for people with Down syndrome and dementia and those who did not care for such service users, with well-being declining as perceived challenging behavior increased. The findings indicate that challenging behavior prevention and reduction may be of benefit to both service users and care staff well-being.

Duncan, C., Wilkinson, E., Jaydeokar, S., & Acton, D.J.
Abstract: This study evaluated the dementia assessment and diagnosis care provided to adults with intellectual disability. The authors selected recommendations from the National Institute for Health and Care Excellence (NICE) standards which could be evidenced in clinical notes and aimed to identify characteristics which may be associated with improved adherence to these recommendations. The study population was adults with an intellectual disability who were diagnosed with dementia between January 2019 and December 2022 by a UK-based intellectual disability service. Data to demonstrate adherence to selected recommendations and demographic and clinical characteristics were extracted from electronic patient records. The authors identified 41 individuals. A mean of six of the eight recommendations were adhered to. There was low adherence with structural imaging to support dementia subtype diagnosis (9 individuals, 22%). This may be linked with the low percentage of people diagnosed with vascular dementia (1 individual, 2%) despite a national figure of 20%. No demographic or clinical characteristics were associated with level of adherence recorded. The authors found incomplete recording of diagnostic clinical coding in electronic patient records. This may disadvantage this population, as they cannot be readily identified for post-diagnostics support or resource allocation. To the best of the authors’ knowledge, this is the first study to examine adherence to these NICE guidelines in this population.

Dunne, P., Reilly, E., Judge, R., Lowe, F. & McCarron, M.
Giving meaning to life - the role of digital life Stories in supporting people with intellectual disability and dementia Journal of Intellectual Disability Research. 2019, 63(8), 643.
Abstract: Over a four-year period individuals with an intellectual disability (ID) in a large service provider in the Republic of Ireland were supported to create their personalized digital life story using selected multi media apps. An easy read survey was distributed to 380 people with ID to gauge their readiness and interest in engaging in digital life story. A bespoke training course was developed to support the introduction of digital life story activities. The use of the personalized digital life stories by the individuals and their support staff, family and social network was captured through recorded observations and individual use was tracked through a developed audit tool. Over the four-year period eighty people with an ID commenced their digital life story. Case studies within this cohort showed a variety of use which varied depending on degree of ID and stage of dementia. In all cases life stories were instrumental to enhanced communication and social interactions. Key factors in uptake and sustainability were staff training, iPAD clubs and champions across the organization Implications: Digital life stories were key in supporting meaningful interactions across the continuum of dementia both in day to day interactions with family, staff, volunteers, and peers and as a tool for social engagement through digital life story clubs.

Eady, N., Sheehan, R., Rantell, K., Sinai, A., Bernal, J., Bohnen, I., Courtney, K., Dodd, K., Gazizova, D., Hassiottis, A., ... Strydom, A.
Abstract: There is little evidence to guide pharmacological treatment in adults with Down syndrome and Alzheimer’s disease. Authors investigated the effect of cholinesterase inhibitors or memantine on survival and function in adults with Down syndrome and Alzheimer’s disease. This was a naturalistic longitudinal follow-up of a clinical cohort of 310 people with Down syndrome diagnosed with Alzheimer’s disease collected from specialist community services in England.
A critical literature review of the effectiveness of various instruments in the diagnosis of dementia in adults with intellectual disabilities

Alzheimer’s and Dementia (Amst), 2016, Jun;30, 4(1), 126-148. doi: 10.1016/j.dadm.2016.06.002. eCollection 2016. Abstract: Currently, there is no consensus on dementia diagnostics in adults with intellectual disabilities (ID). There are three types of assessments available: direct cognitive tests, test batteries, and informant reports. A systematic literature search was conducted in four databases yielding 9840 records. Relevant studies were identified and selected using predefined inclusion and exclusion criteria and then coded and classified according to assessment type. This was completed by two independent researchers, with a third consulted when discrepancies arose. The review collates diagnostic instruments and presents strengths and weaknesses. Overall 47 studies met the search criteria, and 43 instruments were extracted from the selected studies. Of which, 10 instruments were classified as test batteries, 23 were classified as direct cognitive tests, and the remaining 10 were informant reports. This review can recommend that cognitive test batteries can offer the most practical and efficient method for dementia diagnosis in individuals with ID.

Ebbing, K., von Gunten, A., Guinchat, V., Georgescu, D., Bersier, T., Moad, D., & Verloo, H.
Barriers facing direct support professionals when supporting older adults presenting with intellectual disabilities and unusual dementia-related behavior: A multi-site, multi-methods study.

Abstract: Increased life expectancy among people with intellectual disabilities (ID) raises the risk of their diagnosis being superimposed by behavioral and psychological symptoms of dementia (BPSD). The difficulties facing direct support professionals dealing with this is an emerging, under-investigated issue. The study investigates direct support professionals' perceptions and experiences of their daily support for aging people with ID presenting with superimposed BPSD. Twenty-four direct support professionals from long-term care facilities recruited vignettes and attended focus groups conducted to investigate perceptions and lived experiences of the barriers and struggles they faced. Direct support professionals' reactions to vignettes revealed their difficulties recognizing BPSD superimposed on the known challenging behaviors of people with ID. Focus groups highlighted daily struggles with BPSD, the lack of knowledge about detecting and dealing with them, and associated somatic and psychopathological diseases of aging. Improved knowledge transfer about good practices for person-centered support to aging people with ID presenting with BPSD is strongly recommended.

Egan, C., & Dalton, C.T.
An exploration of care-burden experienced by older caregivers of adults with intellectual disabilities in Ireland

Abstract: People with intellectual disabilities are experiencing increased longevity, and in parallel, their family caregivers are also ageing. The literature identifies that these caregivers are at risk of burden. The aim of this study was to measure the level of caregiver burden among older carers of adults with intellectual disabilities in an Irish sample and to analyze the effect of socio-demographic factors upon experiences of caregiver burden. Methods Thirty caregivers completed a survey questionnaire. Data were collected based on participants' self-reports of burden using the Zarit Burden Interview (ZBI) and a socio-demographic questionnaire. Data were analyzed using SPSS version 24. Over 57% of carers indicated a mild-to-moderate level of burden. Analysis indicated that younger caregivers experience significantly higher levels of burden, when compared to older caregivers. This study contributes to our understanding of burden among an Irish population of older caregivers supporting an adult with an intellectual disability. It identified that carers do experience burden. The importance of proactive assessments and supports for these caregivers was revealed. This study highlights a lack of Irish research in this area and may pave the way for future research which could build upon its findings.

Eisner, D.A.
Down’s syndrome and aging: Is senile dementia inevitable?

Abstract: Numerous studies have reported that in elderly Down’s Syndrome individuals there is a high preponderance of senile dementia. An examination of these investigations shows that, while there is accelerated neurological aging, there is not a high incidence of behavioral or overt senile dementia. Changes in cognitive functioning for Down’s Syndrome persons are similar to those found in non-Down’s retarded populations.

Elliott-King, J., Shaw, S., Bandelow, S., Devshi, R., Kassam, S., & Hogervorst, E.
A critical literature review of the effectiveness of various instruments in the experience burden. The importance of proactive assessments and supports for understanding of burden among an Irish population of older caregivers dealing with problem behaviors, helping with activities of daily living, promoting alternative activities) and help for carers.

Engdahl, J. M. K.

36 pp.

Bozeman, Montana: Author [723 South 13th Street, Bozeman, MT 59715] (1995)
Abstract: Training manual developed to provide primary information about care practices for parents and other primary carers of adults with Down syndrome affected by Alzheimer’s disease. Covers, in brief format, recognizing signs and symptoms, diagnostic advice, care management practice (communication, dealing with problem behaviors, helping with activities of daily living, promoting alternative activities) and help for carers.

Englund, A., Jonsson, B., Zander, C.S., Gustafsson, J., & Annerén, G.
Changes in mortality and causes of death in the Swedish Down syndrome population

Abstract: During the past few decades age at death for individuals with Down syndrome (DS) has increased dramatically. The birth frequency of infants with DS has long been constant in Sweden. Thus, the prevalence of DS in the population is increasing. The aim of the present study was to analyze mortality and causes of death in individuals with DS during the period 1969-2003. All individuals with DS that died between 1969 and 2003 in Sweden, and all individuals born with DS in Sweden between 1974 and 2003 were included. Data were obtained from the Swedish Medical Birth Register, the Swedish Birth Defects Register, and the National Cause of Death Register. Median age at death has increased by 1.8 years per year. The main cause of death was pneumonia. Death from congenital heart defects decreased. Death from atherosclerosis was rare but more frequent than reported previously. Dementia was not reported in any subjects with DS before 40 years of age, but was a main or contributing cause of death in 30% of the older subjects. Except for childhood leukemia, cancer as a cause of death was rare in all age groups. Mortality in DS, particularly infant mortality, has decreased markedly during the past decades. Median age at death is increasing and is now almost 60 years. Death from cancer is rare in DS, but death from dementia is common.

ENIDA
Face to face: Respectful coping with dementia in older people with intellectual disability

52 minutes

Working Group on Coping with Dementia in Older People with Intellectual Disability, European Network on Intellectual Disability and Ageing [ENIDA] - c/o Patricia Noonan Walsh, Ph.D., Director, Centre for the Study of Developmental Disabilities, University College Dublin, Belfield, Dublin 4, IRELAND – e-mail: patricia.walsh@ucd.ie (2000)
Abstract: A 52-minute video with an accompanying information booklet, which uses a number of case vignettes from France, Belgium and the Netherlands to illustrate the various symptoms and stages of dementia among older people with intellectual disability. Examples of practices to promote “respectful coping” with dementia, death and dying on the part of direct support professionals and clinicians are presented. Devised for staff training and development, Face to Face may be viewed in short segments. A version with English subtitles and
Ericksson, M., & Sundin, M.
Developing early detection of dementia with people with intellectual and developmental disabilities
Poster presented at the 27th Annual Conference of Alzheimer Europe, Berlin, Germany, October 3, 2017. (PO3.26)
Abstract: In Finland there is a lack of a unified approach to the early detection of dementia for people with intellectual and developmental disabilities (IDD). The aim of this presentation is to describe a currently underway collaboration project, which responds to this challenge. In our opinion, there are two essential elements in early detection of dementia with people with IDD: 1) an overall description of the psychosocial functioning of the person and 2) a screening method for dementia, which developed for persons with IDD. These two components are shortly outlined. Psychosocial functioning: Early detection of neurocognitive disorders in people with IDD is multidisciplinary teamwork. The Finnish Association of Intellectual and Developmental Disabilities (FAIDD) has published two methods for this purpose (the Toimi and the Payto), and we recommend using these methods in the assessment of dementia in people with IDD as well. For example, it is important to distinguish the symptoms of dementia from other possible psychological disorders (like mood disorders and psychotic symptoms). Translation of the National Task Group Early Detection Screen for Dementia (NTG-EDSD): Commonly used assessment methods (like the Cerad and the Mini-Mental) may not be applicable for people with IDD. In our project, we decided to translate the NTG-EDSD into Finnish. The EDSD is already available online in many other languages (www.aadmd.org/ntg/screening). As the authors write, the NTG-EDSD is not an assessment or diagnostic instrument, but an administrative screen that can be used by people who know the client well. The Finnish version of the NTG-EDSD is being introduced into practice in 2017. FAIDD trains staff and other people working with people with IDD in using the screen within the perspective of psychosocial functioning.

Esbensen, A.J.
Health conditions associated with aging and end of life of adults with Down syndrome.
https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3010180/
Abstract: Expectations for the life course of individuals with Down syndrome (DS) have changed, with life expectancy estimates increasing from 12 in 1949 to nearly 60 years of age today. Along with this longer life expectancy comes a larger population of adults with DS who display premature age-related changes in their health. There is thus a need to provide specialized health care to this aging population of adults with DS who are at high risk for some conditions and at lower risk for others. This review focuses on the rates and contributing factors to medical conditions that are common in adults with DS or that show changes with age. The review of medical conditions includes the increased risk for skin and hair changes, early onset menopause, visual and hearing impairments, adult onset seizure disorder, thyroid dysfunction, diabetes, obesity, sleep apnea and musculoskeletal problems. The different pattern of conditions associated with the mortality of adults with DS is also reviewed.

Esbensen, A.J., Boshkoff Johnson, E., Amaral, J.L., Tan, C.M., & Macks, R.
Differentiating aging among adults with Down syndrome and comorbid psychopathology.
Abstract: Differences were examined between three groups of adults with Down syndrome in their behavioral presentation, social life/activities, health, and support needs. We compared those with comorbid dementia, with comorbid psychopathology, and with no comorbid conditions. Adults with comorbid dementia were more likely to be older, have lower functional abilities, have worse health and more health conditions, and need more support in self-care. Adults with comorbid psychopathology were more likely to exhibit more behavior problems and to be living at home with their families. Adults with no comorbidities were most likely to be involved in community employment. Differences in behavioral presentation can help facilitate clinical diagnoses in aging in Down syndrome, and implications for differential diagnosis and service supports are discussed.

Esbensen, A.J., Mailick, M.R., & Silverman, W.
Long-term Impact of parental well-being on adult outcomes and dementia status in individuals with Down syndrome.
Abstract: Parental characteristics were significant predictors of health, functional abilities, and behavior problems in adults with Down syndrome (n = 75) over a 22-year time span, controlling for initial levels and earlier changes in these outcomes. Lower levels of behavior problems were predicted by improvements in maternal depressive symptoms. Higher levels of functional abilities were predicted by prior measures of and improvements in maternal depressive symptoms. Better health was predicted by prior measures of maternal depressive symptoms, paternal positive psychological well-being, relationship quality between fathers and their adult children, and improvements in maternal positive psychological well-being. Dementia status was also predicted by parental characteristics. This study suggests the importance of the family context for healthy aging in adults with Down syndrome.

Esraluw, L., Janicki, M.P., & Keller, S.M.
National Task Group Early Detection Screen for Dementia (NTG-EDSD)
Doi:10.1007/978-3-319-61720-6
Abstract: Early identification of signs and symptoms of cognitive and functional decline associated with dementia is an important first step in managing the course of dementia and providing quality care. Studies show that persons with intellectual disabilities are as susceptible to the causes of dementia as are other adults, with the relative distribution of etiologies mirroring those of other adults. The challenge, as noted by the World Health Organization, is to identify early those adults susceptible and affected so that assessment and diagnostic work-ups can be undertaken. A screening tool can be used to substantiate changes in adaptive skills, behavior and cognition. With early detection, assessment and diagnosis can be carried out to determine whether cognitive changes are the result of a neuropathological process related to disease or trauma to the brain, or attributable to other causes, which may be treatable and reversible. The National Task Group Early Detection Screen for Dementia (NTG-EDSD) is an informant-based rating tool for use with adults with intellectual and developmental disabilities who are suspected of experiencing changes in thinking, behavior, and adaptive skills suggestive of mild cognitive impairment or dementia. This chapter offers background on the NTG-EDSD and describes benefits that may accrue from using a dementia screening instrument.

Adaptacion y validacion del Cambridge Examination for Mental Disorders of Older People with Down's Syndrome and Others with Intellectual Disabilities (CAMDEX-DS) en poblacion espanola con discapacidad intelectual [Adaptation and validation of CAMDEX-DS (Cambridge Examination for Mental Disorders of Older People with Down's Syndrome and Others with Intellectual Disabilities) in Spanish population with intellectual disabilities].
Revisa de Neurologia, 2013 Oct 16, 57(8), 337-346. [In Spanish]
Abstract: Dementia caused by Alzheimer's disease commonly affects the adult population with Down's syndrome. This population presents two characteristic clinical features: a semiotic pattern that differs from the typical Alzheimer's disease, and previous intellectual deficits that may confound the clinical diagnosis. There is a clear need to validate specific instruments adapted to Spanish population. To adapt and to validate CAMDEX-DS (Cambridge Examination for Mental Disorders of Older People with Down's Syndrome and Others with Intellectual Disabilities) in Spanish population, 146 adults with intellectual disability (mild to moderate) were recruited and assessed with CAMDEX-DS, K-BIT I and DMR tests. Test-retest reliability, inter-rater concordance and validity statistic were performed between CAMDEX-DS and clinical diagnosis. This is an observational, multicenter, cross-sectional and validation study. Test-retest and inter-rater reliability achieved kappa coefficient values of 0.92 and 0.91, respectively. Agreement (kappa index) for CAMDEX-DS on clinical diagnosis compared to other clinical criteria was high: CAMDEX-DS vs DSM-IV (kappa = 0.95; p < 0.001); CAMDEX-DS vs ICD-10
Echteld, M.A.  

Development of the DMR (DLD) along with its clinical applications.  

The Dementia Questionnaire for People with Learning Disabilities (DVZ) has been developed by Heleen Evenhuis, the Netherlands. The Vragenlijst voor Zwakzinnigen (DVZ) has been developed by Heleen Evenhuis, H.M.

Mild cognitive impairment and Alzheimer's disease are common in persons with Down's syndrome, but often were not recognized early. In the present group of patients, there was a remarkably high incidence of gait and speech deterioration. Also, the incidence of epileptic seizures and myoclonus was about eightfold, as compared with Alzheimer's disease in persons without Down's syndrome.


Abstract: In a prospective longitudinal study with death as the end point in 17 middle-aged patients with Down's syndrome, dementia was clinically diagnosed in 15 patients, by means of careful observations in daily circumstances. Autopsies were performed in 10 cases: 8 demented patients and 2 nondemented patients. Neuropathologically, Alzheimer-type abnormalities were demonstrated in 9 patients, both demented and nondemented, and combined Alzheimer-type abnormalities with infarctions were demonstrated in 1 patient. In the 14 demented patients who did not show evidence of cerebrovascular or systemic vascular disease, dementia had an early onset and was rapidly progressive (mean age at onset, 51.3 years in the moderately retarded patients and 52.6 years in the severely retarded patients; mean duration of symptoms, respectively, 4.9 and 5.2 years). Cognitive and behavioral decline corresponded to symptoms of dementia of the Alzheimer's type in patients without Down's syndrome, but often were not recognized early. In the present group of patients, there was a remarkably high incidence of gait and speech deterioration. Also, the incidence of epileptic seizures and myoclonus was about eightfold, as compared with dementia of the Alzheimer's type in patients without Down's syndrome.

The Dementia Questionnaire for People with Learning Disabilities (DVZ) can set two key points by the level of intellectual disability on the suspicion of cognitive impairment in people with Down's syndrome.

Evans, E., Bhardwaj, A., Brodaty, H., Sachdev, P., Draper, B., & Trollor, J.N.


Abstract: The population with intellectual disability (ID) is aging, but age-related health concerns such as dementia have received little research attention thus far. We review evidence regarding the prevalence and incidence of dementia in people with ID, and discuss some possible explanations for an increased risk, such as shared genetic risk factors, co-morbid physical and mental disorders, lifestyle factors, trauma, and lowered brain reserve. We discuss practical and theoretical challenges facing researchers in this field, before highlighting the implications of findings to date for future research and clinical care. Research on dementia in this at-risk population has the potential to help us understand dementia in general and to improve services for this group of vulnerable individuals.

Evenhuis H.M.


Abstract: In a prospective longitudinal study with death as the end point in 17 middle-aged patients with Down's syndrome, dementia was clinically diagnosed in 15 patients, by means of careful observations in daily circumstances. Autopsies were performed in 10 cases: 8 demented patients and 2 nondemented patients. Neuropathologically, Alzheimer-type abnormalities were demonstrated in 9 patients, both demented and nondemented, and combined Alzheimer-type abnormalities with infarctions were demonstrated in 1 patient. In the 14 demented patients who did not show evidence of cerebrovascular or systemic vascular disease, dementia had an early onset and was rapidly progressive (mean age at onset, 51.3 years in the moderately retarded patients and 52.6 years in the severely retarded patients; mean duration of symptoms, respectively, 4.9 and 5.2 years). Cognitive and behavioral decline corresponded to symptoms of dementia of the Alzheimer's type in patients without Down's syndrome, but often were not recognized early. In the present group of patients, there was a remarkably high incidence of gait and speech deterioration. Also, the incidence of epileptic seizures and myoclonus was about eightfold, as compared with dementia of the Alzheimer's type in patients without Down's syndrome.

Evenhuis, H.M.


Abstract: To facilitate the diagnosis of dementia in persons with intellectual disabilities (ID), based on observations of caregivers, since 1980 the Dementie Vragenlijst voor Zwakzinnigen (DVZ) has been developed by Heleen Evenhuis, H.M.

Fahey-McCarthy, E., McCarron, M., Connaire, K., & McCallion, P.


Abstract: Generally, staff working in settings that provide care for adults with intellectual disabilities (ID) have not received specific education with respect to extended care for terminal illnesses or late-stage dementia. Equally, staff working in specialist palliative care often are not familiar with the unique issues of supporting persons with intellectual disabilities affected by dementia. To understand care concerns with respect to supporting persons with ID and advanced dementia, and to develop, deliver, and evaluate an educational intervention with staff in ID settings and specialist palliative care services, 14 focus group interviews were conducted with staff across six ID services and one specialist palliative care provider in the greater Dublin (Ireland) area. Qualitative descriptive analysis resulted in the emergence of key themes and formed the development of an educational intervention. Pre- and posttest questionnaires assessed responses to a pilot delivery of the educational intervention. Formal feedback from staff indicated that the educational intervention was highly valued and addressed key training concerns. They agreed that the training supported “aging in place,” and the preparation for a “good death” including support for staff, peers, and family in their grief and bereavement. An educational intervention in the form of a trainer manual was produced to support cross-service system in-service training on issues of advancing dementia in persons with ID.

Fazio, S., Pace, D., Kalmyer, B., & Pike, J.


Abstract: Speaks to the Dementia Care Practice Recommendations that were developed to better define quality care across all care settings and throughout the disease course. Notes that they are intended for professional care providers who work with individuals living with dementia and their families in residential and community based care settings.

Fazio, S., Pace, D., Kalmyer, B., & Maslow, K., & Zimmerman, S.

Alzheimer's Association dementia care practice recommendations. Gerontologist, 2018, Jan 18, 58(suppl1), S1-S9.


Abstract: The Alzheimer's Association 'Dementia Care Practice Recommendations' outline guidance on quality care practices based on a comprehensive review of current evidence, best practice, and expert opinion. The 'Dementia Care Practice Recommendations' were developed to better define quality care across all care settings, and throughout the disease course. They are intended for professional care providers who work with individuals living with dementia and their families in residential and community-based care settings. With the fundamentals of person-centered care as the foundation, the 'Dementia Care Practice Recommendations' posit goals for quality dementia care practice recommendations with emphasis on high-quality, person-centered care in long-term and community-based care settings.
care in the following areas: (a) person-centered care, (b) detection and diagnosis, (c) assessment and care planning, (d) medical management, (e) information, education, and support, (f) ongoing care for behavioral and psychological symptoms of dementia, and support for activities of daily living, (g) staffing, (h) supportive and therapeutic environments, and (i) transitions and coordination of services.

Firth, N.C., Startin, C.M., Hithersay, R., Hamburg, S., Wijeratne, P.A., Mok, K.Y., Hardy, J., Alexander, D.C., LonDownS Consortium, & Strydom, A. Aging related cognitive changes associated with Alzheimer’s disease in Down syndrome. Annals of Clinical and Translational Neurology, 2018, May 20, 5(6), 741-751. doi: 10.1002/acn3.571. eCollection 2018 Jun. Abstract: Individuals with Down syndrome (DS) have an extremely high genetic risk for Alzheimer’s disease (AD), however, the course of cognitive decline associated with progression to dementia is ill-defined. Data-driven methods can estimate long-term trends from cross-sectional data while adjusting for variability in baseline ability, which complicates dementia assessment in those with DS. We applied an event-based model to cognitive test data and informant-rated questionnaire data from 283 adults with DS (the largest study of cognitive functioning in DS to date) to estimate the sequence of cognitive decline and individuals’ disease stage. Decline in tests of memory, sustained attention/working memory coordination, and verbal fluency occurred early, demonstrating that AD in DS follows a similar pattern of change to other forms of AD. Later decline was found for informant measures. Using the resulting staging model, we showed that adults with a clinical diagnosis of dementia and those with APOE 3:4 or 4:4 genotype were significantly more likely to be staged later, suggesting that the model is valid. Our results identify tests of memory and sustained attention may be particularly useful measures to track decline in the preclinical/prodromal stages of AD in DS whereas informant-measures may be useful in later stages (i.e. during conversion into dementia, or postdiagnosis). These results have implications for the selection of outcome measures of treatment trials to delay or prevent cognitive decline due to AD in DS. As clinical diagnoses are generally made late into AD progression, early assessment is essential.

Folin, M., Baigueru, S., Conconi, M.T., Pati, T., Grandi, C., Parnigotto, P.P., & Nussdorfer, G.G. The impact of risk factors of Alzheimer’s disease in the Down syndrome International Journal of Molecular Medicine, 2003, Feb,11(2), 267-270. https://pubmed.ncbi.nlm.nih.gov/12525890/ Abstract: Down syndrome (DS) patients, after the fourth decade of life, display some neuropathological features of the Alzheimer’s disease (AD). Several hypotheses suggested that apolipoprotein protein, an AD risk factor, might promote amyloid formation by stabilizing an aggregated conformation of the beta amyloid protein (Abeta). This peptide is the major proteinaceous component of the senile plaques either in AD or DS, and it is a proteolytic product of the amyloid precursor protein (APP). Both brain and platelets express three APP transcripts of the apparent molecular weight of 106, 110 and 130 kDa. In DS the Abeta deposits may ensue, at least in part, from the overexpression of the Abeta precursor gene located on chromosome 21. Aims of the present study were to evaluate the frequency of apoE4 isoform in DS population, and to ascertain whether the ratio between the 130 and the 106-110 kDa platelet APP isoforms is lower in DS, as seems to occur in AD patients. ApoE4 frequency was significantly lower in DS when compared to AD patients. E4 allele frequency of older DS patients was about half that of younger ones. The 130 to 106-110 kDa APP isoform ratio was similar in young DS and control subjects, and markedly lower in AD patients. Our results indicate that: i) in DS patients the early, selective accumulation of Abeta peptides is independent of the ApoE genotype, but the allele epsilon4 predisposes to various causes of premature death; and ii) platelet APP isoform abnormalities, which can be observed in AD patients, do not occur in young DS patients, suggesting a different processing of APP platelets in DS with respect to AD.

Forster-Gibson, C.J. Behaviour changes in an adult with Down syndrome. Canadian Family Physician, 2019 Apr, 65(Suppl 1), S25-S26. PMID: 31023775; PMCID: PMC6501721 Abstract: A case review of a 43-year-old man with Down syndrome who had a 2-year history of behaviour changes, paranoia, and auditory hallucinations, increased fatigue, and changes in gait and speech, as well as losses in his social, work, and daily living skills formed the basis of this article. Author notes that adults with Down syndrome are at increased risk of depression and hypothyroidism, and Alzheimer dementia (AD) as they age, all of which can cause behaviour changes. Challenges in the primary care of aging adults with developmental disability can include limited information on differences in the aging process; complex health needs; causes of developmental delay; behaviors that challenge; and communication problems. In persons with Down syndrome, pathology studies of the brain have shown the presence of neurofibrillary tangles and plaques typical of AD in 7.5% of teens, 80% of adults in their 30s, and almost all adults older than 40 years of age, although the condition might not be symptomatic. The prevalence of diagnosed AD in adults with Down syndrome has been estimated to be 9.4% for those aged 40 to 49, increasing to 54.5% for those aged 60 to 69. Differentiating between depression and early-stage AD can be very difficult, and depression can be a sign of early AD. Progressive changes that might suggest early-stage AD include loss of language, behaviour changes (including social withdrawal), loss of daily living skills, gait disorder, and, in some, psychosis (including hallucinations and delusions), seizures, and dysphagia. Assessment of behavioural changes in adults with Down syndrome should begin with assessment of medical concerns common to Down syndrome. The diagnostic assessment of adults with Down syndrome for Alzheimer dementia is similar to accepted practice in the general population. Caveats include limitations on the usefulness of neuropsychological assessment tools and neuroimaging, and the common occurrence of hypothyroidism and depression.

Fromage, B., & Anglade, P. [Le vieillissement des sujets trisomiques] [The aging of adults with Down syndrome] Article in French L’Encéphale, 2002, May-June, 28(3 P 1), 212-216. https://pubmed.ncbi.nlm.nih.gov/12091781/ Abstract: The normal aging of adults with Down syndrome is comparable to that observed in individuals who have an equivalent cognitive deficit. However it is earlier and is related to the development of the physical and life story of the person. Before 50 years, there is no significant reduction of memory. After this short-term memory, the speed of information processing and selective attention weaken. These changes are similar to those in older adults with intellectual disability, giving the impression of early ageing in adults with Down syndrome. In terms of autonomy in everyday life, it is possible to establish an average evolutionary profile. From 60 years old, deterioration is estimated at 45% of the score obtained at 40 years, affecting in particular the skills necessary for the carrying out daily tasks (washing, dressing, feeding without assistance.) We have little knowledge of the psychiatric evolution of this people because older handicapped people are a new group in society. In the three fields of cognition, autonomy and mental health, the ageing of adults with Down syndrome is very sensitive to their environment. Pathological aging of adults with Down syndrome is associated with the dementia syndrome that, with varying degrees, combines disorders of the cognitive functions and behavior, modifying the personality. The clinical diagnosis of dementia is difficult to establish in adults with Down syndrome and opinions diverge, also it is important to comply with three rules: 1) to establish an individual base line and to observe, with the help of regular evaluations, a clear change in performance; this must be confirmed by similar modifications in daily conduct; 2) the decline depends not only on the resources of the subject, but also on the demands made by environment; 3) lasting deterioration of capacities must be clearly greater than that observable in normal ageing to signify dementia. As a function of actual age, adults with Down syndrome show early signs of ageing compared to the general population. One notes the presence of pathological anatomic lesions from 36 years old, which are similar to those observable among adults diagnosed with Alzheimer disease. However it seems that about 20% of adults with Down syndrome do not show clinical signs of dementia 20 years later. The diagnosis, which is delicate to establish, requires an ecological process consistent over time, underlining the influence of the context and the human environment on aging among adults with Down syndrome.

Alzheimer's disease occurs at a higher prevalence and an earlier age in individuals with Down syndrome (DS) than typically developing individuals. However, diagnosing dementia in individuals with intellectual disability remains a challenge due to pre-existing cognitive deficits. The aim of this study was to investigate the validity and reliability of the Brazilian version of the Cambridge Examination for Mental Disorders of Older People with Down's syndrome and Others with Intellectual Disabilities (CAMDEX-DS) for individuals with DS. Two psychiatrists, working independently, evaluated 92 adults with DS > 30 years of age. The concurrent validity of the CAMDEX-DS was analyzed in relation to the gold standard of established international criteria. In a subgroup of 20 subjects, the concurrent validity of the CAMDEX-DS was analyzed in relation to an independent objective assessment of cognitive decline over three years. We analyzed the inter-rater reliability of cognitive assessment. The diagnostic accuracy of the CAMDEX-DS compared to the gold standard was 96.7%. CAMDEX-DS-based diagnosis was considered consistent with cognitive decline. The probability of a participant with dementia having cognitive decline was 83%. Inter-rater reliability for the participant assessment was good, with a kappa of > 0.8 for 93% of the CAMDEX-DS items. The CAMDEX-DS can be considered the first valid and reliable instrument for evaluating dementia in adults with DS in Brazil. Its use in such individuals could improve clinical practice and research.


Amnestic and non-amnestic symptoms of dementia: An international study of Alzheimer's disease in people with Down's syndrome


Abstract: The presence of age-related neuropathology characteristic of Alzheimer's disease (AD) in people with Down syndrome (DS) is well-established. However, the early symptoms of dementia may be atypical and appear related to dysfunction of prefrontal circuitry. To characterize the initial informant reported age-related neuropsychiatric symptoms of dementia in people with DS, and their relationship to AD and frontal lobe function. Non-amnestic informant reported symptoms (disinhibition, apathy, and executive dysfunction) and amnestic symptoms from the CAMDEX-DS informant interview were analyzed in a cross-sectional cohort of 162 participants with DS over 30 years of age, divided into three groups: stable cognition, prodromal dementia, and AD. To investigate age-related symptoms prior to evidence of prodromal dementia we stratified the stable cognition group by age. Amnestic and non-amnestic symptoms were present before evidence of informant-reported cognitive decline. In those who received the diagnosis of AD, symptoms tended to be more marked. Memory impairments were more marked in the prodromal dementia than the stable cognition group (OR = 35.07; P < .001), as was executive dysfunction (OR = 7.16; P < .001). Disinhibition was greater in the AD than in the prodromal dementia group (OR = 3.54; P = .04). Apathy was more pronounced in the AD than in the stable cognition group (OR = 34.18; P < .001). Premorbid amnestic and non-amnestic symptoms as reported by informants increase with the progression to AD. For the formal diagnosis of AD in DS this progression of symptoms needs to be taken into account. An understanding of the unique clinical presentation of DS in AD should inform treatment options.

Forbat, L., & Service, K.P.


Abstract: The complexity of the relationship between intellectual disability (ID) and dementia is increasingly acknowledged. In order to operationalize a route towards person-centered care, we introduce the hierarchy model (Pearce, 1999) as a tool to focus the attention of policy and practice on all aspects of caregiving. This tool, which is taken from the family therapy literature, enables practitioners to examine the broad systems that impact on the delivery and receipt of care. In this article, we focus on its utility in scrutinizing end-of-life and later stages of dementia by illustrating its use with three key areas in dementia care. These three areas provide some of the most challenging situations at the end stages, because of the possible treatment options, they are: nutrition, medical interventions, and the location of care provision. This model enables a focused approach to understanding how meaning is created within social interaction. The article draws out implications for practice and policy and has applications for practice internationally.

Fortea, J., Zaman, S.H., Hartley, S., Rafii, M.S., & Head, H.

Maria Carmona-Iragui, M.

Alzheimer's disease associated with Down syndrome: a genetic form of dementia

The Lancet Neurology, 2021 (Nov. 01), 20(11), 930-942. DOI:https://doi.org/10.1016/S1474-4422(21)00245-3

Abstract: Adults with Down syndrome develop the neuropathological hallmarks of Alzheimer's disease at a very high risk of developing early-onset dementia, which is now the leading cause of death in this population. Diagnosis of dementia remains a clinical challenge because of the lack of validated diagnostic criteria in this population, and because symptoms are overshadowed by the intellectual disability associated with Down syndrome. In people with Down syndrome, fluid and imaging biomarkers have shown good diagnostic performances and a strikingly similar temporality of changes with respect to sporadic and autosomal dominant Alzheimer's disease. Most importantly, there are no treatments to prevent Alzheimer's disease, even though adults with Down syndrome could be an optimal population in whom to conduct Alzheimer's disease prevention trials. Unprecedented research efforts in the field of Down syndrome is rapidly changing this bleak scenario that will translate into disease-modifying therapies that could benefit other populations.


Abstract: Alzheimer's disease and its complications are the leading cause of death in adults with Down syndrome. Studies have assessed Alzheimer's disease in individuals with Down syndrome, but the natural history of biomarker changes in Down syndrome has not been established. We characterised the order and timing of changes in biomarkers of Alzheimer's disease in a population of adults with Down syndrome. We did a dual-centre cross-sectional study of adults with Down syndrome recruited through a population-based health plan in Barcelona (Spain) and through services for people with intellectual disabilities in Cambridge (UK). Cognitive impairment in participants with Down syndrome was classified with the Cambridge Cognitive Examination for Older Adults with Down Syndrome (CAMCOG-DS). Only participants with mild or moderate disability were included who had at least one of the following Alzheimer's disease measures: apolipoprotein E allele carrier status; plasma concentrations of amyloid β peptides 1-42 and 1-40 and their ratio (Aβ1-42/1-40), total tau protein, and neurofilament light chain (NFL); tau phosphorylated at threonine 181 (p-tau), and NFL in cerebrospinal fluid (CSF); and one or more of PET with 18F-fluorodeoxyglucose, PET with amyloid tracers, and MNI Cognitively healthy euploid controls aged up to 75 years who had no biomarker abnormalities were recruited from the Sant Pau Initiative on Neurodegeneration. We used a first-order locally estimated scatterplot smoothing curve to determine the order and age at onset of the biomarker changes, and the lowest ages at the divergence with 95% CIs are also reported where appropriate. Between Feb 1, 2013, and June 28, 2019 (Barcelona), and between June 1, 2009, and Dec 31, 2014 (Cambridge), we included 388 participants with Down syndrome (257 [66%] asymptomatic, 48 [12%] with prodromal Alzheimer's disease, and 83 [21%] with Alzheimer's disease dementia) and 242 euploid controls. CSF Aβ1-42/1-40 and plasma NFL values changed in individuals with Down syndrome as early as the third decade of life, and amyloid PET uptake changed in the fourth decade. 18F-fluorodeoxyglucose PET and CSF p-tau changes occurred later in the fourth decade of life, followed by hippocampal atrophy and changes in cognition in the fifth decade of life. Prodromal Alzheimer's disease was diagnosed at a median age of 50-2 years (IQR 47-55-4), and Alzheimer's disease dementia at 53-7 years (49-5-57-2).

Symptomatic Alzheimer's disease prevalence increased with age in individuals with Down syndrome, reaching 90-100% in the seventh decade.
of life. Alzheimer's disease in individuals with Down syndrome has a long preclinical phase in which biomarkers follow a predictable order of changes over more than two decades. The similarities with sporadic and autosomal dominant Alzheimer's disease and the prevalence of Down syndrome make this population a suitable target for Alzheimer's disease preventive treatments.

Foundation for People with Learning Disabilities
Down's syndrome and dementia - Briefing for Commissioners London: The Foundation for People with Learning Disabilities [c/o Mental Health Foundation, 20/21 Cornwall Terrace, London, England NW1 4QL; e/m mhfd@mhfd.org.uk; www.learningdisabilities.org.uk] (February 2001) 8 pp. Abstract: Backgrounder document, written for funders of services in the United Kingdom, outlines the epidemiology of dementia and Down's syndrome and identifies key support services necessary as part of a package of local services to be established for persons affected by dementia and intellectual disabilities (ID). While titled for dementia and Down's syndrome applicable for all persons with ID. Written in brief style, covers main issues and funding considerations and serves as an excellent planning tool for establishing services. Also covers basic clinical diagnostic information and basis for care management decision making. Routes the reader to associated organizations for further information.

Fray, M.T.

Fredericksen, J. & Fabbre, V.
Down syndrome and Alzheimer's disease: Issues and implications for social work. Journal of Gerontological Social Work, 2018, 61(1), 4-10. DOI: 10.1080/01634372.2017.1393480. Owing to recent medical advancements, people with Down Syndrome (DS) are now able to live considerably longer lives and thus experience a variety of complex issues as they age. Alzheimer's Disease (AD) frequently occurs in older adults who have DS, but few practice guidelines exist to inform social work practice with older adults who have this dual diagnosis. This commentary will highlight the connection between these two conditions within a neurobiological framework and discuss implications for practice based on the available literature on this intersection of ability status, cognitive status, and age.

Frederiks, K.S., Nielsen, T.R., Winblad, B., Schmidt, R., Kramerberg, M.G., Jones, R.W., Hort, J., Grimmer, T., Georges, J., Frölichm L., et al. European Academy of Neurology/European Alzheimer's Disease Consortium position statement on diagnostic disclosure, biomarker counseling, and management of patients with mild cognitive impairment European Journal of Neurology, 2021, 28(7), 2147-2155. doi: 10.1111/ene.14688. Abstract: Careful counseling through the diagnostic process and adequate postdiagnostic support in patients with mild cognitive impairment (MCI) is important. Previous studies have indicated heterogeneity in practice and the need for guidance for clinicians. A joint European Academy of Neurology/European Alzheimer's Disease Consortium panel of dementia specialists was appointed. Through online meetings and emails, positions were developed regarding disclosing a syndrome diagnosis of MCI, pre- and postbiomarker sampling counseling, and postdiagnostic support. Prior to diagnostic evaluation, motives and wishes of the patient should be sought. Diagnostic disclosure should be carried out by a dementia specialist taking the ethical principles of “the right to know” versus “the wish not to know” into account. Disclosure should be accompanied by written information and a follow-up plan. It should be made clear that MCI is not dementia. Prebiomarker counseling should always be carried out if biomarker sampling is considered and postbiomarker counseling if sampling is carried out. A dementia specialist knowledgeable about biomarkers should inform about pros and cons, including alternatives, to enable an autonomous and informed decision. Prebiomarker counseling will depend in part on the results of biomarkers. Follow-up should be considered for all patients with MCI and include brain-health advice and possibly treatment for specific underlying causes. Advice on advance directives may be relevant. Guidance to clinicians on various aspects of the diagnostic process in patients with MCI is presented here as position statements. Further studies are needed to enable more evidence-based and standardized recommendations in the future.

Fromage, B., & Anglade, P.
Avancée en âge du sujet atteint d'une trisomie 21 [The aging of Down's syndrome subjects]. Encephale, 2002, 28(3 Pt 1), 212-216. https://www.lissa.fr/rep/articles/12091781 Abstract: The normal ageing of adults with Down syndrome is comparable to that observed in individuals who have an equivalent cognitive deficit. However it is earlier and is related to the former intellectual level and life story of the person. Before 50 years, there is no significant reduction of memory. After this age short-term memory, the speed of information processing and selective attention weaken. These changes are similar to those in older adults with an intellectual disability, giving the impression of early ageing in Down syndrome subjects. In terms of autonomy in everyday life, it is possible to establish an average evolutionary profile. From 60 years old, deterioration is estimated at 45% of the score obtained at 40 years, affecting in particular the skills necessary for the carrying out daily tasks (washing, dressing, feeding without assistance). We have little knowledge of the psychiatric evolution of this people because older handicapped people are a new group in society. In the three fields of cognition, autonomy and mental health, the ageing of Down syndrome subjects is very sensitive to their environment. Pathological aging of adults with Down syndrome is associated with the dementia syndrome that, with varying degrees, combines disorders of the cognitive functions and behavior, modifying their personality. The clinical diagnosis of dementia is difficult to establish in subjects with Down syndrome and opinions diverge, also it is important to comply with three rules: 1) to establish an individual base line and to observe, with the help of regular evaluations, a clear change in performance; this must be confirmed by similar modifications in daily conduct; 2) the decline depends not only on the resources of the subject, but also on the demands made by environment; 3) lasting deterioration of capacities must be clearly greater than that observable in normal ageing to signify dementia. As a function of actual age, adults with Down syndrome show early signs of aging compared to the general population. One notes the presence of pathological anatomic lesions from 36 years old, which are similar to those observable among patients with Alzheimer's disease. However it seems that about 20% of adults with Down syndrome do not show clinical signs of dementia 20 years later. The diagnosis, which is delicate to establish, requires an ecological process consistent over time, underlining the influence of the context and the human environment on the ageing of adults with Down syndrome.

Funaki, Y., Kaneko, F., & Okamura, H.
Study of factors associated with changes in quality of life of demented elderly persons in group homes Scandinavian Journal of Occupational Therapy, 2005, 12(1), 4-9. doi:10.1080/11038120510031725. Abstract: The purpose of the present study was to attempt to identify changes in quality of life (QOL) and factors associated with them shortly after demented elderly residents entered a group home. The subjects were 25 demented elderly persons who had entered a group home within the previous 3 months. Their QOL and factors associated with it were evaluated on two occasions, at baseline and 3 months later. An objective scale for dementia, the Quality of Life Questionnaire for Dementia (QOL-D), was used to evaluate their QOL. The results showed a significant change between the QOL-D scores at baseline and 3 months later, and changes in housekeeping item scores were extracted as factors associated with changes in QOL-D. These findings suggest that the QOL score rises soon after entering a group home, and that the acquisition of roles within the group home may influence the increase in QOL.

Furness, K.A., Loversseed, A., Lippold, T., & Dodd, K.
The views of people who care for adults with Down's syndrome and dementia: A
service evaluation.


Abstract: It is well established that people with Down’s syndrome are more likely to develop dementia than other people and that onset of dementia is likely to occur earlier at an earlier age. The article reports on a specialist service for people with Down’s syndrome and dementia. The service has offered dementia screening and assessment to people with Down’s syndrome for over 10 years and has also developed to offer support and training for carers. Semi-structured interviews were conducted with family carers, relatives and staff about the impact on them of caring for someone with Down’s syndrome and how the dementia service supports them in this role. The resulting data were analyzed using interpretative phenomenological analysis. The responses provide rich insights into the areas of ‘knowledge and information’, ‘coping and support’ and ‘concerns about the future’. Interviewees also identified services they wanted for the future. As a result of this evaluation, a number of changes have been proposed and begun to be implemented within the service. The results have important implications for other health, social care, and voluntary organizations.


Abstract: People with intellectual disabilities (ID) have a higher risk of developing dementia when compared to the general population. The four most common causes of dementia are Alzheimer's disease, Lewy body dementia, vascular dementia and fronto-temporal dementia. The common presentation of dementia in people with ID includes memory loss and deterioration in speech, personality and behavioral changes, disorientation and functional deterioration. The diagnosis of dementia requires evidence of a definitive change in cognitive functioning such as impairment in memory, language ability (aphasia), ability to perform complex tasks (apraxia) and reduced cognitive performance over time and place. A number of medications are used in delaying the progress of dementia as in general population. Evidence base for the use of medication in people with ID is currently limited. Psychotropic medications have only a limited role in the management of neuropsychiatric symptoms in people with ID and dementia.


Abstract: Down syndrome (DS) has been considered a unique model for the investigation of Alzheimer's disease (AD) but intermediate stages in the continuum are poorly defined. Considering this, we investigated the neuropsychiological (i.e., magnetencephalography [MEG]) and neuropsychological patterns of mild cognitive impairment (MCI) and AD in middle-aged adults with DS. The sample was composed of four groups: Control-DS (n = 14, mean age 44.64 ± 3.30 years), MCI-DS (n = 14, 51.64 ± 3.95 years), AD-DS (n = 13, 53.54 ± 6.58 years), and Control-no-DS (healthy controls, n = 14, 45.21 ± 4.39 years). DS individuals were studied with neuropsychological tests and MEG, whereas the Control-no-DS group completed only the MEG session. Our results showed that the AD-DS group exhibited a significantly poorer performance as compared with the Control-DS group in all tests. Furthermore, this effect was crucially evident in AD-DS individuals when compared with the MCI-DS group in verbal and working memory abilities. In the neuropsychological domain, the Control-DS group showed a widespread increase of theta activity when compared with the Control-no-DS group. With disease progression, this increased theta was substituted by an augmented delta, accompanied with a reduction of alpha activity. Such spectral pattern-specifically observed in occipital, posterior temporal, cuneus, and precuneus regions-correlated with the performance in cognitive tests. This is the first MEG study in the field incorporating both neuropsychological and neuropsychiological information, and demonstrating that this combination of markers is sensitive enough to characterize different stages along the AD continuum in DS.


Abstract: The use of anti-dementia medication in people with intellectual disabilities has been controversial and requires additional research to assess the efficacy of such medications. An essential part of this treatment (both in terms of research and clinical practice) is having robust outcome measures to assess the efficacy of these medications for individuals. Currently there is no consensus in the UK regarding which outcome measures, in conjunction with clinical judgement, are effective in informing clinicians’ decision-making regarding anti-dementia medication. This paper aims to present useful outcome measures. A comprehensive literature search was conducted to identify relevant outcome measures. Outcome measures focused on aspects of patients’ presentation such as cognition, activities of daily living, neuropsychiatric presentation or the impact of their presentation (either on themselves, or on others). These outcome measures were critically appraised to ascertain their suitability in informing clinician’s decisions regarding management of anti-dementia medication. The focus of this appraisal was on good quality measures that are practical and accessible and can be easily used within clinical NHS services. This paper provides advice for clinicians on using appropriate outcome measures in anti-dementia medication management and the symptoms of dementia being targeted, that can be used alongside their clinical assessment to enhance their anti-dementia medication management. Two case studies are presented to illustrate the use of such outcome measures. The case for using a range of assessments that are both broad in focus, and those specifically selected to measure the areas of functioning targeted by the anti-dementia medication, is presented.


Abstract: Down syndrome (DS) is characterised by premature aging and an accelerated decline of cognitive functions in the vast majority of cases. As the life expectancy of DS persons is rapidly increasing, this decline is becoming a dramatic health problem. The aim of this study was to thoroughly evaluate a group of 67 non-demented persons with DS of different ages (11 to 66 years), from neuropsychological, neuropsychiatric and psychomotor point of view in order to evaluate in a cross-sectional study the age-related adaptive and neuropsychological features, and to possibly identify early signs predictive of cognitive decline. The main finding of this study is that both neuropsychological functions and adaptive skills are lower in adult DS persons over 40 years old, compared to younger ones. In particular, language and short memory skills, frontal lobe functions, visuo-spatial abilities and adaptive behavior appear to be the more affected domains. A growing deficit in verbal comprehension, along with social isolation, loss of interest and greater fatigue in daily tasks, are the main features found in older, non demented DS persons evaluated in our study. It is proposed that these signs can be alarm bells for incipient dementia, and that neuro-cognitive rehabilitation and psycho-pharmacological interventions must start as soon as the fourth decade (or even earlier) in DS persons, i.e. at an age where interventions can have the greatest efficacy.


Abstract: Adult patients with Down syndrome (DS) are at higher risk of developing Alzheimer-type dementia and epilepsy. The relationship between developing dementia and the risk of developing seizures in DS is poorly characterized to date. In addition, treatment response and medication tolerability have not been rigorously studied. We identified 220 patients with a diagnosis of DS and dementia. Those without a history of developing seizures (DD) were compared to patients with new-onset seizures (DDS) after the age of 35. Electronic records were reviewed for demographics, seizure characteristics, cognitive status, and psychiatric comorbidities. Of the patients included for
analysis, twenty-six out of 60 patients had new-onset seizures or developed seizures during the follow-up period (the DD+S group) with a median onset of 2.4 years after the dementia diagnosis. Generalized tonic-clonic seizures were the most common seizure type (61.5% of DD+S). Sixteen (61.5%) patients were reported to have myoclonus. Levetiracetam was the most commonly used initial medication, with the majority (73%) of patients treated achieving partial or complete seizure control. The DD+S patients tended to have a similar burden of new-onset neuropsychiatric symptoms compared to the DD group. New-onset epilepsy seems to occur early in the course of dementia in DS patients. Patients generally respond to treatment. A great burden of neuropsychiatric symptoms is seen. Future studies need to explore the relationship between ß-amyloid accumulation and epileptiform activity and attend to the care and needs of DS patients with dementia and seizures.

Abstract: Down syndrome (DS) population has a very high prevalence of obstructive sleep apnoea (OSA), but this remains underdiagnosed. Hence, we aimed to evaluate caregiver's knowledge of OSA and related sociodemographic factors that could contribute to OSA screening patterns in this population. An online survey through the LuMind IDSC Foundation focused on OSA diagnosis, treatments and the number of sleep studies performed. Data were compared between subjects born before and after the American Academy of Pediatrics (AAP) recommendations for OSA screening. Of the caregivers, 724 (parents 96.3%), responded to the survey. The median [interquartile (IQR)] age of the subjects with DS was 12 [20;7] years. The majority (84.3%) had sleep apnoea diagnosis, and half of them were initially referred for a sleep study due to disturbed sleep symptoms. Only 58.7% of the responders were aware of the AAP recommendations. This was linked to socioeconomic and/or educational level and to an earlier OSA diagnosis. The median (IQR) age of OSA diagnosis was lower after the AAP guidelines publication compared with before its publication (3 [4;2] years vs. 10 [18;5] years, P < 0.000). Adenotonsillectomy (81.9%) and continuous positive airway pressure (61.5%) were the most commonly prescribed treatments. Few had discussed other new therapies such as hypoglossal nerve stimulation (16.0%). Only 16.0% of the subjects repeated the sleep study to monitor OSA with ageing, and 30.2% had to wait more than 4 years between studies. This study reinforces the need to improve OSA knowledge of caregivers and clinicians of individuals with DS to promote an earlier diagnosis and optimal treatment of OSA in this population.

Gittin, L.N., and Corcoran, M.
Making homes safer: environmental adaptations for people with dementia Alzheimer's Care Quarterly, 2000, 1(1), 50-58
Abstract: Evaluating the safety of the home environment is an important component of clinical care for persons with dementia. This article discusses safety concerns for persons with dementia living at home alone or with family members, specific modifications to the physical environment to address these issues, and guiding principles for implementing environmental changes. A wide range of environmental strategies can be introduced to maximize home safety. Different adaptations may need to be implemented with progressive memory loss thus necessitating periodic reevaluation of the home.

Abstract: Major increases in the survival of people with Down syndrome during the last two generations have resulted in extended periods of adulthood requiring specialist care, which in turn necessitates greater understanding of the nature, timing and impact of comorbidities associated with the disorder. The prevalence of five comorbidities reported as common in adults with Down syndrome, visual impairment, hearing impairment, epilepsy, thyroid disorders and dementia was assessed by decade of life. From early adulthood, people with Down syndrome are at enhanced risk of developing new comorbidities and they may present with multiple conditions. Three specific challenges are identified and discussed: are comorbidities detected in a timely manner, is the clinical progress of the disorder adequately understood, and who is responsible for the provision of care? Further detailed investigations into the development and treatment of comorbidities across the lifespan are needed for a successful longitudinal approach to healthcare in people with Down syndrome. Authors note that implementation of this approach will better inform healthcare providers to ensure continuity of care with advancing age.

Golaz, J., Charnay, Y., Vallet, P., & Bouras, C.
Abstract: The present status of research clearly demonstrates the occurrence of lesions characteristic of Alzheimer's disease in patients suffering from a Down's syndrome or trisomy 21. The senile plaques appear very early in trisomy 21 (from the age of 20) and are constant after 40 or 45 years. In these two illnesses, the beta-amyloid protein or A4 protein (42 kD) leads to deposits in preferential regions of the central nervous system within two compartments: 1) intracellular, contributing to the formation of neurofibrillary tangles and 2) extracellular, making up the amyloid center of senile plaques as well as around the wall of some blood vessels, then corresponding to the amyloid congophilic angiopathies. Unexpectedly, larger proteins including the A4 sequence have been shown to be normally present in several tissues of normal as well as sick individuals and represent possible precursors of the A4 protein. Alzheimer's disease may happen either sporadically or following a familial incidence associated with an autosomic dominant mode of transmission. Moreover, the risk of incidence of trisomy 21 seems to be enhanced for Collins of Alzheimer's disease patients. Since 1987, the use of molecular biology tools has revealed particularly fruitful. A linkage analysis has been undertaken that showed an association of the putative gene for the familial form of Alzheimer’s disease (FAD) with the gene coding for amyloid precursor proteins.

Gomiero, T., Bertelli, M., Deb, S., Weger, E., Marangoni, A., De Bastiani, E., Mantesso, U., & De Vreese, L.P.
Abstract: The USA National Task Group (NTG) guidelines advocate the use of an adapted version of Dementia Screening Questionnaire for Individuals with Intellectual Disabilities (DSQIID) for dementia screening of individuals with Down syndrome (DS) and with other forms of ID (non-DS). In order to meet these guidelines, this study verifies the psychometric properties of an Italian version of the original DSQIID in a population composed of adults aged 40 years and over with DS and non-DS ID. Internal consistency, inter-rater and intra-rater reliabilities, structural validity, convergent validity and known group differences of DSQIID-I were assessed with 200 individuals with ID (mean of 55.2 years; range: 40-80 years) recruited from 15 different centers in Italy. Diagnosis of dementia was done according to IASSID diagnostic criteria and its degree of clinical certainty was defined according to Silverman et al.’s classification (2004). Cronbach’s alpha for the DSQIID-I was 0.94. The ICCs for inter-rater and test-retest reliability were both 0.89. A Principal Component analysis revealed three domains, namely memory and confusion-related items, motor and functional disabilities, depression and apathy, which explained almost 40% of the overall variance. The total DSQIID-I score correlated significantly with DMR and differed significantly among those individuals (n = 34) with cognitive decline from those without (n = 166). Age, gender and severity of ID were unrelated to the DSQIID-I. The present study confirms the cross-cultural value of DSQIID which was proved to be a psychometrically valid and user-friendly observer-rated scale for dementia screening in adults with both DS and non-DS ID.

Goodman, C., Evans, C., Wilcock, J., Froggatt, K., Drennan, V., Sampson, E., Blanchard, M., Bissett, M., & Iliffe, S.
Abstract: [Non-ID population study] Authors reviewed the evidence for end-of-life care for community dwelling older people with dementia (including
those resident in care homes). An integrated review synthesised the qualitative and quantitative evidence on end-of-life care for community dwelling older people with dementia. English language studies that focused on prognostic indicators for end-of-life care, assessment, support/relief, respite and educational interventions for community dwelling older people with dementia were included. A user representative group informed decisions on the breadth of literature used. Each study selected was screened independently by two reviewers using a standardised check list. Sixty eight papers were included. Only 17% (12) exclusively concerned living and dying with dementia at home. Six studies included direct evidence from people with dementia. The studies grouped into four broad categories: Dementia care towards the end of life, palliative symptom management for people with dementia, predicting the approaches to death for people with dementia and dementia decision making. The majority of studies were descriptive. The few studies that developed dementia specific tools to guide end of life care and outcome measures specific to improve comfort and communication, demonstrated what could be achieved, and how much more needs to be done. Research on end-of-life care for people with dementia has yet to develop interventions that address the particular challenges that dying with dementia poses. There is a need for investigation of interventions and outcome measures for providing end-of-life care in the settings where the majority of this population live and die.

Greenwood, N., Pound, C., & Brearley, S. 'What happens when I can no longer care?' Informal carers' concerns about facing their own illness or death: a qualitative focus group study BMJ Open, 2019 Sep 3(9):e030590. doi: 10.1136/bmjopen-2019-030590. Abstract: Older informal carers play an increasingly important role in supporting others with long-term health conditions. This study aimed to explore in depth the perspectives of older carers (70+ years) supporting others with a variety of conditions and disabilities focusing on their thoughts and experiences about when they are unable to continue caring. The design was qualitative with four focus groups and the setting was Greater London in England. Participants were 28 older carers (70+ years) recruited from the voluntary sector and participated in this study. Most of them were women and many were spouses caring for partners with age-related conditions such as dementia, arthritis and visual impairment. Nearly a third were parents of adult children with severe physical or cognitive disabilities. Findings were that the thematic analysis identified two main aspects for carers when contemplating the future-when they are unable to care in the short term or long term if they die or can no longer manage. Themes included the following: the impact of age, health conditions and relationships on future planning; anxiety about future care; carers' ambivalence and challenges in broaching the subject; interventions that might help older carers talk about and plan for the future of those they care for. Authors note that services need to be open to talking about this difficult topic. Our findings suggest that frank discussions about when older carers cannot care and having plans in place, whether these are financial or address other practical issues, makes it easier for all concerned. However, this issue is not easily broached and its timing and ways to access this support must be carefully and individually gauged. Future research with more diverse demographic groups is needed to improve understanding of these carers' perspectives. Research is also needed to develop interventions to support older carers to talk about and plan for the future.

Hahn, H.E., & Cadogan, M.P. Development and Evaluation of a staff training program on palliative care for persons with intellectual and developmental disabilities Journal of Policy and Practice in Intellectual Disabilities, 2011, Mar, 8(1), 42–52 Abstract: Persons with intellectual and developmental disabilities (IDD) face barriers and disparities at end of life. Among thesebarriers are limited educational opportunities and a paucity of targeted training materials on palliative care for staff who provide their day-to-day care. This paper reports on a three-phase project undertaken to develop, implement, and evaluate a palliative care curriculum and educational program that is responsive to the unique learning needs of staff providing services and supports for individuals with IDD living in long-term care settings. Participants’ ratings of their levels of preparation and comfort to provide palliative care improved from pretraining to posttraining. Posttraining use of materials and practice changes in palliative care occurred. When training is developed in partnership with the staff who will use these training resources, it has the potential to sustain use and to alter the care practices to address the palliative care needs of persons with IDD.

Hamadseed O., Elkhidir, I.H., & Skutella, T. Psychosocial risk factors for Alzheimer's disease in patients with Down syndrome and their association with brain changes: A Narrative Review. Neurology and Therapy, 2022 Sep,11(3), 931-953. doi: 10.1007/s40120-022-00361-9. Epub 2022 May 21. Abstract: Several recent epidemiological studies attempted to identify risk factors for Alzheimer’s disease. Age, family history, genetic factors (APOE genotype, trisomy 21), physical activity, and a low level of schooling are significant risk factors. In this review, we summarize the known psychosocial risk factors for the development of Alzheimer’s disease in patients with Down syndrome and their association with neuroanatomical changes in the brains of people with Down syndrome. We completed a comprehensive review of the literature on PubMed, Google Scholar, and Web of Science about psychosocial risk factors for Alzheimer’s disease, for Alzheimer’s disease in Down syndrome, and Alzheimer’s disease in Down syndrome and their association with neuroanatomical changes in the brains of people with Down syndrome. Alzheimer's disease causes early pathological changes in individuals with Down syndrome, especially in the hippocampus and corpus callosum. People with Down syndrome living with dementia showed reduced volumes of brain areas affected by Alzheimer's disease as the hippocampus and corpus callosum in association with cognitive decline. These changes occur with increasing age, and the presence or absence of psychosocial risk factors impacts the degree of cognitive function. Correlating Alzheimer’s disease biomarkers in Down syndrome and cognitive function scores while considering the effect of psychosocial risk factors helps us identify the mechanisms leading to Alzheimer's disease at an early age. Also, this approach enables us to create more sensitive and relevant clinical, memory, and reasoning assessments for people with Down syndrome.


Handen, B.L. The search for biomarkers of Alzheimer’s disease in Down syndrome American Journal of Intellectual and Developmental Disabilities, 2020,125(2), 97-99. doi: 10.1352/1944-7558-125.2.97. Abstract: Adults with Down syndrome are at high risk for Alzheimer’s disease (AD), with most individuals developing clinical dementia by their late 60s. This increased risk for AD has been attributed, at least in part, to triplication and overexpression of the gene for amyloid precursor protein (APP) on chromosome 21, leading to elevated levels of amyloid β peptides. This article offers a brief overview of our current knowledge of AD in the DS population. In addition, the NIA/NIH-funded, multicenter longitudinal study of biomarkers of AD in adults with DS is explored. The Alzheimer’s Biomarkers Consortium—Down Syndrome (ABC-DS) is a longitudinal study of Alzheimer Disease biomarkers in adults with Down syndrome supported by federal grants from the National Institute on Aging (NIA) and the Eunice Kennedy Shriver National Institute of Child Health and Human Development (NICHD). The primary goal of ABC-DS is to understand the factors that moderate the relationship between Aβ, neurodegeneration and dementia in DS and biomarkers for those factors that could be critically important in the design of effective therapeutic trials for AD, not only in DS, but in the general population as well.

Biomarker Consortium-Down Syndrome (ABC-DS).


Abstract: Adults with Down syndrome (DS) are at exceptionally high risk for Alzheimer's disease (AD), with virtually all individuals developing key neuropathological features by age 40. Identifying biomarkers of AD progression in DS can provide valuable insights into pathogenesis and suggest targets for disease modifying treatments. This article describes the development of a multi-center, longitudinal study of biomarkers of AD in DS. The protocol includes longitudinal examination of clinical, cognitive, blood and cerebrospinal fluid biomarkers, brain imaging, magnetic resonance imaging and positron emission tomography measures (at 16-month intervals), as well as genetic modifiers of AD risk and progression. Approximately 400 individuals will be enrolled in the study (more than 370 to date). The methodological approach from the administrative, clinical, neuroimaging, omics, neuropathology, and statistical cores is provided. This represents the largest U.S.-based, multi-site, biomarker initiative of AD in DS. Findings can inform other multidisciplinary networks studying AD in the general population.


Abstract: Primary care integration of Down syndrome (DS)-specific dementia screening is strongly advised. The current study employed principal components analysis (PCA) and classification and regression tree (CART) analyses to identify an abbreviated battery for dementia classification. Scale- and subscale-level scores from 141 participants (no dementia n = 68; probable Alzheimer’s disease n = 73), for the Severe Impairment Battery (SIB), Dementia Scale for People with Learning Disabilities (DLD), and Vineland Adaptive Behavior Scales—Second Edition (Vineland-II) were analyzed. Two principle components (PC1, PC2) were identified with the odds of a probable dementia diagnosis increasing 2.54 times per PC1 unit increase and by 3.73 times per PC2 unit increase. CART analysis identified that the DLD sum of cognitive scores (SCS < 35 raw) and Vineland-II community subscore (< 36 raw) scores best classified dementia. No significant difference in the PCA versus CART area under the curve (AUC) was noted (DLD(65.196) = 0.57683; p = 0.57; CART AUC = 0.87; CART AUC = 0.91). The PCA sensitivity was 80% and specificity was 75%; CART was 100% and specificity was 81%. These results support an abbreviated dementia screening battery to identify at-risk individuals with DS in primary care settings to guide specialized diagnostic referral.


Abstract: Cognitive decline in older people with intellectual disabilities (ID) is often under-recognized. Following the publication of the National Service Framework for Older People and the white paper Valuing People, older people with intellectual disabilities of all aetiologies should have access to a systematic assessment of their cognitive function in order to detect decline in cognition and adaptive skills and implement appropriate treatments as early as possible. The development of a memory clinic for older people with ID is described, including instruments used and characteristics of attendees. Such projects are in line with current UK government policies and can contribute to the improvement of standards of care and support research in this vulnerable group of people.


Abstract: There is a critical need to identify measures of cognitive functioning sensitive to early Alzheimer's disease (AD) pathophysiology in Down syndrome to advance clinical trial research in this at-risk population. The objective of the study was to longitudinally track performance on cognitive measures in relation to necortical and striatal amyloid beta (Aβ) in non-demented Down syndrome. The study included 118 non-demented adults with Down syndrome who participated in two to five points of data collection, spanning 1.5 to 8 years. Episodic memory, visual attention and executive functioning, and motor planning and coordination were assessed. Aβ was measured via [C-11] Pittsburgh Compound-B (PiB) PET. PiB was associated with level and rate of decline in cognitive performance in episodic memory, visual attention, executive functioning, and visuospatial ability in models controlling for chronological age. The Cued Recall Test emerged as a promising indicator of transition from preclinical to prodromal AD.


Abstract: Trisomy 21 causes Down syndrome (DS) and is a recognized cause of early Alzheimer's disease (AD). The current study sought to determine if premorbid intellectual disability level (ID) was associated with variability in age-trajectory of AD biomarkers and cognitive impairments. General linear mixed models compared the age-trajectory of the AD biomarkers PET Aβ and tau and cognitive decline across premorbid ID levels (mild, moderate, and severe/profound), in models controlling trisomy type, APOE status, biological sex, and site. Analyses involved adults with DS from the Alzheimer's Biomarkers Consortium-Down Syndrome. Participants completed measures of memory, mental status, and visuospatial ability. Premorbid ID level was based on IQ or mental age scores prior to dementia concerns. PET was acquired using [11C] PiB for Aβ and [18F] AV-1451 for tau. Cognitive data was available for 361 participants with a mean age of 45.22 (SD = 9.92) and PET biomarker data was available for 154 participants. There was not a significant effect of premorbid ID level by age on cognitive outcomes. There was not a significant effect of premorbid ID by age on PET Aβ or on tau PET. There was not a significant difference in age at time of study visit of those with mild cognitive impairment-DS or dementia by premorbid ID level. Findings provide robust evidence of a similar time course in AD trajectory across premorbid ID levels, laying the groundwork for the inclusion of individuals with DS with a variety of IQ levels in clinical AD trials.


Abstract: Despite the acknowledged increase in the number of older people with intellectual disabilities (ID) in the UK, the age-related health and social care needs of this population have yet to be fully understood and addressed. Although there is some evidence of positive development, the current picture of service provision is characterized by fragmentation and limited choice of resources and specialist care. Policy aims are variably met and inconsistently applied. Research suggests that service planning is often incoherent, that many older people with ID and their carers receive poor quality non-specialist care and that staff are inadequately trained to manage the often multiple and complex needs of this user group. There is a considerable co-joined service development and research challenge in this emerging field. If older people with ID and their carers are to receive quality provision, a coherent and well-funded service planning system is required which is underpinned by articulated agency partnerships, informed by good practice developments in the fields of ID, gerontology and dementia care, and linked to evidence about effective models of care and services. The incorporation of the perspectives of users and carers in the planning process is an essential pre-requisite as is a commitment to the development of effective support across the life course of all individuals with ID.

Head, E., Chen, G., Wang, Z., Guo, C., Li, N., Yun, C., & Zheng, X.

Adults with intellectual disabilities in China: comorbid psychotic disorder and its association with health service utilisation
Abstract: Adults with intellectual disabilities (ID) often have multiple comorbidities. Psychiatric disorders in this population have been poorly studied in developing countries. We aimed to investigate the prevalence of psychiatric disorders in adults with ID and whether comorbid psychiatric disorders were associated with health service utilization. We obtained data from the Second National Sample Survey on Disability, conducted in 31 provinces of China and selected a subsample of 13,631 adults aged 18 years and above with ID. Rates were defined by intelligence quotient score under 70, deficits in two or more adaptive behaviours and age of onset under 18 years. Psychiatric disorders were identified according to the International Statistical Classification of Diseases, Tenth Revision. Logistic regressions were used for data analyses. The prevalence of psychiatric disorders in adults with ID was 16.7%. The most prevalent type of psychiatric disorder was dementia. Older adults, females, being minorities, urban residents, being literate, low-income groups and having severe ID, were associated with elevated risk of psychiatric disorders among adults with ID. Compared with individuals without psychiatric disorders, those with comorbid psychiatric disorders were more likely to use medical service and less likely to use rehabilitation service. The prevalence of psychiatric disorder in adults with ID was strikingly higher than that in the general population. Health service utilisation among Chinese adults with ID remained a big challenge. There is a possibility of diagnostic overshadowing by local clinicians, which may have resulted in overdiagnosis of dementia and underdiagnosis of common mental disorders. This study informs further investigations regarding common mental disorders among people with ID and has implications for public health strategies and health policies to meet health service need for this population.

Head, E., Lott, I.T., Wilcock, D.M., & Lemere, C.A.

Aging in Down syndrome and the development of Alzheimer's disease neuropathology
Abstract: Chromosome 21, implicated in Down Syndrome, contains several genes that are thought to play a critical role in the development of AD neuropathology. The overexpression of the gene for the amyloid precursor protein (APP), on chromosome 21, leads to early onset beta-amyloid (Aβ) plaques in DS. In addition to Aβ accumulation, middle-aged people with DS develop neurofibrillary tangles, cerebrovascular pathology, white matter pathology, oxidative damage, neuroinflammation and neuron loss. There is also evidence of potential compensatory responses in DS that benefit the brain and delay the onset of dementia after there is sufficient neuropathology for a diagnosis of AD. This review describes some of the existing literature and also highlights gaps in our knowledge regarding AD neuropathology in DS. It will be critical in the future to develop networked brain banks with standardized collection procedures to fully characterize the regional and temporal pathological events associated with aging in DS. As more information is acquired regarding AD evolution in DS, there will be opportunities to develop interventions that are age-appropriate to delay AD in DS.

Head, E., Powell, D., Gold, B.T., Schmitt, F.A.

Alzheimer's disease in Down syndrome

Heller, T., Scott, H.M., & Janicki, M.P.

Published online 2018 Jul 10. doi: 10.1016/j.trci.2018.06.002
Abstract: A specially commissioned working group produced a report on caregiving, intellectual and developmental disabilities (IDDs), and dementia for the National Institutes of Health-located National Research Summit on Care, Services, and Supports for Persons with Dementia and Their Caregivers. Experts in caregiving, dementia, and IDDs examined the current state of research, policy, and practice related to caregiving and supports; identified the similarities and dissimilarities between IDD-related care and services and the general population affected by dementia; and considered how these findings might contribute to the conversation on developing a dementia care research and services development agenda. Five major areas related to programs and caregiving were assessed: (1) challenges of dementia; (2) family caregiving interventions; (3) supportive care settings; (4) effects of diversity; and (5) bridging service networks of aging and disability. Recommendations included increasing supports for caregivers of adults with IDDs and dementia; increasing research on community living settings and including caregivers of persons with IDDs in dementia research; acknowledging cultural values and practice diversity in caregiving; increasing screening for dementia and raising awareness; and leveraging integration of aging and disability networks.

Herron, D.L., & Priest, H.M.

Support workers' knowledge about dementia: A vignette study.
Abstract: It is widely acknowledged that people with intellectual disabilities are highly likely to experience mental health problems, but that support workers' knowledge and skill in this area is sometimes lacking. There is little research explicitly exploring knowledge about the mental health of older people with intellectual disabilities and the purpose of this paper is to attempt to fill this gap. In total, 14 support workers completed a questionnaire in which three vignettes presented progressively worsening indicators of dementia in an older person with intellectual disabilities. Participants explained what they thought was happening and what action they would take. Data were analysed using Braun and Clarke's framework. Few participants had undertaken any mental health
training, and only one in relation to older people. They were generally poor at judging early and intermediate indicators of dementia, but were able to identify more overt later signs. However, they believed these advanced indicators to be the onset of dementia. Nonetheless, they would generally take appropriate action, such as observation and referral. Abuse was often considered as a causal factor. The most significant implication is the need for training in the mental health needs of older people and in particular, the general and specific indicators and expected trajectory of dementia in this population.

Herron, D.L., Priest, H.M., & Read, S.
Abstract: There is a need to better understand the experiences and support needs of paid and family carers of people with an intellectual disability and dementia, and the role of Intellectual Disability Dementia Care Pathways (IDDCPs). This study explored the experiences of carers, and IDDCPs and other support structures within those experiences. A constructivist grounded theory methodology was implemented. Data were obtained through 23 semi-structured interviews with two family carers, eight paid carers and eight healthcare professionals. The study aimed to suggest that older people with an intellectual disability may be better served in intellectual disability homes rather than older people homes and that it is an area of research which needs further exploration.

Abstract: People with Down syndrome (DS) typically develop Alzheimer’s disease (AD) neuropathology before age 40, but a lack of outcome measures and longitudinal data have impeded their inclusion in randomized controlled trials (RCTs). A cohort study was employed in which event-based and dose-response Emax models were fitted to longitudinal cognitive data, to stage AD and determine the earliest ages of decline. Results informed sample size estimations for hypothetical RCTs of disease-modifying treatments that reduced decline by 35% or 75%. Seventy-five percent of participants progressed or remained stable in the AD staging model, effect sizes varied by age group and tests. Varied treatment effects could be detected with 50-200 people per arm when using sensitive cognitive outcome measures and targeting recruitment to ages 36 to 45 years. Efficient RCTs of AD preventative treatments can be conducted in the DS population using sensitive outcome measures to monitor early decline. Dose-response models could help tailor future RCTs.

Hithersay, R., Hamburg, S., Knight, B., & Strydom, A.
Abstract: Alzheimer’s disease is most likely universal in older individuals with Down syndrome, due to having three copies of the amyloid precursor protein gene, resulting in amyloid-beta plaque deposition. Down syndrome is an important population in which to consider clinical trials of treatments to prevent or delay the development of dementia. However, assessment of subtle changes in cognitive changes is challenging due to the presence of intellectual disability. Recent research confirmed that older adults with Down syndrome often present with cognitive decline: more than 80% may experience dementia by age 65 years. Efforts have been made to improve and validate neuropsychological assessment and to describe the relationship with comorbidities such as epilepsy and haemorrhagic stroke. There have also been advances in biomarkers such as neuroimaging using amyloid PET. Clinical trials of treatments, particularly in the presymptomatic phase of Alzheimer’s disease, are important to consider in individuals with Down syndrome given their high dementia burden, and may also serve as proof of concept for other forms of Alzheimer’s disease. However, further work is required to improve outcome measures and better understand the biomarkers of progression of disorder and their relationship with symptom development during the presymptomatic period.
Association of dementia with mortality among adults with Down syndrome older than 35 years.


Abstract: This work quantifies the fatal burden of dementia associated with Alzheimer disease in individuals with Down syndrome (DS). To explore the association of dementia associated with Alzheimer disease with mortality and examine factors associated with dementia in adults with DS. Prospective longitudinal study in a community setting in England. Data collection began March 29, 2012. Cases were censored on December 13, 2017. The potential sample consisted of all adults 36 years and older from the London Down Syndrome Consortium cohort with 2 data times and dementia status recorded (N = 300); 6 withdrew from study, 28 were lost to follow-up, and 55 had a single data collection point at time of analysis. The final sample consisted of 211 participants, with 503.92 person-years' follow-up. Dementia status, age, sex, APOE genotype, level of intellectual disability, health variables, and living situation. Crude mortality rates, time to death, and time to dementia diagnosis with proportional hazards of predictors. Of the 211 participants, 96 were women (45.5%) and 66 (31.3%) had a clinical dementia diagnosis. Twenty-seven participants (11 female; mean age at death, 56.74 years) died during the study period. Seventy percent had dementia. Crude mortality rates for individuals with dementia (1191.85 deaths per 10 000 person-years; 95% CI, 1168.49-1215.21) were 5 times higher than for those without (232.22 deaths per 10 000 person-years; 95% CI, 227.67-236.77). For those with dementia, APOE e4 carriers had a 7-fold increased risk of death (hazard ratio [HR], 6.91; 95% CI, 1.756-27.195). For those without dementia, epilepsy with onset after age 36 years was associated with mortality (HR, 9.66; 95% CI, 1.59-58.56). APOE e4 carriers (HR, 4.91; 95% CI, 2.53-9.56), adults with early-onset epilepsy (HR, 3.61; 95% CI, 1.12-11.60), multiple health comorbidities (HR, 1.956; 95% CI, 1.087-3.519), and those living with family (HR, 2.14; 95% CI, 1.08-4.20) received significantly earlier dementia diagnoses. Dementia was associated with mortality in 70% of participants with DS. APOE e4 carriers and/or people with multiple comorbid health conditions were at increased risk of dementia and death, highlighting the need for good health care. For those who died without a dementia diagnosis, late-onset epilepsy was the only significant factor associated with death, raising questions about potentially undiagnosed dementia cases in this group.


Establishing a database for proactive screening of adults with Down’s syndrome: when services work together.


Abstract: This paper describes a service improvement project with two aims: to identify and screen all adults with Down syndrome aged over 30 years in a defined locality using a standardised instrument to establish functional baselines; and to set up a database to facilitate early diagnosis of dementia in this population. An assistant psychologist used a standardised instrument to screen participants who were identified through contact with health, social, and third sector, and housing services. Eligible people were identified and screened using an informant-based measure. Three groups were identified: group 1 showed no significant change; group 2 showed significant change but no signs of dementia; and group 3 showed significant change plus signs of dementia. People with suspected dementia were referred on for further investigation/assessment and supportive services. Terminology is important in engaging families in a screening project, as is the opportunity to provide information. A proactive screening project can be established by employing working partnerships between intellectual disability and older adult services to aid diagnosis. Adults with Down syndrome aged over 30 years in a defined locality can be identified through contact with health, social, and third sector, and housing services. Those identified can be screened using a standardised instrument and a database of screening results established in order to establish baselines against which future re-screening can be conducted. Partnership working between older adult mental health services and intellectual disability services can improve the diagnostic service to adults with Down syndrome.

Hoekman, J., & Maaskant, M.A.

Comparison of instruments for the diagnosis of dementia in individuals with intellectual disability.


Abstract: The authors describe the agreement among the results (dementia/no dementia) of three instruments used for the potential diagnosis of dementia in persons with intellectual disability. The instruments are: the Dementia Questionnaire for Mentally Retarded Persons (DMR), the Checklist with Symptoms of Dementia (CLD) and the Delayed Match-to-Sample Test (DMTS). The results were compared with the expert opinion of a physician/educational specialist/psychologist. The participants were 329 adults affiliated with centres for people with intellectual disability in The Netherlands. It was found that the agreement among the three tools was low (kappa < 0.5). The agreement between the expert opinion and the results of the tests was also found to be low. It was concluded that the instruments do not mutually agree upon which of the adults can be regarded as dementing or not dementing and they also provide inconsistent agreement with the expert opinion when dementia is present. It was further concluded that it is not advisable to use a single instrument when attempting to diagnose dementia in people with intellectual disability.
Abstract: Book chapter that examines the implications of Alzheimer's disease for adults with Down syndrome, including assessment and diagnosis and specialty service provision. Authors note that assigning a tenable diagnosis of Alzheimer disease requires careful and comprehensive data assembly, including medical history, clinical examination, neuropsychological assessment and laboratory investigations. Once the diagnosis is established, effective ongoing management should focus on supporting not only the affected individual (including advocacy for his or her rights) but also the family and professional carers. During the course of the illness various medical, psychiatric and psychological interventions can be helpful as can changes in the environment. A wide range of services for persons with Down syndrome who develop Alzheimer's disease makes it possible for affected individuals, despite deterioration, to remain in the family home or in community residential settings. Authors propose some general suggestions for services and adaptations.

Holland, A.J., Hon, J., Huppert, F.A., & Stevens, F.
Incidence and course of dementia in people with Down's syndrome: findings from a population-based study.
Journal of Intellectual Disability Research, 2000, 44(2), 138-146. DOI: 10.1046/j.1365-2788.2000.00283.x
Abstract: The prevalence rate of Alzheimer's disease (AD) in people with Down's syndrome (DS) increases significantly with age. However, the nature of the early clinical presentation, course and incidence rates of dementia are uncertain. The aims of the present study were to investigate the characteristics of age-related changes and incidence rates for dementia in a population-based sample of people with DS aged 30 years and older at the age of risk for dementia. A modified version of the Cambridge Examination for Mental Disorders of the Elderly informant interview was used to determine the extent and nature of changes in memory, personality, general mental functioning and daily living skill 18 months after a similar assessment. At the time of the first assessment, the initial changes reported were predominately in behavior and personality. At the second assessment, although estimated incidence rates for dementia in people with DS are high (0.24), mainly in the younger groups, with incidence rates of AD, meeting both ICD-10 and DSM-IV criteria, of 0.04 predominately in the older groups. The present authors have hypothesized that the observed personality changes and the high estimated incidence rates of frontal-like dementia in the younger groups may indicate that functions served by the frontal lobes are the first to be compromised with the progressive development of Alzheimer-like neuropathology in people with DS.

Holst, G., Johansson, M., & Ahlstrom, G.
Signs in people with intellectual disabilities: Interviews with managers and staff on the identification process of dementia.
Abstract: The life expectancy of people with intellectual disabilities (ID) has steadily increased, which has been accompanied by an increased risk of dementia. Staff and managers are key resources for safety diagnosis since they deliver information about people with ID behavior every day. The aim of the present study was to explore the identification process employed by staff and managers to detect signs of suspected dementia in people with an ID within intellectual disability services (ID-services). Twenty managers and 24 staff within an ID-service were interviewed and qualitative latent content analysis was applied. A model consisting of three themes on three levels of resources for the identification process of signs of suspected dementia emerged from the analysis. On the first level was the time and continuity in the care relationship, which is crucial for identifying and responding to changes in cognitive ability that indicate dementia. On the second level, the staff identify deficiencies in their own knowledge, seek support from colleagues and managers within their workplace and, on the third level, outside their workplace. Staff and managers expressed a need for early and continuous guidance and support from specialists in dementia and primary healthcare. This finding indicates an urgent need for intervention research and digital support for staff in dementia care. Only when these resources are used will it be possible for staff to identify, and react to, dementia symptoms in a safe and knowledge-based way, initiate examination, design well-adapted care, and, if necessary, consider moving the person to adapted housing. A close collaboration with specialists within dementia care may contribute to a good working environment for staff with skilled guidance and support, continuous education, and the initiation of intervention research concerning the best accommodation form and digital support for people with an ID and staff.

Hom, C.
Neuropsychological subtypes of incident mild cognitive impairment in Down syndrome.
Alzheimer's & Dementia, 2020, 16(S6), https://doi.org/10.1002/alz.043299
Abstract: Past attempts to characterize the earliest cognitive changes as individuals with Down Syndrome (DS) transition from cognitively stable to mild cognitive impairment (MCI) have been equivocal (Garcia-Alba et al., 2019; Lautarescu et al., 2017). Differences identifying MCI in this population are complicated by variability in pre-morbid cognitive abilities, the use of neuropsychological tests that were created for the neuropsychological population, and participants scoring at floor on the baseline assessment (Kinsky-McHale and Silverman, 2013). We examined data from 151 individuals with Down Syndrome (M age=50.25, SD age=6.94). Their pre-morbid level of intellectual impairment ranged from mild to severe. All participants received comprehensive evaluations. Following data collection, the clinical status of each participant was rated at consensus review that considered performance on a core neuropsychological test battery and the clinical data for each participant. Data from the non-demented and MCI groups are examined: Cognitive Stable (N=107, 70.9%) and MCI-DS (N=44, 29.1%). The full battery consists of 27 subtests that were hypothesized a priori to measure five cognitive domains: language, memory, executive function, visuospatial reasoning, and motor coordination. Factor analysis revealed 7 principal components that maximally discriminated between test scores in older adults with DS who have not reached clinical AD status: (1) general intelligence (2) sensorimotor, (3) memory, (4) language comprehension and expression, (5) executive function/speed, (6) attention/language expression, and (7) visuospatial. Cluster analysis for the MCI group produced 3 distinct groups: (1) dysexecutive (n=4), (2) dynamic/visuospatial impaired (n=28), and (3) amnesic/motor impaired (n=12). Author concludes that the neuropsychological battery assesses 7 distinct cognitive functions in older adults with DS. It can also capture cognitive decline, as we were able to empirically identify three distinct neuropsychological subtypes of MCI: amnesic/visuospatial impaired, dysexecutive, and dynamic. These subtypes are generally consistent with those that have been found within the neuropsychological population (Edmonds et al., 2015; Dick et al., 2016), strengthening the evidence that AD has a similar course in the DS population and late onset AD.

Horovitz, M., Kozlowski, A. M., & Matson, J. L.
Compliance training in an adult with dementia of the Alzheimer's type and Down syndrome.
Abstract: The authors describe the treatment of noncompliance and public stripping in a 53-year-old man with Down syndrome and dementia. Based on a review of the relevant literature and a comprehensive functional behavioral analysis, an intervention procedure utilizing contingent reinforcement of compliance was conducted. Compliance with requests to allow assistance getting dressed was reinforced with a combination of verbal praise, physical contact, and edible reinforcers. Noncompliance had decreased by 85.22% at follow-up. Factors responsible for intervention success and implications for clinicians are discussed.

Horvath, S., Garagnani, P., Bacalini, M.G., Pirazzini, C., Salvioli, S., Davide, G., Di Blasio, A.M., Giuliani, C., Tung, S., Vinters, H.V., & Franceschi, C.
Accelerated epigenetic aging in Down syndrome.
Alzheimer's & Dementia, 2020, 16(5), 1-5, eprint. DOI: 10.1111/acel.12325
Abstract: Down syndrome (DS) entails an increased risk of many chronic diseases that are typically associated with older age. The clinical manifestations of accelerated aging suggest that trisomy 21 increases the biological age of tissues, but molecular evidence for this hypothesis has been sparse. Here, we utilize a quantitative molecular marker of aging (known as the epigenetic clock) to demonstrate that trisomy 21 significantly increases the age of blood and brain tissue (on average by 6.6 years, P = 7.0 3 10⁻¹⁸).
Research has shown that there is variability in the quality of life (QOL) outcomes for people with intellectual disabilities who live in group homes. The aim was to examine dimensions of group home culture as predictors of QOL outcomes. The Group Home Culture Scale (GHCS) was used to measure staff perceptions of culture in 23 group homes. QOL data were available from 98 people with intellectual disabilities. Multilevel modelling was used to examine the associations between the GHCS subscales and four QOL-dependent variables. Of the GHCS subscales, Effective Team Leadership and Alignment of Staff with Organizational Values significantly predicted residents' engagement in activities. Supporting Well-Being significantly predicted residents' community involvement. None of the GHCS subscales significantly predicted domestic participation and choice making. The findings suggest that strategies to improve Effective Team Leadership and Supporting Well-Being dimensions of culture may contribute to enhancing certain QOL outcomes.

Huxley, A., Van-Schaik, P., & Witts, P.


Abstract: Research has shown that there is variability in the quality of life (QOL) outcomes for people with intellectual disabilities who live in group homes. The aim was to examine dimensions of group home culture as predictors of QOL outcomes. The Group Home Culture Scale (GHCS) was used to measure staff perceptions of culture in 23 group homes. QOL data were available from 98 people with intellectual disabilities. Multilevel modelling was used to examine the associations between the GHCS subscales and four QOL-dependent variables. Of the GHCS subscales, Effective Team Leadership and Alignment of Staff with Organizational Values significantly predicted residents' engagement in activities. Supporting Well-Being significantly predicted residents' community involvement. None of the GHCS subscales significantly predicted domestic participation and choice making. The findings suggest that strategies to improve Effective Team Leadership and Supporting Well-Being dimensions of culture may contribute to enhancing certain QOL outcomes.

Iacono, T., Bigby, C., Carling-Jenkins, R., & Torr, J.

Abstract: Research has shown that there is variability in the quality of life (QOL) outcomes for people with intellectual disabilities who live in group homes. The aim was to examine dimensions of group home culture as predictors of QOL outcomes. The Group Home Culture Scale (GHCS) was used to measure staff perceptions of culture in 23 group homes. QOL data were available from 98 people with intellectual disabilities. Multilevel modelling was used to examine the associations between the GHCS subscales and four QOL-dependent variables. Of the GHCS subscales, Effective Team Leadership and Alignment of Staff with Organizational Values significantly predicted residents' engagement in activities. Supporting Well-Being significantly predicted residents' community involvement. None of the GHCS subscales significantly predicted domestic participation and choice making. The findings suggest that strategies to improve Effective Team Leadership and Supporting Well-Being dimensions of culture may contribute to enhancing certain QOL outcomes.
in white participants, it was 3.18-fold greater (CI: 1.94-5.2, p < 0.001). In a secondary analysis, compared with MBI-no-psychosis (n = 2,260), MBI-psychosis had a 2.47-fold greater incidence of dementia (CI: 1.69-3.59, p < 0.001). Although psychosis is an infrequently endorsed MCI domain, when present it is associated with substantial risk for dementia. HRs differed between cognitive strata, and these differences were significantly greater when MBI-psychosis emerged in NC as opposed to MCI, emphasizing the importance of cognitive assessment at the time of symptom emergence. In addition, the relationship between MBI-psychosis and incident dementia was stronger in Black participants than in white participants. The emergence of persistent psychotic symptoms in older adults is clinically meaningful, and MBI-psychosis identifies a high-risk group for precision medicine approaches to dementia prevention.


Abstract: The study of sex differences in Alzheimer's disease is increasingly recognized as a key priority in research and clinical development. People with Down syndrome represent the largest population with a genetic link to Alzheimer's disease (>90% in the 7th decade). Yet, sex differences in Alzheimer's disease manifestations have not been fully investigated in these individuals, who are key candidates for preventive clinical trials. In this double-centre, cross-sectional study of 628 adults with Down syndrome (46% females, 44% [34.6; 50.7] years), we compared Alzheimer's disease prevalence, as well as cognitive outcomes and AT(N) biomarkers across age and sex. Participants were recruited from a population-based health plan in Barcelona, Spain, and from a convenience sample recruited via services for people with intellectual disabilities in England and Scotland. They underwent assessment with the Cambridge Cognitive Examination for Older Adults with Down Syndrome, modified cued recall test and determinations of brain amyloidosis (CSF amyloid-β 42 / 40 and amyloid-PET), tau pathology (CSF and plasma phosphorylated-tau181) and neurodegeneration biomarkers (CSF and plasma neurofilament light, total-tau, fluorodeoxyglucose-PET and MRI). We used within-group locally estimated scatterplot smoothing models to compare the trajectory of biomarker changes with age in females versus males, as well as by apolipoprotein ε4 carriership. Our work revealed similar prevalence, age at diagnosis and Cambridge Cognitive Examination for Older Adults with Down Syndrome scores by sex, but males showed lower modified cued recall test scores from age 45 compared with females. AT(N) biomarkers were comparable in males and females. When considering apolipoprotein ε4, female ε4 carriers showed a 3-year earlier age at diagnosis compared with female non-carriers (50.5 versus 53.2 years, P = 0.01). This difference was not seen in males (52.2 versus 52.5 years, P = 0.76). Our exploratory analyses considering sex, apolipoprotein ε4 and biomarkers showed that female ε4 carriers tended to exhibit lower CSF amyloid-β 42/amyloid-β 40 ratios and lower hippocampal volume compared with females without this allele, in line with the clinical difference. This work showed that biological sex did not influence clinical and biomarker profiles of Alzheimer's disease in adults with Down syndrome. Consideration of apolipoprotein ε4 haplotype, particularly in females, may be important for clinical research and clinical trials that consider this population. Accounting for, reporting and publishing sex-stratified data, even when no sex differences are found, is central to helping advance precision medicine.


Abstract: People with Down syndrome have a high risk of developing Alzheimer disease dementia. However, penetration and age at onset are considered variable, and the association of this disease with life expectancy remains unclear because of underreporting in death certificates. The authors assessed whether the variability in symptom onset of Alzheimer disease in Down syndrome is similar to autosomal dominant Alzheimer disease and to assess its association with mortality. This study combines a meta-analysis with the assessment of mortality data from US death certificates (n = 77,347 case records with an International Classification of Diseases code for Down syndrome between 1968 to 2019; 57,900 [49%] female) and from a longitudinal cohort study (n = 889 individually dated years of follow-up) from the Down Syndrome and Alzheimer disease Neuroimaging Initiative (DABNI). A meta-analysis was conducted to investigate the age at onset, age at death, and duration of Alzheimer disease dementia in Down syndrome. PubMed/Medline, Embase, Web of Science, and CINAHL were searched for research reports, and OpenGray was used for gray literature. Studies with data about the age at onset or diagnosis, age at death, and disease duration were included. Pooled estimates with corresponding 95% CIs were calculated using random-effects meta-analysis. The variability in disease onset was compared with that of autosomal dominant Alzheimer disease. Based on these estimates, a hypothetical distribution of age at death was constructed, assuming fully penetrant Alzheimer disease. These results were compared with real-world mortality data. The authors found that in this meta-analysis, the estimate of age at onset was 53.8 years (95% CI, 53.1-54.5 years; n = 2695); the estimate of age at death, 58.4 years (95% CI, 57.2-59.7 years; n = 324); and the estimate of disease duration, 4.6 years (95% CI, 3.7-5.5 years; n = 226). Coefficients of variation and 95% prediction intervals of age at onset were comparable with those reported in autosomal dominant Alzheimer disease. US mortality data revealed an increase in life expectancy in Down syndrome (median IQR, 1 [0.3-16] years in 1968 to 57 [49-61] years in 2019), but with clear ceiling effects in the highest percentiles of age at death in the last decades (99th percentile: 1990, age 93 years; 2019, age 65 years). The mortality rate matched the limits projected by a distribution assuming fully penetrant Alzheimer disease in up to 80% of deaths (corresponding to the highest percentiles). This contrasts with dementia mentioned in 30% of death certificates but agrees with the mortality data in DABNI (78.9%). Important racial disparities persisted in 2019, being more pronounced in the lower percentiles (10th percentile: Black individuals, 1 year; White individuals, 30 years) than in the higher percentiles (90th percentile: Black individuals, 64 years; White individuals, 66 years). These findings suggest that the mortality data and the consistent age at onset were comparable with fully penetrant Alzheimer disease. Lifespan in persons with Down syndrome will not increase until disease-modifying treatments for Alzheimer disease are available.

Jacobs, J., Schwartz, A., McDougle, C.J., & Skotko, B.G.


Abstract: A small percentage of adolescents and young adults with Down syndrome experience a rapid and unexplained deterioration in cognitive, adaptive, and behavioral functioning. Currently, there is no standardized work-up available to evaluate these patients or treat them. Their decline typically involves intellectual deterioration, a loss of skills of daily living, and prominent behavioral changes. Certain cases follow significant life events such as completion of secondary school with friends who proceed on to college or employment beyond the individual with DS. Others develop this condition years after the completion of secondary school with friends who proceed on to college or employment. In this report presents a young adult male with Down syndrome who experienced severe and unexplained clinical deterioration, highlighting specific challenges in the systematic evaluation and treatment of these patients.


Abstract: Dementia disproportionately affects people with intellectual disability. Most qualitative studies explore their experiences by utilizing proxy-reports. A smaller number of studies indicate the potential of exploring perspectives directly from people with intellectual disability and dementia. This systematic review synthesized findings from existing studies (n = 8) that involve people with intellectual disability and dementia as participants to understand their experiences of dementia. Searches were conducted using CINAHL, PsychInfo and Social Services Abstracts. Findings include descriptions of changes in individual functioning, a narrowing of social worlds and of how people made sense of the changes despite often having no knowledge of their dementia diagnosis. Additionally, discussion focuses on how people’s experiences are shaped by their environments. People’s subjective experiences and views on how dementia affected their lives were more ambiguous compared to contextual data and the descriptive portrayal of how people’s lives had changed. Thus, this review highlights the importance of researchers needing consider how to facilitate conversations with people about their experiences and the ethics involved in conducting qualitative research with people with intellectual disability and dementia, particularly how respond to participants not knowing about their dementia diagnosis. Spending time with participants over a longer period, getting to know how people communicate and the use of visual aids or everyday items were examples of approaches that supported research involvement. The review recognized the complexities of speaking to people with intellectual disability and dementia, challenging views that people with intellectual and dementia cannot be involved in research and makes recommendations to support inclusion in future studies.


Abstract: As clinicians often rely on carer reports to identify adults with intellectual disabilities (ID) with early signs of dementia, this study focused on carer-reported symptoms to ascertain whether carer reports of decline in everyday function would be a more effective screening method to detect possible cases of dementia than reports of memory decline in older adults with ID. Subjects were 154 participants who were reassessed along with their carers two to three years after baseline. A questionnaire for carer-reported change in everyday function and the Dementia Questionnaire for Persons with Mental Retardation (DMR) were used to assess carer views of everyday function and memory. The diagnosis of dementia was confirmed by two psychiatrists working independently. Participants who developed dementia displayed both everyday function and memory decline. Overall, decline in everyday function appeared to be the best indicator of new dementia cases. Retrospective carer report of change in everyday function was as good as, if not better than, prospective ratings to identify dementia; however, in those with mild ID, memory change was a better indicator of dementia, while in those with more severe ID, decline in everyday function was a better indicator. Decline in everyday function (whether prospective change from baseline or reported retrospectively by carers) appears to be a better screening method for dementia than memory decline, particularly for participants with moderate/severe ID.


Abstract: Declining health status and comorbidities are often markers for associated mild cognitive impairment or dementia. A group of community-dwelling adults with intellectual disability diagnosed with dementia (along with controls) have been tracked longitudinally with respect to their health and function. With time, the dementia-capable group home residents with dementia are showing significantly varied health status and comorbidities as well as marked behavioral changes. By tracking the health and function longitudinally, outcome information can pinpoint markers that are associated with premonitory dementia and can help health providers maintain surveillance over select functions and health conditions of those adults already affected. Screening instruments, incorporating these markers, can more precisely be used to identify at-risk adults for ADRD and aid providers design remediation programs earlier.


Abstract: Community housing for adults with intellectual disability (ID) and dementia (AD) is becoming more prevalent. An opportunistic longitudinal study (2011-2018) of group homes (GHs) for such adults examined a number of resident and administrative factors. Study followed 2 cohorts of adults with ID, 15 w/dementia and 15 matched controls (CO), over 7 years (8 time intervals). Initial cohort of 15 AD (Xage=59.1; NDS=5) resided in 3 (5-bed each) dementia GHs. COs resided in general GHs/apartments. Over 7 years, 9 AD died, and 7 replacements (Xage=58.7; NDS=3) were added to the study. Instruments captured subject characteristics, behavior/health/function, staffing, time spent on caregiving, and administrative factors. Findings noted differences in health and function factors between the ADs and COs. Deaths occurred at age-norms. Comparative co-morbidities showed AD residents had significantly more classic cognitive and physical health issues associated with dementia and physical debilitation. An ebb and flow was observed of residents affected by dementia when an agency has multiple dementia GHs, as well as variations in staffing patterns and periods of intensity of care during the day. Over time, the 3 dementia GHs, by inter-home transfers and selective new admissions, have trended toward stage/level specific care settings. Findings can help with planning long-term use of such GHs and can providing dementia capable care for adults with ID in a community-based specialized setting. With dementia affecting an increasing number of adults with ID (due to aging) more attention needs to be given to functional GHs providing in-community long-term mid-stage and advanced dementia capable care.


Abstract: Dementia, as a public health challenge, is a phenomenon vexing many care organizations providing specialized residential and family supports for adults with intellectual disabilities. With increasing survivorship to ages where risk is greatest, expectations are that many more adults in service will present with cognitive decline and diagnosed dementia as they grow older. As persons with dementia present with new needs, there is often a call for a reorientation of services. With respect to residential supports, agencies may need to adapt current methods of care, with particular attention to providing care in small group homes. However, dementia-related care also must be quality care and applicable standards need to be met. The author reviewed relevant policy and practice organizational guidelines and applied research literature addressing components of care and service provision that were critical to quality care and that were consistent with professional practice. Examined were the nuances and contributing factors of quality dementia care and it was proposed that quality of care criteria need to be universally applicable and serve as a framework for adapting extant residential environments and make them ‘dementia-capable’. It is proposed that efforts to evaluate dementia-related care provision with respect to quality need to consider quality of care provision components such as (1) clinically relevant early and periodic assessment; (2) functional modifications in the living setting; (3) constructive staff education and functionality for stage-adapted care; and (4) flexible long-term services provision that recognizes and plans for progression of decline and loss of function.


Abstract: The National Task Group on Intellectual Disabilities and Dementia Practices (NTG), organized in 2011, has been actively involved in stimulating development of services for people with intellectual disabilities (ID) affected by dementia. The NTG has created several sets of practice guidelines, a screening and early detection instrument for use by families and agencies, web-based informational materials, and a national curriculum on ID and dementia, and has undertaken the provision of workforce development workshops across the US on dementia and ID. The NTG works to compliment the activities being undertaken under the National Plan to Address Alzheimer’s Disease and
consults with various national organizations focusing on dementia and lifelong disabilities. The goal of the NTG is to continue to affect change and improve the quality of community dementia care provision corresponding with National Plan updates.

Janicki, M.P.

Abstract: Localities are beginning to feel the impact of the growing number of older adults with lifelong intellectual disabilities (ID) who are also affected by dementia. Many local organizations are attempting to adapt their support and residential services to help this group be served more effectively within the community. Yet, questions have been raised as to the models that may most reasonably be used and how to address the discordance between traditional ID service and “dementia-capable” services. Investigated was how localities and organizations have adapted to the onset of dementia and address early stage care demands and determine practices that are effective in promoting “dementia-capable” care. Several studies were conducted to determine how government entities and local providers are adapting services to identify models prevalent in the provider sector, and specifically to identify staff training needs, physical and environmental adaptations, and differential time spent by staff in providing dementia care. Data showed that most US states are not prepared to address growing onset of dementia in select parts of the ID population, that responses to early stage service needs have been mostly handled by local entities and service organizations; that most have not developed extensive training programs for staff and are experimenting with best practice methods to deliver care - primarily via small group homes - and that dementia care takes up a disproportionate amount of staff time in small care settings. To address early stage dementia related services in the most effective manner, a concerted effort needs to be in place to aid local service entities adapt services to dementia-related presentations among ID clientele, set up coordinated training for staff, secure funds for enhancing group homes into “dementia-capable” care, and construction of clinical support services and augmentation of family support services for parents and other kin carers.

Janicki, M.P.

Abstract: Aging persons with intellectual disability (ID) represent a vulnerable population with respect to cumulative neuropathological conditions, including dementia. Adults with Down syndrome (DS), a subset, have a recognized high risk for Alzheimer’s disease. With dementia present, how to provide post-diagnostic supports is challenging. Dementia care group homes (GHs) along with NPIs are emerging as a model for providing out-of-home community supports. Data from a longitudinal study provide insights on what care organizations need to consider when organizing specialty group home care. The study, begun in 2011, followed three co-located homes providing NPIs to 15 adults with dementia. Findings revealed trajectories of changes over time, housing need/function level patterning, and health status outcomes. Key findings noted 3 age-of-admission clusters (X̄=50.5; SD=57.1; X̄=66.8); overall mortality (X̄=deaths=65.4; ID=69.3; DS=56.3) - half of original entrants died within 7 years; age at entry (X̄=59.1); years from entry to death (X̄=5.4 yrs); LOS (X̄=49.4 months/4.12 yrs); morbidity (number of co-morbidities decreased among survivors). In same period, 8/15 deaths in GHs vs 3/15 deaths in Controls. NPI-related practices included daily program activities (adults in mid- to later stages were engaged in regular off-site day activities that agency provided; adults with advanced dementia remained in homes), staffing patterns differed based on level of care – more staff assigned to homes with residents with advanced dementia, and staffing, and differential time spent by staff in providing dementia care GHs can enable provision of in-community group housing and quality care in accord with stage-defined functional changes and needs if structured in a planful way (factoring in dementia-stage, dementia type, mortality expectations, health status, patterns of care needs, dementia-related behaviors, aging-related issues, and probable trajectories of decline of the residents).


Abstract: The growing numbers of individuals with intellectual disabilities affected by Alzheimer disease and related dementias has raised new challenges for community care providers. This paper examines means of providing community group home-based care in a sample of care providers in five different countries. The aim is to identify trends that have emerged. Two samples of group homes for adults with intellectual disabilities affected by dementia were studied to determine: (1) what are the physical characteristics of the homes; (2) what physical environmental adaptations have been made in response to behavioral deterioration expressed by residents with dementia; and (3) what are the demands on staff resulting from dementia care. The first sample of group homes in five countries provided comparative international data on home designs, staffing, costs, and residents. The second sample, drawn from homes in the U.S. and the U.K., provided data on the impact of dementia. Findings revealed staffing and design of homes varied but generally abided by general practices of dementia care; homes relied on existing resources to manage care, programmatic and environmental adaptations were implemented to address progression of dementia; and residents with dementia presented more demands on staff time with respect to hygiene maintenance and behavior management when compared to other residents not affected by dementia.

Janicki, M. P., Heller, T., Seltzer, G., & Hogg, J.

Abstract: The AAMR/IASSID practice guidelines, developed by an international workgroup, provide guidance for stage-related care management of Alzheimer's disease, and suggestions for the training and education of carers, peers, clinicians, and program staff. The guidelines suggest a three step intervention activity process, that includes: (1) recognizing changes, (2) conducting assessments and evaluations, and (3) instituting medical and care management. They provide guidance for public policies in the United States and the UK, and provide support for people with Alzheimer's disease and intellectual disabilities, and avoidance of institutionalization solely because of a diagnosis of dementia. [This report is available also on www.aamr.org at the following URL: http://161.58.153.187/Bookstore/Downloadables/index.shtml]

Supporting people with dementia in community settings. In M.P. Janicki & A.F. Ansello (Eds.), Community Supports for Aging Adults with Lifelong Disabilities. pp. 387-413


Abstract: Due to the ‘greying’ of the nation’s population, dementia associated with Alzheimer’s disease and other causes, has become another challenge for caregivers of people with Alzheimer’s disease and intellectual disability, and avoidance of institutionalization solely because of a diagnosis of dementia. This book chapter, the authors explore the factors, policies, and support structures that can help agencies provide continued “aging-in-place” dementia-capable care, develop “in-place progression” dementia specific programs, or chose alternative care settings. It also explores some features of dementia-related behaviors that may need to be taken into account in program design and makes suggestions for staff training and planning for dementia programs.


Abstract: The number of age-associated pathologies is increasing, with the
increase in the number of elderly persons. One such age-associated condition, Alzheimer's disease and related dementias, affects a significant number of adults with intellectual disability (ID), in particular those with Down syndrome. Many affected adults live in small community group homes or with their families. How to provide sound and responsive community care is becoming a challenge for agencies faced with an increasing number of affected adults. This study reports the outcome of a survey of group homes serving adults with ID and dementia, explores the onset, duration and effects of dementia and their impact on planning for community care of adults with ID. It also examines emerging community care models that provide for “dementia capable” supports and services. Two models, “aging in place,” and “in place progression” are examined with regard to care practices and critical agency decision making. An approach, the ECEPS model, for responding to dementia is offered.

Janicki, M.P. & Dalton A.J.
Care management, diagnostic and epidemiologic considerations in adults with intellectual disabilities and Alzheimer disease
British Journal of Developmental Disabilities, 1996, 42(Supplement), s84
Abstract: Review of the process and outcome of the Invitational International Colloquium on Alzheimer Disease among Persons with intellectual Disabilities held in Minneapolis, Minnesota (USA) and the subsequent development of a set of international practice guidelines and reports on the assessment, epidemiology, and care management of adults with intellectual disabilities affected by dementia.

Janicki, M.P., & Dalton, A.J.
Dementia in developmental disabilities
In N. Bouras (Ed.), Psychiatric and Behavioral Disorders in Developmental Disabilities and Mental Retardation (1999) pp. 121-153
Cambridge: Cambridge University Press
Abstract: This book chapter provides a brief overview of the current status of knowledge about dementia and its relationship to intellectual disability, touching on current developments in the evaluation of possible comorbid psychiatric, medical and age-associated conditions. The clinical presentation of dementia is examined as well as relevant contemporary issues related to diagnosis, assessment, and care management. Lastly, questions of dementia policy and suggestions for training programs on dementia and intellectual disability are addressed.

Janicki, M.P., & Dalton, A.J.
Dementia and public policy considerations
Philadelphia: Brunner-Mazel
Abstract: This book chapter examines a number of the major public policy considerations related to the aging of adults with intellectual disabilities who evidence change due to dementia. Specifically addressed is the changing structure of at-risk adult populations with intellectual disabilities in service systems, the programmatic and policy issues raised by providers attempting to cope with these changes, needs for further training, education and dissemination of information on aging, and lastly, the challenges and policy imperatives to be confronted with the new millennium.

Janicki, M.P., & Dalton, A.J.
Dementia, aging, and intellectual disabilities: A handbook
488pp.
Abstract: 21 chapter text on dementia issues and intellectual disabilities. Six parts: Introduction, Biomedical considerations, Assessment considerations, Clinical considerations, Program considerations, and Education and policy considerations. Text provides most up-to-date information available about Alzheimer’s disease and related dementias as they affect persons with mental disabilities. Text examines biology and physiology of dementia, neurological and medical complications associated with dementia, best practices to meet the needs of aging persons with intellectual disabilities, policy issues raised by the growing number of older adults with ID, and case studies of affected individuals. Contains glossary of terms, and appendices with AAMRI/ASSID practice guidelines for dementia diagnosis and care management in adults with intellectual disabilities, as well as Newroth & Newroth guidelines for coping with Alzheimer’s disease in persons with Down syndrome.

Janicki, M.P., & Dalton, A.J.
Prevalence of dementia and impact on intellectual disability services
Mental Retardation, 2000, 38, 277-289.
Abstract: A statewide survey, conducted to ascertain the administrative prevalence of dementia in adults with an intellectual disability, found a prevalence of about 3% of the adult service population over the age of 40 years (a rate of 29/1000), 6.1% of the population over the age of 60 years, and 12.1% of the population over the age of 80 years (rates of 68.7/1000 and 121.3/1000, respectively). The rate of dementia was consistent with that for adults in the general population, except for those adults with Down syndrome (who made up a third of the overall group) who had a much higher rate: 22.1% among adults age 40 and older and 56.4% among adults age 60 and older. Onset was observed to occur in the mid-60s (early 50s for Down syndrome). Alzheimer-type dementia was the most frequent diagnosis. Late-onset seizures were reported in about 12% of the cases. With the occurrence of dementia expected to rise proportionately with the increase of longevity among adults with an intellectual disability, it is clear that care systems will have to raise the “index of suspicion” among staff and families, adapt to become “dementia capable,” and improve their diagnostic and technical resources, as well as their community-based care management supports.

Janicki, M.P., & Dalton, A.J.
Alzheimer disease in a select population of older adults with mental retardation
Abstract: Twenty New York State Office of Mental Retardation and Developmental Disabilities district offices were surveyed to determine the prevalence of older adults with mental retardation who also had suspected or diagnosed Alzheimer disease (AD). The survey identified 123 individuals in this category, about 1% of an overall case roll of 10,878 people of 40 years of age and older for the 17 reporting offices. Of these 123 individuals, 64% were people with Down syndrome. Typically, a person with suspected Alzheimer disease was identified through staff observation or report of changes in behaviour leading to suspicion of Alzheimer disease. Information on training needs indicated that a workshop on AD was a primary need. Training in various content areas was indicated, including confirming suspicions, diagnosis, programming, and individual program planning.

Janicki, M.P., McCallion, P., Splaine, M., Santos, F.H., Keller, S.M., & Watchman, K.
Consensus statement of the international summit on intellectual disability and dementia related to nomenclature
Abstract: A working group of the 2016 International Summit on Intellectual Disability and Dementia was charged to examine the terminology used to define and report on dementia in publications related to intellectual disability (ID). A review of related publications showed mixed uses of terms associated with dementia or causative diseases. Like general applications, language related to dementia in ID field often lacked precision and could lead to a misunderstanding of the condition(s) under discussion. Most articles related to ID and dementia reporting clinical or medical research generally provided a definition of dementia or related terms; social care articles tended toward term use without definition. Toward terminology standardization within studies/ reports on dementia and ID, the Summit recommended (a) gaining familiarity with dementia-related diagnostic, condition-specific, and social care terms (as identified in the working group’s report), (b) creating a guidance document on accurately defining and presenting information about individuals or groups referenced, and (c) that in reports on neuropathologies or cognitive decline or impairment, definitions are used and data include subjects’ ages, sex, level of ID, residential situation, basis for dementia diagnosis, presence of Down syndrome (or other risk conditions), years from diagnosis, and if available, scores on objective measures of changing function.

Janicki, M.P., Zendell, A., & DeHaven, K.
and intervention, (3) to develop intervention plans that will support both client appropriate diagnosis, (2) to provide a co-ordinated approach to assessment Dementia Care Pathway in 2005. The aims were: (1) to ensure early and unique challenges to both those responsible for delivering services and carers. Someone with a dual diagnosis of intellectual disability and dementia presents

Abstract: As life expectancy improves, increasing numbers of people with
https://www.choiceforum.org/docs/pdem.pdf
University of Glamorgan (Faculty of Health, Sport, and Science), 2008. 78pp.
Adults with learning disabilities presenting with dementia
Jenkins, R., Davies, R., Sardi, I., Llewellyn, P., Northway, R., O’Connor, C., Trudgeon, C. & Keeling, D.
Adults with learning disabilities presenting with dementia [Final Report - May 2009]
University of Glamorgan (Faculty of Health, Sport, and Science), 2008. 78pp. https://www.choiceforum.org/docs/pdem.pdf
Abstract: As life expectancy improves, increasing numbers of people with intellectual (learning) disabilities are affected by dementia. Supporting someone with a dual diagnosis of intellectual disability and dementia presents unique challenges to both those responsible for delivering services and carers. In response to these challenges Gwent Healthcare NHS Trust developed a Dementia Care Pathway in 2005. The aims were: (1) to ensure early and appropriate diagnosis, (2) to provide a co-ordinated approach to assessment and intervention, (3) to develop intervention plans that will support both client and carers, (4) to provide a process for monitoring the person over time, and (5) to support, carers, clients, and professionals via the provision of information and training. The study demonstrated that the stated aims of the Dementia Care Pathway are being met. Stage one results found that the training provided to paid support staff was received enthusiastically and had a positive impact on knowledge, confidence, and competence levels in relation to working with clients who have, or may develop, dementia. In most cases the knowledge was well maintained at six month follow-up. There is some evidence, however, that although staff who attended the training reported it to be ‘interesting’ and ‘very relevant, but they still doubted its usefulness in practical day-to-day situations when caring for their clients. In order for the potential of the pathway to be further maximized attention should be paid to increasing the level of information given to carers, widening contributions to pathway review meetings, and educating carers on the diverse range of interventions possible through the pathway.

Jervis, N., & Prinsloo, L.
Abstract: Much research has identified an increased prevalence of dementia in adults with Down syndrome when compared with the general population. Neuropathological changes associated with Alzheimer’s dementia in the brain have been found in most people with Down syndrome who die over the age of 35 years. Given the limitations of many assessments for dementia in relation to people with Down syndrome for a single completion, it has been recommended that all people with Down syndrome are assessed at least once in early adulthood in order that they have their own baseline which can be compared with in the future if changes in skills and abilities occur. The authors have had many requests from other services enquiring about their project and how a similar initiative could be set up. Therefore, this article focuses on the way the Manchester Learning Disability Partnership approached screening 135 adults with Down syndrome and details the assessments used, practical considerations, what has been learned and future service implications.

Jethwa, H. & Cassidy, G.
https://doi.org/10.5042/amhid.2010.0675
Abstract: Dementia is a condition that involves inevitably progressive deficits in numerous cognitive domains, including thought, language, memory, understanding and judgement. A difference in behavior may be noted, as well as overall loss of skills. Dementia is more than four times as prevalent in people with intellectual disabilities as in the general population. Diagnosis of the condition in people with intellectual disabilities, however, is often difficult due to lack of baseline skill assessment, high staff turnover in supported accommodation and low expectations of capabilities. Current National Institute for Health and Clinical Excellence (NICE) guidelines on anti-dementia medication state that treatment should not be initiated until the condition has reached moderate severity. Determining whether symptoms are at this stage in people with intellectual disabilities is difficult because their skill level is already impaired. An accurate and extensive record of baseline skill levels in people with intellectual disabilities is therefore crucial, and regular comparison with baseline is key to early diagnosis of dementia.

Johannsen, P., & Mai, J.
Abstract: Dementia is seen earlier and more often in patients with Down syndrome (DS) than in the general population. Patients with DS develop neuropathological changes similar to the changes seen in dementia of Alzheimer’s type (DAT). The literature concerning DAT in DS is reviewed. The clinical course and differential diagnoses are discussed. Before the diagnosis of DAT is made, treatable causes of dementia should be excluded.

Johannsen, P., Christensen, J.E.J., & Mai, J.
The prevalence of dementia in Down syndrome Dementia, 1996, 7(4), 221-225.
Abstract: The authors assess the prevalence of clinical dementia in three age groups of persons with Down syndrome in the county of Aarhus, Denmark. Group 1 was composed of 14-16 year olds (n=13), group 2 was composed of 23-29 year olds (n=34), and group 3 was composed of 50-60 year olds (n=25). Of the 85 subjects, 72 (85%) participated. Carers were interviewed and a neurological examination was performed. An EEG was recorded in 50 of the Ss. Definite clinical dementia was defined as a acquired and progressive decline in 4 or more out of 17 items that are considered to indicate dementia in people with Down syndrome. Possible dementia was considered when 1-3 items were affected. Six adults (24%) in group 3 had definite clinical dementia and 6 adults in group 3 and 2 (6%) in group 2 had possible dementia. Authors note that this was the first Danish population-based study of the prevalence of dementia in people with Down syndrome.

Johansson, P.E., & Terenius, O.
Development of an instrument for early detection of dementia in people with Down syndrome
https://doi.org/10.1080/1366825021000029357
Abstract: The successful detection of early signs of dementia in people with Down syndrome could form a basis for useful early support and for drug treatment. This report describes the development and preliminary application of an interview and test instrument for the assessment of dementia among people with intellectual disability, as well as a framework for diagnosis that combines the findings of an interview and a test with the diagnostic criteria of ICD-10, DSM-IV and NINCDS-ADRDA. From among the number of tests and interview questions developed, those showing the most significant differences between participants in three groups of differing levels of intellectual disability and estimated dementia were kept. Reported are the assumptions for the items used, descriptions of the process and items used, and the associations of test items with predicting the presence of dementia. The authors conclude that a protocol combining testing and interview has promise and potential for detecting early signs of dementia in this population and could prove feasible for use in practice.

Johansson, M., Holst,G., & G Ahlström
Signs in people with intellectual disabilities: Interviews with managers and staff on the identification process of dementia
Journal of Intellectual Disability Research, 2019, 63(8), 649.
Abstract: An increasing number of people with intellectual disability (ID) are reaching older ages and an increased risk of dementia diseases. Staff and managers give support in daily living and can deliver information about residents’ changes in behavior. The aim of this Swedish study was to explore the identification process employed by staff and managers to detect signs of suspected dementia in people with ID within intellectual disability services (ID-services). Twenty managers and 24 staff within ID-service were interviewed and qualitative latent content analysis was applied. A model consisting of three themes on three levels of resources for the identification process of signs of suspected dementia emerged from the analysis. On the first level was the time and continuity in the care relationship, which is crucial for identifying and responding to changes in cognitive ability that indicate dementia. On the second level, the staff identifies deficiencies in their own knowledge, seek support from colleagues and managers within their workplace and, on the third level, outside their workplace. Staff and managers expressed needs for guidance and education from specialists in dementia and primary healthcare. This finding indicates an urgent need for intervention research and digital support for staff in dementia care.

Johnson, N., Fahey, C., Choinque, B., Chong, G., & Giteiman, D.
Effects of donepezil on cognitive functioning in Down syndrome
Abstract: This study determined whether donepezil, an acetylcholinesterase inhibitor, would improve cognitive functioning in 19 subjects with Down syndrome and no dementia. They were assigned to either a donepezil or placebo group. Cognitive functioning and caregiver ratings were measured at baseline, 4 weeks, and 12 weeks. With the exception of one area (language), no improvement was noted in any of the cognitive subtests, behavioral scores, or caregiver ratings. Subjects in the donepezil group showed an improvement in language scores compared to subjects in the placebo group. The results suggest that donepezil may improve language performance in subjects with Down syndrome and no dementia, but further studies need to be done on a larger group to confirm this result.

The middle years and beyond: Transitions and families of adults with Down syndrome
Abstract: Normally expected transitions connect the various periods of life. Often these transitions are prompted by life events that require adaptation to a changed circumstance and may challenge both individual and family quality of life. Such transitions may be planned (proactive) or demand (reactive). Little, however, has been written about the nature of such transitions and how they specifically affect older-aged families of adults with Down syndrome. Such families are often predominate lifelong carers of adults with Down syndrome. Drawing on research and experience, the authors examined three transition points from a family perspective. Each of these points of change requires that people adapt and may lead to various outcomes, including at times outcomes that are unexpected, stressful, and challenging. The three points of transition examined include moving away from the parental home, changes occurring within a residential service (e.g., staff changes, relocations), and the reactions to the onset and course of dementia. Vignettes and quotes illustrate the complexities of these transitions and show that, even with planful management, often such transitions can go awry and produce unpredictable outcomes.

Jokinen, N.
The content of available practice literature in dementia and intellectual disability.
Abstract: Adults with intellectual disability are living to ages seen within the general population and they, too, are at risk of developing dementia. This review was to conduct to identify the nature and content of the literature related to adults with intellectual disability and dementia and bring together guidelines for services and staff providing care. The preponderance of work between 1995 and 2004 focuses on the biomedical, diagnosis and assessment aspects of the disease. Although guidelines exist, there is a lack of published literature on the efficacy of practice strategies to guide the provision of daily care. Future research is discussed that could support continued community living and high quality of life during all stages of the disease.

Jokinen, N., Service, K., Marsack-Topolewski, C., & Janicki, M.P.
Support-staging model for caregivers of adults with intellectual disability affected by dementia
Abstract: Adults with intellectual disability (ID) and dementia are a sub-population of persons who are often un- or underserved. Most adults with ID are integrated within the general community (living autonomously, or in apartments/group residences); but significant numbers also reside with their families, particularly adults with Down syndrome. Family help/counseling approaches, such as the New York University-Caregiver Intervention (NYUCI), might benefit from a support-staging model assessment focus on what specific aid a family requires to meet their needs. Patterns of such needs have been identified that can help with providing dementia-capable care. Objective needs include: (a) information on signs and symptoms, (b) diagnostic advice, (c) understanding behavioral changes and managing dementia-related behaviors, (d) adapting homes, (e) determining daily routines most conducive to calming, (f) planning for the future, (g) finding and navigating resources, and (h) responding to end-of-life needs. Subjective needs include: (a) being informed at time of diagnosis and throughout the course of dementia, (b) coping with a profound sense of loss from knowing the diagnosis, (c) fearing the future [including financial concerns], (d) formulating long-term plans, (e) accessing community-based coordinated care, (f) facing difficulties from the medical community, (g) feeling overwhelmed by caregiving demands, (h) feeling a sense of isolation and abandonment, and (i) facing end-of-life issues. A working group emanating from the 2016 Glasgow Summit on Intellectual Disability and Dementia organized a schema encapsulating these concerns into a support-staging model. The schema suggested four fluid stages: Diagnostic (seeking cause of changes in function, abilities, personality), Explorative (exploring dementia capable interventions), Adaptive (coping with and managing the symptoms/changes), and Closure (resolving / relief from responsibilities).
Using this schema, a process (utilizing the NYUCI) is underway to operationalize a support-staging assessment instrument which would enable counseling staff to frame the state of a family’s concerns, build relationships through this knowledge of the caregiver and provide tailored services to them. The outcome will enable systematic coding and organizing both objective and subjective data so that specific interventions and counseling can be adapted to meet both intermittent and continuous caregiver needs.


Guidelines for structuring community care and supports for people with intellectual disabilities affected by dementia


Abstract: To assist families and organizations in their planning for extended care that accompanies the diagnosis of dementia, the National Task Group on Intellectual Disabilities and Dementia Practices (NTG) in the United States adopted a set of practice guidelines covering the period from when suspicions are aroused to when care ends with eventual death. These guidelines are drawn from the research literature as well as clinical experiences and demonstrated best practices. The guidelines delineate what actions should be undertaken and are presented in a manner that reflects the progressive nature of prevalent dementias. To enable the development of the most appropriate and useful services and care management for adults with intellectual disabilities affected by dementia, the NTG adopted the staging model generally accepted for practice among generic dementia services. The staging model follows the flow from a prediagnosis stage when early recognition of symptoms associated with cognitive decline are recognized through to early, mid, and late stages of dementia, and characterizes the expected changes in behavior and function. In keeping with the National Plan to Address Alzheimer’s Disease recommendations for earlier and more widespread efforts to detect possible symptoms, the guidelines cite the application of the NTG-Early Detection Screen for Dementia as a first step in documenting early signs of cognitive and functional changes among people with intellectual disabilities. The guidelines also provide information on nonpharmacological options for providing community care for persons affected by dementia as well as commentary on abuse, financial, managing choice and liability, medication, and nutritional issues.


Perspectives on family caregiving of people aging with intellectual disability affected by dementia: Commentary from the International Summit on Intellectual Disability and Dementia


Abstract: This article, an output of the 2016 International Summit on Intellectual Disability and Dementia, examines familial caregiving situations within the context of a support-staging model for adults with intellectual disability (ID) affected by dementia. Seven narratives offer context to this support-staging model to interpret situations experienced by caregivers. The multidimensional model has two fundamental aspects: identifying the role and nature of caregiving as either primary (direct) or secondary (supportive); and defining how caregiving is influenced by stage of dementia. We propose staging can affect caregiving via different expressions: (1) the “diagnostic phase,” (2) the “explorative phase,” (3) the “adaptive phase,” and (4) the “closure phase.” The international narratives illustrate direct and indirect caregiving with commonality being extent of caregiver involvement and attention to the needs of an adult with ID. We conclude that the model is the first to empirically formalize the variability of caregiving within families of people with ID that is distinct from other caregiving groups, and that many of these caregivers have idiosyncratic needs. A support-staging model that recognizes the changing roles and demands of carers of people with ID and dementia can be useful in constructing research, defining family-based support services, and setting public policy.

Kählin, I., Kjellberg, A., & Hagberg, J-E.

Ageing in people with intellectual disability as it is understood by group home staff


https://doi.org/10.3109/13668250.2015.1094038

Abstract: The number of older residents in group homes for people with intellectual disability (ID) is increasing. This interview study was focused on how group home staff address issues of ageing and being old among people with ID. Twelve members of staff at 4 different group homes in Sweden were interviewed. Findings revealed old age as something unarticulated in the group home. Group home staff felt unprepared to meet age-related changes in residents. The study also revealed that group home staff had a one-track way of describing the process of ageing among people with ID, which was seemingly rooted in a medical paradigm of disability. Based on this study's findings, we suggest that there is a need to raise issues and give guidance related to ageing and ID in disability policy documents to support the development of a formal culture that addresses old age and ID in disability services.

Kalsy, S., McQuillan, S., Adams, D., Basra, T., Konstantinidi, E., Broquard, M., Peters, S., Lloyd, V., & Oliver, C.

A proactive psychological strategy for determining the presence of dementia in adults with Down syndrome: Preliminary description of service use and evaluation


Abstract: The authors describe and assess the experience of providing proactive screening for dementia in older adults with intellectual disabilities (ID) through a dedicated clinical psychology service within the National Health Service in England. Subjects were the first 18 participants who were referred to the clinical service or were identified as showing early signs of probable dementia in a proactive screening strategy. The screening process involved combining neuropsychological, behavioral, and health data with information from a clinical assessment of the presenting problem in a case series approach. The process of psychological assessment and formulation is illustrated together with an outline of the psychological interventions employed for early-, mid-, and late-stage dementia. An appraisal of the service strategy showed that a dedicated psychology service for dementia assessment can be effective when offering a defined and workable psychological response to the increasing presentation of dementia-associated behaviors among people with ID. Ancillary services included supporting carers in contributing to the assessment and intervention process so as to ensure appropriately responsive and respectful care management for the person with ID and dementia. The authors recommend that a multidimensional stage model of intervention founded on direct performance and informant-based assessments (within a framework of differential diagnosis) be employed in supporting people with ID and dementia.

Kalsy, S., Heath, R., Adams, D., & Oliver, C.

Effects of training on controllability attributions of behavioural excesses and deficits shown by adults with Down syndrome and dementia.


Abstract: Whereas there is a knowledge base on staff attributions of challenging behavior, there has been little research on the effects of training, type of behavior and biological context on staff attributions of controllability in the context of people with intellectual disabilities and dementia. A mixed design was used to investigate the effects of three factors on care staff attributions of challenging behavior. Pre- and post-training measures were administered to participants (n = 97) attending training on ageing, dementia and people with intellectual disabilities. Authors found no significant effects of diagnosis or type of behavior on attributions were found. However, there was a significant increase in knowledge after training (P < 0.001), and training was found to significantly decrease the attribution of controllability (P < 0.001). Conclusion was that the results suggest that training that focuses on aspects of change relevant to behavior can favorably influence care staff's knowledge and attributions of controllability within the context of people with Down syndrome and dementia.

Down syndrome: Distribution of brain amyloid in mild cognitive impairment
Alzheimer’s Dementia: Diagnosis, Assessment & Disease Monitoring, 2020, 12(1), e12013. https://doi.org/10.1002/dad.21203

Abstract: Down syndrome (DS) is associated with a higher risk of dementia. We hypothesize that amyloid beta (Aβ) in specific brain regions differentiates mild cognitive impairment in DS (MCI-DS) and test these hypotheses using cross-sectional and longitudinal data. 18F-AV-45 (florbetapir) positron emission tomography (PET) data were collected to analyze amyloid burden in 58 participants clinically classified as cognitively stable (CS) or MCI-DS and 12 longitudinal CS participants. The study confirmed our hypotheses of increased amyloid in inferior parietal, lateral occipital, and superior frontal regions as the main effects differentiating MCI-DS from the CS groups. The largest annualized amyloid increases in longitudinal CS data were in the rostral middle frontal, superior frontal, superior/middle temporal, and posterior cingulate cortices. Authors note that this study helps us to understand amyloid in the MCI-DS transitional state between cognitively stable aging and frank dementia in DS. The spatial distribution of Aβ may be a reliable indicator of MCI-DS in DS.

Keller, S.M., Janicki, M.P., & Esralew, L.

Dementia: screening, evaluation, diagnosis and management

Abstract: The emergent acknowledgment of the increase rate of dementia among adults with intellectual and developmental disability (IDD) has led to the recognition of a lifespan approach to securing and providing services with relevant supports. Consideration of what is most needed in older age has taken on more prominence. Service organizations are responding to the emergence of age-associated neuropsychopathologies as coincident to lifelong conditions and are attempting to adapt their services for continued community care until death. The historical focus on aging among adults with IDD and the recent focus on dementia has heightened awareness of the latter age needs of adults with IDD - and in particular those at-risk of or already affected by dementia. The positive outcome of these public health initiatives is that more dementia-capable services are being developed, technologies are improving, and there is an increased interest in maintaining quality of life through to the end-of-life, irrespective of the nature and complexity of conditions prevalent in older age.

Kenshole AV, Gallichan D, Pahl S, Clibbens J.


Abstract: Lifestyle has previously been associated with the onset of Alzheimer’s disease (AD) in the typically developing population, but research investigating this association in Down syndrome (DS) is limited. Adults with DS and AD (n = 27) were compared to adults with DS without AD (n = 30) on physical activity, diet, weight, where participants currently lived, where participants had lived for the majority of their lives, educational attainment, occupational attainment and cognitive activity. There was a significant difference between samples on where participants currently lived, with the majority of the clinical sample living in institutionalized settings and the majority of the control sample independently living settings. This may reflect a tendency to move people once they start to deteriorate which, if correct, is contrary to clinical recommendations that people with AD should be supported to “die in place.” Further research into the way in which lifestyle factors, particularly living environment, could contribute to the increased risk of AD in adults with DS is required. This may support interventions aimed at preventing or delaying the onset of the disease.

Kerins, G., Petrovic, K., Bruder, M.B., & Gruman, C.

Medical conditions and medication use in adults with Down syndrome: A descriptive analysis.

Abstract: Authors the presence of medical conditions and medication use within a sample of adults with Down syndrome. The author employed a retrospective chart review using a sample of 141 adults with Down syndrome and age range of 30 to 65 years. They identified 23 categories of commonly occurring medical conditions and 24 categories of medications used by adults with Down syndrome. From their work, the authors concluded that approximately 75% of older adults with Down syndrome in their sample experienced memory loss and dementia. Hypothyroidism, seizures, and skin problems also occurred commonly. The prevalence of cancer (i.e., solid tumors) and hypertension was extremely low. Older adults with Down syndrome used anticonvulsants more often than younger adults with Down syndrome. The use of multivitamins and medications such as pain relievers, prophylactic antibiotics, and topical ointments was common.

Kerr, D.

Down’s syndrome and dementia

Abstract: Text providing a comprehensive review of issues and practices relative to adults with Down syndrome affected by Alzheimer’s disease. Covered are a range of topics related to care management, including assessment of need, communication, creating a therapeutic environment, how to maintain skills, and dealing with challenging behaviors. Also covered are specific interventions and supporting carers.

Kerr, D., Cunningham, C., & Wilkinson, H.

Learning disability and dementia Are we prepared?
Journal of Dementia Care, 2006, May/June, 17-19

Abstract: None provided

Kerr, D., Cunningham, C. & Wilkinson, H.

Responding to the pain experiences of people with a learning difficulty and dementia

Abstract: The report explores knowledge and practice in relation to pain recognition and management amongst direct support staff, members of community learning [intellectual] disability teams and general practitioners. It also examines the understanding and experiences of pain amongst people with a learning difficulty [intellectual disability] and dementia. It identifies the dilemmas and obstacles to effective pain management, and reports on examples of good practice. The report found that the pain experiences and management of people with a learning difficulty [intellectual disability] who have dementia mirrored findings in relation to people in the general population. It did, however, identify extra and compounding issues in relation to people with a learning difficulty [intellectual disability]. The authors proffer recommendations for practitioners and service providers.

Kiddle, H., Drew, N., Crabbe, P., & Wigmore, J.

A pilot memory cafe for people with learning disabilities and memory difficulties

Abstract: Memory cafes have been found to normalize experiences of dementia and provide access to an accepting social network. People with learning disabilities are at increased risk of developing dementia, but the possible benefits of attending a memory cafe are not known. This study evaluates a 12-week pilot memory cafe for people with learning disabilities in terms of adaptations required and benefits of attending. Results indicate that affect levels significantly improved across the course of the cafe and that communication, interaction, alertness and participation in other activities improved outside the cafe. Future plans for attendance at memory cafes for people with learning disabilities are discussed.

Kirk, L.J., Hick, R., & Laraway, A.


Abstract: As life expectancy increases for people with intellectual disabilities, the impact of dementia on people with intellectual disabilities and their families, carers and services is becoming more apparent. Psychological services for intellectual disabilities are receiving an increasing number of referrals requesting dementia assessment. Health and social care services are adapting to the
diverse needs of an ageing population with intellectual disabilities. The authors describe a study investigating the relationship between two assessments for dementia in people with intellectual disabilities. Carers of people with intellectual disabilities over the age of 50 (or 40 if the individual had Down syndrome) completed the Dementia Questionnaire for Mentally Retarded People (DMR) and the Adaptive Behavior Scale—Residential and Community (ABS). Overall, the two questionnaire measures showed significant relationships. However, results suggested that both assessments have clinical value in informing individual needs and aiding diagnosis. The authors discuss the implications for both clinical and social care services.

Kirwan, R., Sheerin, F., McGlinchey, E., McCallion, P., & McCarron, M. Functional loss in older adults with intellectual disabilities and dementia. Learning Disability Practice, 2022, 25(5), ??-?? doi: 10.7748/ldp.2022.e2184 Abstract: Diagnosing dementia in people with intellectual disabilities can be challenging due to pre-existing cognitive impairment. In this population, functional loss could be an early indicator of dementia, but the relationship between functional loss and dementia is not well understood. This means there is a risk of delayed diagnosis and therefore a delay in care planning. To identify and compare the prevalence and age distribution of dementia in people with intellectual disabilities, differentiating between those with Down's syndrome and those with an intellectual disability not attributable to Down's syndrome, and to identify and compare functional loss in people with intellectual disabilities who have developed dementia, differentiating between those with Down's syndrome and those with an intellectual disability not attributable to Down's syndrome. This was a secondary analysis of data from waves 1 and 3 of the Intellectual Disability Supplement to the Irish Longitudinal Study on Ageing (IDS-TILDA), a nationally representative study in Ireland of older adults with intellectual disabilities as they age. Functional loss was determined by participants' level of difficulty with activities of daily living (ADLs) and instrumental activities of daily living (IADLs). The prevalence of dementia was higher among participants with Down's syndrome than among those with an intellectual disability not attributable to Down's syndrome, and the age distribution of dementia differed markedly between the two groups. Both groups experienced similar patterns of difficulty with ADLs and IADLs. IADLs posed more difficulty than ADLs, and among ADLs self-care activities posed the most difficulty. It is important to continue to investigate functional loss in older adults with intellectual disability who have developed dementia, as this should lead to earlier assessment and care planning.

Knox, K., Stanley, J., Hendrix, J.A., Hillerstrom, H., Dunn, T., Achenbach, J., Chicoine, B.A., Lai, F., Lott, I., Stanojevic, S., Howlett, S.E., & Rockwood, K. Development of a symptom menu to facilitate Goal Attainment Scaling in adults with Down syndrome-associated Alzheimer's disease: a qualitative study to identify meaningful symptoms. Journal of Patient Reported Outcomes, 2021 Jan 11, 5(1), 5. doi: 10.1186/s41687-020-00278-7. Abstract: As life expectancy of people with Down syndrome (DS) increases, so does the risk of Alzheimer's disease (AD). Identifying symptoms and tracking disease progression is especially challenging whenever levels of function vary before the onset of dementia. Goal Attainment Scaling (GAS), an individualized patient-reported outcome, can aid in monitoring disease progression and treatment effectiveness in adults with DS. Here, with clinical input, a validated dementia symptom menu was revised to facilitate GAS in adults living with Down Syndrome-associated Alzheimer's disease (DS-AD). Four clinicians with expertise in DS-AD and ten caregivers of adults living with DS-AD participated in semi-structured interviews to review the menu. Each participant reviewed 9-15 goal areas to assess their clarity and comprehensiveness. Responses were systematically and independently coded by two researchers as 'clear', 'modestly', 'remove' or 'new'. Caregivers were encouraged to suggest additional items and recommend changes to clarify symptoms. Median caregiver age was 65 years (range 54-77). Most were female (59%) with 15 years of educational (10%); adults with DS-AD had a median age of 58 years (range 52-61) and either a formal diagnosis (6/10) or clinical suspicion (4/10) of dementia. The initial symptom menu consisted of 67 symptoms each with 2-12 descriptors (589 total). The clinicians' adaptation yielded 58 symptoms each with 4-17 descriptors (580 total). Of these 58 descriptors, caregivers identified 37 (6%) as unclear; these were reworded, and one goal area (4 descriptors) was removed. A further 47 descriptors and one goal area were added to include caregiver-identified concepts. The final menu contained 58 goal areas, each with 7-17 descriptors (623 total). A comprehensive symptom menu for adults living with DS-AD was developed to facilitate GAS. Incorporating expert clinician opinion and input from caregivers of adults with DS-AD identified meaningful items that incorporate patient/caregiver perspectives.

Kobayashi, S., Yamamoto-Mitani, N., Nagata, S., & Murashima, S End-of-life care for older adults with dementia living in group homes in Japan Journal of Patient Reported Outcomes, 2008, 5(1), 31-40. doi: 10.1111/j.1742-7924.2008.00097.x Abstract: The aim of this study was to elucidate the components of end-of-life care provided to older adults with dementia who live in group homes (GHs) in Japan. The number of GHs in Japan is rapidly increasing. Although GHs were originally not established to care for elderly people with advanced-stage dementia, many residents remain in the GH even after their stage of dementia advances; thus, end-of-life care is required. Interviews were conducted with seven GH administrators on their experience in providing end-of-life care to their residents. The constant comparative approach was used for data collection and analysis. Four themes emerged as essential components of end-of-life care in the GH setting: (i) maintaining a familiar lifestyle; (ii) minimizing physical and mental discomfort; (iii) proactively utilizing desirable medical care; and (iv) offering emotional and social support. The combination of the four components seems to be a unique characteristic of end-of-life care in GHs in Japan. These findings may be used to establish a framework for end-of-life care at GHs.

Koehl, L.H., Harp, J., Van Pelt, K.L., Head, E., & Schmitt, F.A. Longitudinal assessment of dementia measures in Down syndrome. Alzheimer’s & Dementia (Amst), 2020, Nov 14, 12(1), e12075. doi: 10.1002/dad2.12075. eCollection 2020. Abstract: Early detection of dementia symptoms is critical in Down syndrome (DS) but complicated by clinical assessment barriers. The current study aimed to characterize cognitive and behavioral impairment using longitudinal trajectories comparing several measures of cognitive and behavioral functioning. Measures included global cognitive status (Severe Impairment Battery [SIB]), motor praxis (Brief Praxis Test [BPT]), and clinical dementia informant ratings (Dementia Questionnaire for People with Learning Disabilities [DLD]). One-year reliability was assessed using a two-way mixed effect, consistency, single measurement intraclass correlation among non-demented participants. Longitudinal assessment of SIB, BPT, and DLD was completed using linear mixed effect models. One-year reliability (n = 52, 21 male) was moderate for DLD (0.69 to 0.75) and good for SIB (0.87) and BPT (0.80). Longitudinal analysis (n = 72) revealed significant by diagnosis interactions for SIB (F(2, 115.02) = 6.06, P = .003), BPT (F(2, 85.59) = 4.56, P = .013), and DLD (F(2, 103.56) = 4.48, P = .014). SIB progression (PR) had a faster decline in performance versus no-dementia (ND) (t(159) = -2.87; P = .013). Dementia had a faster decline in BPT performance versus ND (t(112) = -2.46; P = .014). PR showed quickly progressing scores compared to ND (t(128) = -2.86; P = .014). Current measures demonstrated moderate to good reliability. Longitudinal analysis revealed that SIB, BPT, and DLD changed with age depending on diagnostic progression; no change rates were dependent on baseline cognition, indicating usefulness across a variety of severity levels in DS.

Koenig, B.R. Aged and dementia care issues for people with an intellectual disability: Best practices (vol. 2). 80 pp. Brighton, South Australia: MINDA, Inc. (1995) Abstract: Text covering a range of useful topics related to service provision for dementia among persons with intellectual disabilities. Highly detailed chapters cover health issues, physical decline, behavioral changes, and social aspects. Specific remedial information is provided on communication issues and adapting the environment. A chapter also addresses counseling strategies, examining a diverse range of approaches.

Severe enough to meet criteria for a diagnosis of dementia. Consensus criteria even prodromal stages of dementia-causing diseases). A condition called mild dementia, a general term referring to a variety of diseases and conditions causing substantial loss of cognitive ability and functional declines; adults with Down syndrome are at especially high risk. A great deal of recent research on dementia care, as well as those living in a variety of supported living and group settings; and including persons with ID and dementia and their families in research on dementia care.

Kozma, C. Down syndrome and dementia. Topics in Geriatric Rehabilitation, 2008, 24(1), 41–53. doi: 10.1097/01.TGR.0000311405.01555.3b

Abstract: Down syndrome (DS) is one of the most common genetic conditions with an estimated incidence of 1 in 750 in the general population. It results from an extra chromosome 21 with the total chromosome count being 47 instead of the normal 46. The classic features of DS include hypotonia, atypical facial characteristics, an increased incidence of major and minor anomalies, vision and hearing deficits, other health problems, and intellectual disabilities. People with DS are living longer and experiencing premature aging, specifically Alzheimer disease (AD). The incidence of AD among adults with DS varies significantly according to studies averaging between 11% to 22% for people aged 40 to 49 years, 24.3% for people aged 50 to 59 years, and 25.6% to 77% for people older than 60 years. All studies indicate an early onset of AD as well as an exponential increase in prevalence with age. Furthermore, senile plaques and neurofibrillary tangles, the neuropathological characteristics of AD, are seen in the brain of all people with DS. Annual screening for AD should become part of routine medical practice of older adults with DS, because an early diagnosis is important for comprehensive care.

Kozma, C. Down syndrome and dementia. Topics in Geriatric Rehabilitation, 2008, 24(1), 41–53. doi: 10.1097/01.TGR.0000311405.01555.3b

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Lai, F., Mhatre, P.G., Yang, Y., Wang, M.C., Schupf, N., & Rosas, H.D.


Abstract: Adults with Down syndrome (DS) older than 40 have Alzheimer’s disease (AD) neuropathology and high risk for dementia, but little is known about the relationship of sex to AD risk in this population. Using nonparametric methods and Cox proportional hazards models we analyzed differences in incidence of dementia, by sex, presence of an apolipoprotein E (APOE) ?4 or ?2 allele, and dementia duration and decline in 246 adults over 40 with DS. There was no significant sex difference in risk of AD or rate of cognitive decline. APOE ?4 allele significantly increased risk of AD irrespective of sex. No significant interactions were found between sex and APOE status on AD risk. Among those who died, dementia duration was significantly longer in women. This study showed no effect of sex on interactions between sex and APOE for risk of AD in adults with DS; however, women had longer dementia duration.

Lane, A.M., Reed, M.B., & Hawranik, P.


Abstract: Older adults with Down syndrome (DS) and dementia are an emerging sub-population. With much longer life spans than decades ago, issues have arisen as to where these aging adults will live, how nurses and other staff in facilities can provide effective care to these individuals. This article presents a research study that examined the learnings of nurses and staff members working within a western Canadian program for older adults with DS and dementia. These learnings include: the importance of learning from each other; importance of collaboration; how individuals with developmental disabilities communicate; older adults with DS and dementia differ from older adults with dementia; and residents’ impact on staff.

Lautarescu, B.A., Holland, A.J., & Zaman, S.H.


Abstract: Adults with Down syndrome (DS) are at a very high risk of developing early onset Alzheimer’s disease (AD) due to trisomy of chromosome 21. AD is preceded by a prolonged prodromal “pre-clinical” phase presenting with clinical features that do not fulfill the diagnostic criteria for AD. It is important to clinically characterize this prodromal stage to help early detection of the disease as neuropsychopathology of AD is almost universal by the fifth decade in DS. There is a lack of knowledge of the trajectory of decline associated with the onset of dementia in this population and early signs may be overlooked or misdiagnosed, negatively affecting the quality of life of those affected and the use of early pharmacological or psychosocial interventions. The objective of this systematic review was to evaluate the published literature on longitudinal data in order to identify the cognitive and behavioral changes occurring during the prodromal and early stages of AD in this population. Fifteen peer-reviewed articles met the inclusion criteria, including a total number of 831 participants, with a duration between baseline and follow up varying from 1 year to 47 years. Results suggest that, compared to the general population for which short-term (episodic) memory loss is the most common indicator associated with the onset of AD, in people with DS, executive dysfunction and Behavioural and Psychological Symptoms of Dementia (BPSD) are commonly observed during pre-clinical and early stages and may precede memory loss. The review highlights the importance of using a broad spectrum of assessments in the context of heterogeneity of symptoms. Theoretical and practical implications are discussed, as well as the need for further research.

Lazenby, T.


Abstract: Many people with Down’s syndrome (DS) experience eating, drinking, and swallowing (EDS) difficulties, which can potentially lead to life-threatening conditions such as malnutrition, dehydration, and aspiration pneumonia. As the life expectancy of people with DS continues to improve, there is an increasing need to examine how the aging process may further affect these conditions. Published research studies have yet to address this issue; therefore, this article draws on the literature in three associated areas in order to consider the dysphagic problems that might develop in aging people with DS. The areas examined are EDS development in children and adolescents with DS, EDS changes associated with aging, and EDS changes associated with dementia of the Alzheimer’s type (DAT) because this condition is prevalent in older adults with DS. This article concludes that unlike in the general population, the aging process is likely to cause dysphagic difficulties in people with DS as they get older. Therefore, it is suggested that longitudinal studies are needed to examine the specific aspects of EDS function that may be affected by aging and concomitant conditions in DS.


Abstract: Adults with Down syndrome (DS) developmental disorder (AD) pathology by their 5th decade. Compared with the general population, traditional vascular risks in adults with DS are rare, allowing examination of the cerebrovascular disease in this population and insight into its role in AD without the confound of vascular risk factors. We examined in vivo magnetic resonance imaging (MRI)-based biomarkers of cerebrovascular pathology in adults with DS, and determined their cross-sectional relationship with age, beta-amyloid pathology, and mild cognitive impairment or clinical AD diagnostic status. Participants from the Biomarkers of Alzheimer’s Disease in Down Syndrome study (n = 138, 50 ± 7 years, 39% women) with MRI data and a subset (n = 90) with amyloid positron emission tomography (PET) were included. We derived MRI-based biomarkers of cerebrovascular pathology, including white matter hyperintensities (WMH), infarcts, cerebral microbleeds, and enlarged perivascular spaces (PVS), as well as PET-based biomarkers of amyloid burden. Participants were characterized as cognitively healthy (CS), mild cognitive impairment-DS (MCI-DS), possible AD dementia, or definite AD dementia based on in-depth assessments of cognition, function, and health status. There were detectable WMH, enlarged PVS, infarcts, and microbleeds as early as the 5th decade of life. There was a monotonic increase in WMH volume, enlarged PVS, and presence of infarcts across diagnostic groups (CS < MCI-DS < possible AD dementia < definite AD dementia). Higher amyloid burden was associated with a higher likelihood of an infarct. The findings highlight the prevalence of cerebrovascular disease in adults with DS and add to a growing body of evidence that implicates cerebrovascular disease as a core feature of AD and not simply a comorbidity.

Lao, P., Zimmerman, M. E., Hartley, S. L., Gutierrez, J., Keator, D., Igwe, K. C., ... & Brickman, A. M.

Obstructive sleep apnea, cerebrovascular disease, and amyloid in older adults with Down syndrome across the Alzheimer’s continuum. Sleep Advances, 2022 May 5, 3(1):zpac013. doi:
10.1093.sleepadvances/zpac013.

Abstract: We determined the extent to which obstructive sleep apnea (OSA) is associated with increased cerebrovascular disease and amyloid burden, and the relation of the two processes across clinical Alzheimer’s disease (AD) diagnostic groups in adults with Down syndrome (DS). Adults with DS from the Biomarkers of Alzheimer’s Disease in Down Syndrome (ADDS) study were included given available research MRI (n = 116; 50 ± 8 years; 42% women) and amyloid PET scans (n = 71; 50 ± 7 years; 39% women) at the time of analysis. Participants were characterized as cognitively stable (CS; 64%), with mild cognitive impairment-DS (MCI-DS; 23%), with possible AD dementia (5%), or with definite AD dementia (8%). OSA was determined via medical records and interviews. Models tested the effect of OSA on MRI-derived cerebrovascular biomarkers and PET-derived amyloid burden, and the moderating effect of OSA and AD diagnosis on biomarkers. OSA was reported in 39% of participants, which did not differ by clinical AD diagnostic group. OSA was not associated with cerebrovascular biomarkers but was associated with greater cortical amyloid burden. White matter hyperintensity (WMH) volume (primarily in the parietal lobe), enlarged perivascular spaces, and cortical and striatal amyloid burden were greater across clinical AD diagnostic groups (CS<MCI-DS<possible AD<definite AD). OSA increased the differences in WMH volumes across clinical AD diagnostic groups, primarily in the frontal and temporal lobes. Adults with DS and OSA had greater amyloid burden and greater cerebrovascular disease with AD. Importantly, OSA may be a modifiable risk factor that can be targeted for intervention in this population at risk for AD.

Lutarescu, B.A., Holland, A.J., & Zaman, S.H.
The early presentation of dementia in people with Down syndrome: a systematic review of longitudinal studies

Abstract: Adults with Down syndrome (DS) are at a very high risk of developing early onset Alzheimer’s disease (AD) due to trisomy of chromosome 21. AD is preceded by a prolonged prodromal “pre-clinical” phase presenting with clinical features that do not fulfil the diagnostic criteria for AD. It is important to clinically characterise this prodromal stage to help early detection of the disease as neuropathology of AD is almost universal by the fifth decade in DS. There is a lack of knowledge of the trajectory of decline associated with the onset of dementia in this population and early signs may be overlooked or misdiagnosed, negatively affecting the quality of life of those affected and the use of early pharmacological or psychosocial interventions. The objective of this systematic review is to evaluate the published literature on longitudinal data in order to identify the cognitive and behavioural changes occurring during the prodromal and early stages of AD in this population. Fifteen peer-reviewed articles met the inclusion criteria, including a total number of 831 participants, with the duration between baseline and follow up varying from 1 year to 47 years. Results suggest that, compared to the general population for which short-term (episodic) memory loss is the most common indicator associated with the onset of AD, in people with DS, executive dysfunction and Behavioural and Psychological Symptoms of Dementia (BPSD) are commonly observed during pre-clinical and early stages and may precede memory loss. The review highlights the importance of using a broad spectrum of assessments in the context of heterogeneity of symptoms. Theoretical and practical implications are discussed, as well as the need for further research.

Lessov-Schlaggar, C.N., Del Rosario, O.L., Morris, J.C., Ances, B.M., Schlaggar, B.L., & Constantino, J.N.
Adaptation of the Clinical Dementia Rating Scale for adults with Down syndrome

Abstract: Adults with Down syndrome (DS) are at increased risk for Alzheimer disease dementia, and there is a pressing need for the development of assessment instruments that differentiate chronic cognitive impairment, acute neuropsychiatric symptomatology, and dementia in this population of patients. We adapted a widely used instrument, the Clinical Dementia Rating (CDR) Scale, which is a component of the Uniform Data Set used by all federally funded Alzheimer Disease Centers for use in adults with DS, and tested the instrument among 34 DS patients recruited from the community. The participants were assessed using two versions of the modified CDR-a caregiver questionnaire and an in-person interview involving both the caregiver and the DS adult. Assessment also included the Dementia Scale for Down Syndrome (DSDS) and the Raven’s Progressive Matrices to estimate IQ. Both modified questionnaire and interview instruments captured a range of cognitive impairments, a majority of which were found to be chronic when accounting for premorbid function. Two individuals in the sample were strongly suspected to have early dementia, both of whom had elevated scores on the modified CDR instruments. Among individuals rated as having no dementia based on the DSDS, about half showed subthreshold impairments on the modified CDR instruments; there was substantial agreement between caregiver questionnaire screening and in-person interview of caregivers and DS adults. The modified questionnaire and interview instruments capture a range of impairment in DS adults, including subthreshold symptomatology, and the instruments provide complementary information relevant to the ascertainment of dementia in DS. Decline was seen across all cognitive domains and was generally positively related to age and negatively related to IQ. Most importantly, adjusting instrument scores for chronic, premorbid impairment drastically shifted the distribution toward lower (no impairment) scores.

Lethin, C., Rahm Halberg, I., Karlsson, S., & Janlòv, A-C.
Family caregivers' experiences of formal care when caring for persons with dementia through the process of the disease

Abstract: Family caregivers' experiences of formal care when caring for persons with dementia through the process of the disease and interviews. Four focus group interviews were conducted in October 2011 with 23 spouses and adult children of persons with dementia and analyzed with content analysis. The participants' experiences of formal care when caring for a person with dementia were captured in the theme 'Family caregiving requires collaboration with formal care to get support adjusted to needs specific to the stages of dementia'. This can be broken down into the categories ‘The dementia diagnosis - entry into formal care as a novice family caregiver’, ‘Need for expanded collaboration with formal care to continue care at home’ and ‘Being dependent on a nursing home and trying to maintain involvement’. Family caregiving requires collaboration with formal care to get support adjusted to the individual's needs, specific to the stages of dementia. Caregivers experience a transition process with three main turning points: the dementia diagnosis; when they realize increased need for formal care to continue caring at home; and when the person with dementia is moved into a nursing home. The interviewed caregivers experience formal care reactive to their needs and this often promoted unhealthy transitions. Formal care needs to be proactive and deliver available care and support early on in the dementia trajectory. Interventions should focus on facilitating a healthy transition for family caregivers through the trajectory of the dementia disease to ensure their well-being.

Li, R. S. Y., Kwok, H.W.M., Deb, S., Chui, E.M.C., Chan, L.K., & Leung, D.P.K.
Validation of the Chinese version of the dementia screening questionnaire for individuals with intellectual disabilities (DSQIID-CV).

Abstract: An increasing number of people with intellectual disabilities (ID) are at risk of developing age-related disorders such as dementia because of a dramatic increase in life expectancy in this population in the recent years. There is no validated dementia screening instrument for Chinese people with ID. The Dementia Screening Questionnaire for Individuals with Intellectual Disabilities (DSQIID) was reported to be a valid, user-friendly, easy-to-use observer-rated instrument. It was developed in the UK and has good psychometric properties. Validation of a Chinese version of the DSQIID will facilitate its application among the Chinese population. The DSQIID was translated into the Chinese version (DSQIID-CV). By purposive sampling, service users with ID aged 40 years or over were recruited through two large centres serving adults with ID in Hong Kong. Carers who had taken care of the participants continuously for the past 6 months were invited to complete the DSQIID-CV. All participants were examined by qualified psychiatrists to determine the presence or absence of dementia. Two hundred people with ID whose age ranged between 40 and 73

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Abstract: Few studies have investigated in detail which factors influence the needs of people with learning disabilities who develop dementia. Among adults with an intellectual disability (ID), dementia conditions can only explain 3% of the ADL score variation (ß=-9.245, p<0.001) whereas dementia conditions (ß=-15.773, p<0.001), comorbid condition (ß=-4.853, p<0.05) and syndrome (ß=-9.290, p<0.05), severe or profound disability level (ß=-6.725, p<0.05; primary: ß=5.544, p<0.05; junior high or more: ß=8.147, p<0.01), Down’s syndrome resulted in increased emotional exhaustion for carers. The results indicated that 16.5% of the adults with an intellectual disability might have dementia conditions (DSQIID score ≥ 20).

Lin, J-D., Lin, L-P., Hsia, Y-C., Hsai, S-W., Wu, C-L., & Chu, C.M.
A national survey of caregivers’ perspective of early symptoms of dementia among adults with an intellectual disability in Taiwan. A national survey was conducted to recruit 460 community residents of at least 45 years of age with an intellectual disability. The Dementia Screening Questionnaire for Individuals with Intellectual Disabilities (DSQIID) was administered to caregivers to determine the symptoms of dementia in adults with an intellectual disability. The results indicated that 16.5% of the adults with an intellectual disability might have dementia conditions (DSQIID score ≥ 20).

After controlling for other factors in a multiple logistic regression analysis, the older adults with intellectual disability (755 vs. 45–54, OR = 2.594, 95% CI = 1.438–4.679) and those with a comorbid diagnosis of mental illness or neurological disease (with vs. without, OR = 2.926, 95% CI = 1.593–5.012) had a higher risk of dementia than their counterparts. This study suggests that further longitudinal research is needed to investigate the specific aspects of the functions of living and morbidity that might be affected by aging and concomitant conditions in adults with an intellectual disability.

Lin, L.P., Hsu, S.W., Hsia, Y.C., Wu, C.L., Chu, C., & Lin, J.D.

Abstract: Few studies have investigated in detail which factors influence activities of daily living (ADL) in adults with the intellectual disabilities (ID) comorbid with dementia conditions. The objective of the present study was to describe the relation between early onset dementia conditions and progressive loss of ADL capabilities and to examine the influence of dementia conditions and other possible factors toward ADL scores in adults with ID. This study was part of the "Healthy Aging Initiatives for Persons with an Intellectual Disability in Taiwan: A Social Ecological Approach" project. We analyzed data from 459 adults aged 45 years or older with an ID regarding their early onset symptoms of dementia and their ADL profile based on the perspective of the primary caregivers. Results show that a significant negative correlation was found between dementia score and ADL score in a Pearson’s correlation test (r=−0.28, p<0.001). The multiple linear regression model reported that factors of male gender (ß=4.187, p<0.05), marital status (ß=4.79, p<0.05), education level (primary: ß=5.544, p<0.05; junior high or more: ß=8.147, p<0.01), Down’s syndrome (ß=9.290, p<0.05), severe or profound disability level (ß=6.725, p<0.05; primary: ß=15.773, p<0.001), comorbid condition (ß=4.853, p<0.05) and dementia conditions (ß=9.245, p<0.001) were variables that were able to significantly predict the ADL score (R²=0.241) after controlling for age. Disability level and comorbidity can explain 10% of the ADL score variation, whereas dementia conditions can only explain 3% of the ADL score variation in the study. The present study highlights that future studies should scrutinize in detail the reasons for the low explanatory power of dementia for ADL, particularly in examining the appropriateness of the measurement scales for dementia and ADL in aging adults with ID.

Llewellyn, P.
The needs of people with learning disabilities who develop dementia: A literature review


Abstract: People with learning disabilities are living longer and are increasingly developing age related conditions including dementia. If this occurs, their medical and social needs pose many challenges for services. A literature review was undertaken of articles published between 1996—2006. Data was collected relating to the needs of people with learning disabilities and dementia, their carers and their peers. The primary medical need is for timely and accurate diagnosis. There is a multitude of diagnostic tools and advice is available as to which are most suitable for different client groups. The needs of carers are intertwined with those of people with learning disabilities and dementia and meeting their needs for education, training and increased staff numbers, has proved beneficial. Although multiple services will be responsible for the needs of this client group, there is a consensus that learning disability services should be at the heart of service provision.

Lloyd, V., Kalsy, S., & Gatherer, A.
The subjective experience of individuals with Down syndrome living with dementia

Dementia, 2007, 6(1), 63-88.

Abstract: An increasing number of studies have begun to explore the subjective experience of individuals with dementia. However, despite the increased prevalence of dementia in individuals with Down syndrome, no such published research has been undertaken within this population. The aim of this study was to explore the perspectives and subjective experiences of six individuals with Down syndrome and dementia. Semi-structured interview accounts were analyzed using Interpretative Phenomenological Analysis, in order to gain a level of understanding concerning the impact of dementia upon respondents’ lives and sense of self. Five main themes emerged: (1) Self-image, (2) The Relational Self, (3) Making Sense of Decline,(4) Coping Strategies and (5) Emotional Experience. Whilst the process of adjusting to dementia appeared comparable to the general population, the content of this was influenced by multiple levels of context specific to having a concomitant intellectual disability.

Lloyd, V., Kalsy, S., & Gatherer, A.
Impact of dementia upon residential care for individuals with Down syndrome


Abstract: Despite the increased prevalence of dementia in individuals with Down syndrome, relatively little is known about its impact upon care provision. Carers may be familiar with the demands of assisting a person with Down syndrome, but generally have little knowledge about the course or impact of dementia. This dissonance may lead to stress, which can have a detrimental effect on the carer and the quality of care for the recipient. In this exploratory study, the authors examined the objective and subjective impact of dementia upon paraprofessional paid carers of individuals with Down syndrome working in residential settings. The study used the Caregiver Activities Scale—Intellectual Disabilities (CAS-ID), the Caregiver Difficulties Scale—Intellectual Disabilities (CDS-ID), and the Maslach Burnout Inventory (MBI). Responses given for these measures by paraprofessional carers of individuals with Down syndrome and dementia (n = 9) were compared with responses from those caring for recipients with Down syndrome and no additional cognitive decline (n = 11). No significant differences were found in the responses from these sets of carers on measures of objective (CAS-ID) or subjective burden (CDS-ID). However, the MBI revealed that carers of individuals with Down syndrome and dementia reported significantly increased levels of emotional exhaustion. Findings suggested that, while even when there is little difference in the level of caregiving tasks or the subjective difficulties of caregiving, the onset of dementia in individuals with Down syndrome resulted in increased emotional exhaustion for carers. Additional factors not considered within this study, such as challenging behavior, may also be pertinent to carer burden.

Kognition bei Down-Syndrom: Entwicklung über die Lebensspanne und neuropsychologische Diagnostik im Erwachsenenalter [Cognition in Down’s Syndrome: Development across the Life Span and Neuropsychological Assessment in Adults].

Abstract: Down’s syndrome is the most frequent genetic cause of intellectual disability. As the risk for developing Alzheimer’s disease is increased in Down’s syndrome, comprehensive cognitive examination is essential, both in young adults (for baseline evaluation), as well as later for diagnosing dementia. So far, there are only a few recommendations for neuropsychological assessment in Down’s syndrome. Here, we review (1) the development of cognition across the life span, (2) various causes of cognitive change in adults with Down’s syndrome, and (3) procedures available for their evaluation. Furthermore, (4) we provide recommendations for the assessment and interpretation of diagnostic findings in adults with intellectual disabilities. We conclude with recommendations for cognitive assessment in intellectual disability in general.

Lott, J.D.
The rate of decline of social skills across dementing and non-dementing individuals with intellectual disabilities: A longitudinal study.


The author sought to establish rate of decline of adaptive skills in a population of individuals with intellectual disability (ID) with dementia compared to similar adults with ID and without dementia, as well as examining the variability of positive and negative social behaviors across diagnostic classes. The participants were matched for age, sex, Down syndrome, and level of ID. The control group was screened for the presence of dementia with the Early Signs of Dementia Checklist. Rate of decline within groups was assessed by the Vineland Adaptive Behavior Scales and changes in positive and negative behaviors were measured by the Matson Evaluation of Social Skills for the Severely Retarded. (MESSIER) Prior to a diagnosis of dementia groups were equivalent. No significant differences were found for adaptive behaviors. Visual analysis of plotted means supports predicted decline in skills for both groups. Significant differences were found across time for positive social skills. Significant correlations were observed between the VABS and the MESSIER Positive domains. The findings provide support for the diagnostic utility of the MESSIER with dementia. However, no support was observed for different variances of negative behaviors across diagnostic groups. This would suggest that the measure of negative behaviors is not supported as a diagnostic tool at this time.

Lott, I.T., & Head, E.
Dementia in Down’s syndrome: unique insights for Alzheimer disease research


Abstract: Virtually all adults with Down syndrome (DS) show the neuropathological changes of Alzheimer disease (AD) by the age of 40 years. This association is partially due to overexpression of amyloid precursor protein, encoded by APP, as a result of the location of this gene on chromosome 21. Amyloid-ß accumulates in the brain across the lifespan of people with DS, which provides a unique opportunity to understand the temporal progression of AD and the epigenetic factors that contribute to the age of dementia onset. This age dependency in the development of AD in DS can inform research into the age of dementia onset. (2) various causes of cognitive change in adults with Down’s syndrome, and (3) procedures available for their evaluation. Furthermore, (4) we provide recommendations for the assessment and interpretation of diagnostic findings in adults with intellectual disabilities. We conclude with recommendations for cognitive assessment in intellectual disability in general.

Lott, I.T., & Lai, F.
Dementia in Down’s syndrome: observations from a neurology clinic.


Abstract: Clinical manifestations of dementia were reviewed in 15 Down’s syndrome (DS) patients referred to a neurological clinic over a 24-month period for mental deterioration. The ages ranged from 32-64 years. One hundred percent showed personality changes and loss of independent daily living skills, the presenting symptoms in two-thirds of the cases. Other manifestations included seizures (53%), gait deterioration (73%), sphincter incontinence (40%), and pathological release reflexes (67%). All 7 patients with CT-scans showed moderate or severe central and peripheral cortical atrophy. Detailed clinical information is presented for two patients, one of whom showed a temporary remission with imipramine. A characteristic dementia syndrome appears to be present in a subgroup of people with dementia.

Lynngard, H., & Alexander, N.
"Why are my friends changing?" Explaining dementia to people with learning disabilities


Abstract: Many publications seek to explain the causes and effects of dementia to the general population and there is evidence of the benefit of supporting carers and of establishing support groups. However, there is much less published material aimed at people with intellectual disabilities, and little focus on the specific needs of people who share their homes and live with other people with learning disabilities who develop dementia. This article, based on group work, describes residents who had expressed bewilderment at the gradual changes they were witnessing in two of their housemates with dementia with whom they had shared a home and friendships over many years. Employing a wide range of visual aids, equipment, role plays and exercises, we sought to make the explanation of dementia as accessible and concrete as possible. The group also provided a forum for the residents to talk about the effects of living with others who develop dementia. Evaluation showed how a relatively short intervention can result in positive changes for both the people with learning disabilities who develop dementia and their peers.

Lyons, V., Oliver, E., Knifton, & Molesworth, S.
Role of Admiral Nurses in supporting people with learning disabilities and dementia


Abstract: The average age of people with learning disabilities is increasing, meaning that the number of people with learning disabilities and dementia is also rising. The care trajectory for people with learning disabilities and dementia is complex, starting with challenges in obtaining an appropriate diagnosis through to receiving appropriate and high-quality end of life care. The charity Dementia UK recognises the issues that families experience when someone in their family has a learning disability and dementia, and has developed a model of care in which Admiral Nurses, who are specialist dementia nurses, work in learning disability services. This article explores the role of the Admiral Nurse in learning disability services and examines the areas in which these specialist nurses provide tailored support. The article also outlines the expected outcomes of the service provided by these nurses.
MacDonald, S., & Summers, S.J.  
Psychosocial interventions for people with intellectual disabilities and dementia: A systematic review  
*Journal of Applied Research in Intellectual Disabilities*, 27 February 2020  
https://doi.org/10.1111/jar.12722  
Abstract: People with intellectual disability experience a higher prevalence of dementia, at an earlier age, than the general population. The aim of this review was to establish the psychological interventions and outcomes for individuals with intellectual disability and dementia. A search of eight electronic databases and reference lists of all included articles was conducted using PRISMA guidelines. Data were synthesized using an integrative method. Initial searching produced 2,351 papers. Twenty-one studies met the inclusion criteria. Interventions were deductively categorized into behavioural, systemic and therapeutic. All studies reported positive findings for individuals and for the systems which support them, but limited by methodological issues and neglect of the direct experience and impact on individuals themselves. The lack of a synthesis of psychosocial interventions within clinical practice, and the associated evidence, has invariably led to a lack of knowledge in practice. This is clearly evidenced by the omission of psychosocial interventions within established intellectual disability and dementia care pathways. The findings are discussed in relation to the wider literature and evidence base. Future research should aim to adopt methodologically robust designs that are inclusive of the individual experience of people with intellectual disability. The authors also posit that this review will provide essential knowledge for enacting policies that all individuals diagnosed with dementia and their carers have access to meaningful post-diagnostic care, including social and psychological care and support.  

Mahon, A., Tilley, E., Randhawa, G., Pappas, Y., & Vseteckova, J.  
Ageing carers and intellectual disability: a scoping review  
*Quality in Ageing and Older Adults*, 2019, 20(4), 162-178.  
https://doi.org/10.1108/QOAQ-11-2018-0057  
Abstract: Individuals with intellectual disabilities (i.e.) are living longer contributing to an overall increase in the average age of caregivers. The purpose of this paper is to review the literature on the physical, social and psychological needs of ageing carers of individuals with intellectual disability(ies) in the UK. A scoping review framework was used to identify literature from eleven databases, the grey literature and the references lists of relevant studies. Only primary research studies that discussed the needs of non-professional carers, aged 65+ years old, of individuals with intellectual disability(ies) in the UK were included. No date restrictions were applied. Thematic analysis was used to narratively synthesize findings. Six studies were included. Five key themes were identified: Living with fear, lack of information, rebuilding trust, proactive professional involvement and being ignored. Housing and support information is not communicated well to carers. Professionals require more training on carer needs and trust must be rebuilt between carers and professionals. Proactive approaches would help identify carer needs, reduce marginalization, help carers feel heard and reduce the risk of care crisis. Greater recognition of mutual caring relationships is needed.  

Manji, S.W.L.U.  
Aging with dementia and an intellectual disability: A case study of supported empowerment in a community living home  
*Dissertation Abstracts International Section A: Humanities and Social Sciences*, 2009, 70(1-A), 352.  
Abstract: Case study explored the qualitative experience of 4 adults with intellectual disability (ID) and dementia residing in a specializing dementia support group home. Participant observation, daily living log notes, and interviews with family/friend carers, direct-care staff, and administrators were used to obtain data. The three study questions were: (i) how the onset of dementia in people with ID changes their needs, what adjustments have to be made in the support practices, and what service barriers and successes are experienced; (ii) how adults with ID and dementia experience living in a home specializing in dementia support and how stakeholders perceive this model of support; and (iii) what are the ways policymakers can better respond to the changing needs of people with ID and dementia. Two social processes were identified: ‘marginalization’ and ‘supported empowerment’. Marginalization depicted how dementia affected adults with ID as they incurred multiple losses in ability, home, and community. Despite losses, the adults maintained their ‘selfhood’ with good health support, decision-making, self-agency, and autonomy as the home provided an individualized transition process, consistent and person-centered support, and elevated empathy to facilitate freedom of choice. Supported empowerment was found as an empowering social model with micro-practices that harnessed elements of empowerment necessary to support people with dual disabilities. Seven policy considerations that prevent premature placement in nursing homes, enable aging in place, and maintain a participatory life in community were recommended.  

Mann, D.M.  
Alzheimer’s disease and Down’s syndrome.  
Abstract: The neuropathological features of Down's syndrome at middle age is compared with that of Alzheimer’s disease at that age, through a review of the published literature and from the author's personal observations. The pathological changes of Down’s syndrome at middle age, i.e. the form and distribution of senile plaques and neurofibrillary tangles, and the pattern of involvement (atrophy) of neuronal systems are qualitatively the same as those of Alzheimer's disease at that age. Quantitative differences do occur and these may relate to biological or sociological variations inherent to the two parent populations. It is concluded that, in pathological terms, patients with Down's syndrome at middle age do indeed have Alzheimer's disease. Some ways in which a study of patients with Down's syndrome can give insight into the nature and development of the pathological changes of Alzheimer’s disease are put forward and discussed  

Effect of service structure and organization on staff care practices in small community homes for people with intellectual disabilities  
Abstract: An important question in community living is what factors influence the extent to which staff provide ‘active support’. Engagement, care practices and a range of staff and organizational characteristics were studied in 72 residential homes serving 359 adults with intellectual disabilities. Managers in 38 settings were trained in person-centred active support (PCAS). A group comparison design and multivariate analysis was used to investigate the relationship between variables. The PCAS group showed more active support, assistance, other contact from staff and engagement in meaningful activity but no difference in choice-making or assessment of participation in activities of daily living. The PCAS group had more staff with a professional qualification, were more likely to think that challenging behaviour was caused by lack of stimulation, had attitudes more in line with a policy of community care, rated most care tasks as less difficult, and were more organized to deliver active support. The comparison group were more likely to think that challenging behaviour was learned negative behaviour, showed more teamwork and were more satisfied. Multivariate analysis identified a range of staff and organizational variables associated with engagement and active support. The results suggest that some variables which have not hitherto been studied in relation with active support are associated with it. Professional qualification, knowledge and experience appear to be important as do some staff attitudes, clear management guidance, more frequent supervision and team meetings, training and support for staff to help residents engage in meaningful activity.  

Fifteen-year follow-up of 92 hospitalized adults with Down’s syndrome: incidence of cognitive decline, its relationship to age and neuropathology  
Abstract: The clinical and neuropathological features associated with dementia in Down’s syndrome (DS) are not well established. To examine clinico-pathological correlations and the incidence of cognitive decline in a cohort of adults with DS. A total of 92 hospitalized persons with DS were followed up from 1985 to December 2000. At outset, 87 participants were dementia-free, with a median age of 38 years. Assessments included the Prudhoe Cognitive Function Test (PCFT) and the Adaptive Behavior Scale (ABS), to measure cognitive and behavioral deterioration. Dementia was diagnosed from case records and caregivers’ reports. Eighteen (21%) patients developed dementia during follow-up, with a median age of onset 55.5 years (range 45-74). The PCFT demonstrated cognitive decline among those with a
less severe intellectual disability (mild and moderate) but not among the profoundly disabled people (severe and profound). Clinical dementia was associated with neuroanatomical features of Alzheimer's disease, and correlated with neocortical neurofibrillary tangle densities. At the age of 60 years and above, a little more than 50% of patients still alive had clinical evidence of dementia. Authors concluded that clinical dementia associated with measurable cognitive and functional decline is frequent in people with DS after middle age, and can be readily diagnosed among less severely intellectually disabled persons using measures of cognitive function such as the PCFT and behavioral scales such as the ABS. In the more profoundly disabled people, the diagnosis of dementia is facilitated by the use of behavioral and neurological criteria. In this study, the largest prospective DS series including neuropsychology on the original patients, the density of neurofibrillary tangles related more closely to the dementia of DS than senile plaques. In people with DS surviving to middle and old age, the development of dementia of Alzheimer type is frequent but not inevitable, and some people with DS reach old age without clinical features of dementia.


Abstract: The aim of this pilot project was to evaluate the usefulness of establishing a specially designed “Memory Clinic” for the assessment and diagnosis of dementia in people with learning disabilities. This pilot memory clinic was set up by re-organization of existing resources, especially in terms of professional time. The core team members were a psychiatrist, psychologist and a community nurse from the specialist learning disability team with a special interest in dementia and its treatment. Over a period of 8 months a total of 12 assessments were carried out. Seven females and 4 males were assessed, the age range being 41 years – 83 years; one being a repeat assessment. In 3 individuals a definitive diagnosis of dementia was made. Six of these service users also had associated Down’s syndrome. In the Down’s syndrome group, 2 had a definite diagnosis of dementia and for the diagnosis was inconclusive at the initial assessment. Authors note that the professionals involved in the assessment process need to be specially trained in dealing with the issues related to identifying and interpreting the significance of changes in function, behavior, and personality in adults with learning disabilities.


Abstract: This booklet for community carers and agency staff covers some of the fundamentals concerning adults with Down syndrome and Alzheimer’s disease, including information on obtaining diagnoses, approaches to care management, and securing services in the UK. Contains some vignettes and a small glossary and references.


Abstract: Autism spectrum disorder and intellectual disability are associated with psychiatric comorbidities, yet a 2009 study of US child and adolescent psychiatry program directors indicated that psychiatry residents receive insufficient training in autism spectrum disorder/intellectual disability. This follow-up study surveyed child and adolescent psychiatry and general psychiatry program directors to assess (1) the current extent of residency training in autism spectrum disorder/intellectual disability, (2) program director perceptions of educational topics and resident competency in autism spectrum disorder/ intellectual disability, and (3) preferred resources to strengthen autism spectrum disorder/intellectual disability training. As in 2009, many child and adolescent psychiatry program directors reported few lecture hours, although current child and adolescent psychiatry residents saw slightly more patients with autism spectrum disorder but not intellectual disability. General psychiatry program directors reported fewer lecture hours in autism spectrum disorder/intellectual disability and fewer patients with autism spectrum disorder than child and adolescent psychiatry program directors. Both child and adolescent psychiatry and general psychiatry program directors recognized the importance of a range of educational topics pertinent to autism spectrum disorder/intellectual disability. Child and adolescent psychiatry program directors reported higher resident competency, and lecture hours and patients seen moderately correlated with resident competency. Program directors indicated that online videos and other resources would help improve autism spectrum disorder/intellectual disability training in their programs. Collectively, these findings suggest minimal improvements in autism spectrum disorder/intellectual disability training over the past decade and highlight the urgent need to advance psychiatry training in this field through dissemination of resources.


Abstract: Although individuals with intellectual/ developmental disabilities (I/DD) are living longer than in the past, they also are exposed to age-related changes in health and well-being. They are prone to acquire dementia that often manifests earlier and more frequently than in the general population. However, there is sparse knowledge on the daily challenges that affect the quality of life of the individuals with I/DD and comorbid dementia and their family caregivers. This pilot study examined strengths and challenges of individuals with dual diagnoses of I/DD and dementia using the family quality of life (FQOL) framework. Cross-sectional data was gathered from a convenience sample of family caregivers using a web-based electronic survey. This study aims to identify the common and differential elements of the FQL domains among formal and informal services and social supports elucidate a need to empower families with high caregiving needs through research, practice and policy. Providers should be cognizant of the needs of individuals with I/DD and dementia comorbid, as well as the needs of their family caregivers.


Abstract: Neuropsychiatric symptoms (NPS) are significant manifestations of dementia, with important consequences for patients and caregivers. Despite the established genetic link between Down syndrome (DS) and Alzheimer’s disease (AD), studies investigating NPS in individuals with DS and dementia are scarce. The Neuropsychiatric Inventory (NPI) was developed to identify symptoms of dementia and while it is widely used for the assessment of individuals with dementia in the general population, we have found no studies using the NPI in those with DS. The aim of this study is to characterize NPS in a sample with DS with heterogeneous cognitive profiles using the NPI. Participants (N=92) with DS, =30 years of age were assessed with the Cambridge Examination for Mental Disorders of Older People with Down’s Syndrome and Others with Intellectual Disabilities (CAMDEX-DS). They were classified by a psychiatrist into three categories: AD, prodromal dementia, and stable cognition. Another
psychiatrist blinded to the CAMDEX-DS dementia diagnosis, evaluated the participants using the NPI. Chi-square tests were used to check for significant differences in frequency of symptoms. Thirteen participants (14.1%) had AD, 17 (18.4%) were classified as prodromal dementia and 62 (67.5%) were in the stable cognition group. Prevalence of delusion, depression, anxiety, disinhibition, irritability, appetite abnormalities and total NPI did not differ between groups. Anxiety and irritability were common across all groups (~50% of the total), while euphoria was not present in any participant. Hallucination, agitation, apathy, aberrant motor behaviour and night-time behaviour disturbance showed significant difference among the groups (p<0.05), with higher prevalence in the group with AD. Authors note that the results indicate that individuals with DS and AD have some of the symptoms that are characteristic for dementia, with AD being found to be a predictor of AD in comparison to stable cognition (OR for one-point increase in the NPI = 1.342, p = 0.012). Agitation, apathy, nighttime behavior disturbances, and total NPI did not differ between groups. Prevalence of delusion, depression, anxiety, disinhibition, and depression (R2 = 0.627, F(15,76) = 8.510, p < 0.001). NPS observed among people with AD in the general population or may follow a specific NPS pattern. The high frequency of some NPS in individuals with DS and stable cognition should be considered in the diagnostic process in order to reduce the odds of generating a false positive.

Neuropsychiatric symptoms of Alzheimer's disease in Down Syndrome and Its impact on caregiver distress
Abstract: Neuropsychiatric symptoms (NPS) are non-cognitive manifestations common to dementia and other medical conditions, with important consequences for the patient, caregivers, and society. Studies investigating NPS in individuals with Down syndrome (DS) and dementia are scarce. Characterize NPS and caregiver distress among adults with DS using the Neuropsychiatric Inventory (NPI). We evaluated 92 individuals with DS (>30 years of age), divided by clinical diagnosis: stable cognition, prodromal dementia, and AD. Diagnosis was determined by a psychiatrist using the Cambridge Examination for Mental Disorders of Older People with Down's Syndrome and Others with Intellectual Disabilities (CAMDEX-DS). NPS and caregiver distress were evaluated by an independent psychiatrist using the NPI, and participants underwent a neuropsychological assessment with Cambridge Cognitive Examination (CAMCOG-DS). Symptom severity differed between-groups for delusion, agitation, apathy, aberrant motor behavior, nighttime behavior disturbance, and total NPI scores, with NPS total score being found to be a predictor of AD in comparison to stable cognition (OR for one-point increase in the NPI = 1.342, p = 0.012). Agitation, apathy, nighttime behavior disturbances, and total NPI were associated with CAMCOG-DS, and 62% of caregivers of individuals with AD reported severe distress related to NPS. Caregiver distress was most impacted by symptoms of apathy followed by nighttime behavior, appetite/eating abnormalities, anxiety, irritability, disinhibition, and depression (R2 = 0.627, F(15,76) = 8.510, p < 0.001). NPS are frequent and severe in individuals with DS and AD, contributing to caregiver distress. NPS in DS must be considered of critical relevance demanding management and treatment. Further studies are warranted to understand the biological underpinnings of such symptoms.

Amenstic and non-amenstic symptoms of dementia: An international study of Alzheimer's disease in people with Down's syndrome
Abstract: The presence of age-related neuropathology characteristic of Alzheimer's disease (AD) in people with Down syndrome (DS) is well-established. However, the early symptoms of dementia may be atypical and appear related to dysfunction of prefrontal circuitry. The authors sought to characterize the initial informant reported age-related neuropsychiatric symptoms of dementia in people with DS, and their relationship to AD and frontal lobe function. Non-amenstic informant reported symptoms (disinhibition, apathy, and executive dysfunction) and amenstic symptoms from the CAMDEX-DS informant interview were analyzed in a cross-sectional cohort of 162 participants with DS over 30 years of age, divided into three groups: stable cognition, prodromal dementia, and AD. To investigate age-related symptoms prior to evidence of prodromal dementia we stratified the stable cognition group by age. Amenstic and non-amenstic symptoms were present before evidence of informant-reported cognitive decline. In those who received the diagnosis of AD, symptoms tended to be more marked. Memory impairments were more marked in the prodromal dementia than the stable cognition group (OR = 35.07; P < .001), as was executive dysfunction (OR = 7.16; P < .001). Disinhibition was greater in the AD than in the prodromal dementia group (OR = 3.54; P < .04).
Apathy was more pronounced in the AD than in the stable cognition group (OR = 34.18; P < .001). Premorbid amenstic and non-amenstic symptoms as reported by the informant had a stronger progression to AD. Apathy and agitation as symptoms of AD in this DS progression of symptoms needs to be taken into account.
An understanding of the unique clinical presentation of DS in AD should inform treatment options.

Mascarenhas Fonseca, L., Rufino Navatta, A.C., Bottino, C.M.C., & Miotto, E.C.
Cognitive rehabilitation of dementia in adults with Down syndrome: A review of non-pharmacological interventions
Published online 2015 Sep 18. doi: 10.1159/000438858
Abstract: There is a close genetic relationship between Alzheimer's disease (AD) and Down syndrome (DS). AD being the most severe mental disorder affecting aging individuals with DS. The objective of the present study was to evaluate the efficacy of cognitive rehabilitation interventions in DS patients with AD by means of a critical literature review. Because AD is progressive and irreversible, treatment is aimed at delaying and reducing the cognitive and functional decline in order to preserve or improve quality of life. The effects that pharmacological treatments and cognitive interventions have on elderly individuals with AD are well documented. Recent clinical trials have investigated the use of pharmacological treatment in DS patients with AD, generating preliminary results that have been unfavorable. There is a clear lack of studies addressing the efficacy of cognitive rehabilitation interventions in DS patients with AD, and there is an urgent need for studies providing evidence to inform decisions regarding the appropriate choice of treatment strategies.

Mattheys, K., Boustead, I., Doyle, A., & Watchman, K.
`Life through a lens`: understanding the impact of dementia, a participatory action project led by people with intellectual disability
Journal of Intellectual Disability Research, 2019, 63(8), 650.
Abstract: Co-researchers with an intellectual disability are part of a team looking at the effects of non-drug interventions with people who have dementia, including people with Down syndrome. Photovoice is a method of data collection and analysis combining photography with social action supporting the inclusion of people typically excluded from research. Three co-researchers with intellectual disability working on the `Life through a Lens` research project attended training in photovoice methodology and use of the camera, followed by a series of practice exercises. Each engaged in participant observation to understand the impact of non-drug interventions on peers with an intellectual disability and dementia. Photographs were then taken that represented their feelings about the intervention followed by a group discussion with wider research team. Two of the co-researchers believed that their peers benefited from the non-drug interventions. For example, after observing changes to the home environment of one participant, the co-researcher discussed the relaxing and calming effect this created and how it helped her to be safer at home; his photography reflected the security he observed. Such photography can help with understanding of the unique clinical presentation of DS in AD should inform treatment options.
family and guardian considerations, early Alzheimer's disease, mid-stage Alzheimer's disease, feeding and nutrition concerns, and understanding difficult behaviors. Appendix contains a "Level of Capacity Scale," and table outlining implications and treatment suggestions for persons with intellectual disabilities affected by dementia.

Margallo-Lana, M. L., Moore, P. B., Kay, D. W., Perry, R. H., Reid, B. E., Berney, T. P. & Tyrer, S. P.
Abstract: The clinical and neuropathological features associated with dementia in Down's syndrome (DS) are not well established. Aims To examine clinico-pathological correlations and the incidence of cognitive decline in a cohort of adults with DS. A total of 92 hospitalized persons with DS were followed up from 1985 to December 2000. At outset, 87 participants were dementia-free, with a median age of 38 years. Assessments included the Prudhoe Cognitive Function Test (PCFT) and the Adaptive Behavior Scale (ABS), to measure cognitive and behavioural deterioration. Dementia was diagnosed from case records and caregivers' reports. Eighteen (21%) patients developed dementia during follow-up, with a median age of onset 55.5 years (range 45-74). The PCFT demonstrated cognitive decline among those with a less severe intellectual disability (mild and moderate) but not among the profoundly disabled people (severe and profound). Clinical dementia was associated with neuropathological features of Alzheimer's disease, and correlated with neocortical neurofibrillary tangle densities. At the age of 60 years and above, a little more than 50% of patients still alive had clinical evidence of dementia. Clinical dementia associated with measurable cognitive and functional decline is frequent in people with DS after middle age, and can be readily diagnosed among less severely intellectually disabled persons using measures of cognitive function such as the PCFT and behavioural scales such as the ABS. In the more profoundly disabled people, the diagnosis of dementia is facilitated by the use of behavioural and neurological criteria. In this study, the largest prospective DS series including neuropathology on deceased patients, the density of neurofibrillary tangles related more closely to the dementia of DS than senile plaques. In people with DS surviving to middle and old age, the development of dementia of Alzheimer type is frequent but not inevitable, and some people with DS reach old age without clinical features of dementia.

McBrien, J., Whitwham, S., Olverman, K., & Masters, S.
Abstract: Given the now well-recognized risk of Alzheimer's Disease (AD) for adults with Down's Syndrome (DS) as they reach middle age, services for people with learning disability (LD) need to meet this new challenge. Good practice guidance from the Foundation for People with Learning Disabilities recommended that every service for people with learning disability should set up a register of adults with DS, conduct a baseline assessment of cognitive and adaptive functioning before the age of 30 years, develop specialist skills in this area, offer training to other professionals, front-line staff and carers, and seek high-quality co-ordination between agencies. This article reports the progress of one LD service in meeting these challenges, highlighting the successes and difficulties that may guide other teams considering such a development.

McCallion, P.
Maintaining communication
In M. P. Janicki & A. J. Dalton (Eds.), Dementia, Aging, and Intellectual Disabilities pp. 261-277
Abstract: This book chapter is based on the premise that progression of dementia among persons with intellectual disabilities appears to be similar to that in the general population. Therefore, it explores how existing service models and programs may be adapted for the population with intellectual disabilities. A five part program, Maintaining Communication and Independence (MCI), is proposed which adapts an existing program for persons with dementia to better meet the needs of persons with intellectual disabilities. The five parts to MCI are: (1) strengths identification and deficit assessment, (2) environ-mental modification, (3) good communication, (4) memory aids, and (5) taking care of the carer.

McCallion, P., Janicki, M. P.
Intellectual disabilities and dementia (Computer-based Course) 2 CD-Rom set

McCallion, P., McCarron, M., & Force, LT
Abstract: It has been suggested in the literature on family caregiving for persons with Alzheimer's dementia (AD) that levels of objective and subjective burden among carers often predict institutionalization of the persons with AD. There is a paucity of measures to assess whether perceived burden among formal caregivers may also predict movement to more restrictive settings for persons with intellectual disabilities (ID) and AD. This study focused upon the development of a measure of subjective burden, the Caregiving Difficulty Scale – Intellectual Disability (CDS-ID) as a first step in addressing this measurement deficit. An existing caregiver subjective burden scale, the Caregiving Hassles Scale (CHS) was adapted for use with 203 staff caregivers of persons with ID and AD. Preliminary testing of existing CHS items and proposed new items was carried out in two countries, Ireland and the USA. Confirmatory factor analysis with the existing items and exploratory factor analysis with existing and proposed new items for the scale was used to establish the content and test the psychometric properties of a revised scale, the CDS-ID. On the existing CHS items, staff carers appeared to experience greater subjective burden than has been reported for family caregivers. However, the psychometric properties of the CHS found with this population were poor. Factor analysis produced a revised scale, the CDS-ID with three subscales with Cronbach alphas ranging from 0.75 to 0.93 and 38 items overall. This new scale when used with objective burden and other scales offers an opportunity to more systematically measure the difficulties staff experience when caring for persons with ID who present with symptoms of AD.

McCallion, P., Nickle, T., & McCarron, M.
Abstract: There has been increasing concern about the impact of dementia symptoms on the lives and on the care being provided for persons with intellectual disability (ID) in out-of-home settings. One such setting that has received little attention is foster family care homes. These settings in the USA replicate family living and while some supports and resources are provided, they are not designed to meet intensive care needs. As a preliminary step in understanding family experiences and to expand the range of interest in Alzheimer’s disease (AD) in persons with ID beyond traditional out-of-home settings, a pilot study was initiated that included aging persons with ID and symptoms of AD who were living in foster family care settings in two regions of New York State as well as more traditional out-of-home care subjects. Comparisons of matched samples on subjective and objective burden measures suggest that there are few differences in experiences. The limitations of these findings are considered and recommendations made for future, related research.

McCarron, M.
Abstract: Virtually all individuals with Down's syndrome over the age of 35 years have neurological changes characteristic of Alzheimer’s disease. It has become increasingly recognized that people with Down's syndrome and
dementia have very special needs, and those who care for them require specialist knowledge and skills. This paper aims to explore some important issues in caring for persons with this dual disability. It commences with a brief outline on the prevalence of dementia in this population. Diagnostic issues and the clinical presentation of dementia in persons with Down's syndrome are reviewed. In an attempt to help staff respond to the opportunities and challenges they encounter, issues discussed, include: promoting well-being, developing a shared vision on which to build practice, mealtimes – a therapeutic event, reality orientation and validation therapy, communication, activity and entertainment.

McCarron, M., Gill, M., Lawlor, B., & Begley, C.
Time spent caregiving for persons with the dual disability of Down's syndrome and Alzheimer's dementia: Preliminary findings
Abstract: Persons with Down's syndrome (DS) are at increased risk of Alzheimer's type dementia (AD) compared with the general population. Little attention has been paid to the current and future impact of AD on caregivers and clients in residential and community settings. This study sought to test if the Caregiver Activity Survey-Intellectual Disability (CAS-ID) would be useful in measuring time spent by professional caregivers aiding persons with DS and AD. Preliminary findings suggest that caregiving time increases significantly when a person with DS experiences symptoms of dementia. No significant differences were reported in time spent caregiving for subjects at mid-stage versus end-stage dementia; however, the nature and tasks of caregiving change as dementia progresses. This study supports the utility of the CAS-ID in measuring time spent caregiving for persons with DS and AD. Care providers must plan appropriate models of health and social care to effectively address these needs.

McCarron, M., Gill, M., Lawlor, B., & Beagly, C.
A pilot study of the reliability and validity of the Caregiver Activity Survey – Intellectual Disability (CAS-ID)
Abstract: Authors undertook to amend the Caregiver Activity Survey (Davis et al., 1997) and apply it for use with caregivers of persons with intellectual disabilities. Under this study, the CAS-ID was tested with 30 adults and convergent validity was assessed by comparing the CAS-ID with other measures of cognitive and functional impairment of adults with intellectual disabilities. Final version of the CAS-ID contains 8 items: dressing, bathing/showering, grooming, toileting, eating and drinking, housekeeping, nursing care-related activities, and supervision/behavior management. Authors content that the CAS-ID has the potential for identifying and measuring care and resource requirements for people experiencing decline associated with dementia.

McCarron, M., Gill, M., McCallion, P., & Begley, C.
Health co-morbidities in ageing persons with Down syndrome and Alzheimer's dementia.
Abstract: Consideration of the relationship between physical and mental health co-morbidities in ageing persons with Down syndrome (DS) and Alzheimer’s dementia (AD) is of clinical importance both from a care and resource perspective. To investigate and measure health co-morbidities in ageing persons with Down syndrome with and without AD. Recorded physical and mental health needs were ascertained for 124 persons with DS >35 years through a systematic and detailed search of individual medical and nursing case records. Differences in persons with and without AD were investigated, by stage of dementia and by level of intellectual disability (ID). A summed score for health co-morbidities was created and compared using t-tests. Persons with AD had significantly higher co-morbidity scores than persons without AD (t = -8.992, d.f. = 121, P<0.0001). There was also a significant difference in summed co-morbidity scores for persons at end-stage vs. persons at midstage AD (t = -6.429, d.f. = 56, P < 0.0001). No differences were found by level of ID. Increasing health co-morbidities in persons with DS and AD have important implications for care and resources. Appropriate environmental supports combined with competent skilled staff are crucial and will have an important impact on the quality of life for this increasingly at risk population.

McCarron, M., & Lawlor, B.A.
Responding to the challenge of ageing and dementia in intellectual disability in Ireland
Aging and Mental Health, 2003, 7(6), 413-417. doi:10.1080/13607860310001594655
Abstract: The intellectual disability (ID) population in Ireland is ageing and the number of older persons with the dual disability of ID and dementia is increasing. In spite of these demographic trends, as in other countries adequate policy and service provision for this population are lacking. This paper draws upon data available on the population with ID and dementia, reviews both genetic and ID specific literature, considers the policy context and argues for a specific model of service provision. A service model is proposed for the development of multidisciplinary specialist teams within ID, delivered through mobile regional ID dementia clinics.

McCarron, M., Gill, M., McCallion, P., & Begley, C.
Alzheimer's dementia in persons with Down's syndrome: predicting time spent on day-to-day caregiving.
Abstract: The aim of this study was to investigate the amount of time formal caregivers spend addressing activities of day-to-day care activities for persons with Down's syndrome (DS) with and without Alzheimer's dementia (AD). Caregivers completed for 63 persons with DS and AD, and 61 persons with DS without AD, the Caregiving Activity Survey-Intellectual Disability (CAS-ID). Data was also gathered on co-morbid conditions. Regression analysis was used to understand predictors of increased time spent on day-to-day caregiving. Significant differences were found in average time spent in day-to-day caregiving for persons with and without AD. Mid-stage and end-stage AD, and co-morbid conditions were all found to predict increased time spent caregiving. Nature and tasks of day-to-day caregiving appeared to change as AD progressed. The study concluded that staff time to address day-to-day caregiving needs appeared to increase with onset of AD and did so most dramatically for persons with moderate intellectual disability. Equally, while the tasks for staff were different, time demands in caring for persons at both mid-and end-stage AD appeared similar.

McCarron, M., McCallion, P., Fahey-McCarthy, E., Connaire, K., & Dunn-Lane, J.
Supporting persons with Down syndrome and advanced dementia: Challenges and care concerns
Abstract: To understand staff perceptions of critical issues in caring for persons with intellectual disability (ID) and advanced dementia. There has been growing interest in addressing resource, training, and service redesign issues including an increase in collaborative practices in response to the growing incidence of dementia among persons with ID. Most recently this has included consideration of the specific issues in advanced dementia. Thirteen focus group interviews were held involving staff in six ID services and one specialist palliative care provider in Ireland. A qualitative descriptive approach was taken to analysis. Staff identified three key themes: (1) readiness to respond to end of life needs, (2) the fear of swallowing difficulties, and (3) environmental concerns and ageing in place. Four underlying issues that emerged in this study offer clues to solutions: (a) differences in staff preparation associated with settings, (b) lack of understanding and lack of collaboration with palliative care services, © uncertainties about the ability to transfer existing palliative care models to persons with ID and dementia and (d) the need to develop training on end stage dementia and related care approaches.

McCarron, M., McCallion, P., Fahey-McCarthy, E., & Connaire, K.
Staff perceptions of essential prerequisites underpinning end-of-life care for persons with intellectual disability and advanced dementia
Abstract To better address palliative care and end-of-life issues for persons with intellectual disability (ID) and dementia, work was undertaken to understand the perspectives of agency staff in both the ID services and specialist palliative care fields. A qualitative descriptive design composed of 13 focus group interviews involved 50 participants drawn from six ID service providers and seven
McCallon, M., McCallion, P., Reilly, E., & Mulryan, N.

The role and timing of palliative care in supporting persons with intellectual disability and advanced dementia.


Abstract: To better describe the role and timing of palliative care in supporting persons with intellectual disabilities and advanced dementia (AD). Specialist palliative care providers have focused mostly on people with cancers. Working with persons with intellectual disabilities and AD offers opportunities to expand such palliative care to other populations and disease conditions and to better understand the timing and role of palliative care delivery. Thirteen focus group interviews and 12 in-depth interviews involved staff in six intellectual disability services and one specialist palliative care provider in Ireland. A qualitative descriptive approach was taken to analysis. Specialist palliative care staff recognized that person-centered care delivered in intellectual disability services was consistent with palliative approaches, but staff in intellectual disability services did not consider advanced dementia care as ‘palliative’. Both groups were unsure about the role of palliative care at early stage of dementia but appreciated specialist palliative care contributions in addressing pain and symptom management challenges. Successful extension of palliative care principles, philosophy and services to persons with intellectual disabilities and AD will require in-depth understanding of prevailing care philosophies and agreement regarding timing and the unique contributions of specialist palliative care services.

McCallon, M., McCann, P., McCallion, P., Reilly, E., & Mulryan, N.

Responding to the challenges of service developmental to address dementia needs for people with an intellectual disability and caregivers


Abstract: Book chapter addressing the association between intellectual disability and dementia: the experiences of dementia in people with intellectual disabilities; and service planning. This book chapter pertains to the latter. It acknowledges the challenges faced by traditional intellectual disability services and explains the approach adopted by one such service in ROI to supporting and accommodating people with an intellectual disability who develop dementia. It exemplifies the move to an in place progression model in traditional intellectual disability services, whereby services are adapted so that people with an intellectual disability can be supported and accommodated as they progress through dementia.


A prospective 20-year longitudinal follow-up of dementia in persons with Down syndrome.


Abstract: Authors sought to examine dementia characteristics, age at onset and associated co-morbidities in persons with Down syndrome. A total of 77 people with Down syndrome aged 35 years and older were followed up from 1996 to 2015. The diagnosis of dementia was established using the modified ICD 10 Criteria and a combination of objective and informant-based tests. Cognitive tests included the Test for Severe Impairment and the Down Syndrome Mental Status Examination; adaptive behaviour was measured using the Daily Living Skills Questionnaire, and data from the Dementia Questionnaire for People with Intellectual Disabilities have been available since 2005. Over the 20-year period, 97.4% (75 of 77) persons developed dementia with a mean age of dementia diagnosis of 55 years (SD = 7.1, median = 56 years). Clinical dementia was associated with cognitive and function decline and seizure activity. Risk for dementia increased from 23% in those aged 50 years to 80% in those aged 65 years and above. There were no differences by level of ID. The previously reported high risk levels for dementia among people with Down syndrome were confirmed in this data as was the relationship with late onset epilepsy. The value of the instruments utilized in tracking decline and helping to confirm diagnosis is further highlighted.

McCaron, M., & Riley, E.

Supporting persons with intellectual disability and dementia: Quality dementia care standards - A guide to practice

39 pp.

Dublin, Ireland: Trinity College Dublin (2010)

Source: http://www.docservice.ie/includes/documents/Dementia%20Publication%202011.pdf

Abstract: Document contains a series of six standards covering a range of areas concerned with care affecting adults with intellectual disabilities affected by dementia. Drawn from standards affecting the general population, this document groups together focal areas under six main categories reflecting person-centered dementia care. The standards consist of statements, indicators, and criteria for assessing evidence. The standards cover (1) appropriately trained staff and service development, (2) memory assessment services, (3) health and personal care, (4) communication and behavior, (5) promoting well-being and social connectedness, and (6) supporting persons with advanced dementia.

McCaron, M., Reilly, E., & Dunne, P.

Achieving quality environments for person centred dementia care

45 pp.

Dublin, Ireland: Daughters of Charity Service

Abstract: Provides an overview of principles and practices designed to enable the operation of small group homes, including covering the planning process, design of private and public spaces, as well as therapeutic uses. Illustrated by two Daughters of Charity homes established for dementia specific care for people with ID. One home offers care for people with moderate dementia and includes 4 permanent beds and 2 respite beds for people both living with their families in the community and community group homes. Home also has a 6 bed step-down palliative care unit for people with ID in the later stages of dementia. These purpose built facilities were designed to be responsive to the changing needs of persons across the continuum of dementia. The home-like environments support people with dementia and staff to participate and complete tasks together, as well as informal impromptu unplanned activities. The homes are designed so that each resident has his or her own bedroom, with numerous communal areas including sitting rooms and garden areas.

MacDonald, S., & Summers, S.J.

Psychosocial interventions for people with intellectual disabilities and dementia:

A systematic review.


Abstract: People with intellectual disability experience a higher prevalence of dementia, at an earlier age, than the general population. The aim of this review was to establish the psychological interventions and outcomes for individuals with intellectual disability and dementia. A search of eight electronic databases and reference lists of all included articles was conducted using PRISMA guidelines. Data were synthesized using an integrative method. Initial searching produced 2,331 papers. Twenty-one studies met the inclusion criteria. Interventions were deductively categorized into behavioral, systemic, and therapeutic. All studies reported positive findings for individuals and for the systems which support them, but limited by methodological issues and neglect of the direct experience and impact on individuals themselves. The findings are discussed in relation to the wider literature and evidence base. Future research should aim to adopt methodologically robust designs that are inclusive of the individual experience of people with intellectual disability.

McGuire, B. E.; Whyte, N., & Hardardottir, D.

Alzheimer's disease in Down Syndrome and intellectual disability: A review.


Abstract: The authors review the literature on Alzheimer's disease (AD) in persons with general intellectual disabilities and those with Down syndrome. It focuses on the prevalence, clinical manifestations, diagnosis and management
of AD in these populations. The literature indicates that people with Down syndrome have a greatly increased risk of dementia from their mid 40s, while people with general intellectual disabilities have similar rates of AD to the general population. Taking into account the life expectancy of people with intellectual disabilities and those with Down syndrome, guidelines are provided for estimating the proportion of service users in a population that are at risk of developing dementia. The difficulties around diagnosis are reviewed and a particular emphasis is placed on the range of psychometric measures that may contribute to assessment and diagnosis. The management of service users who develop dementia is also reviewed and the implications for service providers are highlighted.

McKenzie, K., Harte, C., Patrick, S., Matheson, E., & Murray, G.C.
Abstract: Article reports study the examined two methods of using the Vineland Adaptive Behavioral Scales (VABS) to measure behavioral change in adults with Down syndrome who were surmised to be at-risk of Alzheimer’s disease. The first approach used the VABS within a semi-structured interview and all areas of behavioral change identified by staff were noted. The second approach used the basal rule of the VABS as indicated in the Scales’ manual. Comanion of the two approaches indicated that using the second approach highlighted significant decline in scores (for adults meeting the criteria for ‘probable Alzheimer’s disease’) on a number of domains between baseline and 12-24 months. One limitation of this approach that was noted was that this scoring method appeared to miss more subtle changes on behavior, which may be indicative of early Alzheimer’s disease – which were picked up by the first approach. Authors recommend flexibility in using the VABS for assessment purposes and caution researchers to be explicit in reporting how the VABS was used in studies assessing dementia.

McKenzie, K., Metcalfe, D., Michie, A., & Murray, G.
Service provision in Scotland for people with an intellectual disability who have, or who are at risk of developing, dementia. Dementia (London), 2020, 19(3), 736-749. doi: 10.1177/1471301218785737
Abstract: This research aimed to identify current national provision by health services in Scotland in relation to proactive screening and reactive assessment for people with an intellectual disability in Scotland who have, or are at risk of developing, dementia. Staff from 12 intellectual disability services, representing the 11 health board areas in Scotland, completed an online questionnaire which asked about proactive screening and reactive assessment for people with intellectual disability who had, or were at risk of developing, dementia as well as suggested areas for improvement. All of the areas provided services for people with intellectual disability who have, or are at risk of developing, dementia, but differed as to whether this was reactive, proactive or both. Nine services offered intervention following diagnosis. The most common elements used across both proactive screening and reactive assessment were conducting a health check, using a general dementia questionnaire designed for people with an intellectual disability and direct assessment with the person. Clinical psychology and community learning disability nurses were the professions most likely to be involved routinely in both proactive screening and reactive assessments. The psychometric properties of the most commonly used assessments of cognitive and behavioral functioning were mixed. The areas of improvement suggested by practitioners mainly related to ways of improving existing pathways. This research represents the first step in providing an overview of service provision in Scotland. There was some inconsistency in relation to the general and specific components which were involved in proactive screening and reactive assessment. Implications for service provision are discussed.

McKenzie, K., Metcalfe, D., & Murray, G.
Abstract: The increasing number of individuals with an intellectual disability who are at risk of developing dementia highlights the need to use measures with strong psychometric properties as part of the screening, assessment and diagnostic process. Searches were made of clinical and good practice guidelines and English language journal articles sourced from Proquest, Web of Science and Scopus databases (up to July 2017) for tools which were designed or adapted for the purpose of helping to diagnose dementia in people with intellectual disability. Based on a detailed review of 81 articles and guidelines, the present authors identified 22 relevant tools (12 cognitive, 10 behaviour). These were reviewed in terms of their psychometric properties. A number of tools were found to be available for use with people with intellectual disability; however, few were specifically standardized for this purpose which also had comprehensive information about reliability and validity.

McLaughlin, K., & Jones, D.
Abstract: A qualitative interview study was undertaken to determine the information and support needs of carers of adults who have Down’s syndrome and dementia. The data were analysed thematically. People who care for someone with Down’s syndrome and dementia were asked about what it was like being a carer. Four of the carers were brothers or sisters of the person who had dementia, and two were paid carers not family members. All carers said that they wanted more information about some of the health and social problems that came with having dementia. Carers said that they wanted to know more information and meet other carers and other people who have dementia. Carers’ information and support needs were seen to change at pre-diagnosis, diagnosis, and post-diagnosis. Helping carers to manage the changing nature of the adult with dementia is seen to be an essential part of the health professional’s role.

McQuillan, S., Kalsy, S., Oyebode, J., Millichap, D., Oliver, C., & Hall, S.
Adults with Down’s syndrome and Alzheimer’s disease Tizard Learning Review, 2003, 8(4), 4-13
Abstract: Adults with Down’s syndrome are at risk of developing Alzheimer’s disease in later life. This paper gives an overview of the current research in the area and discusses the implications it raises for individuals, carers, and service providers. Information on the link between Down’s syndrome and Alzheimer’s disease and prevalence rates are given. The clinical symptoms of Alzheimer’s disease and a stage model documenting the progression of the disease are presented. Attention is drawn to the problems inherent in assessing and diagnosing Alzheimer’s disease in a person with a pre-existing intellectual disability. Also discussed are the management of Alzheimer’s disease, a focus on care management practices, and recommendations for service provision (including guidelines for supporting individuals which include maintaining skills, adapting a person-centered approach, implementing psychosocial interventions, and multi-disciplinary care management). Recommendations for the future include increasing education and awareness, implementing screening services, improving assessment methods, and developing appropriate services.

McVicker, R.W., Shanks, O.E., & McClelland, R.J.
Abstract: The aim of this study was to establish the prevalence of epilepsy in persons with Down’s syndrome aged 19 years and over. A total of 191 adults with Down’s syndrome were identified, giving a prevalence of 0.76/1000 (95% CI 0.75 to 0.77). Of these, 18 had epilepsy, giving a prevalence of 9.4% (95% CI 5.3% to 13.5%). The prevalence of epilepsy increased with age, reaching 46% in those over 50. The neurophysiological (EEG) findings of the epilepsy group were compared with those of a control group of Down’s syndrome adults without epilepsy. Paroxysmal abnormalities consistent with a diagnosis of epilepsy were found in 80% of the epilepsy group, compared with only 13% of controls (P < 0.001). Epilepsy of late onset was associated with diffuse EEG abnormalities and clinical evidence of dementia. The age distribution and EEG findings suggest two independent processes in the causation of epilepsy: late-onset epilepsy associated with clinical evidence of dementia, and early-onset epilepsy in the absence of dementia.

Melville, C.A., Hamilton, S., Hankey, C.R., Miller, S., Boyle, S.
The prevalence and determinants of obesity in adults with intellectual disabilities Obesity Reviews, 2007 (May), 8(3), 223-230.
Abstract: People with intellectual disabilities experience significant health...
inequalities compared with the general population, including a shorter life expectancy and high levels of unmet health needs. Another accepted measure of health inequalities, the prevalence of obesity, has been shown to be higher in adults with intellectual disabilities than in the general population. While the factors contributing to the increased prevalence among adults with intellectual disabilities are not well understood, the high rates of obesity among younger adults highlight the need for further research involving children and adolescents with intellectual disabilities. To take forward the priorities for research and the development of effective, accessible services, there is a need for collaboration between professionals working in the fields of intellectual disabilities and obesity.

Menéndez M. Down syndrome, Alzheimer’s disease and seizures. Brain Development, 2005, 27(4), 246-252. doi:10.1016/j.braindev.2004.07.008. Abstract: Neuropsychologically, Alzheimer-type abnormalities are demonstrated in patients with Down syndrome (DS), both demented and non-demented and more than a half of patients with DS above 50 years develop Alzheimer’s disease (AD). The apolipoprotein E epsilon4 allele, oestrogen deficiency, high levels of Abeta1-42 peptide, elevated expression of BACE2, and valine polymorphism of prion protein gene are associated with earlier onset of dementia in DS individuals. Advanced AD alone may be an important risk factor for new-onset seizures in older adults and age above 60 years is a recognized risk factor for poor outcome from convulsive and nonconvulsive status epilepticus. DS patients aged over 45 years are significantly more likely to develop Alzheimer’s disease than those less than 45 years and up to 84% demented individuals with DS develop seizures. Late-onset epilepsy in DS is associated with AD, while early-onset epilepsy is associated with an absence of dementia. In AD patients with a younger age of dementia onset are particularly susceptible to seizures. DS adults with epilepsy score significantly higher overall on the adaptive behaviour profile. Language function declined significantly more rapidly in AD patients with seizures and there is a good correlation between the severity of EEG abnormalities and cognitive impairment whereas in DS slowing of the dominant occipital rhythm is related to AD and the frequency of the dominant occipital activity decreases at the onset of cognitive deterioration.

Millichap, D., Oliver, C., McQuillan, S., Kalsy, S., Lloyd, V., & Hall, S. Descriptive functional analysis of behavioral excesses shown by adults with Down syndrome and dementia. International Journal of Geriatric Psychiatry, 2003, 18, 844-854. doi: 10.1002/jgps.930. Abstract: The study examined the hypothesis that a functional relationship exists between social environmental events and behavioral excesses in individuals with Down syndrome and dementia. A case-series design was employed (n = 4) using a direct observation-based descriptive functional assessment procedure. Observations were conducted in the natural environments of four participants over periods ranging from 11 to 15.4 hours. Data were collected on non-verbal and verbal behavioral excesses, appropriate engagement and verbal interaction with others. Social environmental events observed including both staff and peer behavior. Analysis of co-occurrence for behavioral excesses and social environmental events indicated significant relationships for some behaviors consistent with operant reinforcement processes. Sequential analysis showed that changes in the probability of social contact occurred in the period directly preceding and following verbal behaviors. Results support the hypothesis that, consistent with literature for older adults with dementia in the general population, some behavioral excesses were functional in nature and not randomly occurring events. No relationship was found between appropriate engagement and staff.

Mircher, C., Cieuta-Walti, C., Marey, I., Rebillat, A-S., Cretu, L., Milenko, E., Conte, M., Sturtz, F., Reithore, M-O., & Ravel, A. Acute regression in young people with Down syndrome. Brain Science, 2017, Jun, 7(6), 57. Published online 2017 May 27. doi: 10.3390/brainsci7060057. Abstract: Adolescents and young adults with Down syndrome (DS) can present a rapid regression with loss of independence and daily skills. Causes of regression are unknown and treatment is most of the time symptomatic. We did a retrospective cohort study of regression cases: patients were born between 1959 and 2000, and were followed from 1984 to now. We found 30 DS patients aged 11 to 30 years old with history of regression. Regression occurred regardless of the cognitive level (severe, moderate, or mild intellectual disability (ID)). Patients presented psychiatric symptoms (catatonia, depression, delusions, stereotypes, etc.), partial or total loss of independence in activities of daily living (dressing, toilet, meals, and continence), language impairment (silence, whispered voice, etc.), and loss of academic skills. All patients experienced severe emotional stress prior to regression, which may be considered the trigger. Partial or total recovery was observed for about 50% of them. In our cohort, girls were more frequently affected than boys (64%). Neurobiological hypotheses are discussed as well as preventative and therapeutic approaches.

Mohan, M., Bennet, C., & Carpenter, P.K. Rivastigmine for dementia in people with Down syndrome Cochrane Systemic Review - Intervention, 2009. 1. https://doi.org/10.1002/14651858.CD007658. Abstract: Alzheimer’s dementia (AD) is the most common form of dementia in people with Down Syndrome (DS). Acetylcholine is a chemical found in the brain that has an important role in memory, attention, reason and language. Rivastigmine is a “pseudo-irreversible” inhibitor of acetylcholinesterase, which is thought to maintain levels of acetylcholine. Rivastigmine can improve cognitive function and slow the decline of AD in the general population over time. It is important to note that people with DS tend to present with AD at a much younger age than the normal population as well as having subtle differences in physiology (e.g. metabolism and heart rate) and may therefore have different requirements from the general population. The authors sought to determine the effectiveness and safety of rivastigmine for people with DS who develop AD by using the following search methods, CENTRAL, MEDLINE, EMBASE, CINHAL, PsycINFO, BIOSIS, SCI, SSCI and the NRR, up to October 2008. They also contacted the manufacturers of rivastigmine as well as experts in the field, to ask about reports of unpublished or ongoing trials. Selection criteria included randomised controlled trials of participants with DS and AD in which treatment with rivastigmine was administered compared with a placebo group. Authors found that no study was identified which met inclusion criteria for this review and concluded that as there are no included trials, recommendations cannot be made about rivastigmine for AD in DS. Well-designed, adequately powered studies are required.

Molnar, F.J., Benjamin, S., Hawkins, S.A., Briscoe, M., & Ehsan, S. One size does not fit all: Choosing practical cognitive screening tools for your practice. JAGS (Journal of the American Geriatrics Society), 2020, 68(10), 2207-2213. DOI: 10.1111/jgs.1671. Abstract: Every year, millions of patients worldwide undergo cognitive testing. Unfortunately, new barriers to the use of free open access cognitive screening tools have arisen over time, making accessibility of tools unstable. This article is in follow-up to an editorial discussing alternative cognitive screening tools for those who cannot afford the costs of the Mini-Mental State Examination and Montreal Cognitive Assessment (see www.dementiascreen.ca). The current article outlines an emerging disruptive “free-to-fee” cycle where free open access cognitive screening tools are integrated into clinical practice and guidelines, where fees are then levied for the use of the tools, resulting in clinicians moving on to other tools. This article provides recommendations on means to break this cycle, including the development of tool kits of valid cognitive screening tools that authors have contracted not to charge for (i.e., have agreed to keep free open access).The PRACTICAL 1 Criteria (PRACTICing Clinician Accessibility and Logistical Criteria Version 1) are introduced to help clinicians select from validated cognitive screening tools, considering barriers and facilitators, such as whether the cognitive screening tools are easy to score and free of cost. It is suggested that future systematic reviews embed the PRACTICAL 1 criteria, or refined future versions, as part of the standard of review. Methodological issues, the need for open access training to insure proper use of cognitive screening tools, and the need to anticipate growing ethnolinguistic diversity by developing tools that are less sensitive to educational, cultural, and linguistic bias are discussed in this opinion piece.

Moss, S., & Patel, P.
Dementia in older people with intellectual disability: symptoms of physical and mental illness, and levels of adaptive behavior.
Abstract: Detailed data on health and functional ability of 101 people with intellectual disability over 50 years of age are presented. Using a combination of informal interviewing, observation, and measurement of cognitive change over a 3-year period, 12 of these individuals were identified as suffering from dementia. Their data are compared to those of the non-dementia sufferers. The people suffering from dementia had a greater number of chronic physical health problems and chronic disability resulting from physical health problems. Their capacity for self-directed activity was lower. The subjects had a reduced capacity for self-directed activity was lower. The question that most physicians feel ill-prepared to answer. The National Task Group on Intellectual Disabilities and Dementia Practices was convened to help formally address this topic, which remains largely under-represented in the medical literature. The task group, comprising specialists who work extensively with adults with I/DD, has promulgated the following Consensus Recommendations for the Evaluation and Management of Dementia in Adults With Intellectual Disabilities as a framework for the practicing physician who seeks to approach this clinical question practically, thoughtfully, and comprehensively.

Moss, S., Schlamb, C., & Harrison, B.
Down syndrome and dementia: Guide to identification, screening, and management.
Abstract: Down syndrome (DS) is an intellectual disability due to the genetic disorder trisomy21. Many individuals with DS are living into middle and older adulthood, experiencing chronic health problems, and are at risk for dementia. This article describes the primary care management of adults with DS, the relationship between DS and Alzheimer’s dementia, and screening protocols for primary care. [Extract follows] The family of the person with DS and AD needs a team to navigate the changes that are inevitable during the course of this disease. An open discussion with both the NP and social worker will assist most families in providing the best quality of life for aging adults with DS. Maintaining social contact with family and friends may be difficult for aging DS adults, but it is vital for well-being. Expectations for self-care and new learning must be readdressed and give way to an emphasis on a positive approach to the adult with DS. Families and caregivers need to proactively create a safe and calming home environment to promote quality of life. As AD progresses, assessment of nonverbal communication is essential in the care of adults with DS, but it may be challenging. Communication may be based on nonverbal cues and gestures. Additional response time is needed to allow for verbal or nonverbal communication of needs. Utilizing communication boards and pictures of common persons or objects is beneficial in this population. Safe relationships grounded in familiarity and trust are paramount in providing a secure environment for adults with DS and AD.

Moss, S., Lambe, L., & Hogg, J.
Physical and mental health
Aging Matters - Pathways for Older People with Learning Disabilities: Manager's Reader. pp. 41-60
Abstract: This unit, one of six that is used for training staff, covers briefly some of the key issues related to physical and mental health, and touches on dementia. Although not specifically developed for care management of adults with dementia, the text, in total, can be a useful resource for staff working in care settings when one or more of the adults in the setting are affected by dementia.

Mullins, D., Daly, E., Simmons, A., Beacher, F., Foy, C.M.L., Lovestone, S., Hallahan, B., Murphy, K.C., & Murphy, D. G.
Dementia in Down’s syndrome: an MRI comparison with Alzheimer’s disease in the general population.
Abstract: Down’s syndrome (DS) is the most common genetic cause of intellectual disability. People with DS are at an increased risk of Alzheimer’s disease (AD) compared to the general population. Neuroimaging studies of AD have focused on medial temporal structures; however, to our knowledge, no in vivo case–control study exists comparing the anatomy of dementia in DS to people with AD in the general population. We therefore compared the in vivo brain anatomy of people with DS and dementia (DS+) to those with AD in the general population. Using MRI in 192 adults, we compared the volume of whole brain matter, lateral ventricles, temporal lobes and hippocampus in DS subjects with and without dementia (DS+, DS−), to each other and to three non-DS controls. These included one group of individuals with AD and two groups of controls (each age-matched for their respective DS and general population AD cohorts). AD and DS+ subjects showed significant reductions in the volume of the whole brain, hippocampus and temporal lobes and a significant elevation in the volume of the lateral ventricle, compared to their non-demented counterparts. People with DS+ had a smaller reduction in temporal lobe volume compared to individuals with AD. DS+ and AD subjects have a significant reduction in volume in the same brain regions. We found preliminary evidence that DS individuals may be more sensitive to tissue loss than others and have less ‘cognitive reserve’.

Muralidhara, M., Spectorb, A., Huia, E.K., Liua, L., & Alia, A.
A systematic review of psychosocial interventions for people with intellectual disabilities and dementia.
Aging & Mental Health, 2023, 9 Oct. DOI: 10.1080/13607863.2023.2265322
Abstract: As the life expectancy of individuals with intellectual disabilities (ID) continues to increase, there is an increased risk of developing dementia. While psychosocial interventions are gaining prominence, evidence is limited for people with both dementia and ID. This review discusses the effectiveness of direct psychosocial interventions and adaptations to facilitate delivery within this population. The review followed the PRISMA guidelines. Five electronic databases, grey literature, and reference lists of included articles were searched for relevant studies. 10 eligible studies were appraised and analyzed by narrative synthesis. Ten distinct interventions were identified and categorized as (1) behavioral, (2) structured (manualized), and (3) personalized interventions, based on their purpose and delivery. All interventions were beneficial in improving a range of outcomes, though some studies were of low quality and most had small samples. Common adaptations included simplification of tasks and material, higher staff-to-client ratio, and alternative communication methods. There is emerging evidence for several psychosocial interventions for people with ID and dementia, though further research is required on effectiveness and generalizability. The adaptations discussed may guide implementation into routine care and contribute to current policies and guidelines on improving ID and dementia care.

Nagdee, M.
Dementia in intellectual disability: a review of diagnostic challenges.
African Journal of Psychiatry (Johannesburg), 2011, 14, 194-199. doi: 10.4314/ajps.v14i3.1
Abstract: The evaluation of dementia in individuals with intellectual disability, which will guide subsequent intervention, care and management depends on the systematic review of a number of factors: (1) the individual historical context, obtained from multiple sources, (2) evaluation of the pre-existing cognitive, behavioral, psychiatric, medical and adaptive skill profile, (3) the constellation, and pattern of evolution, of presenting signs and symptoms, (4) results of focused investigations, and (5) refinement of the differential diagnosis. In patients with ID, standard clinical methods need to be supplemented by careful,
longitudinal behavioral observations, and individually tailored assessment techniques. Co-morbidity, multiple biological, psychological and socio-environmental factors, and complex interactions among events, are the reality for many ageing people with ID. Determining the various influences is often a formidable clinical task, but should be systematically carried out using medical, cognitive, behavioral, neuropsychiatric and psycho-social frameworks.


Abstract: This guideline covers diagnosing and managing dementia (including Alzheimer's disease). It aims to improve care by making recommendations on training staff and helping carers to support people living with dementia.

National Task Group on Intellectual Disabilities and Dementia Practices. My thinker's not working: A national strategy for enabling adults with intellectual disabilities affected by dementia to remain in their community and receive quality supports. 42pp.

National Task Group on Intellectual Disabilities and Dementia Practices [www.aadmd.org/nlq]. (2012). Abstract: My Thinker's Not Working is the short title for the 42-page summative report issued by the National Task Group on Intellectual Disabilities and Dementia Practices, a planning and advocacy group organized to produce a national plan on dementia and intellectual disabilities. The report offers 20 recommendations for the improvement of services nationally and locally and suggests that its findings and recommendations be considered and integrated into the reports and plans being developed by the federal Advisory Council on Alzheimer's Research, Care, and Services -- under the National Alzheimer's Project Act. The document reviews the main issue facing adults with intellectual disabilities as they age when they are affected by dementia, as well as their families and provider organizations. The document is composed of 7 sections (Change and Purpose, The Population, Challenges Facing the Population, Community Services, Education and Training, Financing, and Possible Solutions) and the National Dementia and Intellectual Disabilities Action Plan.

NAMHI Alzheimer's Dementia in persons with intellectual disabilities: Some common questions and concerns

NAMHI, 5 Fitzwilliam Place, Dublin 2, Ireland

Abstract: 28 page booklet with 18 sections/question areas outlining basic information about Alzheimer's disease and people with ID, diagnostic resources, and service to help cope with the course of the disease. Developed by Dr. Mary McCarron of Trinity College Dublin.


Abstract:A comprehensive baseline of emotional functioning was established for adults with Down syndrome. Five emotional factors were studied using groups of (a) adults with Down syndrome (n = 30), (b) clinical control subjects with dementia of the Alzheimer type (n = 18), and (c) elderly control subjects without intellectual disability (n = 25). Results of planned statistical comparisons showed indifference, pragmatism, and inappropriateness as primary emotional factors separating Down syndrome and Alzheimer disease groups from elderly control subjects without intellectual disability. Indifference was also shown to co-vary with cognitive mental state, whereby increased levels of indifference were associated with decreased levels of cognitive functioning. There is the possibility of noncognitive variables signaling dementia of the Alzheimer type in individuals with Down syndrome.


Abstract: Study examined emotional changes in adults with Down Syndrome (DS) over time and to determine whether changes in these psychological variables were associated with brain atrophy on MRI scan and the presence of pathological reflexes on the neurological examination. Participants were 26 adults with DS and their caregivers. Caregivers completed a measure of emotional functioning about individuals with DS at two different time points (1 year apart). Levels of cognitive functioning were measured and neurological and MRI examinations were performed on all subjects at initial testing. Significant group effect separated those with and without pathological findings on MRI and neurological exam across three different scales: depression, indifference, and pragmatic language functioning. Problems of poor pragmatic language functioning appeared later in the course of suspected Alzheimer's disease (AD), as demonstrated by a significant group effect at time 2, but not at initial testing. In these subjects, the primary emotional change was a decline in social discourse (e.g. conversational style, literal understanding, verbal expression in social contexts). These emotional levels were stable over time, regardless of degree of cognitive decline. Specific emotional changes occur during the course of AD which were associated with abnormal findings from MRI and from neurological examination. These results, along with abnormalities in brain imaging and the presence of pathological reflexes, suggested that frontal lobe dysfunction is likely to be an early manifestation of Alzheimer's Disease in Down Syndrome.

New York State Developmental Disabilities Planning Council

Dementia and people with intellectual disabilities – What can we do? 23 minutes


Abstract: Sweden has closed all institutions and imposed legislation to ensure health promotion and disease control strategies. We investigated patterns and risk of mortality among older adults with ID in Sweden. This retrospective cohort study compared older adults aged 55 years and older with ID with a control population. Participants were followed during 2002-2015 or death, and censored if they moved out of Sweden. Individuals with ID were identified from two national registers: one covering all specialist health-care visits (out-patient visits and hospitalization) and the other covering people accessing social/support services. Individuals with ID (n = 15,289) were matched with a control population.
by sex, birth year, and year of first hospitalisation/out-patient visit/access to LSS services. Cause-of-death data were recorded using International Classification of Diseases, Tenth Revision. Cox proportional hazards regression were conducted to assess if overall and cause-specific mortality rate among individuals with ID was higher than in the Swedish population. The overall mortality rate among individuals with ID was 2483 per 100,000 people compared with 810 in the control population. Among those who died, more individuals with ID were younger than 75 years and unmarried. Leading causes of death among individuals with ID were circulatory diseases (34%), respiratory diseases (17%) and neoplasms (15%). Leading causes of death in a sub-sample with Down syndrome (DS) were respiratory diseases (37%), circulatory diseases (28%) and mental/behavioural disorders (11%). Epilepsy and pneumonia were more common among individuals with ID than controls. Alzheimer’s disease was common in the control population and individuals with DS, but not among those with ID when DS was excluded. Individuals with ID had a higher overall mortality risk (hazard ratio [HR] 4.1, 95% confidence interval [CI] 4.0-4.3) and respiratory disease death risk (HR 12.5, 95% CI 10.9-14.2) than controls. Older adults with ID in Sweden carry a higher mortality risk compared with the general population, mainly attributable to respiratory, nervous and circulatory diseases. Care for this group, particularly during the terminal stage of illness, needs to be tailored on understanding of their main health problem.

Nieuwenhuis-Mark, R.E.

Diasising Alzheimer’s dementia in Down syndrome: Problems and possible solutions.


Abstract: It is widely accepted that people with Down syndrome are more likely than the general population to develop Alzheimer’s dementia as they age. However, the diagnosis can be problematic in this population for a number of reasons. These include: the large intra-individual variability in cognitive functioning, the different diagnostic and methodological procedures used in the field and the difficulty in obtaining baseline levels of cognitive functioning in this population with which to assess cognitive and behavioral change. Recent researchers have begun to suggest ways around these difficulties. This review explores these recent developments and provides recommendations which may aid clinicians in their attempts to diagnose Alzheimer’s dementia in the early stages in the Down syndrome population.

Noelker, E.A. & Somple, L.C.


Abstract: Book chapter providing a brief summary of significant assessment and care issues affecting adults with Down syndrome who have Alzheimer’s disease. Noted are the needs for education of carers and families, as well as specialist care provision and community services.

Northway, R., Holland-Hart, D., & Jenkins, R.


Abstract: Older people with intellectual disabilities often experience high levels of health needs and multiple morbidities but they may be supported by residential care staff with little or no previous experience of identifying and meeting health needs. Little is known regarding how they undertake this health-related role and this exploratory study seeks to address this gap. A purposive sample of 14 managers of supported living accommodation in Wales were interviewed in 2014 to determine their experiences of supporting tenants in relation to age-related health needs. The semi-structured interviews were transcribed and thematically analysed. Three of the emerging themes are reported in this paper: meeting health needs, the consequences of ageing and relationships. Findings indicate that residential care staff support older people with intellectual disabilities with complex and multiple health needs: they monitor health status, support access to healthcare, provide additional support arising from changing health needs and advocate for tenants in the context of healthcare. However, their role is often not understood by healthcare professionals. The importance of staff having a long-term relationship with those they support was identified as being important to identifying any health-related changes. The need to develop effective relationships with healthcare staff was also noted. It is concluded that there is a need for better understanding among health staff of the role of residential social care workers and for further research regarding health-related communication.


Eine deutschsprachige Fassung der Cambridge Examination for Mental Disorders of Older People with Down’s Syndrome and Others with Intellectual Disabilities: Ein Diagnoseverfahren zur Erfassung von Demenz bei Menschen mit einem Down-Syndrom (A German version of the Cambridge examination for mental disorders of older people with Down’s syndrome and others with intellectual disabilities: A diagnostic procedure for detecting dementia in people with Down's syndrome).


Abstract: Although people with Down’s syndrome (DS) are at a high risk of developing an Alzheimer type dementia (AD) due to a duplication of the amyloid precursor gene, there are practically no internationally available test procedures to detect cognitive deficits in this at risk population in the German language. The aim was to provide a German translation and intercultural adaptation of the Cambridge examination for mental disorders of older people with Down’s syndrome and others with intellectual disabilities (CAMDEX-DS), which is available in English and Spanish. This instrument for diagnostics and monitoring consists of a psychological test examination (CAMCOG-DS) and a caregiver interview. The translation and adaptation of the CAMDEX-DS were achieved through a multistep translation process, whereby two independent forward and back translations were provided by professional translators and a consensus version was finalized and tested. The final version of the caregiver interview was applied to 11 subjects and the CAMCOG-DS was conducted with 28 patients. The German version of the CAMDEX-DS proved to be easily administered. The CAMCOG-DS could be fully administered to 21 out of 28 patients (75%). The CAMCOG-DS values were much lower for older patients aged ≥45 years than for younger patients (46/109 vs. 73.5/109; p = 0.033). The German version of the CAMDEX-DS provides an internationally recognized tool for the diagnostics and monitoring of cognitive decline in Down’s syndrome. Furthermore, the German version can standardize medical care of these patients. In particular it provides a means of participation in international research trials for this at risk population.


Proteomic profiles of incident mild cognitive impairment and Alzheimer’s disease among adults with Down syndrome Alzheimer’s & Dementia: Diagnosis, Assessment & Disease Monitoring, 12, 1, doi: https://doi.org/10.1002/dad2.12033

Abstract: We sought to determine if proteomic profiles could predict risk for incident mild cognitive impairment (MCI) and Alzheimer’s disease (AD) among adults with Down syndrome (DS). In a cohort of 398 adults with DS, a total of n = 186 participants were determined to be non-demented and without MCI or AD at baseline and throughout follow-up; n = 103 had incident MCI and n = 81 had incident AD. Proteomics were conducted on banked plasma samples from a previously generated algorithm. The proteomic profile was highly accurate in predicting incident MCI (area under the curve [AUC] = 0.92) and incident AD (AUC = 0.88). For MCI risk, the support vector machine (SVM)-based highflow cut-point yielded an adjusted hazard ratio (HR) = 6.46 (P < .001). For AD risk, the SVM-based high/low cut-point score yielded an adjusted HR = 8.4 (P < .001). On the current results provide support for our blood-based proteomic profile for predicting risk for MCI and AD among adults with DS.

O’Caoimh, R., Clune, Y., & Molloy, D.W.

Screening for Alzheimer’s disease in Downs syndrome Alzheimer’s & Dementia: Diagnosis, Assessment & Disease Monitoring, 2013, S7; 001.

http://dx.doi.org/10.4172/2161-0460.S7-001

Abstract: Down syndrome (DS), is associated with an increased incidence of...
Alzheimer’s disease (AD). Although pathological changes are ubiquitous by 60 years of age, prevalence rates are lower. The diagnosis of AD in persons with DS is challenging, complicated by atypical presentations, baseline intellectual disability and normal age associated cognitive decline. Effective screening is limited by a paucity of diagnostic criteria, cognitive screening instruments and screening programs. Both observer-rated questionnaires and direct neuropsychological testing are suggested to screen for cognitive impairment, each with different strengths and weaknesses. This paper reviews commonly used screening instruments and explores the unique challenges of screening for AD in persons with DS. It concludes that single, one-dimensional screening tools and opportunistic evaluations are insufficient for detecting dementia in this population. These should be replaced by batteries of tests, incorporating informant questionnaires, direct neuropsychological testing, assessment of activities of daily living and behaviors, measured at baseline and reassessed at intervals. Developing these strategies into organized screening programs should improve diagnostic efficiency and management.

O’Dwyer, M., Finnerty, S., Henman, M., Carroll, R., McCallion, P., & McCarron, M.
Abstract: High rates of dementia have been reported among older adults with intellectual disability (ID), particularly those with Down Syndrome. As the use of dementia drugs in this patient group lacks an evidence base, their rates of use are of interest. Incidence and prevalence rates were determined using a combined dementia variable for three waves of the IDS-TILDA study, a nationally representative study of older adults with ID in Ireland. Incidence of dementia was defined as participants newly reporting a diagnosis and/or newly receiving dementia drug(s) at each wave. Prevalence of dementia was defined as those who had reported a diagnosis at a previous wave and/or received dementia drug(s) at a previous wave. Drugs for dementia were included as a proxy for dementia diagnosis, in those with no diagnosis. Dementia incidence remained similar across Waves: 5.0% at Wave 1, 4.3% at Wave 3. Prevalence increased, 5% at Wave 1 to 9.6% by Wave 3. Those receiving dementia drug(s) decreased, from 54.1% of those with dementia at Wave 1 to 28.8% at Wave 3. Three dementia drugs were reported: donepezil, memantine and rivastigmine. It was found that use of drugs for dementia decreased, despite an increased incidence. Further research into efficacy of use of a drugs is needed.

O’Leary, L., Cooper, S.-A., & Hughes-McCormack, L.
Abstract: Death of people with intellectual disabilities is considered to be earlier than for the general population. Databases were searched for key words on intellectual disabilities and death. Strict inclusion/exclusion criteria were used. Information was extracted from selected papers, tabulated and synthesized. Prospero registration number: CRD42015020161. Of 19,111 retrieved articles, 27 met criteria. Death was earlier by 20 years. It has improved in recent decades; however, the same inequality gap with the general population remains. More severe intellectual disabilities, and/or additional comorbidities rendered it shortest. Standardized mortality ratios showed a greater inequality for women than men. Respiratory disease and circulatory diseases (with greater congenital and lesser ischemic disease compared with the general population) were the main causes of death. Cancer was less common, and cancer profile differed from the general population. Some deaths are potentially preventable. All research is from high-income countries, and cause of death is surprisingly little investigated. Authors concluded that improved health care, including anticipatory care such as health checks, and initiatives addressing most relevant lifestyle behaviors and health risks are indicated.

Oliver, C., Adams, D., Holland, A.J., Brown, S.S.G., Ball, S., Dodd, K., & Carr, J.
Acquired mild cognitive impairment in adults with Down syndrome: Age-related prevalence derived from single point assessment data normed by degree of intellectual disability.

Abstract: Individuals with Down syndrome (DS) are at significant risk for early onset Alzheimer’s disease (AD), likely due to the triplexation of genes on chromosome 21 that facilitate AD neuropathology. To aid the effective early diagnosis of dementia in DS, we demonstrate the strategy of using single point assessment of cognitive performance with scoring normed for degree of intellectual disability to generate age related prevalence data for acquired mild cognitive impairment (AMCI). Four hundred and twelve adults with DS were assessed using the Neuropsychological Assessment of dementia in adults with Intellectual Disability. Normative data, banded by degree of intellectual disability, allowed identification of AMCI by atypical deviation from expected performance. AMCI was evident in approximately 20% of adults with DS aged 40 and under, 40%, aged 41-50 and 45% aged 51 and over. Relative risk increased significantly in those aged 46 and over. Analysis of prevalence by 5-year age bands revealed two peaks for higher prevalence of AMCI. Psychometric data indicate single point assessment of AMCI is possible for the majority of adults with DS. Two peaks for age-related prevalence of AMCI suggest the risk for onset of AD conferred by trisomy of chromosome 21 is moderated by another factor, possibly ApoE status.

Oliver, C., & Holland, A.J.
Abstract: Neuropathological change found in nearly all individuals with Down syndrome over the age of 35 years closely resembles that of Alzheimer’s disease. The extent to which dementia occurs as a result of this change is unclear, and the studies which have investigated presumed cognitive deficits are reviewed. The theories put forward to explain the association between these two disorders and their possible significance to the understanding of the aetiology of Alzheimer’s disease are discussed.

Oliver, C., Crayton, L., Holland, A., & Hall, S.
Cognitive deterioration in adults with Down syndrome: effects on the individual, caregivers, and service use American Journal on Mental Retardation, 2000, 103, 455-465
Abstract: Individuals with Down syndrome (N = 49) who had participated in serial neuropsychological assessments were assigned to one of three groups comparable in level of premorbid intellectual disability: (1) those showing cognitive deterioration, (2) those comparable in age but not showing cognitive deterioration and (3) those not showing cognitive deterioration but younger. Those experiencing cognitive deterioration were less likely to receive day services, had more impoverished life experiences, and required more support compared to groups without cognitive deterioration. When age was controlled for, cognitive deterioration was significantly positively associated with carer difficulties and service use and negatively associated with life experiences for the individual. Results suggest a potential role for carer difficulties in influencing life experiences of adults with Down syndrome showing cognitive decline.

Oliver, C., Kaisy, S., McQuillan, S., & Hall, S.
Abstract: Informant-based assessment of behavioral change and difference in dementia in Down syndrome can aid diagnosis and inform service delivery. To date few studies have examined the impact of different types of behavioral change. The Assessment for Adults with Developmental Disabilities (AADS), developed for this study, assesses behavioral excesses (11 items) and deficits (17 items) associated with dementia. Inter-informant reliability, internal consistency and concurrent validity were evaluated and found to be robust. A comparison of the AADS subscale scores for three groups (n = 12) of adults with Down syndrome demonstrated more frequent deficits and excesses and greater management difficulty and effects on the individual in a dementia group than age comparable and younger groups. The AADS is a promising dementia specific measure for people with intellectual disability. Further research should evaluate change as dementia progresses and the nature of management difficulty and effects on the individual.
Olsen, R.V., Ehrenkranz, E., & Hutchings, B.L.  
Creating the movement-access continuum in home environments for dementia care.  
Topics in Geriatric Rehabilitation, 1996, 12(2), 1-8.  
10.1097/00013614-199612000-00003  
Abstract: The majority of people with Alzheimer's disease receive some care at home, the environment of that home must be safe and supportive. In-depth interviews of 90 "seasoned" caregivers identified tactics for creating these settings through home modifications and technology. A successful modification strategy follows a three-stage movement-access continuum that responds to the disease course – assistance, restriction with compensation, and wheelchair accessibility. Approaching home modifications along this continuum encourages independence and movement when appropriate while providing safety and control. With a sensitive and ongoing modification strategy, the home environment can become an asset rather than a liability for caregiving.

Olsen, R.V., Ehrenkranz, E., & Hutchings, B.  
Creating supportive environments for people with dementia and their caregivers through home modifications.  
Technology and Disability, 1993, 2(4), 47-57  
Abstract: Article examines what caregivers did to enhance or modify their homes when a spouse or other family member had dementia. Authors address controlling access (using locking techniques, blocking access with gates and partial doors), and the like, as examining modifications to kitchens, bathrooms, and furniture. Data showed that many built ramps, double railings, hand grips, as well as extending landings for ease of wheelchair use, reducing riser heights, removing steps, and installing electric chair lifts. Home owners also reconfigured space and rooms. Authors conclude that home owners modified spaces to increase access and independence in some life areas and to limit or curtail access in others. Article is a good source of information for how the process and outcome of families tackle home modifications.

Olsen, R.V., Ehrenkranz, E., & Hutchings, B.  
Homes that help: Advice from caregivers for creating a supportive home (Alzheimer's and Related Dementias).  
77 pp.  
Newark, New Jersey: New Jersey Institute of Technology [Architecture and Building Science Research Group, School of Architecture, NJIoT, University Heights, Newark, New Jersey 07102-1982] (1993)  
Abstract: Manual that details examples of how to adapt a home for persons affected by dementia, covering care management techniques, physical adaptations, and personal monitoring strategies.

Owens, D., Dawson, J. C., & Losin, S.  
Alzheimer's disease in Down's syndrome.  
Abstract: Although neuropathologists describe Alzheimer's changes in the brains of all victims of Down's syndrome over 35 yr. of age, only 3 cases of clinical dementia in such individuals are described in the literature. In order to establish clinical correlates of Alzheimer's disease, psychiatric and neurologic findings obtained from a middle-aged group were compared to those of Down's syndrome patients in their early 20s. The older group exhibited significantly greater incidence of abnormality in (a) object identification, (b) snout reflex, (c) Babinski sign, and (d) palmaromental sign. Both groups displayed mild hypertonia rather than hypotonia, and a moderate increase in the number of patients tested in 75% of the population. While dementia is uncommon, subtle neurological changes reflect neuropathological findings present in aging sufferers of Down's syndrome.

Paiva, A.F., Nolan, A., Thumser, C., & Santos, F.H.  
Screening of cognitive changes in adults with intellectual disabilities: A systematic review.  
Brain Sciences, 2020, 10(11), 848; https://doi.org/10.3390/brainsci10110848  
Abstract: Screening and assessment of cognitive changes in adults with Intellectual Disabilities (ID), mainly Down Syndrome (DS), is crucial to offer appropriate services to their needs. Authors present a systematic review of the existing instruments assessing dementia, aiming to support researchers and clinicians' best practice. Searches were carried out in the databases Web of Science; PubMed; PsyCNFO in March 2019 and updated in October 2020. Studies were selected and examined if they: (1) focused on assessing age-related cognitive changes in persons with ID; (2) included adults and/or older adults; (3) included scales and batteries for cognitive assessment. Forty-eight cross-sectional studies and twenty-seven longitudinal studies were selected representing a total sample of 6451 participants (4650 DS and 1801 with other ID). In those studies, we found 39 scales, questionnaires, and inventories, and 13 batteries for assessing cognitive and behavioural changes in adults with DS and other ID. It was noted that the most used instrument completed by an informant or carer was the Dementia Questionnaire for Learning Disabilities (DLD), and its previous versions. The authors explore the strengths and limitations of the instruments and outline recommendations for future use.

Pape, S.  
Dementia diagnostic criteria in persons with IDD.  
Journal of Intellectual Disability Research, 2019, 63(8), 641.  
Abstract: Dementia is a common clinical presentation among older adults with IDD, particularly those with Down syndrome. The presentation of dementia may differ compared with typical Alzheimer's disease, and criteria thus require validation in IDD populations. Data from memory assessments in individuals with Down syndrome were presented to expert raters who rated the case as dementia or no dementia using ICD-10, DSM-IV-TR and DSM-5 criteria and their own clinical judgment. Estimates were then made of the concurrent validity and reliability of clinicians' diagnoses of dementia against these guidelines. Validity of clinical diagnoses were explored by establishing the stability of diagnoses over time. Similar data from previous studies in other individuals with intellectual disabilities were compared. It was found that clinical diagnoses of dementia in Down syndrome were valid and reliable and could be used as the standard against which new criteria such as the DSM-5 are measured. Criteria had good inter-rater reliability but concurrent validity varied. Author cautions that clinicians should consider the reliability and validity of dementia diagnostic criteria when applying these in clinical settings.

Pape, S.E., Baksh, R.A., Startin, C., Hamburg, S., Hithersay, R., & Strydon, A.  
The association between physical activity and CAMDEX-DS changes prior to the onset of Alzheimer's disease in Down syndrome.  
Journal of Clinical Medicine, 2021, Apr 27, 10(9), 1882. doi: 10.3390/jcm10091882.  
Abstract: People with Down syndrome are at ultra-high-risk of developing Alzheimer's dementia. At present, there are no preventative or curative treatments. Evidence from sporadic Alzheimer's disease literature suggests that lifestyle factors including physical activity may help maintain cognitive and functional skills and reduce dementia risk. Our study aimed to explore the association between regular exercise undertaken by participants with Down syndrome and changes in dementia-related domains of cognition and function. This was to consider whether physical activity may be a protective measure to delay cognitive decline and dementia in Down syndrome. Demographic, lifestyle, and health information was collected at baseline and at a two year follow up from 214 adults with Down syndrome without dementia, who also underwent assessment using the Cambridge Examination for Mental Disorders of Older People with Down Syndrome and Others with Intellectual Disabilities (CAMDEX-DS) and genetic analysis. Logistic regression models were used to examine the potential associations between decline in CAMDEX-DS domains and exercise whilst controlling for key variables. At baseline, engaging in moderate intensity exercise was associated with a 47% reduced risk of everyday skills decline and engaging in high intensity exercise was associated with a 62% reduced risk of decline in personality and behaviour. At follow-up, high levels of exercise were associated with an 87% reduced risk of decline in personality and behaviour. Moderate intensity exercise at baseline was associated with a 62% reduction in risk of decline during the follow-up period in memory and orientation. Based on our data it appears that regular moderate and high intensity exercise could reduce the risk of clinically detectable decline in a Down syndrome population with possible long-term benefits. People with Down syndrome may engage in less physical activity than their peers, and barriers remain which can prevent people with Down syndrome engaging in exercise. Our work highlights how important it is that people with Down syndrome are supported to be physically active, and to promote exercise as part of a healthy ageing plan. Clinical trials in this area would be justified to determine if engaging in exercise can lead to realistic improvements in maintaining functioning and delaying dementia onset in Down syndrome and to help develop guidance in this area.
Abstract: Authors sought to gain insight into caregivers’ understanding of the causes of behaviours they find problematic in people with Alzheimer’s disease in order to inform the development of educational strategies. A qualitative, semi-structured interview was used. Participants were 205 caregivers for a person with Alzheimer’s disease, all of whom were aware of the diagnosis and who had been recruited as part of a larger longitudinal study. Participants were from inner-city and suburban London/semi-rural Essex. The main outcome measures were caregivers’ understanding of: the cause of problematic behaviour; the ability of the person with dementia to control this behaviour; the prognosis of the illness. Most carers attribute the cognitive, behavioural and psychological symptoms of dementia to causes other than dementia; many believe that the person with dementia has control over their behaviour and substantial numbers believe the person with dementia will return to normal. This study suggests that providing facts about the illness to caregivers is not enough, as caregivers may not understand that the symptoms they observe are related to the diagnosis. Education by clinicians should focus on the understanding of caregivers and in particular explore the caregivers’ attributions of the symptoms which are present in the person for whom they care.

Patti, P., Amble, K. & Flory, M.

Abstract: Aging adults with Down’s syndrome (DS) experience more relocations and other life events than adults with intellectual disabilities aged 50 and older without DS. Age-related functional decline and the higher incidence of dementia were implicated as the contributing factors that led to relocation and nursing home placement. A retrospective study of adults with intellectual disabilities who were born prior to the year 1946 was conducted to analyze the number of relocations experienced over a 5- and 10-year period. The cohort consisted of 140 individuals (61 with DS between ages 50–71 years, and 79 without DS between ages 57–89 years) who had been referred to a diagnostic and research clinic. Analyses revealed the number of relocations over a 5- and 10-year period were significantly greater in the DS group. Placement in a nursing home for end of life care was significantly higher in the DS group whereas the majority (90%) in the non-DS group remained in a group home setting. Mortality was significantly earlier in the DS group with the mean age at death to be 61.4 years compared with 73.2 years in the non-DS group. The authors concluded that the present results suggest that aging adults with DS encounter more relocations, and are more likely to have their final placement for end of life care in a nursing home. In contrast, the adults without DS were subjected to less relocation and remained in the same group home setting.

Pary, R.J.

Abstract: (non provided - extract from article) Breif overview of examining adults with Down syndrome (DS) for functional decline. Provides brief summary of Alzheimer’s disease and the prevalence and occurrence of AD in persons with DS. Provides two tables, Table 1: Disorders which are common in DS and Miscellaneous cerebral conditions with no apparent predilections for individuals with DS; Table 2: Work-up for functional decline in DS. Discusses complicating factors such as thyroid disease, major depression, sensory impairment, infection, and other conditions. Provides brief description of factors comprising a diagnostic workup. References (n-49) provide historical glimpse into extant topical literature prior to 1992.

Pendi, D., Glatz, M., & Gasteiger-Klicpera, B.
Intellectual disabilities and dementia: New tasks and experiences of Austrian formal caregivers.
battery (DTB). Participants were adults with DS living in New York (N = 29; average age 46 years). We asked three questions. 1. Does having any type of heart disease affect the association between DTB scores and chronological age? 2. Does thyroid status affect the association between heart disease and DTB scores? 3. Are the E4 or E2 alleles of apolipoprotein E (APOE) associated with DTB scores or with heart disease? The study was retrospective, pilot, and exploratory. It involved analysis of information in a database previously established for the study of aging in DS. Participants had moderate intellectual disability on average. Information for each person included: sex, age, a single DTB score obtained by combining results from individual subscales of the DTB, the presence or absence of heart disease, thyroid status (treated hypothyroidism or normal), and APOE genotype. Trends were visualized by inspection of graphs and contingency tables. Statistical methods used to evaluate associations included Pearson correlation analysis, Fisher’s exact tests (2-tailed), and odds ratio analysis. P values were interpreted at the 95% confidence level without Bonferroni correction. P values > 05<1 were considered trends. The negative correlation between DTB scores and age was significant in those with heart disease but not in those without. Heart disease was significantly associated with DTB scores >1 SD below the sample mean; there was a strong association between heart disease and low DTB scores in those with treated hypothyroidism but not in those with normal thyroid status. The APOE genotype was weakly associated with heart disease (E4, predisposing; E2, protective) in males. On the basis of the potentially important findings from the present study, large prospective studies are warranted to confirm and extend the observations. In these, particular heart conditions or diseases and other medical comorbidities in individuals should be documented.


Abstract: Down syndrome (DS) occurs due to triplication of chromosome 21. Individuals with DS face an elevated risk for development of Alzheimer’s disease (AD) due to increased amyloid beta (Aβ) resulting from the over-expression of the amyloid precursor protein found on chromosome 21. Diagnosis of AD among individuals with DS poses particular challenges resulting in an increased focus on alternative diagnostic methods such as blood-based biomarkers. The aim of this review was to evaluate the current state of the literature of blood-based biomarkers found in individuals with DS and particularly among those also diagnosed with AD or in prodromal stages (mild cognitive impairment (MCI)). A systematic review was conducted utilizing a comprehensive search strategy. Twenty-four references were identified, of which 22 fulfilled inclusion criteria (E2, protective) in males. On the basis of the potentially important findings from the present study, large prospective studies are warranted to confirm and extend the observations. In these, particular heart conditions or diseases and other medical comorbidities in individuals should be documented.


Abstract: The management of dementia in Alzheimer’s disease has dramatically changed since the development of anti-dementia drugs. However, there is limited information available regarding the bio-medical aspects of the differing types of dementia in adults with Down syndrome. The review highlights the important pharmacological and clinical aspects of donepezil, rivastigmine, galantamine and memantine and supports the view that such drugs play an important part in the management of dementia in adults with intellectual disability. Future clinical and research issues are discussed.


Abstract: An association between Down syndrome and dementia type is well established. This study demonstrates that late onset seizures in people with Down syndrome are a strong indicator of a dementing process. Further, late onset seizures in people with Down syndrome may be used as a prognostic indicator, indicating life expectancy of less than a further 2 years, probable death within 3 years and death almost invariably within 5 years of onset.


Abstract: The diagnosis of dementia in Alzheimer’s disease remains at times problematic in adults with intellectual disability. The analysis of 5-year consecutive data developed a researched-based clinical screening tool for
of dementia, treating any reversible additional factors, consideration of but remain limited. The management of DAD in the DS population can be at symptoms, and sleep difficulty. Antidementia medications have a role to play The management of dementia is holistic. This includes treating reversible literature review aimed to look at the management of DAD in people with DS. make the management of DAD more challenging in older adults with DS. This with intellectual disabilities. However, the underlying intellectual disability can treatment of DAD is not too dissimilar in the general population and in people with Down syndrome (DS), with characteristically an earlier onset. The Developmental Disabilities in Ontario 4th Ed., Toronto, Ontario, Canada: Delphi Graphic Communications. https://oadd.org/publications/textbook/ Abstract: Chapter provides an introduction to the topic of dementia in persons with developmental disabilities. Dementia, as a worldwide public health concern, is increasing in prevalence markedly because the world’s population is living longer and aging in greater numbers. Chapter 4 covers the physiology of dementia, options for services, mechanisms for multidisciplinary management, and advances in advocacy, dementia prevention, and dementia research. Proveda, B., & Broxholme, S. Assessments for dementia in people with learning disabilities: Evaluation of a dementia battery developed for people with mild to moderate learning disabilities Learning Disability Practice, 2016, 19(1), 31-40. doi.org/10.7748/lidp.19.1.31.s23 Abstract: An intellectual disabilities’ dementia battery was developed to assess cognitive abilities in individuals referred to the intellectual disabilities service because of concerns of possible dementia. The present study aimed to establish concurrent validity with previously validated measures of cognitive ability and its clinical effectiveness in detecting dementia in this population. Fifty-five individuals aged 29 and over (range: 29 to 71), received a baseline and a follow-up assessment using the dementia battery between 2000 and 2010. Fifty-five individuals aged 29 and over (range: 29 to 71), received a baseline and a follow-up assessment using the dementia battery between 2000 and 2010. Differences in performance between individuals allocated to ‘probable’, ‘unsure’ and ‘no’ dementia groupings were investigated at domain and subtest level, as well as overall performance. Results on the battery were compared with clinically relevant measures of dementia also included in the local assessment protocol. Significant differences in overall performance were found between the ‘probable’ and ‘no’ dementia groups as well as cognitive domain-specific differences. No differences were found at subtest level. Good concurrent validity was found between the battery and comparable measures of change within the dementia assessment protocol, namely the VABS, DMR and BPIVS II. The intellectual disabilities’ dementia battery appears to be a good measure, which can be used longitudinally, to detect change in individuals and help establish a psychological and behavioral management, use of antidementia medication, and a multidisciplinary team approach.
diagnosis of dementia. It is also comparable with other measures of change incorporated in the dementia assessment protocol. Subtests included in the language domain appear to be the most relevant at detecting significant changes between baseline and follow up. Future studies should attempt to standardize this measure and establish cut-off scores.

**PTAC [PASRR Technical Assistance Center]**

How does a categorical determination for dementia and intellectual disability affect the PASRR process? **PTAC, April 10, 2018;**

https://www.pasrrassist.org/resources/How-does-a-categorical-determination-for-dementia-and-intellectual-disability-affect-the-PASRR-process%3F

Abstract: 42 CFR 483.130(h) provides that the State intellectual disability authority may make categorical determinations that individuals with dementia, which exists in combination with an intellectual disability (ID) or a related condition, do not need specialized services. A categorical determination for dementia and ID can be applied at the Level I screening, but the categorical must be determined at the Level II evaluation phase of PASRR. Beyond the specialized services determinations, there is no basis for ending a Level II evaluation for an individual with an intellectual disability diagnosis, as the evaluation still determines if nursing facility services are needed. Under 42 CFR § 483.126(j), findings must be issued in the form of an abbreviated written evaluative report which—(1) Identifies the name and professional title of the person applying the categorical determination and the data on which the application was made; (2) Explains the categorical determination(s) that has (have) been made and, if only one of the two required determinations can be made categorically, describes the nature of any further screening which is required; (3) Identifies, to the extent possible, based on the available data, NF services, including any mental health or specialized psychiatric rehabilitative services, that may be needed; and (4) Includes the bases for the report's conclusions. A categorical determination for dementia and intellectual disability also requires the issuing of a written determination notice per the following 483.130(k)(l) requirements: (k) Notice of determination. The State mental health or intellectual disability authority must notify in writing the following entities of a determination made under this subpart: 1. The evaluated individual and legal representative; 2. The admitting or retaining NF; 3. The individual or resident's attending physician; and 4. The discharging hospital, unless the individual is exempt from predetermination screening as provided at §483.106(b)(2). (l) Contents of notice. Each notice of the determination made by the State mental health or intellectual disability authority must include: 1. Whether a NF level of service is needed; 2. Whether specialized services are needed; 3. The placement options that are available to the individual consistent with these determinations; and 4. The rights of the individual to appeal the determination made. **Puri, B.K., Ho, K.W., & Singh, I.**


Abstract: In a cohort of 68 adults (35 males and 33 females) with Down's syndrome aged 29-83 years, a history of seizures was found in 26.5%. The overall mean age of onset of seizures was 37 years, males (22 years) being significantly younger than females (51 years). The age of onset was bimodally distributed, with the first peak occurring in the first two decades, and a late-onset peak occurring in the fifth and sixth decades. A strong association between Alzheimer's disease and seizures was confirmed. Of those with a history of seizures, those aged over 45 years were significantly more likely to develop Alzheimer's disease than those younger than 45. It is suggested that late-onset epilepsy in Down's syndrome is associated with Alzheimer's disease, while early-onset epilepsy is associated with an absence of dementia.


Language skills as a predictor of cognitive decline in adults with Down syndrome. *Alzheimers Dement (Amst)*, 2020, Aug 25, 12(1), e12080. doi: 10.1002/dad2.12080. eCollection 2020. Abstract: Adults with Down syndrome (DS) are at high risk for early onset Alzheimer's disease (AD), characterized by a progressive decline in multiple cognitive domains including language, which can impact social interactions, behavior, and quality of life. This cross-sectional study examined the relationship between language skills and dementia. A total of 168 adults with DS (mean age = 51.4 years) received neuropsychological assessments, including Vineland Communication Domain, McCarthy Verbal Fluency, and Boston Naming Test, and were categorized in one of three clinical groups: cognitively stable (CS, 57.8%); mild cognitive impairment (MCI-DS, 22.6%); and probable/definite dementia (AD-DS, 19.6%). Logistic regression was used to determine how well language measures predict group status. Vineland Communication, particularly receptive language, was a significant predictor of MCI-DS. Semantic verbal fluency was the strongest predictor of AD-DS. Assessment of language skills can aid in the identification of dementia in adults with DS. Clinically, indications of emerging language problems should warrant further evaluation and monitoring.

Rafii, M.S.


Abstract: Alzheimer's disease (AD) pathology and early-onset dementia develop almost universally in Down syndrome (DS). AD is defined neuropathologically by the presence of extracellular plaques of aggregated amyloid ß protein and intracellular neurofibrillary tangles (NFTs) of aggregated hyperphosphorylated tau protein. The development of radiolabeled positron emission tomography (PET) ligands for amyloid plaques and tau tangles enables the longitudinal assessment of the spatial pattern of their accumulation in relation to symptomatology. Recent work indicates that amyloid pathology develops 15–20 years before neurodegeneration and symptom onset in the sporadic and autosomal dominant forms of AD, while tau pathology correlates more closely with symptomatic stages evidenced by cognitive decline and dementia. Recent work on AD biomarkers in DS illustrates similarities between DS and sporadic AD. It may soon be possible to apply recently developed staging classifications to DS to obtain a more nuanced understanding of the development AD in DS and to provide more accurate diagnosis and prognosis in the clinic.

Rafii, M.S.


Abstract: Individuals with Down syndrome (DS) are at high risk for developing Alzheimer's disease (AD) pathology and this has provided significant insights into our understanding of the genetic basis of AD. The present review summarizes recent clinical, neuropathologic, imaging, and fluid biomarker studies of AD in DS (DSAD), highlighting the striking similarities, as well as some notable differences, between DSAD and the more common late-onset form of AD (LOAD) in the general population, as well as the much rarer, autosomal-dominant form of AD (ADAD). There has been significant progress in our understanding of the natural history of AD biomarkers in DS and their relationship to clinically meaningful changes. Additional work is needed to clearly define the continuum of AD that has been described in the general population, such as the preclinical, prodromal, and dementia stages of AD. Multiple therapeutic approaches, including those targeting not only ß-amyloid but also tau and the amyloid precursor protein itself, require consideration. Recent developments in the field are presented within the context of such efforts to conduct clinical trials to treat and potentially prevent AD in DS.

Rafii, M., & Santoro, S.L.


Abstract: The median life expectancy for a child with Down syndrome (DS) born in the 1950s was less than 10 years of age, with congenital heart defects being the main cause of death. With advances in medical care and improvements in the overall health of individuals with DS, life expectancy has increased dramatically; for children with DS born in 2010, median life expectancy is estimated to be 65 years. However, along with this longer lifespan comes the prospect of a considerable increase in the risk of developing dementia associated with Alzheimer disease (AD), with a prevalence of nearly 80% for those with DS who are older than 65 years.

Rafii, M.S., Ances, B.M., Schupf, N., Krinsky-Mchale, S.J., Mapstone, M., Silverman, W., Lott, I. ... O’Bryant, S.

Abstract: The National Institute on Aging in conjunction with the Alzheimer’s Association (NIA-AA) recently proposed a biological framework for defining the Alzheimer’s disease (AD) continuum. This new framework is based upon the key AD biomarkers (amyloid, tau, neurodegeneration, AT(N)) instead of clinical symptoms and represents the latest understanding that the pathological processes underlying AD begin decades before the manifestation of symptoms. By using these same biomarkers, individuals with Down syndrome (DS), who are genetically predisposed to developing AD, can also be placed more precisely along the AD continuum. The AT(N) framework is therefore thought to provide a more objective method by which to select and enrich samples for clinical trials. This new framework is highly flexible and allows the addition of newly confirmed AD biomarkers into the existing AT(N) groups. As biomarkers for other pathological processes are validated, they can also be added to the AT(N) classification scheme, which will allow for better characterization and staging of AD in DS. These biological classifications can then be merged with clinical staging for an examination of factors that impact the biological and clinical progression of the disease. Here, we leverage previously published guidelines for the AT(N) framework to generate such a plan for AD among adults with DS.


Abstract: To gain further knowledge on the preclinical phase of Alzheimer’s disease (AD), we sought to characterize cognitive performance, neuroimaging and plasma-based AD biomarkers in a cohort of non-demented adults with Down syndrome (DS). The goal of the down syndrome biomarker Initiative (DSBI) pilot is to test feasibility of this approach for future multicenter studies. We enrolled 12 non-demented participants with DS between the ages of 30-60 years old. Participants underwent extensive cognitive testing, volumetric MRI, amyloid positron emission tomography (PET; 18F-florbetapir), fluorodeoxyglucose (FDG) PET (18F-fluorodeoxyglucose) and retinal amyloid imaging. In addition, plasma beta-amyloid (Aβ) species were measured and Apolipoprotein E (APOE) genotyping was performed. Results from our multimodal analysis suggest greater hippocampal atrophy with amyloid load. Additionally, we identified an inverse relationship between amyloid load and regional glucose metabolism. Cognitive and functional measures did not correlate with amyloid load in DS but did correlate with regional FDG PET measures. Biomarkers of AD can be readily studied in adults with DS as in other preclinical AD populations. Importantly, all subjects in this feasibility study were able to complete all test procedures. The data indicate that a large, multicenter longitudinal study is feasible to better understand the trajectories of AD biomarkers in this enriched population.

Rafii, M.S., Zaman, S., & Handen, B.L.


Abstract: The NIH-funded Alzheimer’s Biomarker Consortium Down Syndrome (ABC-DS) and the European Horizon 21 Consortium are collecting critical new information on the natural history of Alzheimer’s Disease (AD) biomarkers in adults with Down syndrome (DS), a population genetically predisposed to developing AD. These studies are also providing key insights into which biomarkers best represent clinically meaningful outcomes that are most feasible in clinical trials. This paper considers how these data can be integrated in clinical trials for individuals with DS. The Alzheimer’s Clinical Trial Consortium - Down syndrome (ACTC-DS) is a platform that brings expert researchers from both networks together to conduct clinical trials for AD in DS across international sites while building on their expertise and experience.

Raj, S., Stanley, M., Mackintosh, S., & Fryer, C.


Abstract: Dementia in adults with Down syndrome causes a progressive decline in daily occupations impacting both persons with Down syndrome and their informal caregivers. This study aimed to explore the scope of occupational therapy practice for adults with both Down syndrome and dementia and their informal caregivers living in their homes. A survey was conducted with occupational therapists having clinical experience in providing interventions for adults with Down syndrome. A web-based survey was developed to explore occupational therapy practice for this group of people with Down syndrome and their informal caregivers. Responses to closed-ended questions were analysed descriptively, and inductive content analysis was used for open-ended questionnaire responses. Therapists from eight sites across the United Kingdom and the United States of America participated in the survey. Two-thirds were from the United Kingdom, most of whom were employed in the public sector and had at least 10 years of clinical experience. Over 90% of respondents received one or more referrals in a typical month for adults with Down syndrome having dementia, 68% of which were for a decline in activities of daily living. Home environment and activities of daily living were frequently assessed areas, and the commonest interventions were compensatory strategies and environmental modifications. Only half the respondents provided interventions for informal caregivers. Risk and safety and manual handling were frequently addressed domains. Collaboration and developing clinical expertise were the two key perceived enablers for providing effective occupational therapy services. Fragmentation of services and a lack of client-centred care were the common perceived barriers. Occupational therapists often address decline in activities of daily living for individuals with both Down syndrome and dementia. To support participation in meaningful occupations for these people and support the needs of their informal caregivers, it is essential that services are offered in a collaborative approach.

Raj, S.E., Mackintosh, S., Kernot, J., Fryer, C., & Stanley, M.


Abstract: This paper describes the development of a home-based occupational therapy intervention program for people with Down syndrome who experience early on-set dementia causing a decline in their performance skills and increasing care dependency on their informal caregivers. A six-step methodological process adapted from the Medical Research Council framework for developing and evaluating complex interventions was formulated to develop an evidence-based occupational therapy program for people with both Down syndrome and dementia and their informal caregivers. The first two steps gathered evidence through systematic reviews of the literature and determined the scope of current occupational therapy practice. The gathered evidence was synthesised in step three to develop a client-centred occupational therapy intervention program for persons with both Down syndrome and dementia and their informal caregivers. In steps four and five, opinions were sought from occupational therapists working in this area of practice on the content of the developed program and its feasibility within the Australian disability services context. The final testing step can be conducted in the future using a single-case experimental design study. It is important to use rigorous frameworks and gather comprehensive evidence using multiple methods to develop interventions for small heterogeneous populations. The developed occupational therapy program for persons with both Down syndrome and dementia and their informal caregivers appears feasible to be implemented within the Australian disability services; however, funding limitations imposes barriers for its implementation in clinical practice.

Rebillett, A-S., Hiance-Delahaye, A., Falguero, S., Radice, G., & Sacco, S.


Abstract: People with Down Syndrome (DS) are at an increased risk of developing Alzheimer’s Disease (AD) relatively early in life. The dementia screening questionnaire for individuals with intellectual disabilities (DSQID) has been developed for people with intellectual disabilities and was shown to have
high discriminative power to distinguish between people with and without dementia. The objective of this study was to verify if the French version of the DSQIID (DSQIID-F) had a good diagnostic specificity and to determine the optimal cut-off for screening people with DS for dementia. This was a single-centre, retrospective, medical chart review study in people with DS aged ≥40 years. Demographics, level of intellectual disability, DSQIID-F data and clinical assessment of dementia were extracted from medical records. Sensitivity and specificity for different DSQIID-F cut-offs were calculated to determine the optimal cut-off. Some 151 people with DS were included with a median age of 51 years. The optimal DSQIID-F cut-off was 19, sensitivity was 0.940 (95% CI: 0.830; 0.985) and specificity was 0.941 (95% CI: 0.873; 0.975). Results were comparable to those for the English DSQIID (cut-off: 20; sensitivity: 0.92; specificity: 0.97). However, the psychometric qualities of the DSQIID-F, used for clinical follow-up, have not been verified. The DSQIID-F has good discriminative power and represents a useful tool to screen people with DS for dementia

Reppermund, S., & Trollor, J.N.
Successful ageing for people with an intellectual disability
Abstract: Successful ageing has not yet been defined in people with an intellectual disability. The purpose of this review is to discuss and define successful ageing in the context of intellectual disability and to propose strategies to improve health and wellbeing for this population. People with an intellectual disability experience higher rates of diabetes, hypertension, obesity and cardiovascular disease, and higher rates of mental disorders than people without an intellectual disability. People with an intellectual disability engage in more passive leisure activities because many active leisure activities require the participation of or assistance by others. Health promotion programmes tailored to people with an intellectual disability consisting of exercise and health education can result in more positive attitudes toward exercise and improvements in psychosocial outcomes. With modifications for people with an intellectual disability, the concept of successful ageing can be used as a template for development of strategies to improve health and wellbeing for people with an intellectual disability as they age. Targeted programmes focused on health promotion and prevention of age-related morbidities is required. There is a need for policies addressing positive ageing, including social participation and maximizing community participation. Appropriate and ongoing education for people with an intellectual disability and their carers on healthy living in areas of physical, social, and cognitive activity, nutrition and avoidance of risk factors is essential.

Reid, A. H., & Aungle, P. G.
Abstract: Review of literature on dementia and Down syndrome to date [no published abstract]

Robertson, J., Hatton, C., Emerson, E., Baines, S.
Prevalence of epilepsy among people with intellectual disabilities: A systematic review.
Seizure, 2015, 29, 46-62. doi: 10.1016/j.seizure.2015.03.016
Abstract: Epilepsy is more common in people with intellectual disabilities than in the general population. However, reported prevalence rates vary widely between studies. This systematic review aimed to provide a summary of prevalence studies and estimates of prevalence based on meta-analyses. Studies were identified via electronic searches using Medline, Cinahil and PsycINFO and cross-citations. Information extracted from studies was tabulated. Prevalence rate estimates were pooled using random effects meta-analyses and subgroup analyses were conducted. A total of 48 studies were included in the tabulation and 46 studies were included in meta-analyses. In general samples of people with intellectual disabilities, the pooled estimate from 38 studies was 22.2% (95% CI 19.6-25.1). Prevalence increased with increasing level of intellectual disability. For samples of people with Down syndrome, the pooled estimate from data in 13 studies was 12.4% (95% CI 9.1-16.7), decreasing to 10.3% (95% CI 8.4-12.6) following removal of two studies focusing on older people. Prevalence increased with age in people with Down syndrome and was particularly prevalent in those with Alzheimer’s/dementia. Epilepsy is highly prevalent in people with intellectual disabilities. Services must be equipped with the skills and information needed to manage this condition.

Robinson, A., Spencer, B., & White, L.
Understanding difficult behaviors: Some suggestions for coping with Alzheimer’s disease and related illnesses
Abstract: In this study, we examined normative data and diagnostic accuracy of a pictorial screening test to detect memory impairment for mild cognitive impairment (MCI) and Alzheimer’s disease (AD) in Spanish-speaking adults with intellectual disability (ID). A total of 94 volunteers with ID (60 controls, 17 MCI, and 17 AD), were evaluated by neuropsychological tests including the PmIS-ID in a cross-sectional validation study. Discriminative validity between the MCI, AD, and control group was analyzed by the area under the ROC curve. A cut-off score of 4.5 on the immediate recall trial had a sensitivity of 69% and a specificity of 80% to detect memory impairment (AUC = 0.685; 95% CI = 0.506-0.863) in the AD group. The PmIS-ID is a useful screening test to rule out a diagnosis of memory decline in people with moderate level of ID and AD, and it shows good psychometric properties.

Rodríguez-Hidalgo E, García-Alba J, Buxó M, Novell R, Estebá-Castillo S.
Abstract: The aim of this study is to adapt and validate the global deterioration scale (GDS) for the systematic tracking of Alzheimer’s disease (AD) progression in a population with Down syndrome (DS). A retrospective dual-center cohort study was conducted with 83 participants with DS (46.65 ± 5.08 years) who formed the primary diagnosis (PD) group: cognitive stability (n = 48), mild cognitive impairment (n = 24), and Alzheimer’s disease (n = 11). The proposed scale for adults with DS (GDS-DS) comprises six stages, from cognitive and/or behavioral stability to advanced AD. Two neuropsychologists placed the participants of the PD group in each stage of the GDS-DS according to cognitive, behavioral and daily living skills data. Inter-rater reliability in staging with the GDS-DS was excellent (ICC = 0.86; CI: 0.80-0.93), and the agreement with the diagnosis categories of the PD group ranged from substantial to excellent with k values of 0.82 (95% CI: 0.73-0.92) and 0.85 (95% CI: 0.72. 0.99). Performance with regard to the CAMCOG-DS total score and orientation subtest of the Barcelona test for intellectual disability showed a slight progressive decline across all the GDS-DS stages. The GDS-DS scale is a sensitive tool for staging the progression of AD in the DS population, with special relevance in daily clinical practice.

Ross, W.T., & Olsen, M.
Care of the adult patient with Down syndrome
doi:10.14423/SMJ.0000000000000193
Abstract: Individuals with Down syndrome have an increased risk for many conditions, including cardiovascular disease, cancer, infections, and osteoporosis, and endocrine, neurological, orthopedic, auditory, and ophthalmic
disorders. They also are at increased risk for abuse and human rights violations and receive fewer screenings and interventions than the population without Down syndrome. In this literature review, the most common health conditions associated with Down syndrome are examined, along with the topics of sexual abuse, menstrual hygiene, contraception, and human rights. Clinical guidelines for this population are summarized in an effort to assist practicing physicians in improving their provision of health care to the adult patient with Down syndrome.


Abstract: Assessment of age-associated disorders has become increasingly important. In a clinical setting, people with intellectual disability with and without dementia were assessed retrospectively using the Neuropsychological Test Battery (NTB) and the Dementia Questionnaire for People with Learning Disabilities (DLD) at two different times to analyse neuropsychological changes and diagnostic validity. One group (n = 44) was assessed with both instruments, while the DLD was applied in 71 patients. In the NTB (n = 44), only patients with dementia (n = 26) showed a decrease in the NTB total score and three subscales. Receiver operating characteristic analysis revealed a diagnostic sensitivity of .67, a specificity of .81, and an area under the curve (AUC) of .767. In the DLD group (n = 71), only those with dementia displayed a decrease in the cognitive and social scale; diagnostic sensitivity and specificity values were low (.61/63) and the AUC was .704. Neuropsychological assessment was sensitive to detect cognitive changes over time. Sensitivity values of both instruments suggest a reassessment at a later time point.


Abstract: This article provides a timely response in regard to the UK’s Department of Health’s current initiative to financially reward general practitioners (GPs) to prioritise and undertake dementia screening for people with learning disabilities over the age of 50 years and for people with Down syndrome over the age of 40 years. Whilst GPs are becoming increasingly aware of their responsibility to care for the complex needs of people with learning disabilities, the implementation of dementia screening poses a multitude of challenges. Research has continued to suggest how difficult it is to detect the early and often ambiguous signs of dementia for someone who has pre-existing cognitive impairments and may present with atypical symptomology. And it continues to be a difficult process even for those who specialise within this area. However, GPs who choose to opt into this financially incentivised scheme will now be offering dementia screening. This article outlines the main GP aims within the dementia screening process and the difficulties that may be encountered, with specific focus upon (1) offering a dementia screen, (2) obtaining consent, (3) undertaking screening within the most appropriate setting, (4) choosing a dementia screening tool and developing a baseline, and (5) detecting early dementia signs.


Abstract: We describe prevalence and incidence of dementia and AD in DS in a full Medicaid population of adults with DS in Wisconsin from 2008 through 2018. We assessed Medicaid claims for adults (>21 years) who ever had 2 DS claims over their lifetime (based on International Classification of Diseases, Ninth Revision and Tenth Revision codes) on 2 separate days during Medicaid enrollment. Dementia claims were extracted from codes for any dementia (with AD as a subset) from the Centers for Medicare & Medicaid Services Chronic Conditions Data Warehouse. We required 3 or more years of Medicaid enrollment for adults with DS to ensure validity of dementia claims, therefore, beneficiaries entered the cohort at any point between 2008 and 2015. We categorized age at first and last claims (<40 years, 40-54 years, and ≥55 years) to account for confounding by age. A total of 2,968 individuals were included, of whom 1,507 (51.8%) were male. The median (interquartile range) age at first claim was 39 (25-48) years. In the category of individuals aged 55 years or older, 490 of 938 had dementia claims (52.2%), 307 of 938 had AD claims (32.7%), and dementia incidence was 102 (95% CI, 87-119) cases per 1000 person-years. Among individuals aged 40 to 54 years, 190 of 1013 had dementia claims (18.8%), and dementia incidence was 49 (95% CI, 44-53) cases per 1000 person-years. The probability of an incident dementia claim was 40% (95% CI, 41%-47%) over 11 years of enrollment for adults with DS who were aged 40-54 years at cohort entry and 67% (95% CI, 60%-74%) for those 55 years and older at cohort entry (Figure). There were no sex differences for dementia among individuals younger than 40 years (prevalence ratio, 1.07 [95% CI, 0.63-1.81]) or among those 55 years and older (prevalence ratio, 0.94 [95% CI, 0.69-1.29]). Dementia prevalence was higher in female individuals than male individuals aged 40 to 54 years (prevalence ratio, 1.23 [95% CI, 1.02-1.50]). Findings from a statewide health system confirm that both dementia and AD in individuals with DS present in claims data at rates similar to those ascertained from clinical samples. The hypothesized causative mechanism and similar eligibility requirements between state Medicaid programs for people with DS likely mean that other state Medicaid systems experience high incidence and prevalence of dementia and AD in individuals with DS. Dementia and AD prevalence and incidence in Medicaid beneficiaries with DS highlight the need to identify prodromal presentations and develop dementia services and supports for DS as they age and continue to rely on Medicaid and Medicaid-funded assisted living or skilled nursing facilities.


Abstract: In keeping with worldwide demographic changes and an ageing population, people with intellectual disabilities are living longer and all the evidence suggests that this trend will continue. This ‘new’ population of older people and their carers will pose challenges for health and social care providers. This paper presents a review of the literature on key issues influencing caregiving for older people with intellectual disabilities and their ageing family carers. The review was undertaken using a framework adapted from the NHS Centre for Reviews and Dissemination. Papers were identified through the use of databases including CINAHL, ScienceDirect, PsychInfo, Blackwell Synergy, the Cochrane Library and MEDLINE. The key themes which emerged from the literature and which consequently form the basis of this review include: ageing family carers, future planning and support services. In the context of family caregiving, older people with intellectual disabilities represent a unique group insofar as they are unlikely to be married and therefore have no spouse or dependents to care for them in later life. As a result, parents (usually mothers) have to continue caring for their son or daughter with an intellectual disability as they both grow older, often resulting in a mutually dependent relationship. The caregiving situation is further complicated by poor emergency and future planning and by a lack of appropriate services for this group of individuals. In light of the emergence of a ‘new’ population of older people with intellectual disabilities, there is an urgent need to develop services and support structures which will enable these individuals and their ageing carers to ‘age in place’ and when this is no longer possible, to have appropriate alternatives that recognise the duality of their needs as older people and as people with intellectual disabilities.


Abstract: Living with dementia is challenging, but poses unique difficulties for adults with an intellectual disability. The demands of dementia are also challenging for family, carers, and friends. The authors explored the impact of dementia on direct care staff using a focus group methodology. Thematic analysis was used to investigate the staff narratives. There were four key themes that emerged: (a) the difficulty of recognizing symptoms of dementia in people with intellectual disability, (b) the process of diagnosis, (c) the
of dementia for the person, (d) the emotional impact of dementia for other people. The authors concluded that the themes identified a number of important potential targets for supporting staff and peers when dementia is present in an adult with an intellectual disability.


Abstract: This article provides an overview of the current literature on person-centered planning, health action planning and associated concepts such as facilitation and advocacy. It explores how flexible approaches to person-centered planning and health action planning were introduced and adapted to meet the individual daily changing needs of Anna, an older person with Down syndrome and dementia.


Abstract: Little is known of paid carers’ perspectives when caring for people with intellectual disabilities at the end-of-life. Sixty four individuals from intellectual disability services took part in 12 focus groups. Interviews were analysed using framework analysis. Participants wanted to provide palliative care and felt the experience enriched practice. However, they were inadequately prepared to meet need and this often led to staff stress. A number of issues appeared to heighten stress: situations when end-of-life care decision making was challenging, when staff felt ‘pushed out’ by relatives and when staff did not have sufficient support or time to provide care or mourn the loss of service users. The study describes issues which contribute to the development of staff stress when providing palliative care and draws attention to areas where strategies should be developed in order to improve the quality of care provided to people with intellectual disabilities.


Abstract: Approximately 40-80% of persons with Down syndrome (DS) develop Alzheimer's disease (AD)-like dementia by the fifth to sixth decade of life, a much younger age than is typically seen in sporadic AD. The onset of dementia symptoms in DS parallels the development of classic brain neuropathological lesions (i.e., amyloid plaques) similar to that evident in AD. Both disorders appear to have a similar genetic linkage, which is supported by the triplication of the gene that codes for amyloid beta (A4) precursor protein (APP) in persons with DS and an extra copy of the APP gene causes familial AD in persons without DS. Despite an overlap in the genetics of these disorders, the clinical presentation of dementia differs between persons with DS and AD. Whereas 'forgetfulness' is a typical symptom of early-phase dementia for persons in the general population, behavioral problems and personality changes are early signs of dementia for persons with DS. There are indications that amyloid burden begins in the frontal cortex before spreading to other brain regions in those with DS-AD, something that is not always the case in sporadic AD suggesting pathological variances between these disorders. These differences beg the question: Are the genetic and neuropathological commonalities found in DS- and AD-related dementia an associated similarity or do these disorders share a common pathogenesis? To address this query, we briefly review the clinical, histopathological, and genetic research supporting a putative link between dementia in DS and AD.


Abstract: In developed countries the population of elderly people with Down syndrome expands resulting in an increasing incidence of age-related diseases, including dementia. The assessment of dementia in individuals with intellectual disability is often complicated due to large intra-individual variability in cognitive functioning prior to dementia and to lack of standardised measures to detects dementia. Structured observations of symptoms of dementia and assessment techniques tailored for people with intellectual disability are increasingly needed.


Abstract: Specialty centers improve care for patients with Down syndrome. The cohort of adults with Down syndrome is increasing, but the capacity for specialty centers to meet their medical care needs is unknown. Electronic survey of staff of specialty clinics for adults with Down syndrome was conducted. Review of online clinic list ings, and calculation of the number of adults with Down syndrome were performed. Analysis identified the percent of adults with Down syndrome who could have their medical care needs met in a current specialty clinic. Fourteen specialty clinics report providing care for 4038 adults with Down syndrome. Respondents reported gaps in care including: limitations of existing clinics, need for additional clinics, and know edgeable health professionals in Down syndrome. Survey-respondent clinic capacity would meet needs of 3% of adults with Down syndrome. Twenty-five clinics for adults with Down syndrome were listed online with capacity to care for 6517 adults with Down syndrome meeting the needs of adults with Down syndrome. Sixty-four individuals from national specialty clinics agreed to meet the needs of adults with Down syndrome. Survey of existing clinics provides guidance to create additional clinics, including: must-have team members, current sources of clinic financial support, and gaps in current clinical care.


Abstract: In October of 2016, an interdisciplinary group representing North and South American and European countries met in Glasgow, Scotland, to scrutinize universal issues regarding adults with intellectual disability (ID) affected by dementia and to produce recommendations and guidelines for public policy, practice, and further research. The aim of this paper is to apprise relevant outcomes of the Summit targeting Brazilian researchers, clinicians, and nongovernmental organizations in the field of ageing and dementia that are committed to developing the Brazilian national dementia plan. Three core themes were covered by the Summit: i) human rights and personal resources, ii) personalized services and caregiver support, and iii) advocacy and public impact. The exploration of the themes highlighted variations across countries, and revealed consensus views on matters such as international networks, guidance for practices, and advocacy on behalf of both people with ID affected by dementia, and their families. The authors outline the challenges Brazil must confront regarding ageing and dementia and proffer recommendations to address the needs of adults with ID affected by dementia within this scenario; both of which would help in developing the Brazilian national dementia plan.


Abstract: Intellectual disability (ID) and dementia reflect disturbed cortical function during and after developmental age, respectively. Due to the wide heterogeneity of ID population the decline in cognitive and adaptive skills may be different in distinct genetic subgroups. Using the British Present Psychiatric State-learning Disabilities assessment (PPS-LD) questionnaire the dementia signs were screened in 62, 22 and 44 individuals (> 35 year of age) with Down (DS, OMIM number 190685), Williams (WS, OMIM number, 194050), and Fragile X syndrome (FXS, 6% of the population. Additional clinic capacity is needed to meet the needs of adults with Down syndrome. Survey of existing clinics provides guidance to create additional clinics, including: must-have team members, current sources of clinic financial support, and gaps in current clinical care.
Dementia that was confirmed by brain imaging, manifested as Alzheimer's disease and as moyamyat disorder associated with vascular dementia in DS and as vascular dementia in WS. This survey suggests that the risk of dementia varies depending on the cause of ID and that the severity of ID in adolescence does not predict the development of dementia at a later age. Consequently, the ID and dementia should be understood as separate clinical entities that need to be taken into account in the health management of intellectually disabled people. This is important for the arrangement of appropriate and timely interventions, which can be expected to delay the need for institutionalization.

Schaap, F.D., Dijkstra, G.J., Finnema, E.J., & Reijneveld, S.A.
The first use of dementia care mapping in the care for older people with intellectual disability: a process analysis according to the RE-AIM framework Aging & Mental Health, 22(7), 912-919. DOI: 10.1080/13607453.2017.1401582
Abstract: The ageing of the population with intellectual disability (ID), with associated consequences as dementia, creates a need for evidence-based methods to support staff. Dementia Care Mapping (DCM) is perceived to be valuable in dementia care and promising in ID-care. The aim of this study was to evaluate the process of the first use of DCM in ID-care. DCM was used among older people with ID and care-staff in 12 group homes of six organisations. We obtained data on the first use of DCM in ID-care via focus-group discussions and face-to-face interviews with: care-staff (N = 24), managers (N = 10), behavioural specialists (N = 7), DCM-ID mappers (N = 12), and DCM-trainers (N = 2). We used the RE-AIM framework for a thematic process-analysis. All available staff (94%) participated in DCM (reach). Regarding its efficacy, staff considered DCM valuable; it provided them new knowledge and skills. Participants intended to adopt DCM, by continuing and expanding its use in their organisations. DCM was implemented as intended, and strictly monitored and supported by DCM-trainers. As for maintenance, DCM was further tailored to ID-care and a version for individual ID-care settings was developed, both as standards for international use. To sustain the use of DCM in ID-care, a multidisciplinary, interorganisational learning network was established. DCM tailored to ID-care proved to be an appropriate and valuable method to support staff in their work with ageing clients, and it allows for further implementation. This is a first step to obtain an evidence-based method in ID-care for older clients.

Schaap, F.D., Dijkstra, G.J., Stewart, R.E., Finnema, E.J., & Reijneveld, J.S.A.
Abstract: The ageing of people with intellectual disability, accompanied with consequences like dementia, challenges intellectual disability-care staff and creates a need for supporting methods, with Dementia Care Mapping (DCM) as a promising possibility. This study examined the effect of DCM on the quality of life of older people with intellectual disability. We performed a quasi-experimental study in 23 group homes for older people with intellectual disability in the Netherlands, comparing DCM (n = 113) with care-as-usual (CAU; n = 111). Using three measures, we assessed the staff-reported quality of life of older people with intellectual disability. DCM achieved no significantly better or worse quality of life than CAU. Effect sizes varied from 0.01 to -0.22. Adjustment for covariates and restriction of analyses to people with dementia yielded similar results. The finding that DCM does not increase quality of life of older people with intellectual disability contradicts previous findings and deserves further study.

Schaap, F.D., Finnema, E.J., Dijkstra, G.J., & Reijneveld, M.
What can we learn from dementia care in the care of older people with intellectual disability? Journal of Intellectual Disability Research, 2019, 63(8), 645-646.
Abstract: The ageing of people with intellectual disability (ID) increases rates of dementia, starting earlier and are more prevalent than in the general population. ID-care staff call for methods, knowledge, and skills to support their older residents. Person-centred methods derived from dementia care can fill this gap, but are often used unsystematically, and not adapted to ID-care. Moreover, their effectiveness in ID-care is not yet clear. One person-centred method adapted to ID-care, is Dementia Care Mapping (DCM). The aim of this study is to examine the experiences of care staff with DCM. We assessed this after two applications of DCM in twelve group homes for older people with ID, with a qualitative study (N = 24) and a quantitative study on care-staff (N = 136). Our study showed that DCM provided better understanding of the behaviour of their residents with and without dementia, more reflection and awareness of their own professional behaviour, and new knowledge and (dementia-care) skills. Furthermore, relating the needs and interpretation of the behaviour of residents to the theory of person-centred care provided care-staff a rationale and significance in daily care. Finally, DCM led to more team coordination of care. Authors concluded that evidence from dementia care can improve the quality of care for older people with ID, if adequately embedded in ID-care.

Schäper, S. & Graumann, S.
Abstract: In the coming years, a growing number of people with an intellectual disability will reach retirement age. In line with the change of paradigms, the leading ideas of participation, inclusion and self-determination have become the principles of the ideological and conceptual framework in social services for people with disabilities. However, in many places convincing concepts and arrangements of support for elderly people with intellectual disabilities are lacking, particularly beyond institutionalized concepts. The research project "Lebensqualität inklusiv(e)" (quality of life included) tried to bridge this gap. On the base of an estimation of the demographic development for this group of people, models of best practice have been documented and evaluated focusing on living conditions and the special requirements for elderly people with intellectual disabilities in order to gather ideas for the development of arrangements of support. The results show that an interdisciplinary cooperation is indispensable.

Schlamb, C.D., & Moriconi, C.D.
Betsy: A case study of a client with Down’s syndrome and dementia Advancing Care Excellence for Seniors. 8pp. https://digitalcommons.wcupa.edu/cgi/viewcontent.cgi?article=1000&context=nu rs_facpub
Abstract: This case study is about an aging woman experiencing Down syndrome (DS) and dementia. People with Down syndrome are living longer than ever before. Since the 1980s their life expectancy has doubled and many now live into their 60s, most likely because of advances in medical treatment and improved living conditions. Adults with DS and dementia typically experience several residential relocations during their lifetime and these may be traumatic events for these individuals. This study explores the complex needs of aging clients with intellectual disabilities. Target students for this teaching strategy have completed medical-surgical or geriatric nursing.

Schupf, N., Kapell, D., Lee, J.H., Ottman, R. & Mayeux, R.
Abstract: Most adults with Down’s syndrome (DS) develop neuropathology characteristic of Alzheimer’s disease (AD) by the age of 40. Most of the non-disjunction events in DS are of maternal origin. We postulated therefore that a shared genetic susceptibility to DS and AD would be associated with an increased frequency of AD among mothers, but not fathers, of individuals with DS. We further hypothesised that the shared susceptibility could involve an accelerated ageing process, leading to the birth of a child with DS to a relatively young mother and to an increased risk of dementia in the mother and her relatives. Families of 96 adults with DS and of 80 adults with other forms of mental retardation were ascertained through the New York State Developmental Disabilities services network. A semi-structured interview was used to obtain information on the presence or absence of non-stroke-related dementia and other disorders in parents. There was an increase in risk of dementia among mothers of DS probands compared with control mothers (risk ratio 2.6 [95% CI 0.9-7.3]). The risk of dementia among mothers who were 35 or younger when their DS children were born was 5 times that of control mothers (4.9 [1.6-15.4]). There was no increase in risk of dementia among mothers who were older (> 35 years) at the proband’s birth (0.8 [0.2-3.4]). There was no difference in risk of dementia between fathers of DS cases and fathers of controls (1.2 [0.4-3.9]) and...
no discernible influence of age on this risk. Familial aggregation of dementia among mothers of adults with DS supports the hypothesis of a shared genetic susceptibility to DS and AD.

Schupf, N., Kapell, D., Nightingale, B, Rodriguez, A., Tycko, B., & Mayeux, R.


Abstract: Virtually all individuals with Down syndrome (DS) have neuropathologic changes characteristic of Alzheimer’s disease (AD) beginning at 40 years of age. Few studies have examined factors that influence age at onset of AD in DS. We investigated whether sex differences in age at onset and risk of AD among adults with DS are similar to those observed in the general population and whether the effect of sex on risk of AD is modified by apolipoprotein E (APOE) genotype. A community-based sample of 111 adults with cytogenetically confirmed DS (34 to 71 years of age) was ascertained through the New York State Developmental Disabilities system. A semi-structured interview with caregivers and review of medical records was used to ascertain the presence or absence of AD. APOE genotyping was carried out without knowledge of the subject's medical history or clinical diagnosis. Both male gender and the presence of an APOE 4 allele were associated with an earlier onset of AD. Compared with women, men with DS were three times as likely to develop AD. Compared with those with the APOE 3/3 genotype, adults with DS with the 3/4 or 4/4 genotypes were four times as likely to develop AD. No individual with an APOE 7 allele developed AD. No evidence of interaction of sex and APOE genotype was found in risk of AD. The higher risk of AD in men may be related to differences in hormonal function between men and women with DS that are distinct from those in the general population.

Schupf, N., Winsten, S., Patel, B., Pang, D., Ferin, M., Zigman, W.B., Silverman, W., & Mayeux R.

Bioavailable estradiol and age at onset of Alzheimer’s disease in postmenopausal women with Down syndrome.


Abstract: Several lines of evidence suggest that loss of estrogen after menopause may play a role in the cognitive declines associated with Alzheimer’s disease (AD). Women with Down syndrome (DS) experience early onset of both menopause and AD. This timing provides a model to examine the influence of endogenous estrogen deficiency on risk of AD. We hypothesized that low serum levels of bioavailable estradiol (E2) would be associated with increased risk of AD. One hundred and nineteen postmenopausal women with DS, 42-69 years of age, were ascertained through the New York State Developmental Disability service system and followed at 15-month intervals. Information from cognitive assessments, caregiver interviews, medical record review and neurological examination was used to establish the diagnosis of dementia. Women with DS who developed AD had lower levels of bioavailable E2, lower levels of total estradiol, higher levels of sex-hormone binding globulin, and lower levels of dehydroepiandrosterone sulfate at baseline than women who remained nondemented free over the course of follow-up. Women who had low levels of bioavailable E2 at baseline were four times as likely to develop AD (HR=4.1, 95% CI: 1.2-13.9) and developed AD, on average, 3 years earlier, than those with high levels of bioavailable E2, after adjustment for age, level of mental retardation, ethnicity, body mass index, history of hypothyroidism or depression and the presence of the apolipoprotein varepsion-4 allele. Our findings support the hypothesis that reductions in estrogen following menopause can contribute to the cascade of pathological processes leading to AD.


Onset of dementia is associated with age at menopause in women with Down's syndrome.


Abstract: Women with Down’s syndrome experience early onset of both menopause and Alzheimer’s disease. This timing provides an opportunity to examine the influence of endogenous estrogen deficiency, indicated by age at menopause, on risk of Alzheimer’s disease. A community-based sample of 163 postmenopausal women with Down’s syndrome, 40 to 60 years of age, was ascertained through the New York State Developmental Disability service system. Information from cognitive assessments, medical record review, neurological evaluation, and caregiver interviews was used to establish ages for onset of menopause and dementia. We used survival and multivariate regression analyses to determine the relation of age at menopause to age at onset of Alzheimer’s disease, adjusting for age, level of mental retardation, body mass index, and history of hypothyroidism or depression. Women with early onset of menopause (46 years or younger) had earlier onset and increased risk of Alzheimer’s disease (AD) compared with women with onset of menopause after 46 years (rate ratio, 2.7; 95% confidence interval [CI], 1.2-5.9). Demented women had higher mean serum sex hormone binding globulin levels than nondemented women (86.4 vs 56.6 nmol/L, p = 0.02), but similar levels of total estradiol, suggesting that bioavailable estradiol, rather than total estradiol, is associated with dementia. Our findings support the hypothesis that reductions in estrogen after menopause contribute to the cascade of pathological processes leading to AD.

Schweber, M.S.

Alzheimer's disease and Down syndrome


Abstract: This report contains a summary of an extensive survey of autopsy data for persons with intellectual disability. Among adults with Down syndrome (DS), the brain neuropathology of AD was universal in those age 37 and over; claimed exceptions were indefensible. The behavioral evaluations of the DS adults, however, could be classified into three divisions: 1. "quiescent" (neither seizures nor dementia, 2) "partial" (seizures but no dementia), and 3) "active" (dementia +/- seizures). Thus, it is reasonable to argue that all persons with DS develop AD itself upon aging. However, DS cannot be used uncritically as an AD model since no increased incidence of active AD was found in DS with aging beyond the critical threshold age (mid-30’s). Improved accurate quantification of Southern blots produced 100% accuracy in decoding blind samples of DS and non-DS samples. Using this system, DNA levels similar to those of DS have been demonstrated for all categories of AD at a small subsection of chromosome 21 near to, or within the DS DNA location on chromosome 21. Increased amounts of a complete, structural gene sequence were not found (or expected). The results provide evidence for a unitary hypothesis for DS and all forms of AD.

Scottish Down’s Syndrome Association

What is dementia? - A booklet about dementia for adults who have a learning disability.

14pp
Edinburgh: Scottish Down's Syndrome Association [158-160 Balgreen Road, Edinburgh, Scotland EH11 3AU; e-mail: info@sdlsa.org.uk; www.sdlsa.org.uk] [n.d.]

[Source: http://www.rrcadd.co.uk/T/A/Dementia_Care_Resources/Info.html]

Abstract: Written for the Scottish Down's Syndrome Association by Diana Kerr and Mo Innes this A4 size booklet is designed to explain dementia and its nuances to persons with intellectual disabilities (termed "learning disabilities in Scotland"). Using drawings and easy language this booklet covers many of the symptoms and behaviors classically associated with Alzheimer’s disease.

Service, K.P.

Considerations in care for individuals with intellectual disability with advanced dementia


Abstract: A number of physical, psychosocial, or ethical issues related to the care of the individual with advanced dementia are reviewed and related to individuals with intellectual disabilities. The sources used include the published literature and illustrations drawn from personal observations. The author notes that through anticipation and early planning, advanced directives and service planning (which looks to adaptation of services and other care management interventions), can effectively impact care at the end. Areas that need to be addressed include technical information, including a review of and, as appropriate, adaptation of general advanced dementia resources, relief, rest, support, reassurance, receipt of on-going information, participation in planning, a sense of humor, and appreciation. Also noted, are the differences experienced because of the presence of paid staff as carers and residence outside of the family home. It is concluded that, although the goals of quality care is the same for all people with advanced dementia, the process by which to reach these goals often needs further consideration and adaptation for people with intellectual disabilities.
Service, K.P., & Clifford, C.J.  
What do I really need? Assessment of caregiver supports for people with intellectual and developmental disability and dementia.  
https://doi.org/10.1002/alz.047106  
Abstract: The increased needs of people with an intellectual and developmental disability (IDD) and a dementia related disorder can strain caregivers and existing community support systems. The project team, comprised of IDD and Aging experts and funded, in part, through a federal grant, conducted a needs assessment on the awareness and use of typical community-based resources such as senior centers. The assessment consisted of both telephone interviews and home visits with the home, shared living, and family caregivers. Nurse Practitioner (NP) conducted a total of 95 interviews with 54 site visits, and the evaluator completed 40 interviews with caregivers of people with IDD and dementia diagnosis. Analysis included both qualitative and quantitative data. Caregivers were asked about the following: functional and health status of person with IDD since dementia diagnosis, receipt of dementia specific caregiving training, care confidence levels, perceived barriers to care, and access to community-based aging resources. Caregiver's most frequent concerns included lack of suitable day programming, planning for the person's future, and caregiver burnout and stress. 78% reported feeling confident providing care currently and 68% were confident about providing care in the future. Most caregivers are aware of local community resources such as senior centers, Alzheimer's Association Counseling and an on-line training series on aging, but rarely used the resources. Authors note that caregivers generally relied on support from provider agencies indicating a need for increased collaboration across the IDD and Aging human service support systems. Trainings, delivered to both aging advocates and caregivers of people with IDD, and designed to improve communication and collaboration focused on dementia capable care, state systems, and available community resources. In addition, a series of web-based resources were developed with a focus on IDD and dementia. Results of the assessment will continue to guide resource and training development to improve collaboration and support the relationship between the Aging and IDD communities.

Service, K.P., Lavoie, D, Herlihy, J.E.  
Coping with losses, death and grieving  
Abstract: This book chapter uses a composite case to demonstrate strategies to address the issues related to losses and death for people with intellectual disability and the diagnosis of dementia and for their families and staff. Dealing with the diagnosis and the changes are explained in the framework of the stages of death and dying as developed by Kubler-Ross. The responses to the losses of dementia which are manifested by affected individuals and their families concern included lack of suitable day programming, planning for the person's future, and caregiver burnout and stress. 78% reported feeling confident providing care currently and 68% were confident about providing care in the future. Most caregivers are aware of local community resources such as senior centers, Alzheimer's Association Counseling and an on-line training series on aging, but rarely used the resources. Authors note that caregivers generally relied on support from provider agencies indicating a need for increased collaboration across the IDD and Aging human service support systems. Trainings, delivered to both aging advocates and caregivers of people with IDD, and designed to improve communication and collaboration focused on dementia capable care, state systems, and available community resources. In addition, a series of web-based resources were developed with a focus on IDD and dementia. Results of the assessment will continue to guide resource and training development to improve collaboration and support the relationship between the Aging and IDD communities.

Sheehan, R., Ali, A., & Hassiotis, A.  
Dementia in intellectual disability  
doi: 10.1097/YCO.000000000000032.  
Abstract: Dementia is emerging as a significant condition in the population with intellectual disability. This review is aimed at clinicians working in the field. We revisit what is known on the subject and expand on this with results from recent research. The emphasis of this review is on the clinical research rather than laboratory or molecular research. Research has encompassed all aspects of dementia in intellectual disability, from epidemiology, assessment and diagnosis, through to management. There remains a lack of evidence concerning both pharmacological and nonpharmacological treatment of dementia in people with intellectual disability. Recent research has tended to focus on dementia in Down syndrome. More research is necessary in order to translate improvements in the understanding of the neuropathology of intellectual disability and dementia into effective treatments. There is also a need to investigate the optimum environment in which to provide holistic care for individuals affected.

Dementia diagnostic criteria in Down syndrome.  
Abstract: Dementia is a common clinical presentation among older adults with Down syndrome. The presentation of dementia in Down syndrome differs compared with typical Alzheimer's disease. The performance of manualised dementia criteria in the International Classification of Diseases (ICD)-10 and Diagnostic and Statistical Manual of Mental Disorders-IV-Text Revision (DSM-IV-TR) is uncertain in this population. We aimed to determine the concurrent validity and reliability of clinicians' diagnoses of dementia against ICD-10 and DSM-IV-TR diagnoses. Validity of clinical diagnoses were also explored by establishing the stability of diagnoses over time. The authors used clinical data from memory assessments of 85 people with Down syndrome, of whom 64 (75.3%) had a diagnosis of dementia. The cases of dementia were presented to expert raters who rated the case as dementia or no dementia using ICD-10 and DSM-IV-TR criteria and their own clinical judgement. The authors found that clinician's judgement corresponded best with clinically diagnosed cases of dementia, identifying 84.4% cases of clinically diagnosed dementia at the time of diagnosis. ICD-10 criteria identified 70.3% cases, and DSM-IV-TR criteria identified 56.3% cases at the time of clinically diagnosed dementia. Over time, the proportion of cases meeting ICD-10 or DSM-IV-TR diagnoses increased, suggesting that experienced clinicians used their clinical knowledge of dementia presentation in Down syndrome to diagnose the disorder at an earlier stage than would have been possible had they relied on the classic description contained in the diagnostic systems. The authors concluded that clinical diagnosis of dementia in Down syndrome is valid and reliable and can be used as the standard against which new criteria such as the DSM-5 are measured.

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DSM-IV-TR diagnoses. Validity of clinical diagnoses were also explored by establishing the stability of diagnoses over time. The authors used clinical data from memory assessments of 85 people with Down syndrome, of whom 64 (75.3%) had a diagnosis of dementia. The cases of dementia were presented to expert raters who rated the case as dementia or no dementia using ICD-10 and DSM-IV-TR criteria and their own clinical judgement. The authors found that clinician’s judgement corresponded best with clinically diagnosed cases of dementia, identifying 84.4% cases of clinically diagnosed dementia at the time of diagnosis. ICD-10 criteria identified 70.3% cases, and DSM-IV-TR criteria identified 56.3% cases at the time of clinically diagnosed dementia. Over time, the proportion of cases meeting ICD-10 or DSM-IV-TR diagnoses increased, suggesting that experienced clinicians used their clinical knowledge of dementia presentation in Down syndrome to diagnose the disorder at an earlier stage than would have been possible had they relied on the classic description contained in the diagnostic systems. The authors concluded that clinical diagnosis of dementia in Down syndrome is valid and reliable and can be used as the standard against which new criteria such as the DSM-5 are measured.

Sheth, A.J.
Abstract: The purpose of this paper was to improve understanding of environmental influences on participation in routine and familiar activities for people with intellectual disability and dementia from first-person and caregiver perspectives. The methodology involved four adults with intellectual disability and dementia participating in 2 nominal group technique sessions and 12 family and staff caregivers participating in 5 standard focus groups. Transcripts were analyzed utilizing thematic analysis centering the findings from nominal group technique sessions and an ecological systems lens. The findings revealed that participants with intellectual disability and dementia identified six important themes: activity access, caregiver assistance, social interactions, responsibilities, privacy, and health and wellness. Their perspectives focused primarily at an immediate environment level, while caregiver input added additional understandings from broader ecological systems levels. This study provides a beginning point to establishing a framework for creating supports and addressing barriers to participation for adults with intellectual disability and dementia based on direct input from potential service consumers and their caregivers. People with intellectual disabilities and dementia provide valuable insights into their experiences through engagement in accessible research.

"It's not the same without you" exploring the experience and perception of transition for people with intellectual disabilities and dementia. British Journal of Learning Disabilities, 2021 (Sep), 49(3), 365-372.
https://doi.org/10.1111/bld.12412
Abstract: For people with intellectual disabilities and dementia, transitions are likely to become increasingly common as they age. While transitions experienced by people with intellectual disabilities in young adulthood are frequently studied, less is known about transitions in older adults, including residential, vocational and leisure changes. This article aims to explore the experiences of transition from the perspectives of people with intellectual disabilities and dementia, including the impact on their daily lives. Three women with intellectual disabilities and dementia living in residential settings participated in participant observations and informal interviews across a variety of environments and activities. Field notes and interview transcripts underwent a thematic analysis focusing on transitions. Participants experienced the impact of transitions in their residential placements, day programming, leisure activities and relationships. Themes related to their experiences of transitions included making sense of transitions, utilising peer care networks for support and tackling the looming threat of loss and transition. Peer care networks and friendships are crucial in supporting people during and after transitions. Receiving effective supports to maintain relationships, roles and activities, even in seemingly minor ways, is an important right for people with intellectual disabilities and dementia, particularly as care needs increase.

Shirai, Y., Bishop, K., & Kushner, M.
https://doi.org/10.1352/1934-9556-59.5.422
Abstract: With a growing need for specialized training for direct caregivers and support staff of persons with intellectual and developmental disabilities (IDD) affected by dementia, the National Task Group on Developmental Disabilities and Dementia Practices (NTG) developed a comprehensive evidence-informed Dementia Capable Care Training (DCCT). To overcome the challenge of the training length and cost, and to extend its dissemination, the Sonoran Center developed a shorter version of the NTG-DCCT while retaining its core components, and implemented it in seven cities in the U. S. Southwest (N = 368). The pre- and post-training evaluation (n = 260) demonstrated that the short version of the NTG-DCCT is effective in significantly improving participants' knowledge and/or confidence in dementia capable care. The follow-up semi-structured interviews of participants (n = 7) provide some insights.

Shooshtari, S., Stoess, B.M., Udell, L., Fenez, L., Nik, N., Burchill, C., Sachs, E., Menec, V.
Aging with intellectual and developmental disabilities and dementia in Manitoba: Advances in Mental Health and Intellectual Disabilities, 11(4), 134-144.
https://doi.org/10.1108/AMHID-03-2017-0007
Abstract: Information on the risk of dementia in aging persons with intellectual and/or developmental disability (IDD) in Manitoba, Canada is lacking. The purpose of this paper is to estimate dementia prevalence in adults with IDD. Anonymized population-level health and non-health administrative data (1979-2012) contained in the Population Health Research Data Repository of the Manitoba Centre for Health Policy (MCHP) were linked to identify adults with IDD, and estimate the prevalence of dementia based on the presence of ICD codes. Prevalence of dementia was estimated for persons aged 18-55 years and 55+ years, and was reported by sex, type of residence, region of residence, neighbourhood income quintiles, and IDD diagnostic category. Of the 8,655 adults with IDD identified, 8.1 per cent had an indication of dementia in their medical records; an estimate three times greater than that found for those without IDD (2.6 per cent). More than 17 per cent of Manitobans with IDD aged 55+ years had an indication of dementia, which was nearly twice the rate reported previously. Of those with IDD and dementia, 34.7 per cent lived in long-term care facilities. Health and social support services are typically available to individuals with dementia aged 65+ years; thus, younger adults with IDD and dementia may not be eligible for those supports. To promote equity in health and access to care, dementia screening and increased supports for aging individuals with IDD are recommended.

Shultz, J.M., Aman, M.G., & Rojahn, J.
Abstract: Forty elderly persons with mental retardation were assessed by their care providers on a modified version of the Short Informant Questionnaire on Cognitive Decline in The Elderly (IQCODE) an instrument designed to quantify cognitive decline in elderly people in the general population. They were also assessed for IQ, aberrant behavior, and current mental status; test-retest and interrater reliability were evaluated as well. Internal consistency, as assessed by coefficient alpha, was moderately high (alpha = .86). Test-retest reliability was mediocre and interrater reliability levels did not reach statistical significance. The Short IQCODE was not correlated with a variety of demographic features or with behavior ratings, showing evidence of divergent validity. However, the Short IQCODE was only weakly (nonsignificantly) correlated with a measure of current mental status, which challenges its concurrent validity. The Short IQCODE probably needs to be modified further for satisfactory psychometric performance in people with mental retardation. However, some features of this study may have resulted in suboptimal estimates of the Short IQCODE’s psychometric characteristics.

Sigal, M.J., & Levine, N.
Abstract: Individuals with Down’s syndrome (DS) who live to be 40 years of age will demonstrate neuropathological changes that are consistent with Alzheimer’s disease (AD). Due to modern medical intervention, we are now observing an aging DS population. Middle-aged Down’s syndrome adults are actually...
considered to be "very old," and it is not uncommon to observe a progressive loss of cognitive function and a decline in the ability to perform daily tasks consistent with that seen in Alzheimer's disease. At this stage, the DS individual will not be able to perform daily preventive dental care and may be unable to cooperate for professional dental care. Clinicians who care for DS adults must be aware of this problem when preparing their dental treatment plans, which must emphasize preventive care prior to the onset of dementia and the maintenance of that program during their patients' cognitive decline. In the latter stages of AD, it may be necessary to extract all the remaining teeth due to the inability of the individual or care giver to provide adequate oral hygiene to prevent dental caries or periodontal disease. Simard, M., & van Reekum, R. Dementia with Lewy bodies in Down's syndrome. International Journal of Geriatric Psychiatry, 2001, Mar;16(3), 311-20. Abstract: The association between Down's syndrome (DS) and Alzheimer's disease is well established. This paper presents a review of the literature, suggesting a possible association between DS and the more recently recognized dementia with Lewy bodies (DLB). Patients with DLB frequently present with changes in affect and behavior, and in particular with psychiatric symptoms. The literature suggests a possible role for atypical neuroleptics in the management of psychosis in DLB.

Sinai, A., Mokrysz, C., Bernal, J., Bohnen, I., Bonell, S., Courtney, K., Dodd, K., ... Strydom, A. Predictors of age of diagnosis and survival of Alzheimer's disease in Down syndrome. Journal of Alzheimer's Disease, 2018, 61(2), 717-728. doi: 10.3233/JAD-170624. Abstract: People with Down syndrome (DS) are an ultra-high risk population for Alzheimer's disease (AD). Understanding the factors associated with age of onset and survival in this population could highlight factors associated with modulation of the amyloid cascade. This study aimed to establish the typical age at diagnosis and survival associated with AD in DS and the risk factors associated with these. Data was obtained from the Aging with Down Syndrome and Intellectual Disabilities (ADSID) research database, consisting of data extracted from clinical records of patients seen by Community Intellectual Disability Services (CIDS) in England. Survival times when considering different risk factors were calculated. The mean age of diagnosis was 55.80 years, SD 6.29. Median survival time after diagnosis was 3.78 years, and median age at death was approximately 60 years. Survival time was associated with age of diagnosis, severity of intellectual disability, living status, anti-dementia medication use, family history of epilepsy. Age at diagnosis and treatment status remained predictive of survival time following adjustment. This study provides the best estimate of survival in dementia within the DS population to date, and is in keeping with previous estimates from smaller studies in the DS population. This study provides important estimates and insights into possible predictors of survival and age of diagnosis of AD in adults with DS, which will inform selection of participants for treatment trials in the future.

Silverman W, Krinsky-McHale SJ, Kovacs C, Lee JH, Listwan T, Pang DI, Zigman WB, Schupf N. Individualized estimated years from onset of Alzheimer's disease-related decline for adults with Down syndrome. Alzheimers Dement (Amst). 2023 Jun 27;19(2):e12444. doi: 10.1002/dad.212444. Abstract: Adults with Down syndrome (DS) are at increased risk for Alzheimer's disease (AD) and vary in their age of transition from AD preclinical to prodromal or more advanced clinical stages. An empirically based method is needed to determine individual "estimated years from symptom onset (EYO)," the same construct used in studies of autosomal dominant AD. Archived data from a previous study of > 600 adults with DS were examined using survival analysis methods. Age-specific prevalence of prodromal AD or dementia, cumulative risk, and EYOs were determined. Individualized EYOs for adults with DS ranging in age from 30 to 70+ were determined, dependent upon chronological age and clinical status. EYOs can be a useful tool for studies focused on biomarker changes during AD progression in this and other populations at risk, studies that should contribute to improved methods for diagnosis, prediction of risk, and identification of promising treatment targets. Silverman, W., Krinsky-McHale, S.J., Lai, F., Rosas, H.D., Hom, C., Doran, E., Pulsifer, M., Lott, I., Schupf, N., and Alzheimer's Disease in Down Syndrome (ADDS) Consortium. Evaluation of the National Task Group-Early Detection Screen for Dementia: Sensitivity to 'mild cognitive impairment' in adults with Down syndrome Journal of Applied Research in Intellectual Disabilities, 2020, Dec 13. doi:10.1111/jar.12849. Abstract: The accuracy of the National Task Group-Early Detection Screen for Dementia (NTG-EDSD) was evaluated in a sample of 185 adults with Down syndrome (DS), emphasizing 'mild cognitive impairment (MCI-DS). Knowledgeable informants were interviewed with the NTG-EDSD, and findings were compared to an independent dementia status rating based on consensus review of detailed assessments of cognition, functional abilities and health status (including physician examination). Results indicated that sections of the NTG-EDSD were sensitive to MCI-DS, with one or more concerns within the 'Memory' or 'Language and Communication' domains being most informative. The NTG-EDSD is a useful tool for evaluating dementia status, including MCI-DS. However, estimates of sensitivity and specificity, even for detecting frank dementia, indicated that NTG-EDSD findings need to be supplemented by additional sources of relevant information to achieve an acceptable level of diagnostic/screening accuracy.

Silverman, W., Schupf, N., Zigman, W., Devenny, D., Miezejeski, C., Schubert, R., & Ryan, R. Dementia in adults with mental retardation: Assessment at a single point in time. American Journal of Mental Retardation, 2004, 109, 111-125. doi:10.1352/0895-8017(2004)109<111:AJAMR2.0.CO;2 Abstract: Dementia status of 273 adults with mental retardation was rated based upon two extensive evaluations conducted 18 months apart. Overall, 184 individuals did not have dementia, 33 had possible or definite dementia, and 66 had findings suggesting uncertain or questionable status. These ratings were compared to binary classifications (dementia vs. no dementia) generated from the Dementia Questionnaire for Persons With Mental Retardation (Evenhuis, 1995) and the IBR Mental Status Examination (Wisniewski & Hill, 1985). When performance was referenced to IQs (established earlier in adulthood), quantitative criteria effectively distinguished between individuals with and without dementia based upon assessment at a single point in time. Findings suggest that procedures of this type could soon contribute to more accurate and rapid diagnoses of dementia. Smith, D., & Chicoine, B. Difficulties of diagnosing and managing dementia in people with Down syndrome British Journal of Psychiatry, 2018, Nov, 213(5), 668-669. doi:10.1192/bjp.2018.199. Abstract: In our experience as the directors of Down syndrome clinics for adults, the big issue is how the diagnosis of dementia is made. Clinicians tend to easily apply the diagnosis or Alzheimer's dementia without looking at all of the potential causes of pseudodementia in this population. They often assume that loss of ability is due to dementia because of a study published in 1985 that showed plaques and tangles in the brain tissue of all people with Down syndrome over the age of 35. Wisniewski & Rabe subsequently wrote that there was a discrepancy between neuropathology and the occurrence of dementia in people with Down syndrome. Just as the population of typically developed older adults the diagnosis of Alzheimer's dementia in people with Down syndrome should be made only after evaluation for causes of pseudodementia. Snyder HM, Bain LJ, Brickman AM, Carrillo MC, Esbensen AJ, Espinosa JM, Fernandez F, Fortea J, Hartley SL, Head E, Hendrix J, Kishnani PS, Lai F, Lao P, Lemere C, Moblew W, Mufson EJ, Potter H, Zaman SH, Granholm AC, Rosas HD, Strydom A, Whitten MS, Rafii MS. Further understanding the connection between Alzheimer's disease and Down syndrome. Alzheimers Dement. 2020 Jul;16(7):1065-1077. doi: 10.1002/alz.12112. Epub 2020 Jun 16 Abstract: Improved medical care of individuals with Down syndrome (DS) has led to an increase in life expectancy to over the age of 60 years. In conjunction, there has been an increase in age-related co-occurring conditions including Alzheimer's disease (AD). Understanding the factors that underlie symptom and age of clinical presentation of dementia in people with DS may provide insights into the mechanisms of sporadic and DS-associated AD (DS-AD). In March
The role of prospective screening in the diagnosis of dementia in people with Down's syndrome


Abstract: People with Down's syndrome are at increased risk of developing early onset Alzheimer's disease. It has been recommended that all adults with Down's syndrome receive baseline neuropsychological testing for dementia. In certain areas prospective screening of people with Down's syndrome takes place to ensure the early diagnosis of the condition. However, little has been published on the value of this type of screening. The purpose of this paper is to report on a prospective screening programme and asks whether the programme is effective in identifying dementia-related changes in people with Down's syndrome and whether the current screening intervals are appropriate. All adults with Down's syndrome in Plymouth (UK) are identified and offered a comprehensive test battery at baseline at the age of 20 and then have testing biennially from 40 to 50 and annually after 50. All individuals diagnosed with dementia between 2001 and 2013 were identified and their case notes examined. The symptoms at the time of diagnosis were identified and whether these symptoms had been identified through the screening programme or by other routes were recorded. Prevalence data and age at diagnosis were also recorded. In total, 26 people were diagnosed with dementia during the study period. Of these, the diagnosis of dementia followed concerns being identified during the routine screening programme in 54 per cent of cases. In the younger age group (age 40-49) 63 per cent of people were identified through the screening programme. At the time of diagnosis a mean of 5.5 areas of concern were in evidence. This paper adds to the growing evidence base around the value of prospective dementia screening in people with Down's syndrome. It is also one of a few studies exploring the frequency of screening. Additionally, it adds further data about prevalence of dementia in people with Down's syndrome.

Starin, C.M., Rodger, R., Fodor-Wynne, L., Hamburg, S., & Strydom, A.
Abstract: Down syndrome (DS) is the most common genetic cause of intellectual disability (ID). Abilities relating to executive function, memory, and language are particularly affected in DS, although there is a large variability across individuals. People with DS also show an increased risk of developing dementia. While assessment batteries have been developed for adults with DS to assess cognitive abilities, these batteries may not be suitable for those with more severe IDs, dementia, or visual / hearing difficulties. Here we report the development of an informant rated questionnaire, the Cognitive Scale for Down Syndrome (CS-DS), which focuses on everyday abilities relating to executive function, memory and language, and is suitable for assessing these abilities in all adults with DS regardless of cognitive ability. Complete questionnaires were collected for approximately 128 individuals with DS. After final question selection we found high internal consistency scores across the total questionnaire and within the executive function, memory and language domains. CS-DS scores showed a wide range, with minimal floor and ceiling effects. We found high interrater (n = 55) and test retest (n = 36) intraclass correlations. CS-DS scores were significantly lower in those aged 41+ with significant cognitive decline compared to those without decline. Across all adults without cognitive decline, CS-DS scores correlated significantly to measures of general abilities. Exploratory factor analysis suggested five factors within the scale, relating to memory, self-regulation / inhibition, self-direction / initiation, communication, and focussing. The CS-DS therefore shows good interrater and test retest reliability, and appears to be a valid and suitable informant rating tool for assessing everyday cognitive abilities in a wide range of individuals with DS. Such a questionnaire may be a useful outcome measure for intervention studies to assess improvements to cognition, in addition to detecting dementia-related cognitive decline. The CS-DS may also be a useful tool for other populations with ID.


Abstract: Reviewed are the common characteristics and co-morbidities of individuals with Down syndrome (DS), his epidemiology of AD and clinical presentation of AD in this cohort, and discussed are practical considerations for diagnosis and treatment as well as strategies to maximize support for caregivers. Consistent with the National Plan to Address Alzheimer’s Disease, we will focus on the importance of the early detection of AD and what it means to patients, caregivers, and the healthcare system. The most commonly used screening tool for evaluating the early indicators of dementia in DS is the National Task Group - Early Detection Screen for Dementia (NTG-EDSD). The NTG-EDSD scoring scale helps clinicians compare patients with DS to their own baseline as the questions are scored always been the case, always but worse, new symptom in the past year, or does not apply. The framework of analysis reflects transitions from “clinically stable” to concern for MCI-DS to suspicion for AD and additional considerations of individuals with DS. Pharmacotherapy and non-pharmacologic approaches to treat co-existing depression and anxiety, and improve quality of life are important considerations in the care of patients with AD and DS. Caring for and supporting patients with DS and MCI-DS and AD is best managed from an interprofessional team approach to achieve the best outcomes. Social workers, case managers, nurses and occupational therapists can evaluate the needs of the patient and caregiver and provide recommendations to support patient participation in activities and routines which can improve patient overall health and well-being as well as reduce unwanted behaviors. The goal of the primary care physician is to recognize and manage the medical and socioeconomic disparity that exists for patients with DS, especially as they age.


Abstract: The purpose of this review was twofold: (1) to gain insight into what is known from the literature about home modifications for people with Alzheimer's disease (AD) and (2) to identify gaps in the literature that could lead to opportunities for research. A systematic scoping review of peer-reviewed articles published from 1994 through 2014 explored home modifications and AD. Seventeen articles met the inclusion criteria. The three major findings pertain to (1) the caregiver role and caregiver training, (2) a client-centered collaborative approach to assessment and intervention, and (3) modifications for safety and function. Home modifications involved the physical and social environments as well as cognitive strategies at the task level. Opportunities exist for the development of assessment procedures, the exploration of home modifications in the later stages of AD, and the study of home modification needs of people with dementia who live alone.


Abstract: Adults with Down syndrome (DS) have neuropathological features identical to individuals with sporadic Alzheimer’s disease (AD) and this discovery played an important role in the identification of the amyloid precursor protein gene on chromosome 21. Individuals with DS have a lifetime risk for dementia in excess of 90% and DS is now acknowledged to be the most common genetic cause of AD, but this group is often excluded from AD medication trials. This review will discuss primary and secondary prevention trials for AD in DS and the potential barriers and solutions to such trials. It will include a brief overview of the epidemiology, diagnosis and outcome measurement issues pertinent to prevention trials, as well as important aging-related co-morbidities that need to be considered in the design of such trials. Described is the work of the European-wide Horizon21 Consortium and other efforts to establish DS clinical trials networks, as well as to consider the methodological issues for trials to prevent or delay AD in DS. It was noted that individuals with DS could benefit from treatments to prevent or delay AD. Improved knowledge of pathogenic processes and their clinical consequences in DS will hopefully lead to new clinical trials.


Abstract: There is a need for simple and reliable screening instruments for dementia in the intellectual disability (ID) population that can also be used to follow their progress, particularly if they are being treated with anti-dementia drugs. Commonly used tests for the general population such as the Mini Mental State Examination (MMSE) are not appropriate for many people with ID. This paper is a literature review of alternative instruments that have been used in research or recommended by experts since 1991 and have the potential to be used as screening instruments. Two types of tests have been identified: those administered to informants, and those that rely on direct assessment of the individual. The most promising informant rated screening tool in most adults with ID including Down syndrome (DS) diagnosis is the Dementia Questionnaire for Persons with Mental Retardation (DMR). However, sensitivity in single assessments is variable and cut-off scores need further optimization. In those with DS, the Dementia Scale for Down Syndrome (DSDS) has good specificity but mediocre sensitivity. The Test for Severe Impairment and Severe Impairment Battery are two direct assessment tools that show promise as screening instruments, but need further evaluation.


Abstract: People with intellectual [learning] disabilities, particularly Down’s syndrome, are at increased risk of dementia. At present, services and care tailored to people with both dementia and a intellectual disability are unsatisfactory. This article reviews the literature specific to dementia in people with learning disabilities, including: comprehensive screening, diagnosis, management, environmental considerations, end of life care and training issues for nursing staff. Recommendations for best practice and service improvement are made to improve the quality of life for individuals with dementia and learning disabilities, pre and post-diagnosis.
Prevalence of dementia in older adults with intellectual disability without Down syndrome


Abstract: The aim of this study was to determine the prevalence of dementia in older adults with intellectual disability (ID) without Down syndrome. The authors identified the total population of adults with ID aged 60+ in the five London boroughs served through local social services registers, ID teams and residential services providers and then screened the Ss with a simple object memory task, information about functional status, and the Dementia Questionnaire for Persons with Mental Retardation (DMR). Screen positives on the DMR, or those with unexplained functional decline or memory deficits underwent detailed examination. Full assessment of cognitive and physical function was undertaken and additional information was collected from informants and medical records. All information was summarized to determine dementia status with ICD-10, DSM-IV, and DC-LD criteria. The authors identified 264 adults with ID and 222 (84%) participated in the study. One in four screened positive. The prevalence rate for ICD-10 or DSM-IV was 12%. Prevalence differed between those with mild and severe ID, and between diagnostic criteria. The authors concluded that dementia is common in older adults with ID without DS, but prevalence in severe ID deviated from prediction and the use of diagnostic criteria needs to be reviewed.

Mental health and social care needs of older people with intellectual disabilities


Abstract: Older people with intellectual disabilities (ID) are a growing population but their age-related needs are rarely considered and community services are still geared towards the younger age group. We aimed to examine the mental health and social care needs of this new service user group. We identified all adults with ID without Down syndrome (DS) aged 65+ living in the London boroughs of Camden and Islington. The Psychiatric Assessment Schedule for Adults with a Developmental Disability (PASAD) checklist was used to detect psychiatric disorder, the Vineland Behavior Scale (maladaptive domain) for problem behaviors and the Dementia Questionnaire for Persons with Mental Retardation (DMR) to screen for dementia. Carers reported health problems and disability. Needs were measured with the Camberwell Assessment of Need for adults with Intellectual Disabilities (CANDID-S). A total of 23 older people with ID (13 had mild ID and nine more severe ID) and their carers participated in the survey. In which, 74% had one or more psychiatric symptoms; 30% were previously known with a diagnosis of mental illness. One-third of the older people screened positive for dementia (range: 17.44%, depending on sensitivity of DMR scores used). Three quarters of the group had physical health problems, 74% had poor sight, 22% had hearing loss and 30% had mobility problems. Carers rated unmet needs for accommodation (22%), day activities, and eyesight and hearing. The people with ID rated unmet needs to be social relationships (44%), information and physical health. Authors concluded that older people with ID without DS have considerable prevalence of health problems and psychiatric disorders, including symptoms of functional decline and dementia. Such symptoms are often not recognized and further research into their needs is a priority.

Prevalence of dementia in intellectual disability using different diagnostic criteria.


Abstract: Diagnosis of dementia is complex in adults with intellectual disability owing to their pre-existing deficits and different presentation. To describe the clinical features and prevalence of dementia and its subtypes and to compare the concurrent validity of dementia criteria in older adults with intellectual disability. The Becoming Older with Learning Disability (BOLD) memory study is a two-stage epidemiological survey of adults with intellectual disability without Down syndrome aged 60 years and older, with comprehensive assessment of people who screen positive. Dementia was diagnosed according to ICD–10, DSM–IV and DC–LD criteria. The DSM–IV dementia criteria were more inclusive. Diagnosis using ICD–10 excluded people with even moderate dementia. Clinical subtypes of dementia can be recognized in adults with intellectual disability. Alzheimer’s dementia was the most common, with a prevalence of 6.6% (95% CI 5.2–13.0), almost three times greater than expected. Dementia is common in older adults with intellectual disability, but prevalence differs according to the diagnostic criteria used. This has implications for clinical practice.

The relationship of dementia prevalence in older adults with intellectual disability (ID) to age and severity of ID.


Abstract: Previous research has shown that adults with intellectual disability (ID) may be more at risk of developing dementia in old age than expected. However, the effect of age and ID severity on dementia prevalence rates has never been reported for adults with ID. We investigated the predictions that older adults with ID should have high prevalence rates of dementia that differ between ID severity groups and that the age-associated risk should be shifted to a younger age relative to the general population. A two-staged epidemiological survey of 281 adults with ID without Down syndrome (DS) aged 60 years; participants who screened positive with a memory task, informant-reported change in function or with the Dementia Questionnaire for Persons with Mental Retardation (DMR) underwent a detailed assessment. Diagnoses were made by psychiatrists according to international criteria. Prevalence rates were compared with UK prevalence and European consensus rates using standardized morbidity ratios (SMRs). Dementia was more common in this population (prevalence of 18.3%, SMR 2.77 in those aged 65 years). Prevalence rates did not differ between mild, moderate and severe ID groups. Age was a strong risk factor and was not influenced by sex or ID severity. As predicted, SMRs were higher for younger age groups compared to older age groups, indicating a relative shift in age-associated risk. Criteria-defined dementia is 2-3 times more common in the ID population, with a shift in risk to younger age groups compared to the general population.

Incidence rates from a longitudinal study of dementia in older adults with ID in the general population. A two-staged epidemiological survey of 281 adults with ID without DS aged 60 years; participants who screened positive with a memory task, informant-reported change in function or with the Dementia Questionnaire for Persons with Mental Retardation (DMR) underwent a detailed assessment. Diagnoses were made by psychiatrists according to international criteria. Prevalence rates were compared with UK prevalence and European consensus rates using standardized morbidity ratios (SMRs). Dementia was more common in this population (prevalence of 18.3%, SMR 2.77 in those aged 65 years). Prevalence rates did not differ between mild, moderate and severe ID groups. Age was a strong risk factor and was not influenced by sex or ID severity. As predicted, SMRs were higher for younger age groups compared to older age groups, indicating a relative shift in age-associated risk. Criteria-defined dementia is 2-3 times more common in the ID population, with a shift in risk to younger age groups compared to the general population.

Prevalence of dementia in older adults with ID without Down syndrome aged 60 years and older were followed up an average of 2.9 years. The authors conducted a comprehensive review of the published literature from 1997–2008 with a specific focus on: (1) epidemiology of dementia in ID in general as well as in specific genetic syndromes; (2) presentation; and (3) diagnostic criteria for dementia. The review drew upon a combination of searches in electronic databases Medline, EMBASE, and PsycINFO for original research papers in English, Dutch, or German. The authors report that varied methodologies and inherent challenges in diagnosis yield a wide range of reported prevalence rates of dementia. Rates of dementia in the population with intellectual disability not because of Down syndrome (DS) are comparable with or higher than the general population. Alzheimer’s disease onset in DS appears earlier and the prevalence increases from under 10% in the 40s to more than 30% in the 50s, with varying prevalence reported for those 60 and older. Incidence rates increase with age. Few studies of dementia in other genetic syndromes were identified. Presentation differs in the ID population compared with the general population; those with DS present with prominent behavioral changes believed to be because of frontal lobe deficits. Authors recommend large-scale collaborative studies of high quality to further knowledge on the epidemiology and clinical presentation of dementia in this population.

Incidence of dementia in older adults with intellectual disabilities


Abstract: Dementia may be more common in older adults with intellectual disability (ID) than in the general population. The increased risk for Alzheimer’s disease in people with Down syndrome (DS) is well established, but much less is known about dementia in adults with ID who do not have DS. We estimated incidence rates from a longitudinal study of dementia in older adults with ID without DS and compared them to general population rates. 222 participants with ID without DS aged 60 years and older were followed up an average of 2.9 years later to identify those who had declined in functional or cognitive abilities.
Those who screened positive had a comprehensive assessment for dementia, diagnosed using ICD 10 and DSM IV criteria. 134 participants who did not have dementia at initial assessment were alive and interviewed at follow up; 21 (15.7%) were diagnosed with dementia. Overall incidence rate for those aged = 60 was 54.6/1000 person years (95% CI 34.1-82.3). The highest incidence rate (97.8/1000 person years) was in the age group 70-74. Standardised incidence ratio for those aged = 65 was 4.98 (95% CI 1.62-11.67). Incidence of dementia in older people with intellectual disabilities are up to five times higher than older adults in the general population. Screening may be useful in this population given the high incident rates, particularly as more effective treatments become available. Studies to explore the underlying aetiological factors for dementia associated with intellectual disability could help to identify novel protective and risk factors.

Abstract: The discovery that adults with Down syndrome (DS) have neuropathological features identical to individuals with sporadic Alzheimer’s disease (AD) played a key role in the identification of the amyloid precursor protein gene on chromosome 21 and resulted in the amyloid cascade hypothesis. Individuals with DS have a lifetime risk for dementia in excess of 90%, and DS is now acknowledged to be a genetic form of AD similar to rare autosomal-dominant causes. Just as DS put the spotlight on amyloid precursor protein mutations, it is also likely to inform the impact of manipulating the amyloid pathway on treatment outcomes in AD. Ironically, however, individuals with DS are usually excluded from AD trials. This review will discuss primary and secondary prevention trials for AD in DS and the potential barriers and solutions to such trials and describe the Europe-wide Horizon21 Consortium to establish a DS-AD prevention clinical trials network.

Stuart, C. Estimating the number of people with Down’s syndrome in Scotland and the cohort at elevated risk of early onset dementia. 
Abstract: The purpose of this study was to ascertain the size of the population of adults with Down syndrome in Scotland and provide a basis for estimating their number and age ranges with dementia. Data were requested from all general practitioners (GP) in Scotland on people with an identified READ code denoting Down syndrome. A statistical weighting model was then applied to account for non-response bias. Findings were that there were 3,261 adults with Down estimated by the application of a statistical weighting model. Of these, 1,118 (34%) were aged between 40 and 59. This age group includes those adults with the highest incidence of early onset dementia. It was not possible to apply a benchmark to the percentage of observed data so as to determine the accuracy of the estimates. Adults with Down have an elevated risk of developing dementia significantly earlier than the general population and require specific age appropriate supports and services to meet their needs both pre- and post-diagnosis. The reality of this is currently not fully realized in either standard practice or national policy concerning the issue.

Abstract: To update the 2011 Canadian guidelines for primary care of adults with intellectual and developmental disabilities (IDD). Family physicians and other health professionals experienced in the care of people with IDD reviewed and synthesized recent empirical, ecosystem, expert, and experiential knowledge. A system was developed to grade the strength of recommendations. Adults with IDD are a heterogeneous group of patients and have health conditions and factors affecting their health that can vary in kind, manifestation, severity, or complexity from those of others in the community. They require approaches to care and interventions that are adapted to their needs. These guidelines provide advice regarding standards of care. References to clinical tools and other practical resources are incorporated. The approaches to care that are outlined here can be applied to other groups of patients that have impairments in cognitive, communicative, or other adaptive functioning. As primary care providers, family physicians play a vital role in promoting the health and well-being of adults with IDD. These guidelines can aid their decision making with patients and caregivers.

Abstract: Patterns of symptoms associated with depression and dementia were examined in 3 aging adults with Down syndrome. A case study approach (Yin, 1994) was employed to identify and link these symptoms. Results of the case analyses provide further insight into distinguishing between depression and dementia in older persons with Down syndrome.

Svanelov, E. An observation study of power practices and participation in group homes for people with intellectual disability. 
Abstract: This study explored how participation constitutes and is constituted by practices of power in group homes for people with intellectual disability. The study used disciplinary power as theoretical perspective and was based on 50 h of observation in two group homes with a total of 15 residents. The analysis identifies practices of power and their relationship to individual agency and participation. The results show that institutional structures construct practices of power that define codes of conduct for the group home residents and their possibility for participation. This study offers implications for the daily lives of residents in group homes for people with intellectual disability.

Abstract: [Non-ID population] The aim of this study was to investigate the effects of group living homes on quality of life and functioning of people with dementia. The study had a quasi-experimental design with a baseline measurement on admission and an effect measurement six months later. Participants were 67 residents in 19 group living homes and 97 residents in seven traditional nursing homes. DQOL and QUALIDEM measured quality of life, functional status was examined with MMSE, IADD, RMBPC, NPI-Q and RISE from RAI. Use of psychotropic drugs and physical restraints was also assessed. Linear and logistic regression analyses analyzed the data. After adjustment for differences in baseline characteristics, residents of group living homes needed less help with ADL and were more socially engaged. There were no differences in behavioral problems or cognitive status. Also after adjusting, two of the 12 quality of life subscales differed between the groups. Residents of group living homes had more sense of aesthetics and had more to do. While there were no differences in prescription of psychotropic drugs, residents of group living homes had less physical restraints. Group living homes had some beneficial effects on its residents, but traditional nursing homes performed well as well. Possible study limitations included the baseline differences between the study groups and the use of different informants on T0 and T1. Future nursing home care may very well be a combination of the best group living care and traditional nursing home care to offer.
Working in group living homes for older people with dementia: The effects on job satisfaction and burnout and the role of job characteristics
Abstract: [Non-ID population] Group living homes are a fast-growing form of nursing home care for older people with dementia. This study seeks to determine the differences in job characteristics of nursing staff in group living homes and their influence on well-being. We examined the Job Demand Control Support (JDCS) model in relation to 183 professional caregivers in group living homes and 197 professional caregivers in traditional nursing homes. Multilevel linear regression analysis was used to study the mediator effect of the three job characteristics of the JDCS-model (demands, control and social support) on job satisfaction and three components of burnout (emotional exhaustion, depersonalization and decreased personal accomplishment). Demands were lower in group living homes, while control and social support from co-workers were higher in this setting. Likewise, job satisfaction was higher and burnout was lower in group living homes. Analysis of the mediator effects showed that job satisfaction was fully mediated by all three psychosocial job characteristics, as was emotional exhaustion. Depersonalization was also fully mediated, but only by control and social support. Decreased personal accomplishment was partially mediated, again only by job characteristics, control and support. This study indicates that working in a group living home instead of a traditional nursing home has a beneficial effect on the well-being of nursing staff, largely because of a positive difference in psychosocial job characteristics.

Taggert, L., Truesdale-Kennedy, M., Ryan, A., McConkey, R.
Examining the support needs of ageing family carers in developing future plans for a relative with an intellectual disability
Abstract: Planning for the future care of adults with an intellectual disability after the main family carer ceases their care, continues to be a sensitive and difficult time posing challenges for service providers internationally. Limited research has been undertaken on this topic because until recently, people with intellectual disability usually pre-deceased their parents. This study examined ageing carers’ preferences for future care and the support systems required to make such future plans. The study was conducted in one region of the United Kingdom with a high proportion of family carers. A mixed methods design was employed. In Stage 1, a structured questionnaire was used to collate information on the health, caregiving demands and future planning preferences of 112 parent and sibling carers; aged 60-94 years. In Stage 2, 19 in-depth semistructured interviews were undertaken with a sample of carers to explore a range of issues around future planning. Over half of the carers were lone carers, mainly female, with many reporting a wide range of health problems. A third of these carers reported that their caregiving resulted in high levels of anxiety. The main preference of the carers was for the person to remain in the family home, with either the family and/or paid staff to support them. A minority of parent carers preferred the person to move into the home of a sibling, although some favored the person moving to a residential facility with other people with intellectual disabilities. The majority of carers did not want their relative to move into an older people’s residential/nursing facility. In the qualitative data, four main themes were identified around future planning: unremitting appraisal, the extent of planning, obstacles encountered and solutions for future planning. Avoidance, lack of guidance and a lack of appropriate residential provision were cited as obstacles to making future plans compounded by the emotional upset experienced by carers in thinking about the future. Findings of this study clearly identify the emotional, informational and practical supports required by these ageing family carers. These findings have national and international relevance in influencing how governments and service providers support parent and sibling carers to proactively plan for the future, and in the development of both in-home and out-of-home options when a family carer can no longer provide care. This is more urgent than ever given the growing numbers of older persons with intellectual disabilities in future decades.

Takenoshita, S., Terada, S., Kuwano, R., Inoue, T., Kurozumi, T., Yamada, N., & Suemitsu, S.
Prevalence of dementia in people with intellectual disabilities without Down syndrome in Japan
Abstract: There are few large-scale studies of the prevalence of dementia in people with intellectual disability who do not have Down syndrome, although Down syndrome is well known to be related with Alzheimer’s disease. We investigated 1831 adults with intellectual disability but without Down syndrome who were residents of a facility for people with intellectual disability in Japan. The caregivers answered a questionnaire, and physicians directly examined the participants suspected of cognitive decline. Of the 1831 patients, 118 were diagnosed with dementia and 51 with mild cognitive impairment (MCI). The prevalence of dementia was 5.6% for the 55 to 64 age group, 13.2% for the 65 to 74 age group, 22.2% for the 75 to 84 age group, and 42.3% for the 85 to 94 age group. The prevalence of MCI was 2.1% for the 55 to 64 age group, 7.4% for the 65 to 74 age group, and 11.5% for the 85 to 94 age group. Sixty-five patients (55.1%) had not been diagnosed with dementia before the survey. People with intellectual disability without Down syndrome may develop dementia at an earlier age than those without intellectual disability. Our nationwide survey suggested there are many undiagnosed dementia patients in the community of people with intellectual disability.

Takenoshita, S., Kuwano, R., Inoue, T., Kurozumi, T., Choju, A., Suemitsu, S., & Yamada, N.
Prevalence of dementia in people with intellectual disabilities: Cross-sectional study
Abstract: There are only a few studies of the prevalence of dementia in people with intellectual disability (ID) without Down syndrome (DS), and there is a large difference in the prevalences between reported studies. Moreover, the prevalence of mild cognitive impairment (MCI) in ID has not been reported. We aimed to evaluate the prevalence of dementia in adults of all ages and the prevalence of MCI in people with ID. Furthermore, we tried to clarify the differences depending on the various diagnostic criteria. The survey included 493 adults with ID at 28 facilities in Japan. The caregivers answered a questionnaire, and physicians directly examined the participants who were suspected of cognitive decline. Dementia and MCI were diagnosed according to ICD-10, DC-LD, and DSM-5 criteria. The prevalence of dementia was 0.8% for the 45 to 54 years old group, 3.5% for the 55 to 64 years old group, and 13.9% for the 65 to 74 years old group in people with ID without DS. The prevalence of MCI was 3.1% for patients 45 to 54, 3.5% for patients 55 to 64, and 2.8% for patients 65 to 74 with ID without DS. DSM-5 was the most inclusive in diagnosing dementia and MCI in people with ID. People with ID without DS may develop dementia and MCI at an earlier age and higher rate than the general population. Among the diagnostic criteria, DSM-5 was the most useful for diagnosing their cognitive impairment.

Takenoshita, S., Terada, S., Kuwano, R., Inoue, T., Kurozumi, T., Choju, A., Suemitsu, S., & Yamada, N.
Validation of the Japanese version of the Dementia Screening Questionnaire for Individuals with Intellectual Disabilities.
Abstract: Dementia in people with intellectual disabilities (IDs) is difficult to detect because of preexisting cognitive deficits. An effective screening method is required. The Dementia Screening Questionnaire for Individuals with Intellectual Disabilities (DSQIID) was developed as an observer rating tool to screen dementia in people with ID. The aim of this study was to verify the
screening accuracy of the DSQIID for Japanese people with ID. Four-hundred ninety-three subjects with ID participated in this study. Caregivers who had observed the participants for more than 2 years scored the Japanese version of the DSQIID (DSQIID-J) of the participants. Three doctors examined participants directly and diagnosed dementia using the Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition criteria. To identify the key screening items that predict dementia, the specificities of a single and pairs of items with 100% sensitivity were evaluated relative to the dementia diagnosis. Of 493 participants, 34 were people with Down syndrome (DS), and 459 were people without DS. Seventeen participants were diagnosed with dementia. The suitable cut-off score of the DSQIID-J was 10/11 (sensitivity 100% and specificity 96.8%) for screening dementia. The inter-rater reliability, test-retest reliability and internal consistency of the DSQIID-J were excellent. Regarding key items, there was no single item with 100% sensitivity, and the best two-item combination was the pair of ‘Cannot dress without help’ and ‘Walks slower’ (sensitivity 100% and specificity 93.5%). We identified several important question items of the DSQIID-J related to the diagnosis of dementia in people with ID. The DSQIID-J is a useful screening tool for dementia in adults with ID.

Abstract: The behavioral and emotional changes associated with Alzheimer's disease (AD) are compared for individuals with Down syndrome and AD and individuals with AD from the general population (AD-only). The primary caregivers of 30 people with Down syndrome and AD and 30 people with AD-only completed the BEHAVE-AD and the Apathy subscale of the CERAD. As well, behavioral observations at adult day programs were undertaken with selected participants (n=26). The Down syndrome group experienced fewer delusions and had lower total scores on the BEHAVE-AD, indicating fewer problem behaviors overall. Day program observations suggested that the AD-only group were more likely to be sedentary and observe the activities of others, while the Down syndrome group were more physically active. Improving our understanding of the similarities and differences between these two groups may help facilitate the integration of individuals with Down syndrome into adult day programs for the general population.

Abstract: More years of education have been found to be associated with a lower rate of Alzheimer disease (AD) in individuals without intellectual disability. It has been proposed that education reflects greater 'synaptic reserve' and that greater synaptic reserve may defer the development of AD. The present study compared individuals with Down's syndrome (DS) who were found to have symptoms of dementia with those who remained symptom-free to determine if the two groups differed in their level of education, employment, recreational activities, years in an institution or overall level of cognitive functioning. Thirty-five adults with DS aged between 29 and 67 years were assessed. The participants were recruited from a community health facility and included individuals with a wide range of ability levels. Neuropsychological testing, caregiver report and the Dementia Scale for Down Syndrome (Gedye 1995) were used to identify decline in participants over periods of 6 months to 3 years. After the effect of age was statistically removed, multiple regression analyses revealed that level of cognitive functioning was significantly associated with decline such that a higher level of cognitive functioning predicted less decline. None of the environmental variables (i.e. educational level, years in an institution and employment) were directly associated with decline; however, a post hoc regression using level of cognitive functioning as the outcome variable revealed that level of cognitive functioning itself was associated with these environmental variables. A higher level of cognitive functioning was associated with fewer cases of dementia in individuals with DS, and level of cognitive functioning appears to be associated with environmental factors such as level of education, years in an institution and employment. The present findings suggest that environmental interventions aimed at improving level of cognitive functioning may also be useful in deferring the onset of dementia.

Thalen, M., Volkers, K.M., van Oorsouw, W.M.W.J., & Embregts, P.J.C.M. Psychosocial interventions for older people with intellectual disabilities and the role of support staff. A systematic review. Journal of Applied Research in Intellectual Disabilities, 2022 Mar;35(2):312-337. doi: 10.1111/jar.12953. Epub 2021 Nov 15. Abstract: The life expectancy of people with intellectual disabilities has increased. The implications of ageing have resulted in changes in their support needs and challenges to support staff. Access to evidence-based strategies for support staff providing care to elderly with intellectual disabilities remains scarce. A systematic review was conducted to provide an overview of available psychosocial interventions. Four databases were searched and assessed according to the PRISMA guidelines. A narrative, integrative method of analysis was conducted to synthesize quantitative and qualitative data. The 36 studies included in the review reported on interventions aimed at either identifying and meeting the needs or perceptions of older individuals or at improving their behavior and skills. Furthermore, the role of support staff in the implementation of interventions was either active, assisting or undefined. This overview of studies could contribute to the existing body of knowledge and help to optimize psychosocial support for a growing population.

Thase, M.E., Tigner, R., Smeltzer, D.J., & Liss, L. Age-related neuropsychological deficits in Down's syndrome Biological Psychiatry, 1984 Apr; 19(4), 571-585. https://pubmed.ncbi.nlm.nih.gov/6234031/ Abstract: Down's syndrome (DS) has been suggested as a high-risk condition for Dementia Alzheimer's type (DAT). In the present study, neuropsychological variables were assessed in 165 DS subjects and 163 matched controls with intellectual disability. Overall, DS subjects had lower scores for orientation, digit span, visual memory, object-naming, and general knowledge, as well as more "released" reflexes. Impairments were most evident in DS subjects greater than 50 years old. These findings provide further support for an association between aging and DAT in Down's syndrome. Methodological issues and areas for future research are discussed.


Thompson, D.J., Ryrie, I., & Wright, S. People with intellectual disabilities living in generic residential services for older people in the UK Journal of Applied Research in Intellectual Disabilities, 2004, 17(2), 101-108. https://doi.org/10.1111/j.1365-2322.2004.00187.x Abstract: As part of a UK program of work focusing on older people with ID, the circumstances of those who reside in generic services for older people were investigated. Some 215 people with ID were identified living in 150 homes. They were significantly younger than other residents and were placed in these homes more because of organizational change or the aging/death of family
carers, rather than due to their own needs. Of the residents, 24 adults had Down syndrome, 8 of whom were noted to have dementia. Of the 215, 45 had dementia. Average age of people with DS upon entry was 60 and those remaining at the homes was about 65.

Abstract: Authors estimated dementia incidence rates, life expectancies with and without dementia, and percentage of total life expectancy without dementia. Authors studied 3605 members of Group Health (Seattle, WA) aged 65 years or older who did not have dementia at enrollment to the Adult Changes in Thought study between 1994 and 2008. We estimated incidence rates of Alzheimer's disease and dementia, as well as life expectancies with and without dementia, defined as the average number of years one is expected to live with and without dementia, and percentage of total life expectancy without dementia. Dementia incidence increased through ages 85 to 89 years (74.2 cases per 1000 person-years) and 90 years or older (105 cases per 1000 person-years). Life expectancy without dementia and percentage of total life expectancy without dementia decreased with age. Life expectancy with dementia was longer in women and people with at least a college degree. Percentage of total life expectancy without dementia was greater in younger age groups, men, and those with more education. Efforts to delay onset of dementia, if successful, would likely benefit older adults of all ages.

Torr, J., & Davis, R.
Abstract: Increasing numbers of people with intellectual disability are now living well into old age. This paper will review the recent literature pertaining to the mental health of older people with intellectual disability. Overall, the prevalence of mental health problems is high in adults of all ages with intellectual disability. A major epidemiological study did not report sufficient detail to examine the effect of ageing on specific disorders or the differential effects of ageing and early mortality in people with Down's syndrome. At least a third of people with Down's syndrome can expect to develop Alzheimer's disease in middle age whilst for other people with intellectual disability, Alzheimer's disease is probably no more common than in the general population. Diagnosis and management of dementia is complicated by the high rates of comorbid physical and mental health problems. Overall, mental health problems in older people with intellectual disability are similar to younger people with intellectual disability, however there are more cases of dementia and physical health problems. Further research is needed to improve our understanding of the effects of ageing on the mental health and care needs of older people with intellectual disability.

Torr, J., Strydom, A., Patti, P. & Jokinen N.
Abstract: The life expectancy of adults with Down syndrome has increased dramatically over the last 30 years, leading to increasing numbers of adults with Down syndrome now living into middle and old age. Early-onset dementia of the Alzheimer type is highly prevalent in adults with Down syndrome in the sixth decade, and this has overshadowed other important conditions related to aging among adults with Down syndrome. The authors’ aim was to update and summarize current knowledge on these conditions, and examine causes of morbidity and mortality in older people with Down syndrome by conducting a systematic review of the published literature for the period: 1993–2008. They reviewed English-language literature drawn from searches in the electronic databases Medline, CINAHL, and PsycINFO, as well as supplementary historical papers. The authors conclude that functional decline in older adults with Down syndrome cannot be assumed to be due only to dementia of the Alzheimer type (which is not inevitable in all adults with Down syndrome). Functional decline may be the result from a range of disorders, especially sensory and musculoskeletal impairments. Given the high rates of early-onset age-related disorders among adults with Down syndrome, programmatic screening, monitoring, and preventive interventions are required to limit secondary disabilities and premature mortality. With respect to assessment and treatment, in the absence of specialist disability physicians, geriatricians have a role to play.

Tsang, W., Oliver, D., & Triantafyllopoulou, P.
Abstract: Adults with intellectual disabilities are an at-risk group of developing dementia. In the absence of a cure for dementia, emphasis on treatment is the promotion of Quality of life (QoL). The aim of this review is to identify and describe QoL tools for people with intellectual disabilities and dementia. A systematic review was carried out using 10 databases and papers from up to March year 2021. Two instruments were identified and examined. The QoL in late-stage dementia, which showed evidence of good levels of internal consistency, intra-rater reliability, test-retest reliability, and convergent validity. The Dementia Quality of Life - proxy was also used; however, its psychometric properties have yet to be studied within the intellectual disabilities population. Due to the degenerative nature of dementia involving cognitive impairment such as memory and language difficulties, there can be significant barriers in capturing the subjective experience of QoL in a person with dementia. Therefore, it may not be possible to capture the patient’s perceived QoL from self-report measures. Interpretation of concepts can be problematic, especially when life situation of the respondent is vastly different to that of a researcher. Also, the conceptualisation of QoL can vary in different stages of dementia in the general population, such that in the early stages of dementia enjoyment of daily activities is a relevant domain for QoL, but may no longer seem to be applicable in severe or advanced dementia. It is recommended instruments should be developed and psychometrically tested specifically for adults with intellectual disabilities and dementia to help inform policy makers, measure outcomes of interventions and personal outcomes.

Tse, M.M., Kwan, R.Y, & Lau, J.L.
Abstract: The increasing longevity of people with intellectual disability is testimony to the positive developments occurring in medical intervention. Nonetheless, early-onset age-related issues and concerns cause deterioration of their overall wellbeing. This paper aimed to explore the issues and concerns about individuals with intellectual disability as they age. Articles that discussed people older than 30 years with an intellectual disability and those that identified ageing health issues and concerns were included. Only studies reported in English from 1996 to 2016 were included. We searched PubMed, Google Scholar, and Science Direct using the terms 'intellectual disability', 'ageing', 'cognitive impairment', 'health', and 'screening'. Apart from the early onset of age-related health problems, dementia is more likely to develop by the age of 40 years in individuals with intellectual disability. Geriatric services to people with intellectual disability, however, are only available for those aged 60 years and older. Cognitive instruments used for the general population are not suitable for people with intellectual disability because of floor effects. In Hong Kong, the
Adverse effects of phenytoin given for late-onset seizures in adults with Down syndrome.


Summary (no abstract provided): A brief report that indicates the adverse effects of therapeutic levels of phenytoin and the improvement observed when phenytoin was replaced with other antiepileptics in 17 adults with DS, Alzheimer disease (AD) and late-onset seizures (LOS). The reported deterioration in the patients’ condition was found to be due to the adverse effects from phenytoin and not to AD. It was suggested that practitioners avoid prescribing phenytoin to treat LOS in persons with DS and AD. If phenytoin is already prescribed, it should be replaced with another anticonvulsive agent.


Medical care of adults with Down syndrome: A clinical guideline


Abstract: Down syndrome is the most common chromosomal condition, and average life expectancy has increased substantially, from 25 years in 1983 to 60 years in 2020. Despite the unique clinical comorbidities among adults with Down syndrome, there are no clinical guidelines for the care of these patients. To develop an evidence-based clinical practice guideline for adults with Down syndrome. The Global Down Syndrome Foundation Medical Care Guidelines for Adults with Down Syndrome Workgroup (n=13) developed 10 Population/ Intervention/ Comparison/Outcome (PICO) questions for adults with Down syndrome addressing multiple clinical areas including mental health (2 questions), dementia, screening or treatment of diabetes, cardiovascular disease, obesity, osteoporosis, atlantoaxial instability, thyroid disease, and celiac disease. These questions guided the literature search in MEDLINE, EMBASE, PubMed, PsycINFO, Cochrane Library, and the TRIP Database, searched from January 1, 2000, to February 26, 2018, with an updated search through August 6, 2020. Using the GRADE (Grading of Recommendations, Assessment, Development, and Evaluation) methodology and the Evidence-to-Decision framework, in January 2019, the 13-member Workgroup and 16 additional clinical and scientific experts, nurses, patient representatives, and a methodologist developed clinical recommendations. A statement of good practice was made when there was a high level of certainty that the recommendation would do more good than harm, but there was little direct evidence. From 11295 literature citations associated with 20 PICO questions, 20 relevant studies were identified. An updated search identified 2 additional studies, for a total of 22 included studies (3 systematic reviews, 19 primary studies), which were reviewed and synthesized. Based on this analysis, 14 recommendations and 4 statements of good practice were developed. Overall, the evidence base was limited. Only 1 strong recommendation was formulated: screening for Alzheimer-type dementia starting at age 40 years. Four recommendations (managing risk factors for cardiovascular disease and stroke prevention, screening for obesity, and evaluation for secondary causes of osteoporosis) agreed with existing guidance for individuals without Down syndrome. Two recommendations for diabetes screening recommend earlier initiation of screening and at shorter intervals given the high prevalence and earlier onset in adults with Down syndrome. These evidence-based clinical guidelines provide recommendations to support primary care of adults with Down syndrome. The lack of high-quality evidence limits the strength of the recommendations and highlights the need for additional research.

Tuffrey-Wijne I.


Abstract: How to tell bad news to people with dementia? First assess the person’s current framework of knowledge, then give the new chunks of information one by one, then finally check and reassess the person’s...
knowledge. This might be useful for bereaved people with dementia to fill the gap.

**Tuffrey-Wijne, I., & Watchman, K.**
**Breaking bad news to people with learning disabilities and dementia**
DOI:10.7748/ldp.18.7.16.e1672
Abstract: People with learning disabilities are now enjoying a longer life expectancy than ever before as a result of enhanced medical and social interventions and improved quality of life. Some, particularly individuals with Down’s syndrome, are susceptible to dementia at a significantly younger age than the average age of onset in the rest of the population. Currently, there is limited guidance on how to talk to people with learning disabilities about dementia and, until such information is shared, individuals cannot be positioned as an authority on their own condition. The new model presented here suggests a way of supporting staff and families to have enabling conversations about dementia that center on the person’s current situation, level of understanding and capacity.

**Tuffrey-Wijne, I., & Watchman, K.**
**Sharing the diagnosis of dementia: Breaking bad news to people with intellectual disabilities.**
Abstract: Not available - see Tuffrey-Wijne & Watchman 2015

**Turky, A., Felce, D., Jones, G., & Kerr, M.**
A prospective case control study of psychiatric disorders in adults with epilepsy and intellectual disability
Epilepsia, 2011 July, 52(7), 1223-1230.
https://doi.org/10.1111/j.1528-1167.2011.03044.x
Abstract: No study to date has prospectively investigated the impact of epilepsy on psychiatric disorders among adults with an intellectual disability (ID). This study aimed to determine prospectively the influence of epilepsy on the development of psychiatric disorders in adults with ID. Psychiatric symptoms were measured prospectively over a 1-year period among 45 adults with ID and active epilepsy and 45 adults with ID without epilepsy, matched on level of ID. The 1-year incidence rate (IR) of common Axis I psychiatric disorders was compared with and without controlling for possible confounding factors. Total psychiatric symptom scores over the period were compared between the two groups using repeated-measures analysis of covariance. Adults with epilepsy and ID had a more than seven times increased risk for developing psychiatric disorders, particularly depression and unspecified disorders of presumed organic origin, including dementia, over a 1-year period compared to those with ID only. Comparison of the psychiatric scores showed the epilepsy group to have significantly higher unspecified disorder and depression symptom scores. The findings point to an increased risk of depression and unspecified disorders, including dementia, among adults with ID and epilepsy. Further exploration of the nature and treatment of these unspecified disorders may help the care of people with epilepsy and ID.

**Tyler, C.V., & Shank, J.C.**
Dementia and Down syndrome
Abstract: Case report of a 43-year-old woman with Down syndrome and progressive decline over three years that was attributed to dementia of the Alzheimer's type. Authors describe the medical conditions evident during decline, whilst living with her family. Identifies typical features associated with decline for persons with Down syndrome and defines areas for concern during examinations by physicians.
dementia. The 16 papers range from the theoretical to the practical.

Urv, T.K., Zigman, W.B., & Silverman, W.

Abstract: Changes in psychiatric symptoms related to specific stages of dementia were investigated in 224 adults 45 years of age or older with Down syndrome. Findings indicate that psychiatric symptoms are a prevalent feature of dementia in the population with Down syndrome and that clinical presentation is qualitatively similar to that seen in Alzheimer's disease within the general population. Psychiatric symptoms related to Alzheimer's disease vary by the type of behavior and stage of dementia, but do not seem to be influenced by sex or level of premorbid intellectual impairment. Some psychiatric symptoms may be early indicators of Alzheimer's disease and may appear prior to substantial changes in daily functioning. Improvements in understanding the progression of dementia in individuals with Down syndrome may lead to improved diagnosis and treatment.

van Hoof, J., & Kort, H.

Abstract: The vast majority of older adults want to remain living independently at home, with or without a sufficient amount of professional home care, even when overall health is starting to decline. The ageing of society and the increase in the number of very old elders goes together with an increase in the number of people with dementia. About two thirds of the diagnosed people in the Netherlands live at home. Dementia has severe implications to the quality of daily life, in particular to independent functioning. This sets extra demands to living environments. Older adults with dementia and their partners ask for living environments that support independence, compensate for declining vitality, and lower the burden of family care. For this purpose, a first concept of a design for a dementia dwelling is presented in this paper, which incorporates modifications in terms of architecture, interior design, the indoor environment, and technological solutions. These design features were derived from literature search and focus group sessions. Current design guidelines are frequently based on practical experience only, and therefore, more systematic field research should be carried out to find evidence for the various design modifications. Also, it needs to be studied how the design features of the dementia dwelling can be incorporated into the existing housing stock.

van Hoof, J., Kort, H.S.M., van Waarde, H., & Blom, M.M.

Abstract: In Western societies, the vast majority of people with dementia live at home and wish to remain doing so for as long as possible. Aging in place can be facilitated through a variety of environmental interventions, including home modifications. This article provides an overview of existing design principles and design goals, and environmental interventions implemented at home, based on literature study and additional focus group sessions. There is a multitude of design principles, design goals, and environmental interventions available to assist with activities of daily living and functions, although few systematic studies have been conducted on the efficacy of these goals and interventions. The own home seems to be a largely ignored territory in research and government policies, which implies that many problems concerning aging in place and environmental interventions for dementia are not adequately dealt with.

van Hoof, J., Kort, H.H., Duijnstee, M.S., Rutten, P.P., & Hensen, J.J.

Abstract: There are currently about 6 million - mainly older - people with dementia in the European Union. With ageing, a number of sensory changes occur. Dementia syndrome exacerbates the effects of these sensory changes and alters perceptual stimuli. People with dementia have an altered sensitivity for indoor environmental conditions, which can induce problematic behavior with burdensome symptoms to both the person with dementia and the family carer. This paper, based on literature review, provides an overview of the indoor environmental parameters, as well as the integrated design and implementation of relevant building systems. The overview is presented in relation to the intrinsic ageing of senses, the responses of older people with dementia and the impact on other relevant stakeholders through the combined use of the International Classification of Functioning, Disability and Health, and the Model of Integrated Building Design. Results are presented as indicators of the basic value, functional value and economic value, as well as a synthesis of building-related solutions. Results can help designers and building services engineers to create optimal environmental conditions inside the living environments for people with dementia, and can be used to raise awareness among health care professionals about the influence of the indoor environment on behavior of the person with dementia.

Vaughan, R.M., McGee, C., Guerin, S., Tyrrell, J., & Dodd, P.

Abstract: This study analysed retrospective data on twenty adults with Down’s syndrome (DS), who are clients of a specialist service in Dublin. The aim was to compare the practice of this service on diagnosis and treatment of dementia, with the consensus recommendations of the Royal College of Psychiatry, British Psychological Society and National Institute for Health and Care Excellence. Further aims were to establish average time to make a diagnosis and commence pharmacotherapy, and to describe tolerability to acetylcholinesterase inhibitors. It was found that screening for dementia did not take place before the age of 30yrs with the mean age for first assessment being 48yrs and average age at diagnosis being 51yrs. Average length of time from first identified symptoms to diagnosis was 1.3yrs. Of those diagnosed, 83% were prescribed acetylcholinesterase inhibitors but the authors were concerned at the continued use of the drug even when there appeared to be no benefit to the person. They found that a wide range of assessments were in use and that longitudinal assessment and follow up were not practised. The authors recommend the streamlining of assessment tools and repeating assessment on a longitudinal basis.


Abstract: There is large cross-national variation in the characteristics of small, domestic-style care settings which emphasize normalized living. However, a systematic overview of existing types is lacking. This study provides an international comparison of the care concepts which have adopted a homelike philosophy in a small-scale context. Insight into their characteristics is vital for theory, planning and implementation of such dementia care settings. A literature search was performed using various electronic databases, including PubMed, Medline, CINAHL and PsycINFO. In addition, “gray” literature was identified on the internet. Concepts were analyzed according to five main characteristics: physical setting, number of residents, residents’ characteristics, domestic characteristics and care concept. 75 papers were included covering 11 different concept types in various countries. Similarities among concepts reflected a focus on meaningful activities centered around the daily household. Staff have
integrated tasks and are part of the household, and archetypical home-style features, such as kitchens, are incorporated in the buildings. Differences among concepts were found mainly in the physical settings, numbers of residents and residents' characteristics. Some concepts have become regular dementia care settings, while others are smaller initiatives. The care concepts are implemented in various ways with a changing staff role. However, many aspects of these small, home-like facilities remain unclear. Future research is needed, focusing on residents' characteristics, family, staff and costs.


Trajectories of decline in cognition and daily functioning in preclinical dementia Alzheimer's & Dementia, 2016, Feb, 12(2), 144-153. doi: 10.1016/j.jalz.2015.08.001. Epub 2015 Sep 9. Abstract: [Note: this article refers to adults in the general population] Although preclinical dementia is characterized by decline in cognition and daily functioning, little is known on their temporal sequence. We investigated trajectories of cognition and daily functioning in preclinical dementia, during 18 years of follow-up. In 856 dementia cases and 1712 controls, we repetitively assessed cognition and daily functioning with memory complaints, mini-mental state examination (MMSE), instrumental activities of daily living (IADL), and basic activities of daily living (BADL). Dementia cases first reported memory complaints 16 years before diagnosis, followed by decline in MMSE, IADL, and finally BADL. Vascular dementia related to earlier decline in daily functioning but later in cognition, compared with Alzheimer's disease. Higher education related to larger preclinical cognitive decline, whereas apolipoprotein E (APOE) e4 carriers declined less in daily functioning. These results emphasize the long hierarchical preclinical trajectory of functional decline in dementia. Furthermore, they show that various pathologic, environmental, and genetic factors may influence these trajectories of decline.


Longitudinal clinical and cognitive changes along the Alzheimer disease continuum in Down syndrome. JAMA Netw Open. 2022 Aug 1;5(8):e2225573. doi: 10.1001/jamanetworkopen.2022.25573. PMID: 39530282; PMCID: PMC9356319. Abstract: Alzheimer disease (AD) is the main medical problem in adults with Down syndrome (DS). However, the associations of age, intellectual disability (ID), and clinical status with progression and longitudinal cognitive decline have not been established. To examine clinical progression along the AD continuum and its related cognitive decline and to explore the presence of practice effects and floor effects with repeated assessments, authors used a single-center cohort study of adults (aged >18 years) with DS with different ID levels and at least 6 months of follow-up between November 2012 and December 2021. The data are from a population-based health plan designed to screen for AD in adults with DS in Catalonia, Spain. Individuals were classified as being asymptomatic, having prodromal AD, or having AD dementia. The main outcome was clinical change along the AD continuum. Cognitive decline was measured by the Cambridge Cognitive Examination for Older Adults With Down Syndrome and the modified Cued Recall Test. Authors assessed a total of 632 adults with DS (mean [SD] age, 42.6 [11.4] years; 292 women [46.2%]) with 2847 evaluations (mean [SD] follow-up, 28.8 [18.7] months). At baseline, there were 436 asymptomatic individuals, 69 patients with prodromal AD, and 127 with AD dementia. After 5 years of follow-up, 17.1% (95% CI, 12.5%-21.5%) of asymptomatic individuals progressed to symptomatic AD in an age-dependent manner (0.6% [95% CI, 0%-1.8%] for age <40 years; 21.1% [95% CI, 8.0%-32.5%] for age 40-44 years; 41.4% [95% CI, 23.1%-55.3%] for age 45-49 years; 57.5% [95% CI, 38.2%-70.8%] for age >75 years; P < .001), and 94.1% (95% CI, 84.6%-98.0%) of patients with prodromal AD progressed to dementia with no age dependency. Cognitive decline in the older individuals was most common among those who progressed to symptomatic AD and symptomatic individuals themselves. Importantly, individuals with mild and moderate ID had no differences in longitudinal cognitive decline despite having different performance at baseline. This study also found practice and floor effects, which obscured the assessment of longitudinal cognitive decline. This study found an association between the development of symptomatic AD and a high risk of progressive cognitive decline among patients with DS. These results support the need for population health plans to screen for AD-related cognitive decline from the fourth decade of life and provide important longitudinal data to inform clinical trials in adults with DS to prevent AD.

Visser, F.E., Aldenkamp, A.P., van Huggelen, A.C., Kuilman, M., Overweg, J., & van Wijk, J.

Prospective study of the prevalence of Alzheimer-type dementia in institutionalized individuals with Down syndrome American Journal on Mental Retardation, 1997, Jan, 101(4), 400-412. https://pubmed.ncbi.nlm.nih.gov/9017086/ Abstract: Institutionalized patients with Down syndrome (n = 307) were monitored for 5 to 10 years prospectively to determine prevalence of Alzheimer-type dementia. Clinical signs, cognitive functioning, and EEGs were
assessed. When possible, postmortem neuropathological examinations were conducted. Progressive mental and physical deterioration was found for 56 of the residents. **Mean age at onset of dementia was 56 years.** Prevalence increased from 11% between ages 40 and 49 to 77% between 60 and 69. All patients 70 and over had dementia. Neuropathological findings were consistent with clinical diagnosis. Use of a dementia checklist, cognitive skills inventory, and EEG reliably detected Alzheimer-type dementia at an early stage.

Walaszek, A., Schroeder, M., Krainer, J., Pritchett, G., Wilcenski, M., Endicott, S., Albrecht, T., Carlsson, C.M., & Mahoney, J. Effectively training professional caregivers to screen and refer persons with dementia and intellectual/developmental disability


Abstract: By age 40, almost all people with Down syndrome, the most common cause of intellectual/developmental disability (IDD), have neuropathological changes consistent with Alzheimer’s disease; by age 60, about half have dementia. Detecting dementia in persons with IDD can be challenging because baseline cognitive impairment can be severe and because persons with IDD may have difficulty reporting symptoms. The National Task Group Early Detection Screen for Dementia (NTG-EDSD) was developed to aid detection of cognitive impairment in adults with IDD. We implemented an educational curriculum to increase the ability of professional caregivers to screen for dementia in persons with IDD using the NTG-EDSD. In November 2018 to April 2019, we held five training sessions for professional caregivers of persons with IDD, partnering with various managed care organizations (MCO), aging and disability resource centers, adult day programs, and adult family homes. We assessed knowledge and attitudes at baseline, immediately after training, and one week, one month and six months after training. Participants (N=154) included direct care workers, case managers, healthcare providers, and other social services staff. Participants reported a marked increase in confidence in their ability to detect changes associated with mild cognitive impairment or dementia (p<0.001), decline in activities of daily living (p=0.02), and changes in behavior and affect (p<0.001). Satisfaction with the training was very high, and 94.0% of participants agreed or strongly agreed they could use the NTG-EDSD tool with their clients. Following the training, one MCO we partnered with, serving 82 of 72 counties in Wisconsin, made the NTG-EDSD a standard part of its assessment of adults with DS starting at age 40. Authors note that a wide variety of social services and healthcare professionals can be effectively trained to screen for dementia in persons with IDD using a standardized screening tool, the NTG-EDSD. Satisfaction with the training was high, and use of the NTG-EDSD was thought to be feasible. This educational intervention led to change in practice at a systems level within an MCO. Next steps could include assessing impact of such training on the quality of life and healthcare outcomes of persons with IDD.


Abstract: By age 60, 60% of adults with Down syndrome (DS) have dementia. Detecting dementia in persons with intellectual disability (ID) can be challenging because their underlying cognitive impairment can confound presentation of dementia symptoms and because adults with ID may have difficulty reporting symptoms. The National Task Group Early Detection Screen for Dementia (NTG-EDSD) was developed to aid detection of report of cognitive impairment in adults with ID. We implemented an educational curriculum using the NTG-EDSD and evaluated the impact of the intervention on professional caregivers’ self-assessed capacity to identify persons with ID and dementia. We held five in-person training sessions for professional caregivers of persons with ID, partnering with various managed care organizations and social services agencies. We assessed knowledge and attitudes at baseline; immediately after training; and 1 week, 1 month, and 6 months after training. A total of 154 direct care workers, case managers, health-care providers, and other social services staff attended the trainings. Satisfaction with the NTG-EDSD training was high; 94% of attendees agreed or strongly agreed that they could use the NTG-EDSD with their clients. After training, attendees reported a marked increase in confidence in their ability to track various health circumstances and detect functional decline in their clients, although some gains were not sustained over time. As a result of the training, one managed care organization made the NTG-EDSD a standard part of its assessment of adults with DS starting at age 40. Social services and health-care professionals can learn to document signs of cognitive decline in adults with ID using the NTG-EDSD. Attendees were highly satisfied with the training, experienced an increase in confidence in their care of persons with ID, and found the NTG-EDSD feasible to use. Because not all gains were sustained over time, booster trainings may be necessary.

Walker, B., MacBryer, S., Jones, A., & Law, J. Interinformant agreement of the dementia questionnaire for people with learning disabilities


Abstract: Because of difficulties with neuropsychological assessments for dementia in people with intellectual disabilities, professionals in clinical practice have relied heavily on carer interviews, one of the most widely used being the Dementia Questionnaire for People with Learning Disabilities (DLD – Evenhuis et al. 2006 Dementia questionnaire for people with intellectual disabilities manual (second edition). Amsterdam, Netherlands, Harcourt Test Publishers). Because dementia is indicated by the magnitude of changes in scores between longitudinal assessments, interinformant agreement is paramount. We carried out the DLD interview independently with two carers for each of 26 people with Down syndrome. Only 15% of pairs of carers achieved ‘good’ agreement. Levels of agreement varied widely across the DLD subscales and individual questions. Interinformant agreement was better for less able people with Down syndrome than for more able individuals.

Walker, C.A., & Walker, A. Uncertain Futures: people with learning difficulties and their ageing family carers


Abstract: This monograph provides an overview of research, policy and practice relating to service responses to adults with learning difficulties living at home with older family carers in the UK. The authors’ premise is that as life expectancy increases, a growing proportion of people with learning difficulties continues to live with family members, most frequently parents, whose caring role is being extended into their own advanced old age. Highlighted are some of the issues raised by service users, carers and service providers, including care for someone with diminishing abilities. The text argues that there is urgent need for the paid service sector to work with families to provide the necessary support and planning to take the uncertainty out of the future.


Abstract: Individuals with Down syndrome (DS) are at high risk for dementia, specifically Alzheimer’s disease (AD). However, many measures regularly used for the detection of AD in the general population are not suitable for individuals
with DS. Some measures, including the Severe Impairment Battery (SIB), Brief Praxis Test (BPT), and Dementia Questionnaire for Persons with Intellectual Disabilities (DMR), have been used in clinical trials and other research with this population. Validity research is limited, however, particularly regarding identification of predementia symptoms in the DS population. The current project presents baseline cross-sectional SIB, BPT, and DMR performance in order to characterize their ability to discriminate normal cognition, possible AD, and probable AD in DS. Baseline SIB, BPT, and DMR performances from 117 individuals were analyzed as part of a large longitudinal cohort of aging individuals with DS. Receiver operating characteristic (ROC) curves were calculated to investigate accuracy in differentiating levels of dementia status. In comparing no/possible AD vs. probable AD, the SIB and BPT exhibited fair discrimination ability (AUC = .78 and .79, respectively). In comparing no/possible AD vs. probable AD, the DMR exhibited good discrimination ability (AUC = .89), with qualitatively similar performance of the DMR-Cognitive and DMR-Social subscales (AUC = .89 and .83, respectively). In comparing no AD vs. possible AD, the SIB and BPT failed to differentiate these groups (AUC = .53 and .55, respectively), whereas the DMR exhibited good differentiation (AUC = .80). Authors note that the results suggest that the SIB, BPT, and DMR are able to reasonably discriminate consensus dementia diagnoses in individuals with DS, supporting their continued use in the clinical assessment of dementia in DS. Specifically, the DMR, based on informant ratings of social and cognitive behaviors of daily living, outperformed the SIB and BPT, tests of cognitive performance, in discriminating no/possible AD vs. probable AD as well as no AD vs. possible AD. Such findings suggest that the DMR is better equipped to identify symptoms of overt dementia as well as predementia in this population. Findings reinforce the importance of including informant behavior ratings in assessment of this population.

Abstract: Individuals with Down syndrome (DS) are at high risk for dementia, specifically Alzheimer's disease. However, many measures regularly used for the detection of dementia in the general population are not suitable for individuals with DS due in part to floor effects. Some measures, including the Severe Impairment Battery (SIB), Brief Praxis Test (BPT) and Dementia Scale for People with Learning Disabilities (DLD), have been used in clinical trials and other research with this population. Validity research is limited, particularly regarding the use of such tools for detection of prodromal dementia in the DS population. The current project presents baseline cross-sectional SIB, BPT and DLD performance in order to characterise their predictive utility in discriminating normal cognition, possible dementia and probable dementia in adult DS. Baseline SIB, BPT and DLD performances from 100 individuals (no dementia = 68, possible dementia = 16 & probable dementia = 16) were examined from a longitudinal cohort of aging individuals with DS. Receiver operating characteristic curves investigated the accuracy of these measures in relation to consensus dementia diagnoses, diagnoses which demonstrated high percent agreement with the examining neurologist's independent diagnostic impression. The SIB and BPT exhibited fair discrimination ability for differentiating no/possible versus probable dementia [area under the curve (AUC) = 0.61 and 0.66, respectively]. The DLD exhibited good discrimination ability for differentiating no versus possible/probable dementia (AUC = 0.75) and further demonstrated better performance of the DLD Cognitive subscale compared with the DLD Social subscale (AUC = 0.77 and 0.67, respectively). Results suggest that the SIB, BPT and DLD are able to reasonably discriminate consensus dementia diagnoses in individuals with DS, supporting their continued use in the clinical assessment of dementia in DS. The general performance of these measures suggests that further work in the area of test development is needed to improve on the AUCs for dementia status discrimination in this unique population. At present, however, the current findings suggest that the DLD may be the best option for reliable identification of prodromal dementia in this population, reinforcing the importance of including informant behaviour ratings in assessment of cognition for adults with DS.

Walsh, D.M., Doran, E., Silverman, W., Tournay, A., Movsesyan, N., & Lott, I.T.
Rapid assessment of cognitive function in down syndrome across intellectual level and dementia status
Abstract: Adults with Down syndrome (DS) are at risk of developing dementia and cognitive assessment is a fundamental part of the diagnostic process. Previously, we developed a Rapid Assessment for Developmental Disabilities (RADD), a brief, broadly focused direct test of cognition. In the current report, we assess whether the RADD is sensitive to dementia in DS and the degree to which it compares with other cognitive measures of dementia in this population. In a sample of 114 individuals with DS, with dementia diagnosed in 62%, the RADD was compared with the Dementia Questionnaire for Mentally Retarded Persons (DMR), the Bristol Activities of Daily Living Scale, Severe Impairment Battery (SIB), and the Brief Praxis Test (BPT). The RADD showed predicted effects across intellectual disability (ID) levels and dementia status ($p < 0.001$). Six-month test-retest reliability for the subset of individuals without dementia was high ($r(41) = 0.95, p < 0.001$). Criterion-referenced validity was demonstrated by correlations between RADD scores and ID levels based upon prior intelligence testing and clinical diagnoses ($r (114) = 0.67, p = 0.001$) and with other measures of cognitive skills, such as the BPT, SIB, and DMR-Sum of Cognitive scores (range 0.84 through 0.92). Using receiver operating characteristic curves for groups differing in pre-morbid severity of ID, the RADD exhibited high sensitivity (0.87) and specificity (0.81) in discriminating among individuals with and without dementia, although sensitivity was somewhat lower (0.73) for the subsample of dementia cases diagnosed no more than 2 years prior to their RADD assessment. Taken together, findings indicated that the RADD, a relatively brief, easy-to-administer test for cognitive function assessment across ID levels and dementia status, would be a useful component of cognitive assessments for adults with DS, including assessments explicitly focused on dementia.

Dementia in people with severe/profound intellectual disabilities
https://doi.org/10.1111/jar.12917
Abstract: In people with severe or profound intellectual disabilities, it is difficult to diagnose dementia. As timely identification and diagnosis of dementia allows for a timely response to changing client wishes and needs, this study examined symptoms, and diagnosis of dementia in practice. Family members and professionals were invited to fill out a survey about symptoms and diagnosis of dementia in people with severe or profound intellectual disabilities. Results of the survey were further explored within semi-structured interviews with professionals having experience with signaling and diagnosing dementia in these people. Symptoms found in the survey and transcripts of the interviews were qualitatively analyzed, using thematic analyses based on a developed symptom-matrix. The survey was filled out completely by 14 family members and 90 professionals with different backgrounds. Results showed that behavioral changes were recognized more frequently than cognitive decline. Compared to those without dementia, epilepsy and motor decline were more present in case of dementia. Fifteen interviews (until saturation) with professionals provided an in-depth view into the symptoms, and how to identify them, again stressing behavioral alterations and to a lesser extent cognitive symptoms.

Waninge, A., Wissing, M., Hobbelen, J., Fokkens, A., Dekker, A., & De Deyn, P.
Abstract: Life expectancy of people with severe or profound intellectual disability (SPID) increases, which contributes to the risk of developing dementia. However, early detection and diagnosing dementia is complex, because of their low-level baseline functioning. Therefore, the aim is to identify observable dementia symptoms in adults with SPID in available literature. A systematic literature search, in line with PRISMA guidelines, was conducted in PubMed, PsycINFO and Web of Science using a combination of search terms for SPID, dementia/aging and aged population. In total, fifteen studies met inclusion criteria. Cognitive, behavioral and psychological symptoms (BPSD) and a decline in the ability to perform activities of daily living as well as neurological and physical changes were found. This presentation gives an overview of reported symptoms of (possible) dementia-related symptoms in SPID. Despite growing attention for dementia in people with ID in literature, only very few studies have studied dementia symptoms in SPID. Given the complexity of signaling and diagnosing dementia in SPID, dedicated studies are required to unravel the natural history of dementia in SPID, specifically focusing on observable symptoms for caregivers of (early) dementia in this population.

Ward, S., Opie, J., O’Connor, D.W.
Family carer’s responses to behavioural and psychological symptoms of dementia
Abstract: The author sought to describe the responses of family carers to the behavioural and psychological symptoms associated with dementia. Thirty family carers of people with dementia were identified in a survey of mental disorder in general practice. Another 20 were referred by local aged mental health services. Carers were interviewed using the Manchester and Oxford University Scale for the Psychopathological Assessment of Dementia (MOUSEPAD) which rates behavioural and psychological disturbances. Carers’ customary responses to current symptoms were recorded verbatim and categorised using a structured typology. Symptom frequency increased in line with dementia severity. Disturbances were generally well tolerated. Most were ignored where possible, except for wandering from home. Other common responses included avoiding triggers, providing reassurance, reality orientation, diversion, and collusion with false beliefs. Restrictive or punitive responses were uncommon. Few carers articulated clear strategies to deal with behavioural and psychological symptoms. For most, tolerance proved more effective and less distressing than arguments and reprimands. Carers’ responses are likely to be influenced by social and cultural factors and may differ in other settings.

Wark, S., Hussain, R., & Parmenter, T.
Down syndrome and dementia: Is depression a confounder for accurate diagnosis and treatment?
Abstract: The past century has seen a dramatic improvement in the life expectancy of people with Down syndrome. However, research has shown that individuals with Down syndrome now have an increased likelihood of early onset dementia. They are more likely than their mainstream peers to experience other significant co-morbidities including mental health issues such as depression. This case report also describes a phenomenon in which three individuals with Down syndrome and dementia are described as experiencing a rebound in their functioning after a clear and sustained period of decline. It is hypothesized that the relationship is not actually a reversal of the expected dementia trajectory but is an undiagnosed depression exaggerating the true level of functional decline associated with the dementia. The proactive identification and treatment of depressive symptoms may therefore increase the quality of life of some people with Down syndrome and dementia.

Abstract: Alzheimer's disease (AD) occurs in aging adults with Down syndrome (DS) at a higher prevalence and an earlier age than in typical aging adults. As with the general aging adult population, there is an urgent need to understand the preclinical and early phases of AD progression in the adult population with DS. The aim of this scoping review was to synthesize the current state of the evidence and identify gaps in the literature regarding functional activity performance and falls and their significance to disease staging (i.e., mild, moderate, and severe defined staging criteria) in relation to Alzheimer's disease and related dementias (ADRD) in adults with DS. This scoping review included six electronic databases (e.g., PsycINFO, Academic Search Complete, CINAHL, COCHRANE Library, MEDLINE, and PubMed). Eligible studies included participants with DS ≥25 years of age, studies with functional measures and/or outcomes (e.g., activities of daily living, balance, gait, motor control, speech, behavior, and cognition; falls; and fall risks), and studies that investigated AD pathology and implications. Fourteen eligible studies were included and categorized through thematic analysis into the following themes: (1) physical activity and motor coordination (PAMC), (2) cognition, (3) behavior, and (4) sleep. The studies indicated how functional activity performance and engagement may contribute to early identification of those at risk of cognitive decline and AD development and/or progression. There is a need to expand the research regarding ADRD pathology relative to functional outcomes in adults with DS. Functional measures related to disease staging and cognitive impairment are essential to understanding how AD progression is characterized within real-world settings. This scoping review identified the need for additional mixed-methods research to examine the use of assessment and intervention related to function and its detection of cognitive decline and AD progression.

Watchman, K., Kerr, D., & Wilkinson, H.
Supporting Derek: A new resource for staff working with people who have a learning difficulty and dementia. 58 pp.
York, United Kingdom: Joseph Rountree Foundation (2010)

Abstract: This resource pack published by the Joseph Rowntree Foundation in partnership with the University of Edinburgh, is aimed at staff supporting people with intellectual disability who develop dementia. Its focus is on helping care staff and training officers from intellectual disability and dementia care settings, as well as community, housing and health care staff. The pack is composed of 10 topic area (chapters), including basics on dementia, understanding behavior, development care environments, pain, communication, meaningful activities, friends with dementia, nutrition and hydration, night-time care, and palliative care. The pack includes a DVD and training materials which cover many of the key issues related to diagnosing and responding to dementia in people with intellectual disabilities. A short drama included on the DVD (acted by people with an intellectual disability) provides an insight into the reality of dementia and how it might feel to the individual affected.

Watchman, K.
Critical issues for service planners and providers of care for people with Down’s syndrome and dementia. 

Abstract: This discussion paper raises critical issues that need to be addressed along with suggestions as to how they may be met with. Author notes that the role of service planners and providers of care is one that cannot be understated while considering the future needs of people with Down’s syndrome and dementia. Discussed are appropriateness of accommodations, care management, diagnosis, and training.

Watchman, K.
Why wait for dementia? 

Abstract: Adults with Down syndrome living in supported accommodation, who develop dementia, may also experience other preventable difficulties caused by the environment in which they live. This can result in their enforced move to a different accommodation. Yet it is known that it is beneficial for people with intellectual disabilities and dementia to remain in familiar surroundings for as long as possible. This article puts forward a new set of guidelines suggesting the modification of the living environment of adults with Down syndrome before they develop dementia. The guidelines are discussed along with possible barriers to their implementation.

Watchman, K.
Practitioner-raised issues and end-of-life care for adults with Down syndrome and dementia 

Abstract: The author interviewed a small group of practitioners working in intellectual disability and palliative care settings about their perceptions of a number of end-of-life issues related to people with Down syndrome who were affected by dementia. The study, which took place in Scotland, identified a number of issues and perceptions expressed by the subjects as well as gaps in services and practice. Key among the findings were the need for people with Down syndrome to be more involved in planning for their own end-of-life care; a lack of communication between those persons working in palliative care and intellectual disability settings; identification of a “care culture clash;” deficits in training programs for staff involving dying, death, and bereavement; and that end-of-life care for people with Down syndrome and dementia is a neglected area of research. The author highlights the lack of uniform practice when working with people with Down syndrome in the end stages of dementia and provides some recommendations for further discourse and research.

Watchman, K.
Changes in accommodation experienced by people with Down syndrome and dementia in the first five years after diagnosis 

Abstract: Research that has tracked living situation changes is lacking for people with Down syndrome post-diagnosis of dementia. Extant studies have not considered reasons for a move, the stage at which it happened, and how involved in the decision the person with Down syndrome was. To study this, a postal questionnaire was used with 35 carers of persons with Down syndrome who had been diagnosed with dementia during the previous five years. Results showed that there are fewer accommodation changes in the early stages of dementia among people with Down syndrome than have previously been suspected and that confusion exists over the interpretation of existing care models. Findings also revealed that adults with Down syndrome were often denied the opportunity to take part in discussions about their future accommodations and there was a lack of forward planning on the part of carers.

Watchman, K.
People with a learning disability and dementia: reducing marginalisation
Abstract: The awareness that people with a learning disability, particularly Down’s syndrome, are at risk of dementia at a younger age brings an associated need for clarity over service planning and delivery. In order to record changes and developments in approaches, research literature documents the changing history of people with a learning disability and, separately, people with dementia. We do not have the same knowledge about the most appropriate ways of supporting individuals who have both a learning disability and dementia. People will already experience social exclusion due to society’s interpretation of their learning disability and this review identifies from the literature factors that have contributed to the further marginalization of this group. The review highlights the need for accurate data and statistics, an individualized approach to sharing information about the diagnosis, general and specialist training, an increased use of adapting methods of communication as dementia progresses and a consistent staff approach across care settings.

Watchman, K.  
Intellectual Disability and Dementia: Research into Practice. 336 pp.  
Abstract: In 16 chapters, this edited text offers a balanced appraisal of the evidence base on people with intellectual disabilities who develop dementia. It includes a range of resources, and is split into three sections that address the following: (1) The association between intellectual disabilities and dementia: what do we know? (2) Experiences of dementia in people with intellectual disabilities: how do we know?, and (3) Service planning: what are we going to do? Section one explores issues such as defining and diagnosing dementia in people with intellectual disabilities, prevalence and incidence and treatment options. The authors explain the differing theories about why people with Down’s syndrome are more likely to experience dementia, which provides a useful foundation for discussions about the use of medication. Section two explores the perspectives of people with learning disabilities and their families and the experiences of families via case studies. This section also explores some checklists for use with family members to help plan for the future. Section three focuses on service planning by describing a framework that can be used by practitioners for discussing diagnosis and prognosis of dementia. This section also explores the issues related to ageing in place and dementia-specific services and suggests that training is important for staff supporting those with learning disabilities and dementia.

Watchman, K., Janicki, M.P., and the members of the International Summit on Intellectual Disability and Dementia  
The intersection of intellectual disability and dementia: Report of the international summit on intellectual disability and dementia  
Abstract: An International Summit on Intellectual Disability and Dementia, held in Glasgow, Scotland (October 13-14, 2016) drew individuals and representatives of numerous international and national organizations and universities with a stake in issues affecting adults with intellectual disability (ID) affected by dementia. A discussion-based consensus process was used to examine and produce a series of topical reports examining three main conceptual areas: (1) human rights and personal resources (applications of the Convention for Rights of People with Disabilities and human rights to societal inclusion, and perspectives of persons with ID), (2) individualized services and clinical supports (advancing and advanced dementia, post-diagnostic supports, community supports and services, dementia-capable care practice, and end-of-life care practices), and (3) advocacy, public impact, family caregiver issues (nomenclature/terminology, inclusion of persons with ID in national plans, and family caregiver issues). Outcomes included recommendations incorporated into a series of publications and topical summary bulletins designed to be international resources, practice guidelines, and the impetus for planning and advocacy with, and on behalf of, people with ID affected by dementia, as well as their families. The general themes of the conceptual areas are discussed and the main recommendations are associated with three primary concerns.

Watchman, K., Janicki, M.P., Splaine, M., Larsen, F.K., Gomiero, T., & Lucchino, R.  
International summit consensus statement: Intellectual disability inclusion in national dementia plans.  
https://doi.org/10.1177/1533317517704082  
Abstract: The World Health Organization (WHO) has called for the development and adoption of national plans or strategies to guide public policy and set goals for services, supports, and research related to dementia. It called for distinct populations to be included within national plans, including adults with intellectual disability (ID). Inclusion of this group is important as having Down’s syndrome is a significant risk factor for early-onset dementia. Adults with other ID may have specific needs for dementia-related care that, if unmet, can lead to diminished quality of old age. An International Summit on Intellectual Disability and Dementia, held in Scotland, reviewed the inclusion of ID in national plans and recommended that inclusion goes beyond just description and relevance of ID. Reviews of national plans and reports on dementia show minimal consideration of ID and the challenges that carers face. The Summit recommended that persons with ID, as well as family carers, should be included in consultation processes, and greater advocacy is required from national organizations on behalf of families, with need for an infrastructure in health and social care that supports quality care for dementia.

Watchman, K., Janicki, M.P., Udell, L., Hogan, M., Quinn, S., Beránková, A.  
Consensus statement of the International Summit on Intellectual Disability and Dementia on valuing the perspectives of persons with intellectual disability.  
Abstract: The International Summit on Intellectual Disability and Dementia covered a range of issues related to dementia and intellectual disability, including the dearth of personal reflections of persons with intellectual disability affected by dementia. This article reflects on this deficiency and explores some of the personal perspectives gleaned from the literature, from the Summit attendees and from the experiences of persons with intellectual disability recorded or scribed in advance of the two-day Summit meeting. Systemic recommendations included reinforcing the value of the involvement of persons with intellectual disability in (a) research alongside removing barriers to inclusion posed by institutional/ethics review boards, (b) planning groups that establish supports for dementia and (c) peer support. Practice recommendations included (a) valuing personal perspectives in decision-making, (b) enabling peer-to-peer support models.

Watchman, K., Mattheys, K., & McKernon, M.  
Effects of the implementation of non-drug Interventions on behaviour and psychosocial symptoms of dementia in people who have an Intellectual disability.  
Journal of Intellectual Disability Research, 2019, 63(8), 647.  
Abstract: This three-year study investigates if non-drug interventions result in positive changes in behaviour associated with dementia in people with intellectual disability. People with intellectual disability are involved as advisors (n = 1) and co-researchers (n = 4) in both cycles. Cycle 1 (concluded) included 7
participants with intellectual disability in the early stage of dementia (4 with Down syndrome) and 12 support staff. Cycle 2 (ongoing) includes participants who have a more profound intellectual disability, and/or are experiencing advanced dementia. In both cycles, a goal-setting tool firstly helped to identify individualised non-drug interventions. In Cycle 1, a pre- and post-behaviour change tool (NPI-Q), was completed alongside semi-structured interviews, a bespoke tool to measure 'in the moment' changes, intervention diaries, and photovoice. Cycle 1 interventions included reminiscence, life story, music playlists, cookery, aromatherapy, environmental design change, exercise and cognitive games. Of 239 separate intervention over a 6-month period in Cycle 1, 193 resulted in positive behaviour change with 75% of goals being achieved or exceeded. The study offers insight into the support of people with intellectual disability and dementia. Use of non-drug supports in response to distress has led to cultural change within participating organisations with less reliance on medication as a first response.

Watchman, K., Mattheys, K., McKernon, M., Strachan, H., Andreis, F., & Murdoch, J.
A person-centred approach to implementation of psychosocial interventions with people who have an intellectual disability and dementia-A participatory action study.
Abstract: Numbers of people with an intellectual disability and dementia present a global health and social challenge with associated need to reduce stress or agitation and improve quality of life in affected individuals. This study aimed to identify effectiveness of psychosocial interventions in social care settings and, uniquely, explore use of photovoice methodology to develop dialogue about dementia. This mixed-method participatory action study used individualized goal-setting theory with 16 participants with intellectual disability and dementia, and 22 social care staff across 11 sites. Five co-researchers with intellectual disability were part of an inclusive research team collecting data using existing and bespoke tools including photovoice. Analysis used descriptive and inferential statistics and framework analysis. Seventy four percentage of individual goals met or exceeded expectations with some "as required" medication. Qualitative findings include themes of enabling care and interventions as tools for practice. Photovoice provided insight into previously unreported fears about dementia. Individualized psychosocial interventions have potential to reduce distress or agitation.

Watchman, K., McKernon, M., Boustead, L., & Doyle, A.
Intellectual disability and dementia: Understanding the effectiveness of psychosocial interventions
Abstract: The study aim was to identify effectiveness of psychosocial interventions with people who have an intellectual disability and dementia. This mixed-method participatory action study used goal-setting theory with 16 participants with intellectual disability and dementia, and 22 social care staff across 11 sites. Five core researchers with intellectual disability were part of an inclusive research team collecting data using existing and bespoke tools, including photovoice. Psychosocial interventions included: music playlists, reminiscence, animal therapy, robotic animals, and design changes. Analysis used descriptive and inferential statistics and framework analysis. Found that 74% of individual goals met or exceeded expectations with reduction in some "as required" medication. Qualitative findings include themes of enabling care and interventions as tools for practice. Photovoice provided insight into previously unreported fears about dementia. This poster combines an easy-read and pictorial summary of the study. Recommendations are made to maximise wellbeing and ensure the perspectives of people with dementia are heard: medication review, design changes to the home of a person with dementia, staff training, and talking about dementia more with people who have an intellectual disability. Individualized psychosocial interventions have potential to reduce distress or agitation in persons with intellectual disability and dementia, and to increase quality of life.

 WARNER, M.L.
The complete guide to Alzheimer's-proofing your home.
470 pp.
West Lafayette, Indiana: Purdue University Press (1998)
Abstract: General text on adapting homes and living environments for persons with dementia; applicable to home and other residential situations for adults with intellectual disabilities and dementia.

Webber, R., Bowers, B., & Bigby, C
Confidence of group home staff in supporting the health needs of older residents with intellectual disability
Abstract: Increased life expectancy for people with intellectual disability is accompanied by increased age-related health concerns. People ageing with intellectual disability experience more health conditions and are relocated to aged care earlier than their age peers. Group home staff were surveyed about their (a) training and confidence in 11 health conditions and 7 health procedures, and (b) attitude to relocating residents with health needs to aged care. Staff training in each of 10 health conditions and 7 health procedures was positively associated with increased confidence in supporting residents with those health issues. Higher staff confidence in caring for residents with 9 conditions and requiring 4 procedures was negatively associated with a likelihood of recommending that a person with those health needs should relocate to aged care. Targeted training of staff in age-related health issues may contribute to better health care and delay residents relocating to aged care.

Webber, R., Bowers, B. & McKenzie-Green, B.
Staff responses to age-related health changes in people with an intellectual disability in group homes.
Abstract: The purpose of this study was to explore how supervisors in group homes caring for people with intellectual disability responded to the development of age-related health changes in their residents. Ten group home supervisors working in the disability sector were interviewed once. Data were analyzed using Dimensional Analysis. The study identified several factors related to whether a resident could stay ‘at home’ or would need to be moved to residential aged care (nursing home) including: nature and extent of group home resources, group home staff comfort with residents’ health changes, staff skill at navigating the intersection between the disability and ageing sectors, and the supervisor’s philosophy of care. The ability of older people with an intellectual disability to ‘age in place’ is affected by staff knowledge about and comfort with age-related illnesses, staff skills at navigating formal services, staffing flexibility, and the philosophy of group home supervisors. Despite the growing international concern for the rights of people with disability, particularly in relation to decision making, questions about the older person’s choice of residence and participation in decision making about what was best for them, were almost nonexistent. Rather, decisions were made based on what was considered to be in ‘the best interest.”
Whitaker, S., & Read, S.
The prevalence of psychiatric disorders among people with intellectual disabilities: An analysis of the literature
Abstract: It has often been stated that the prevalence of psychiatric disorders in people with intellectual disabilities is greater than it is in the population as a whole. The epidemiological studies on psychiatric disorders in people with intellectual disabilities were reviewed. There is evidence that the prevalence of psychiatric disorder is greater in children with intellectual disabilities, compared with children with normal IQs, that it is higher in both adults and children with severe intellectual disabilities compared with people with mild or no intellectual disability and that the rate of problematic behavior is higher in both adults and children having intellectual disabilities, than in their non-disabled peers. There is no sound evidence that the prevalence of psychiatric disorders in adults with mild intellectual disability is greater than in the population as a whole.

Whitehouse, R., Chamberlain, P., & Tunna, K.
Dementia in people with learning disability: a preliminary study into care staff knowledge and attributions
Abstract: This paper describes the findings of a pilot study funded by the NHS Executive Primary and Community Care Research Initiative Small Projects Scheme that investigated the knowledge and attributions of dementia held by care staff who work with older adults with learning disability. Meetings took place with 21 members of care staff identified as “keyworkers” to older adults with learning disability living in residential homes provided by Solihull Healthcare NHS Trust, Solihull, UK. The results suggest that staff have knowledge of ageing at a similar level to that of college students. Forgetfulness was the sign that they would most expect to see if they thought someone was suffering from dementia. When a change in behavior was attributed to dementia, it was most likely to be viewed as ‘stable, uncontrollable’ with staff feeling pessimistic about being able to change the behavior.

Whittick, J.E.
Dementia and mental handicap: attitudes, emotional distress and caregiving
Abstract: Against the current climate of hospital closure programs and community care, attitudes to caregiving were examined in three groups of carers, namely mothers caring for a mentally handicapped child, mothers caring for a mentally handicapped adult and daughters caring for a parent with dementia. An ‘attitude questionnaire’ was developed by the author and administered, postally, to the three groups. Daughters were found to be more likely than the mothers to see their caring role in a negative way and were more inclined to favor institutional care. Possible reasons for this are discussed. The relationship between attitudes and emotional distress (as measured by the GHQ-30) were also examined for the sample as a whole. Negative and pro-institutional attitudes towards the caregiving situation were associated with elevated levels of emotional distress. Implications at both a local and a national level for all those involved with carers are discussed in the light of these findings.

Whitwham, S., McBrien, J., & Broom, W.
Should we refer for a dementia assessment? A checklist to help know when to be concerned about dementia in adults with Down syndrome and other intellectual disabilities.

Abstract: The aim of this research was to develop a simple screening checklist to help carers and professionals know when to make a referral for a dementia assessment. A checklist was completed for all new referrals to a dementia service for people with intellectual disabilities. The obtained scores were compared to the diagnostic outcome of a comprehensive dementia assessment. The data (n = 159) indicate a higher score on the checklist correlates significantly with a subsequent diagnosis of dementia. Cut-off scores are explored. The checklist appears to be a useful tool to prompt referrals for a full dementia assessment. By helping the referrer to know when to be concerned about dementia, it may reduce the number of people referred late or not at all.

Wiener, J.M., & Pazzaglia, F.
Ageing- and dementia-friendly design: theory and evidence from cognitive psychology, neuropsychology and environmental psychology can contribute to design guidelines that minimise spatial disorientation.
Abstract: Many older people, both with and without dementia, eventually move from their familiar home environments into unfamiliar surroundings, such as sheltered housing or care homes. Age-related declines in wayfinding skills can make it difficult to learn to navigate in these new, unfamiliar environments. To facilitate the transition to their new accommodation, it is therefore important to develop retirement complexes and care homes specifically designed to reduce the wayfinding difficulties of older people and those with Alzheimer’s disease (AD). Residential complexes that are designed to support spatial orientation and that compensate for impaired navigation abilities would make it easier for people with dementia to adapt to their new living environment. This would improve the independence, quality of life and well-being of residents, and reduce the caregivers’ workload. Based on these premises, this opinion paper considers how evidence from cognitive psychology, neuropsychology and environmental psychology can contribute to ageing- and dementia-friendly design with a view to minimising spatial disorientation. After an introduction of the cognitive mechanisms and processes involved in spatial navigation, and the changes that occur in typical and atypical ageing, research from the field of environmental psychology is considered, highlighting design factors likely to facilitate (or impair) indoor wayfinding in complex buildings. Finally, psychological theories and design knowledge are combined to suggest ageing- and dementia-friendly design guidelines that aim to minimize spatial disorientation by focusing on residual navigation skills.

Wiese, M., Stancliffe, R.J., Dew, A., Balandin, S., & Howarth, G.
What is talked about? Community living staff experiences of talking with older people with intellectual disability about dying and death
Abstract: This study explored what community living staff talked about and did with people with intellectual disability (ID) to assist them to understand dying and death. Guided by grounded theory methodology, focus groups and one-to-one interviews were conducted with 22 staff who had talked about any topic relating to dying and death with their clients. There was little evidence that staff talked with, or did things with clients to assist understanding of the end of life, both prior to and after a death. Prior to death staff assisted clients in a limited way to understand about determining wishes in preparation for death, and what dying looks like by observance of its passage. Following a death staff offered limited assistance to clients to understand the immutability of death, and how the dead can be honored with ritual, and remembered.
The findings have implications for why people with ID have only partial understanding of the end of life, the staff skills required to support clients’ understanding, and when conversations about the end of life should occur.

Abstract: A panel of experts attending a 3-day meeting held in Edinburgh, UK, in February 2001 was charged with producing a set of principles outlining the rights and needs of people with intellectual disability (ID) and dementia, and defining service practices which would enhance the supports available to them. The Edinburgh Principles, seven statements identifying a foundation for the design and support of services to people with ID affected by dementia, and their carers, were the outcome of this meeting. The accompanying guidelines and recommendations document provides an elaboration of the key points associated with the Principles and is structured toward a four-point approach: (1) adopting a workable philosophy of care; (2) adapting practices at the point of service delivery; (3) working out the coordination of diverse systems; and (4) promoting relevant research. It is expected that the Principles will be adopted by service organizations worldwide, and that the accompanying document will provide a useful and detailed baseline from which further discussions, research efforts and practice development can progress.

Wilkinson, H., Kerr, D., & Rae, C.
Abstract: With people with a learning disability living longer, more of them are developing dementia. In planning the services they need, an important first step is to ask them what they think. Authors report information from surveying a group of older adults with intellectual disabilities.

Wilkinson, H., Kerr, D., & Cunningham, C.
Abstract: The knowledge, experiences and skills of direct care staff working in care home settings are essential in ensuring a good quality of life and care for a person with an intellectual disability (ID) who develops dementia. Drawing on the findings of a wider study, the issues of training, support and the wider needs of staff when trying to support a resident who develops dementia are explored, specifically as relating to the role played by staff and the need to determine their experiences and related training needs. Following an introduction to the policy and practice context for working with people with an ID and dementia, and a brief description of the research method, the authors discuss the attitudes and practices of staff; supportive changes at an organizational level; and the knowledge and training needs of staff and specific gaps in knowledge. The authors argue that, within the policy and practice context of aiming to support residents to ‘age in place’, support for staff is a crucial aspect of ensuring that such an approach is effective and provides a coordinated approach to planning, resourcing and support.

Wilson, B., Jones, K.B., Weedon, D., & Bilder, D.
Abstract: Down syndrome (DS) is a genetic disorder involving excess genetic material from chromosome 21. The incidence of DS is increasing, and the life expectancy for individuals with DS has increased to a median age of 55 years. Adults with DS are at increased risk of several conditions, including significant neurologic, cardiovascular, pulmonary, gastrointestinal, musculoskeletal, endocrine, psychiatric, hematologic, and social comorbidities, and additional screening or monitoring may be needed. Additional preventive measures for patients with DS include regular screening for thyroid dysfunction, hearing loss, eye disorders, heart disease, osteoporosis, and dementia, and one-time vaccination with the polyclonal pneumococcal polysaccharide vaccine (PPV23). Quality of life should be the main focus of treatment, with patients being involved in medical decisions as much as possible.

Wisniewski, K.E., Wisniewski, H.M., & Wen, G.Y.
Abstract: One hundred brains of patients with Down’s syndrome (DS) who died in institutions for chronic care were examined for clinico-pathological correlation of Alzheimer’s disease. Fifty-one were below and 49 were above age 30 years at death. Tissues from the right, prefrontal, and hippocampal cortices were processed for microscopy using H&E and Bodian-periodic acid-Schiff impregnation. Morphometric evaluations of plaques and tangles were carried out. Plaques or plaques and tangles were found in the brains of 56 patients with DS, 7 below age 30 and 49 above that age. A history of dementia was evident in the medical records of 15 of these patients; of these only 2 were below the age of 30. The brains of the patients with DS who also had clinical dementia had more than twenty plaques or plaques and tangles per 1.5 X 10(6) micron 2 of cortex. The numbers of plaques and tangles found in the brains of the patients with DS above the age of 30 greatly increased with age but varied from brain to brain. These observations suggest a correlation among dementia, the density of plaques and tangles, and age. All 100 brains studied showed early arrest of brain growth and brain atrophy, a condition that may have been due to prenatal arrest of neurogenesis mainly in the granular cell layers, prenatal and postnatal arrest of synaptogenesis, and early aging. Plaques and tangles developed twenty to thirty years earlier and dementia was clinically detected at least three times more frequently (20 to 30%) in DS than it is known to occur in the non-DS population.
Abstract: Dementia is increasingly prevalent in people with severe/profound intellectual disabilities. However, early detection and diagnosis of dementia is complex in this population. This study aimed to identify observable dementia symptoms in adults with severe/profound intellectual disabilities in available literature. A systematic literature search was conducted in PubMed, PsycINFO and Web of Science with an exhaustive search string using a combination of search terms for severe/profound intellectual disabilities and dementia/ageing. Eleven studies met inclusion criteria. Despite the few small-sized studies, a range of dementia symptoms were identified, subdivided in cognitive decline (e.g., memory loss, for getfulness, deterioration in speech, losses of social skills), decline in activities of daily living (e.g., self-cares skills, everyday functioning/skills), BPSD (e.g., apathy, aggression, irritability, altered eating/drinking behaviour) as well as neurologic and other physical symptoms (e.g., incontinence, (late-onset) epilepsy, hypotonia, gait deterioration). Cognitive decline, behavioral and psychological alterations, decline in activities of daily living as well as neurologic and physical changes were found. Only a very limited number of studies reported symptoms ascribed to dementia in adults with severe/profound intellectual disabilities. Given the complexity of signaling and diagnosing dementia, dedicated studies are required to unravel the natural history of dementia in this population.

Dementia in people with severe/profound intellectual (and multiple) disabilities: applicability of items in dementia screening instruments for people with intellectual disabilities  
https://doi.org/10.1080/19315864.2022.2111737  
Abstract: Diagnosing dementia in people with severe/profound intellectual (and multiple) disabilities (SPI(M)D) is complex. Whereas existing dementia screening instruments as a whole are unsuitable for this population, a number of individual items may apply. Therefore, this study aimed to identify applicable items in existing dementia screening instruments. Informant interviews about 40 people with SPI(M)D were conducted to identify applicable items in the Dementia Scale for Down Syndrome, Behavioral and Psychological Symptoms of Dementia in Down Syndrome II scale, Dementia Questionnaire for persons with Mental Retardation and Social competence Rating scale for people with Intellectual Disabilities. Among 193 items, 101 items were found applicable, categorized in 55 domains: behavioral and psychological functioning (60 items), cognitive functioning (25), motor functioning (6), activities of daily living (5) and medical comorbidities (5). Identifying applicable items for people with SPI(M)D is an essential step in developing a dedicated dementia screening instrument for this population.

Dementia in people with severe/profound intellectual (and multiple) disabilities: practice-based observations of symptoms  
https://doi.org/10.1080/19315864.2022.2061092  
Abstract: Observable dementia symptoms are hardly studied in people with severe/profound intellectual (and multiple) disabilities (SPI(M)D). Insight in symptomatology is needed for timely signaling/diagnosis. This study aimed to identify practice-based observations of dementia symptoms in this population. Care professionals and family members were invited to complete a survey about symptoms. Quantitatively analyzed survey data were further deepened through semi-structured interviews with care professionals having vast experience in signaling/diagnosing dementia in this population. Symptoms were categorized using a symptom matrix. Survey respondents and interviewees frequently observed a decline in activities of daily living (ADL) functioning and behavioral and psychological changes, like increased irritability, anxiety, apathy and decreased eating/drinking behavior. Cognitive symptoms were particularly recognized in persons with verbal communication and/or walking skills. To lesser extent motor changes and medical comorbidities were reported. Increased insight in dementia symptoms contributes to developing a dedicated screening instrument for dementia in people with SPI(M)D.

Woods, R.T., Moniz-Cook, E., Liliffe, S., Campion, P., Vernooij-Dassen, M., Zanetti, O., & Franco, M.  
Journal of the Royal Society of Medicine, 2003, 96, 320-324.  
Abstract: Generic article about the need for quality and accurate screening and assessment of adults suspected of showing signs of Alzheimer’s disease and the need for psychosocial interventions and family carer supports. Authors note need for better training of medical practitioners who may be screening for dementia, indicating that there is a need for timely detection and diagnosis that will prevent crises, facilitate adjustment and provide access to treatments and supports.

Yang, Q., Rasmussen, S. A., & Friedman, J. M.  
https://doi.org/10.1016/s0140-6736(02)08092-3  
Abstract: : Down syndrome is the most frequently identified cause of mental retardation, but information about mortality and comorbidity in people with Down syndrome is limited. We used data from US death certificates from 1983 to 1997 to calculate median age at death and standardized mortality odds ratios (SMORs) for common medical disorders in people with Down syndrome. Of 17897 people reported to have Down syndrome, median age at death increased from 25 years in 1983 to 49 years in 1997, an average increase of 1.7 years per year studied (p<0.0001). Median age at death was significantly lower in black people and people of other races than in white people with Down syndrome. As expected, death certificates with a diagnosis of Down syndrome were more likely to list congenital heart defects (SMOR 29.1, 95% CI 27.8-30.4), dementia (21.2, 19.6-22.7), hypothyroidism (20.3, 18.5-22.3), or leukemia (1.6, 1.4-1.8) than were those that did not report Down syndrome. By contrast, malignant neoplasms other than leukemia were listed on death certificates of people with Down's syndrome less than one-tenth as often as expected (0.07, 0.06-0.08). A strikingly low SMOR for malignancy was associated with Down syndrome at all ages, in both sexes, and for all common tumor types except leukemia and testicular cancer. Identification of factors responsible for the racial differences recorded could facilitate further improvement in survival of people with Down syndrome. Reduced exposure to environmental factors that contribute to cancer risk, tumor-suppressor genes on chromosome 21, or a slower rate of replication or higher likelihood of apoptosis in Down syndrome cells, could be possible reasons for paucity of cancer in people with Down syndrome.

Zaman, S., & Fortea, J.  
The crucial history of Down syndrome.  
PMID: 35182507.  
Full article: In 1866, John Langdon Down was the first clinician to provide a detailed account of Down syndrome. A decade later, Fraser and Mitchell highlighted the early onset of senility in this population. The observations of
neuropathological features of Alzheimer’s disease in Down syndrome were published in 1929 by Struwe. In 1948, Jervis noted, “with the exception of the age of onset, the clinical and pathological manifestations are those of senile dementia”. In 1984, using samples taken from the brain of people with Down syndrome, Glenner and Wong biochemically characterized amyloid β (Aβ), predicted that the gene for Alzheimer’s disease would be on chromosome 21, and proposed a pathogenic role for Aβ. Using the Aβ peptide sequence, the locus of the APP gene was discovered on chromosome 21. Genetic linkage studies later identified the first mutations in APP, PSEN1, and PSEN2. Furthermore, Down syndrome studies have shown the importance of endosomal misprocessing of APP in Alzheimer’s disease. Down syndrome is now regarded as a genetically determined form of Alzheimer’s disease, similar to autosomal dominant Alzheimer’s disease. Clinical and biomarker changes show a parallel natural history in these two forms. These genetic types of Alzheimer’s disease provide the strongest evidence supporting the amyloid cascade hypothesis and give researchers a unique opportunity to study pathophysiology and its temporality, and to undertake preventive trials.

Abstract: Authors evaluated the feasibility of using the German-language version of a recently developed screening tool for dementia for persons with intellectual disability (ID): the National Task Group – Early Detection Screen for Dementia (NTG-EDSD). Some 221 paid carers of ageing persons with ID were asked to use the NTG-EDSD and report back on its utility and on feasibility dimensions, and to provide detailed feedback on aspects deemed critical or missing. All feasibility dimensions were rated good to very good, and 80% of respondents found the NTG-EDSD useful or very useful for the early detection of dementia. This highlights a high acceptability of this instrument by the main target group. The positive feasibility evaluation of the NTG-EDSD indicates the usability and adequacy of this instrument for application of early detection of dementia in persons with ID.

Zeilingher, E.L., Komenda, S., Znic, I., Franken, F., & Woditschka, K.
Informant-based assessment instruments for dementia and their measurement properties in persons with intellectual disability: systematic review protocol *BMJ Open*, 2020, 10(12), 1136. https://bmjopen.bmj.com/content/10/12/e040920
Abstract: Persons with intellectual disability (ID) are at a higher risk of developing dementia than persons without ID, with an expected earlier onset. Assessment methods for the general population cannot be applied for persons with ID due to their pre-existing intellectual and functional impairments. As there is no agreed-upon measure to assess dementia in persons with ID, multiple instruments for this purpose have been developed and adapted in the past decades. This review aimed to identify all available informant-based instruments for the assessment of dementia in persons with ID, to evaluate and compare them according to their measurement properties, and to provide a recommendation for the most suitable instruments. Additionally, an overview of the amount and quality of research on these instruments was provided.

Abstract: Dementia in people with intellectual disability (ID) is frequent but hard to recognize. Evidence-based recommendations for suitable instruments are lacking. The present study set out to evaluate informant-based dementia assessment instruments and to provide evidence-based recommendations for instruments most suitable in clinical practice and research. A systematic review was conducted across ten international electronic databases. The COSMIN-based Standards for the selection of health Measurement Instruments (COSMIN) guidelines, including a risk of bias assessment, was applied to extract information and to evaluate measurement properties and the quality of available evidence. In total, 42 studies evaluating 18 informant-based assessment instruments were analysed. For screening purposes, we recommend the Behavioral and Psychological Symptoms of Dementia in Down Syndrome Scale (BPSD-DS), the Cognitive Scale for Down Syndrome (CS-DS), and the Dementia Screening Questionnaire for Individuals with Intellectual Disabilities (DSQIID). For a more thorough dementia assessment, we recommend the Cambridge Examination for Mental Disorders of Older People with Down Syndrome and Others with Intellectual Disabilities (CAMDEX-DS).

Our study informs clinicians and researchers about adequate, well-evaluated dementia assessment instruments for people with ID, and highlights the need for high quality studies, especially regarding content validity.

Abstract: This work describes an extensive systematic literature review on assessment instruments for dementia in persons with intellectual disability (ID). Existing instruments for the detection of dementia in persons with ID were collected and described systematically. This allows a direct and quick overview of available tools. Additionally, it contributes to the availability and usability of information about these instruments, thus enhancing further developments in this field. A systematic literature search in five databases (CINAHL, PsycInfo, PubMed, Scopus, and Web of Science) was conducted. In order to include gray literature an invisible college approach was used. Relevant studies were identified and selected using defined inclusion and exclusion criteria. After the selection process all instruments were coded and classified. It was determined which concepts they assess, whether they were especially developed or adapted for persons with ID, and whether they were designed to assess dementia. The selection of relevant papers, as well as the coding of instruments was done independently by two researchers. In total, 97 records met the search criteria. Out of these, 114 different instruments were extracted. There were 79 instruments to be completed by the person with ID, and 35 informant-based instruments. Additionally, four test batteries were found. Some of these instruments were neither designed for the assessment of dementia, nor for persons with ID. There are a variety of different tools used for the assessment of dementia in ID. Nevertheless, an agreed-upon approach or instrument is missing. Establishing this would improve the quality of assessment in clinical practice, and benefit research. Data collected would become comparable and combinable, and allow research to have more informative value.

Zigman, W.B.
Abstract: At present, there may be over 210,000 people with Down syndrome (DS) over the age of 55 in the United States (US) who have significant needs for...
augmented services due to circumstances related to ordinary and/or pathological aging. From 1979 through 2003, the birth prevalence of DS rose from 9.0 to 11.8 (31.1%) per 10,000 live births in 10 representative US regions. This increase, largely due to women conceiving after age 35, portends an ever-growing population of people with DS who may be subject to pathogenic aging. Whereas Trisomy 21 is one of the most widespread genetic causes of intellectual disability (ID), it still is one of the least understood of all genetic ID syndromes. While longevity in people with DS has improved appreciably in as modest a period as 30 years, age-specific risk for mortality still is considerably increased compared with both other people with ID or with the typically developing population. The penetrance of the phenotype is widely distributed, even though a consistent genotype is assumed in 95% of the cases. Some, but not all body systems, exhibit signs of premature or accelerated aging. This may be due to both genetic and epigenetic inheritance. We now know that the long-term outcome for people with DS is not as ominous as once contemplated; a number of people with DS are living into their late 60s and 70s with few if any major signs of pathogenic aging. Alzheimer's disease (AD), a devastating disease that robs a person of their memory, abilities and personality, is particularly common in elder adults with DS, but is not a certainty as originally thought, some 20% to 30% of elder adults with DS might never show any, or at least, mild signs of AD. DS has been called a mature well-understood syndrome, not in need of further research or science funding. We are only beginning to understand how epigenetics affects the phenotype and it may be feasible in the future to alter the phenotype through epigenetic interventions. This chapter is divided into two sections. The first section will review typical and atypical aging patterns in somatic issues in elder adults with DS; the second section will review the multifaceted relationship between AD and DS.

Zigman, W.B., & Lott, I.T.
Abstract: Down syndrome (DS) is characterized by increased mortality rates, both during early and later stages of life, and age-specific mortality risk remains higher in adults with DS compared with the overall population of people with mental retardation and with typically developing populations. Causes of increased mortality rates early in life are primarily due to the increased incidence of congenital heart disease and leukemia, while causes of higher mortality rates later in life may be due to a number of factors, two of which are an increased risk for Alzheimer's disease (AD) and an apparent tendency toward premature aging. In this article, we describe the increase in lifespan for people with DS that has occurred over the past 100 years, as well as advances in the understanding of the occurrence of AD in adults with DS. Aspects of the neurobiology of AD, including the role of amyloid, oxidative stress, Cu/Zn superoxide dismutase (SOD-1), as well as advances in neuroimaging are presented. The function of risk factors in the observed heterogeneity in the expression of AD dementia in adults with DS, as well as the need for sensitive and specific biomarkers of the clinical and pathological progressing of AD in adults with DS is considered.

Abstract: Down syndrome is associated with increased mortality rates due to congenital cardiac defects and leukemia early in life, and with Alzheimer's disease and a tendency toward premature aging later in life. Alzheimer's disease was once considered an inexorable result of growing old with Down syndrome, but recent data indicate that risk does not reach 100%. Although some individuals exhibit signs and symptoms of Alzheimer's disease in their 40s, other individuals have reached the age of 70 without developing dementia. This chapter presents a wealth of data from a longstanding longitudinal study with the overall objective of understanding and recounting the mechanisms responsible for these substantial individual differences.

Zigman, W.B., Schupf, N., Devenny, D., Miezejeski, C., Ryan, R., Urv, T.K., Schubert, R., & Silverman, W.
Abstract: Rates of dementia in adults with mental retardation without Down syndrome were equivalent to or lower than would be expected compared to general population rates, whereas prevalence rates of other chronic health concerns varied as a function of condition. Given that individual differences in vulnerability to Alzheimer's disease have been hypothesized to be due to variation in cognitive reserve, adults with mental retardation, who have long-standing intellectual and cognitive impairments, should be at increased risk. This suggests that factors determining intelligence may have little or no direct relationship to risk for dementia and that dementia risk for individuals with mental retardation will be comparable to that of adults without mental retardation unless predisposing risk factors for dementia are also present.

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