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To cite this article: Malvika Muralidhar, Aimee Spector, Esther K. Hui, Lisa Liu & Afia Ali (09 Oct 2023): A systematic review of psychosocial interventions for people with intellectual disabilities and dementia, Aging & Mental Health, DOI: 10.1080/13607863.2023.2265322

To link to this article:  https://doi.org/10.1080/13607863.2023.2265322
A systematic review of psychosocial interventions for people with intellectual disabilities and dementia

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ABSTRACT

Objectives: 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.

Methods: 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 analysed by narrative synthesis.

Results: Ten distinct interventions were identified and categorised 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.

Conclusion: There is emerging evidence for several psychosocial interventions for people with ID and dementia, though further research is required on effectiveness and generalisability. The adaptations discussed may guide implementation into routine care and contribute to current policies and guidelines on improving ID and dementia care.

Introduction

Increasing life expectancy of people with intellectual disabilities (ID) is accompanied by an increased risk of several health conditions in the ageing ID population (Coppus, 2013). Individuals with ID tend to be at greater risk of developing dementia, and at an earlier age, compared to the general population (Strydom et al., 2009). According to the cognitive reserve hypothesis (Stern, 2002), higher reserves (greater cognitive capabilities and rich environment) should act as a protective factor in delaying dementia onset, whereas lower reserves would increase vulnerability to it. Considering this rationale, ID comprises significant functional and cognitive impairments which may therefore, increase the risk of developing dementia, and at an early onset (Silverman et al., 2013).

The management of ID and dementia includes substantial focus on long-term goals, and psychological and environmental interventions are recommended, especially for managing non-cognitive symptoms (NICE, 2018). Similarly, medication is suggested only in combination with psychosocial interventions to manage behaviour that challenges in people with ID (NICE, 2015).

Psychosocial interventions aim to improve symptomatology, emotional and psychological wellbeing, and overall quality of life (MacDonald & Summers, 2020). Research has examined a range of psychosocial interventions for dementia and there is increasing evidence for the effectiveness of cognitive stimulation therapy, cognitive rehabilitation therapy, occupational therapy and group reminiscence therapy, as suggested by the NICE guidelines for management of dementia (NICE, 2018). While psychosocial interventions are progressively being implemented, research has predominantly focused on individuals with dementia in the general population, maintaining uncertainty regarding the effectiveness and applicability of these interventions for people with ID.

The combined diagnosis and distinct presentations may render some interventions as effective for one but not the other. Furthermore, individual, service, and system level barriers to treatment (Willner, 2009) question how acceptable psychosocial interventions for dementia would be among the ID population. To abide by person-centred treatment, adaptations for people with ID need to be considered. Widely practised adaptations include simplifying tasks and language, using flexible methods, and considering developmental level (Whitehouse et al., 2006). Two systematic reviews have presented evidence on psychosocial interventions for people with ID and dementia; MacDonald and Summers (2020) examined current interventions and their outcomes for people with ID and dementia; however, these included staff and carer focused (i.e. non-direct) interventions and discussed the lack of direct interventions tested and outcomes measured in this population. Moreover, the paper did not discuss intervention adaptations. Thalen et al. (2022) reviewed the focus of psychosocial interventions for older people with ID and the role of staff support in applying these interventions. This review discussed factors that could optimise interventions for older people with ID but limited insight was given on dementia in ID or assessing intervention effectiveness for this population.
Therefore, this systematic review aims to extend current knowledge, with the following questions:

1. What is the evidence for the effectiveness of direct psychosocial interventions in ID and dementia?
2. What specific adaptations have been made to facilitate the use of these interventions with this population?

Methods

This review followed the Preferred Reporting Items for Systematic reviews and Meta-Analyses (PRISMA) checklist (Page et al., 2021).

Study protocol

The review protocol was published online on the PROSPERO website (CRD42022314605).

Inclusion criteria

Studies were included if: (1) participants had a diagnosis of ID and dementia, of any severity and duration, (2) they evaluated a direct psychosocial intervention delivered to people with ID and dementia (in clinical or non-clinical settings, including community and online modes, to account for COVID-19), (3) outcomes related to cognition, behaviour, psychiatric symptoms, adaptive functioning, and quality of life (all measured by validated scales), and (4) the study was published in English.

There was no limit on: (1) participants age, as people with ID may have an earlier onset of dementia, (2) country of study, (3) study design, and (4) publication date.

Exclusion criteria

Papers were excluded if: (1) participants (a) did not have a clinical diagnosis of ID and dementia or (b) had sub-clinical cognitive impairment; (2) the outcomes measures were collected only from carer/staff/others; (3) they were protocols, conference abstracts, and books.

Search strategy

Five electronic databases, Medline (OVID), Embase (OVID), PsychInfo (OVID), Web of Science (Core), and Google Scholar were used. Grey literature including unpublished studies and PhD theses (ProQuest Dissertations and Theses Global) were included. An initial search was conducted between March and April 2022 by the main researcher MM (Table 1). Additionally, reference lists of included full-text articles and the two systematic reviews (MacDonald & Summers, 2020; Thalen et al., 2022) were checked for further relevant studies.

The database searches were re-run in May 2022 before the final analyses.

Data extraction and synthesis

The database search results were deduplicated and papers were imported into Endnote. The first reviewer MM screened titles and abstracts based on the inclusion and exclusion criteria. Full texts were retrieved and checked for relevance alongside a second reviewer.

Data extraction examined key characteristics including information on the author, year, country, sample size, study design, demographics, and intervention details (type, modality, duration, number of planned sessions, length of session, treatment period, control condition, dropout rates, and retention), outcomes and findings. This was combined into a table, and narrative synthesis was carried out by summarising the studies, comparing their similarities and differences, and assessing quality.

Risk of bias assessment

Studies were assessed for quality using the Mixed Methods Appraisal Tool (MMAT), a critical appraisal tool suitable for systematic reviews that include quantitative, qualitative, and mixed-method studies (Hong et al., 2018b). It has five sets of methodological quality criteria for (1) qualitative (2) quantitative randomised controlled trials (3) quantitative non-randomised (4) quantitative descriptive, and (5) mixed-methods studies, with each criterion covering a specific attribute of the respective study design. Each category has five questions, with ‘Yes,’ ‘No,’ and ‘Can’t tell’ as responses. Two independent reviewers were involved in the quality assessment, and disagreements were resolved with discussion.

Results

Characteristics of included studies

A total of 2126 papers were identified from the database search and 1967 were screened after deduplication. Subsequently, 165 full texts were retrieved and assessed for eligibility. 10 papers met the eligibility criteria and were included in the review. The PRISMA flowchart (Figure 1) shows the process of study selection.

Four quantitative (one RCT, one non-randomised, and two descriptive), four qualitative (case studies) and two mixed-methods studies were included (Table 2). The studies were carried out between 2008 and 2019, across five different countries. Sample characteristics varied in terms of gender distribution and ethnicity, sample sizes ranged from 1 to 97 participants. The studies included ID and dementia of mild to profound/advanced severities, with four focusing on Down Syndrome and

Table 1. Search strategy for all databases.

<table>
<thead>
<tr>
<th>Intellectual disability</th>
<th>Dementia</th>
<th>Psychosocial</th>
<th>Intervention</th>
</tr>
</thead>
<tbody>
<tr>
<td>intellectual disability* OR learning disability* OR learning difficulty* OR developmental disability* OR mental retardation* OR intellectual impairment* OR down's syndrome</td>
<td>dementia OR Alzheimer* OR vascular dementia OR Parkinson* dementia* OR PDD OR Lewy bod* OR LBD OR DLB OR mixed dementia</td>
<td>psychosocial OR non-pharmacological OR non-drug OR drug OR cognitive stimulation OR CST OR cognitive rehabilitation* OR reminiscence* OR occupational therapy* OR OT OR creative therap* OR art therap* OR music therap* OR music-oriented OR music-based OR drama therap* OR movement OR movement-based</td>
<td>intervention OR treatment OR therap*</td>
</tr>
</tbody>
</table>
the remaining either had a sample with diverse aetiologies or did not specify the type of ID.

A range of outcomes were examined, including quality of life, behavioural changes, functioning, cognition, and symptom deterioration.

**Quality assessment**

Table 3 depicts the quality assessment. A low-quality study met 0%–20%, medium quality met 40%–60%, and high-quality met 80%–100% of the criteria under the respective study design in the MMAT. Three studies were rated as high quality—Watchman et al. (2021; quality score 100%), De Vreese et al. (2012; 80%), and Ali et al. (2022; 80%); three were of medium quality—Crook et al. (2016; 60%), Hawkes et al. (2019; 60%), and Lifshitz and Klein (2011; 40%); one of low quality (Kiddle et al., 2016; 20%). Finally, three studies (Horovitz et al. 2010; Nichols, 2011; Vogl & Rapp, 2011) did not satisfy the screening criteria, so further appraisal was not conducted (i.e. total quality score N/A).
### Table 2. Summary of included studies.

<table>
<thead>
<tr>
<th>Author, Country</th>
<th>Design</th>
<th>Sample Details</th>
<th>Intervention Details</th>
<th>Control</th>
<th>Outcomes</th>
<th>Findings</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Watchman et al. (2021), UK</td>
<td>Mixed-method participatory action study</td>
<td>16 participants with ID and dementia (aged 40–77, 10 females, 6 males); 13 with Down Syndrome; 3 with other ID; duration of dementia &lt;1 year to 6 years. 22 social care staff across 11 social care sites.</td>
<td>Relevant psychosocial interventions, including reminiscence, creative activities, animal therapy. Participants received 2–6 interventions (M = 5). Number and timing of sessions varied according to goals and intervention used. Overall, interventions implemented over six months.</td>
<td>Not specified</td>
<td>Qualitative</td>
<td>- Behaviour changes (bespoke behaviour change tool) - Neuropsychiatric symptoms (NPI-Q) - Quality of life (QUAliD) - Goal attainment (pictorial) scale</td>
<td>- 74% of individual goals were met or exceeded expectations.</td>
</tr>
<tr>
<td>Kiddle et al. (2016), UK</td>
<td>Service evaluation</td>
<td>97 individuals across the 12-weeks (weekly attendance: 5–11 individuals), all from four local residential settings. Mild to moderate ID (3 with Down Syndrome); 3 diagnosed with moderate dementia, 4 undergoing assessments.</td>
<td>12-week memory café held at a local youth centre, with a different theme every week. Several physical, cognitive, and creative group activities, alongside individual activities when needed. Weekly talks for carers by a member of the Community Learning Disability Team.</td>
<td>None</td>
<td>Qualitative</td>
<td>- Semi-structured interviews with social care staff - Photovoice with co-researchers with ID</td>
<td>Attendance - Measured weekly - Affect ratings - Affect scale from an observational tool designed by Jarrott et al. (2002) Feedback - Weekly feedback questionnaires - Verbal feedback from group during last café week - Postal questionnaires (6 weeks after café).</td>
</tr>
<tr>
<td>Horovitz et al. (2010), USA</td>
<td>Case study</td>
<td>1 53-year-old Caucasian male; Alzheimer's-type dementia and profound ID secondary to Down Syndrome. Presenting complaints: noncompliant behaviour, inappropriate sexual behaviours, and disruptive behaviours.</td>
<td>Compliance training (initially by therapist): sessions that contingently reinforced compliance (edible reinforcers, verbal praise and physical contact). 30 weeks (baseline = weeks 1–13, intervention = weeks 14–30). Where needed, modifications were made to number of therapists, transition to staff members, and varying number of trials within session and frequency of sessions.</td>
<td>None</td>
<td>Qualitative</td>
<td>Incidence of behavioural noncompliance recorded on behavioural report forms.</td>
<td>Overall, 61.11% decrease in average noncompliance. Reduced incidences (mean per week): Baseline: M = 8.49 per week; post-intervention: M = 3.29. 85.22% decrease from baseline at 1-month follow up (M = 1.25).</td>
</tr>
<tr>
<td>Vogel and Rapp (2011), USA</td>
<td>Case study</td>
<td>1 52-year-old White female; Down Syndrome and early-onset Alzheimer's dementia. Presenting complaints: loitering, wandering, stealing.</td>
<td>DRO procedure with extinction - Component 1 = non-resetting 30-minute DRO for loitering (choice of alternative activities for up to 15 minutes, contingent on her absence from the entryway in the 30-minute interval prior). - Component 2 = extinction of tangible reinforcement of stealing items (counting number of items at the start of day, and ‘approved’ items at the end of the day; unapproved items removed on discretion of individual service-planning team). 11 days of baseline followed, 11 days of the intervention. 2 additional, non-consecutive follow-up days.</td>
<td>None</td>
<td>Quantitative</td>
<td>- Partial interval recording of occurrence of being present in hallway - Behavioural product measure (number of unapproved items in participant's cubby at the end of each day) recorded on data collection sheet - Staff reports at 2 and 3-month follow ups.</td>
<td>Baseline Presence in entryway (M = 5, range = 1–12) Unapproved items in cubby (M = 1, range = 0–4) Post-intervention Presence in entryway (M = 1.45, range = 0–4) Unapproved items in cubby (M = 0.27, range = 0–1) Follow-up Both at zero.</td>
</tr>
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(Continued)
Table 2. Continued.

<table>
<thead>
<tr>
<th>Author, Country</th>
<th>Design</th>
<th>Sample</th>
<th>Intervention</th>
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<tbody>
<tr>
<td>Crook et al. (2016), UK</td>
<td>Randomised single case series experiment</td>
<td>Five participants; Down Syndrome and dementia (3 females, 2 males; aged 55-63 years). Duration of dementia diagnosis: 3 months to 2 years.</td>
<td>3 experimental conditions, each lasting 30 minutes: (1) life story book (2) rummage box (3) no-intervention. 2 baseline periods of no-intervention followed by intervention phase; participants took part in each of the 3 experimental conditions thrice (total = 9 occasions), delivered over consecutive days in a randomised order.</td>
<td>None (excluding no-intervention condition)</td>
<td>Dementia Care Mapping (DCM). Within DCM: - Wellbeing, including mood and engagement (Mood-Engagement Value; ME)- Behaviour (Behaviour Category Code; BCC)</td>
<td>Both reminiscence conditions (life story book and rummage box): higher well-being scores (P &lt; .05) for all five participants. Both reminiscence conditions associated with increased communicative, expressive and intellectual behaviours. No significant difference between reminiscence conditions.</td>
<td>***</td>
</tr>
<tr>
<td>Nichols (2011), UK</td>
<td>Case study</td>
<td>4 individuals (3 males and 1 female; aged 52-62); ID and differing severities of dementia, living at an ID service.</td>
<td>Personalised technology- P1: Bedroom Door Sensor (to alert staff when P1 left the room to prevent unsafe use of kitchen equipment); Talking Photo Album (to prompt memories about himself and his life)- P2: Fingerprint Lock (to tackle tendency of losing keys)- P3 and P4: Being Alert trial (transmitters which use RFID Technology, to enable independence while minimising wandering into high-risk areas).</td>
<td>None</td>
<td>Individualised outcomes relating to independence, helping participants make meaningful choices and decisions, improving safety, and overall wellbeing.</td>
<td>Observed benefits - P1: improved safety; interactive photobook integrated into personal plan, enabling him to maintain an active and fulfilling life; - P2: Retained independence and could remain in his self-contained flat safely.- P3 and P4: Maintain familiarity of routine and surroundings, reduced restrictions, and improved independent living.</td>
<td>Screening criteria unmet</td>
</tr>
<tr>
<td>De Vreese et al. (2012), Italy</td>
<td>Prospective cohort study</td>
<td>60 participants (50% male in each group, mean age = 53.4 years); ID and varying stages of dementia (ranging from MCI to early stages of dementia).</td>
<td>Special care unit (SCU; N = 14)- Staff-oriented interventions- Environment-oriented interventions- Miscellaneous environmental factors (surveillance system, music therapy, animal-assisted therapy)- Client-oriented interventions: internal and external activities offered regularly (individual and/or group).</td>
<td>TAU: (1) Day Care (DC; N = 22)(2) Nursing Home (NH; N = 24)</td>
<td>Primary outcome: Collective rate of decline in cognition, daily functioning, and behaviour of participants (validated Italian version of DMR). Secondary Outcome: Number of DC attendees transferred to NH due to dementia-related increased need for assistance, with or without behavioural problems (compared to SCU residents over the 3 years).</td>
<td>3 deaths during 3-year period before follow-up (2 DC and 1 NH). SCU significantly improved DMR/SCS scores across 3-year period (compared to DC and NH), only when participants had Down’s Syndrome (p = .003). After 3 years in SCU, daily functioning remained stable in experimental group, but worsened in control groups.</td>
<td>****</td>
</tr>
<tr>
<td>Ali et al. (2022), UK</td>
<td>Randomised controlled trial</td>
<td>40 dyads (individuals with ID and dementia, and their carers); mean age = 60; 65% male, 95%. White British, 60% with Down Syndrome, 65% with Alzheimer’s disease, 55% with mild dementia.</td>
<td>40 sessions of manualised ICST, delivered by carers, twice a week for 30 minutes, over 20 weeks. Outcome assessments at 11 weeks (midpoint) and 21 weeks (end of intervention) with individual and their carer.</td>
<td>Waiting list (TAU)</td>
<td>Cognition (Orientation, language, attention, praxis, and abstract thinking (CAM-COG-DS); The Modified Memory for Objects tests from Neuropsychological assessment of Dementia in Intellectual Disabilities Battery)- Proxy measures (Executive functioning, memory, and language (CSDS); Ability to carry out daily activities (ADCS-ADL); Quality of life (QOL-AD)- Care (Caregiver burden (Care Giving Burden Scale); Competence (SCIDS); Depression and anxiety (HADS))</td>
<td>At 21 weeks, QoL was significantly higher after ICST, with a large effect size (adjusted mean difference: 3.11; 95% CI: -2.44 to 8.67). No differences in other outcome measures (underpowered). 87.5% and 97.5% completed midpoint and endpoint assessments, respectively. One dyad dropped out. 70% completed at least 20 sessions; high satisfaction with ICST.</td>
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Table 2. Continued.

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<tr>
<td>Lifshitz and Klein (2011), Israel</td>
<td>Case Study</td>
<td>1 54-year-old male; Down Syndrome and early-stage Alzheimer's disease.</td>
<td>Tailored program with the five parameters of MISC, relating to cognitive (i) focusing, time focusing, and orientation in space (ii) cognitive expansion, and expanding the communication chain, emotional (iii) providing opportunity for making choices (iv) rewarding with explanation, and behavioural (v) regulation of behaviour) components.</td>
<td>None</td>
<td>- Deterioration (DMR)- Staff observation</td>
<td>Memory difficulties remained; improvements in daily life (capacity and motivation to learn new skills). DMR score increased by only four points.</td>
<td>**</td>
</tr>
</tbody>
</table>
| Hawkes et al. (2019), New Zealand | Mixed methods repeated action research cycle (qualitative observations + quantitative surveys) | 4 individuals living in residential care; ID (3 with Down Syndrome, 1 with autism) and dementia (2 advanced, 1 moderate, 1 probable). | PIE programme (4 meetings, 3 observation rounds), activity resources, and life template books. First cycle: baseline observation, intervention, and repeat observation (conducted by two observers simultaneously for at least two 2-hour periods). Actions reviewed and followed by another round of observation. Second cycle focused on presence of continuous change after introducing activity resources (in meeting 3). | None | - Staff survey to rate self-efficacy, project usefulness, and degree of benefit to themselves and residents (survey adapted from Clinical Excellence Commission, alongside additional comments section). | Increased 'enriching' events with increased engagement, better communication and movement, and more resident interaction. Engagement was lasting, and more positive expressions (smiling, laughing, positive comments) were observed. Staff reports supported benefits to residents and themselves and discussed the positive effects of the variety of activity resources. | *** 

**Reminiscence.** Reminiscence therapy (RT) involves generally past activities, events, and experiences focusing on psychopathology (Butler, 1963). RT has shown beneficial effects on people with dementia (Butler, 1963). RT has been used in several studies to enhance reminiscence therapy, therapy, and care. Reminiscence therapy has been shown to improve mood, increase social interaction, and decrease agitation.

**Structured interventions.** Structured interventions all of which varied in content and method.

**Memory café.** Memory cafés have commonly facilitated open conversations for people with dementia and their carers and helped them feel less isolated (Probst et al., 2022). Kiddie et al. (2016) piloted a memory café facilitated by professionals with expertise in memory cafés, focusing on psychopathology (Butler, 1963). RT has shown beneficial effects on people with dementia (Butler, 1963). RT has been used in several studies to enhance reminiscence therapy, therapy, and care. Reminiscence therapy has been shown to improve mood, increase social interaction, and decrease agitation.
Cognitive stimulation therapy (CST). CST is a group psychosocial intervention implemented as a structured program incorporating mental stimulation, reminiscence, and orientation activities (Morley & Cruz-Oliver, 2014). Individual cognitive stimulation therapy (iCST) aims to improve accessibility where group therapy may not be feasible or preferred (Orrell et al., 2017). While CST and iCST were originally developed for people with dementia, Ali et al. (2022) adapted iCST for people with ID and dementia and conducted a feasibility RCT where carers were trained to carry out 40 sessions of manualised iCST over 20 weeks. Outcomes including cognition, quality of life, and caregiver wellbeing, were measured. Compared to usual care, quality of life was significantly higher in the iCST group at 21 weeks, with a large effect size (Cohen’s $d = 0.89$). However, no significant difference was found on other outcomes, and the fidelity assessment indicated that the sessions were not delivered to the intended standard.

Occupational therapy. Occupational therapy aims to promote wellbeing and overall functioning and is a suggested intervention for mild to moderate dementia (NICE, 2018). Hawkes et al. (2019) examined the effectiveness of a person-centred Person, Interaction, Environment, or PIE programme (Young et al., 2011) that included a manual, observation tool, and self-reflection form. Experiences were measured as ‘enriching’, ‘depriving’, or ‘neutral’, in context to (1) staff knowledge of the person, (2) staff interaction with the person, and (3) the physical environment and culture of care. A range of activities and life story books were offered alongside the PIE programme, and four cycles of observations and reflections were carried out (repeated action research cycle) over two months, with a mixed-methods approach. Compared to baseline, there were more ‘enriching’ events for residents, with increased engagement, better communication and movement, and more resident interaction post-intervention.

Combined psychosocial interventions. One study (De Vreese et al., 2012) investigated the effectiveness of a multicomponent nonpharmacological approach in a special care unit (SCU) compared to treatment as usual in day centre (DC) and nursing home (NH) groups. The components included staff-oriented, environment-oriented, miscellaneous environmental considerations, and client-oriented interventions. Post-intervention, they found some improvement in cognition and stabilisation of everyday functioning and behaviour in SCU residents, and deterioration in the control groups. The improvement, however, was only significant when the participants had Down Syndrome.
**Personalised interventions**

These are multicomponent interventions tailored to individual needs, rather than a manualised intervention delivered uniformly. Three studies, one mixed-method and two case studies, examined the effectiveness of individualised interventions.

Watchman et al. (2021) carried out a mixed-methods study on person-centred approaches to care, examining the impact of individualised goal-setting on behaviour and quality of life for people with ID and dementia. Participants set goals themselves, largely focused on improving mood, reducing anxiety, wanting to feel more like themselves, engagement in activities, and personal safety. They chose interventions relevant to their needs from a range of offered interventions; data collected over 10 months found that 74% of individual goals were met or exceeded expectations and qualitative interviews with staff echoed these positive effects. However, no significant change was found on individual measures of symptom severity and quality of life.

One case study (Nichols, 2011) discussed the efficacy of personalised technology in promoting independent and safe living in a local residential care home. Four distinct technological adaptations or interventions were implemented for individual cases; these included bedroom door sensors, talking photo album, fingerprint lock, and Being Alert (mark low and high-risk areas to prevent wandering off site into unsafe areas). Being Alert, specifically, was trialled for 12 weeks with two individuals, who often left the site without support staff. While there was no quantitative finding or statistical analysis conducted, all participants benefitted from personalised technology, primarily retaining their independence and leading a fulfilling life with minimal restrictions. Broader effects included economic benefits and family wellbeing, due to the reassurance of their relatives’ safety.

Mediational Intervention for Sensitizing Caregivers (MISC) was originally developed as an early intervention to improve child-caregiver relations, subsequently enhancing the child’s cognitive, behavioural, and emotional functioning (Klein, 2000). Another case study (Lifshitz & Klein, 2011), of an individual with Down Syndrome and early Alzheimer’s Disease dementia, investigated whether MISC mediation parameters were applicable for caregivers of people with ID and Alzheimer’s, and how it may affect the quality of interactions between the caregiver and individual. A tailored program with five parameters was implemented with the given case, and effects were measured through DMR scores (higher the score, greater the deterioration) and staff observations. Although memory difficulties remained, there were improvements in daily life and capacity and motivation to learn new skills, and deterioration was minimal. However, specific outcomes were not measured.

**What specific adaptations have been made to facilitate the use of these interventions with this population?**

Adapting techniques is crucial when delivering interventions for people with ID, to meet individual cognitive abilities, subjective presentations, and ensure that they can gain and retain treatment benefits. Five out of the 10 included studies discussed adaptations made at varying stages of the study.

Photovoice is a participatory research approach wherein photographs are used to depict the research questions; participants observe the study procedure, take photographs that are meaningful to them, and elaborate on the observations through interviews (Povee et al., 2014). Watchman et al. (2021)’s study included five co-researchers with ID, who attended seven workshops over 10 months to learn about dementia, photovoice, using cameras, and data analysis. Participatory action research helped address one of their research questions on enabling co-researchers with ID to reflect on personal experiences and build a novel and inclusive dialogue on dementia; here, it included themes on peer support, future planning, and fear of dementia and its progression.

Furthermore, Kiddie et al. (2016) piloted an adapted memory café for people with ID by simplifying cognitive tasks and broadening the time period for reminiscence, considering the possibility of early onset dementia. Other adaptations included higher ratio of staff to attendees to maximise one-to-one support, inclusion, and engagement of attendees in each activity. The staff was skilled in alternative communication methods such as Makaton, which aids communication by combining signs and symbols with speech (Tuffrey-Wijne & McEnhill, 2008). This knowledge promoted inclusion, alongside being beneficial in managing challenging behaviour. Lastly, the information provided at the café was specific to people with ID. Ali et al. (2022)’s study on iCST simplified some activities or substituted them with alternative ones to better suit people with ID, while aiming to retain the original themes. This involved consultations from a speech and language therapist, alongside a focus group consisting of health and social care professionals, carers, and individuals with ID, who reviewed the activities and provided feedback. The adapted manual was piloted, and feedback sought (Ali et al., 2018).

To promote inclusivity during recruitment, Crook et al. (2016) included a short demonstration of reminiscence during recruitment and provided red- and green-coloured cards to indicate agreement or disagreement. However, on trying this process and discussing with staff, they concluded that none of the participants had capacity; consultees were therefore invited. Hawkes et al. (2019) supplemented their manualised intervention with activity resources and explained that staff were ‘encouraged’ to use flexible methods and adapt the activity to person-centred goals. While no further detail was provided, flexible methods are commonly used when implementing psychotherapy techniques with individuals with ID (Whitehouse et al., 2006).

**Discussion**

This review aimed to summarize the evidence for effectiveness of direct psychosocial interventions for ID and dementia, while discussing adaptations that facilitate their use. Based on the ten included studies of distinct interventions, there is emerging but limited evidence on the effectiveness of psychosocial interventions for this population. Given the small sample sizes in all studies, and absence of large-scale RCTs, caution is needed while interpreting effectiveness. Behavioural interventions may reduce occurrences of stealing, loitering, and inappropriate sexual behaviours. Studies of structured interventions suggested improvements in several outcomes, including affect, communication, engagement, cognition, daily functioning, and quality of life. Personalised interventions were beneficial for achieving personal goals, largely pertaining to improving mood, increasing engagement, and ensuring safety and independence. Furthermore, several interventions were adapted.
through reasonable adjustments and flexible delivery. Other adaptations promoted accessibility by collaborating with co-researchers with ID and promoting participant autonomy and individual decision-making during recruitment, where possible.

To our knowledge, this is the first review focused on direct interventions and outcomes for people with ID and dementia, and specific adaptations that facilitate these interventions, therefore alleviating limitations identified by previous reviews.

Prior research has largely focused on either ID or dementia. DRO has been effective in reducing challenging behaviour in adults with ID (Lloyd & Kennedy, 2014) while other functional-analysis-based behavioural therapy for dementia has effectively reduced wandering (Douglas et al., 2004). Reminiscence has shown positive effects on mood and cognitive abilities in people with dementia (Cotelli et al., 2012), with suggestions of including it in routine care (Huang et al., 2015). Similar acceptability was found among people with ID, though less common in practice (Stueber & Hassiotis, 2012). There is substantial evidence of the effectiveness of CST and iCST in improving cognition and quality of life in dementia in individuals without ID (Gibbor et al., 2021), with recommendations of group CST for mild to moderate dementia (NICE, 2018). Similar improvements in cognition were observed among participants with Down Syndrome (Shanahan, 2014), though these were not significant compared to control groups, and research on CST for people with ID is limited. Furthermore, multicomponent interventions may target multiple modifiable risk factors and slow cognitive decline in dementia (Chalfont et al., 2020), while tailored interventions have been effective in improving quality of life for people with dementia living at home (Cooper et al., 2012).

**Implications**

This review provides some evidence for the benefits of psychosocial interventions in this population, which could be implemented in clinical practice. While behavioural interventions may improve behavioural problems, structured interventions, including multicomponent interventions, target several areas and may therefore, provide multiple benefits. Individualised interventions seem to have notable potential in meeting complex needs, including sensory impairments, which require adaptations and tailoring interventions to individual needs. However, the issue of access persists as most community ID teams do not provide these interventions. Lack of resources and staff who are trained to provide dementia-specific interventions are common barriers to person-centred care in health and social care settings (Martin et al., 2020).

The benefits of psychosocial interventions discussed in this review may furthermore, contribute towards alleviating increased antipsychotic prescription in people with ID and dementia (Sheehan et al., 2015), and support measures of reducing overmedication within ID and dementia populations, respectively.

Future research needs to focus on samples of people with both ID and dementia. RCTs or other experimental designs may help establish a cause-and-effect relationship, while follow-ups would indicate whether the intervention’s benefits are maintained (Llewellyn-Bennett et al., 2016). Co-production in adapting interventions may further provide insight from lived experiences (Acton et al., 2022), through increased involvement and engagement in feedback and consultation, as opposed to researcher-led methods. Larger sample sizes and comparative research would increase validity and reduce bias. Previous studies have acknowledged small sample sizes, recruitment difficulties, need for multi-site involvement, and longer recruitment periods (Ali et al., 2022; De Vreese et al., 2012; Watchman et al., 2021). Implementing these measures is however, accompanied by logistical issues; feasible methods that explore the acceptability of the intervention, while maintaining cost-effectiveness, therefore need to be discussed.

**Conclusion**

This review has analysed direct psychosocial interventions for people with ID and dementia, alongside adaptations to facilitate their implementation. While some interventions have emerging evidence supporting their potential to address complex individual needs, large RCTs and follow-ups are required to counter methodological limitations. Suggestions have been made for future research, with implications for accessibility in clinical practice and national initiatives on ID.

**Strengths and limitations**

Including grey literature reduced publication bias and improved the comprehensiveness of the review (Paez, 2017). On screening, none of these papers met the inclusion criteria for the final review however, as they either did not include participants with both ID and dementia or did not assess a psychosocial intervention or outcomes. Screening and quality assessment of papers was undertaken by two independent researchers, which ensured that relevant papers were not missed. Another strength is the systematic mixed studies review design, which includes quantitative, qualitative, and mixed-methods studies. While quantitative research is useful to determine the effectiveness of interventions, and the magnitude and direction of the relationship between the intervention and outcome, qualitative findings may explain how, why, and in what context an intervention is effective (Cerigo & Quesnel-Vallée, 2020). Considering the limited evidence on interventions for the ID and dementia population, broadening the study designs provides a more holistic view of current interventions.

However, there are several limitations to consider. Only including English language papers may increase bias. Furthermore, there was no co-production or consultation with service users. While six out of ten papers were of high or medium quality, three papers did not satisfy the screening criteria, thus questioning credibility surrounding the interventions’ effectiveness. Large RCTs, studies with medium to long-term follow-ups, and studies examining outcomes related to cost-effectiveness were all absent. Moreover, inadequate data collection methods form a source of methodological bias. For example, staff observations may not be a sensitive measure of change. However, most quantitative studies used adapted measures relevant to the population or cultural context. Behavioural interventions were investigated through low-quality case studies, which tend to have low generalisability and are difficult to make causal inferences from (Simon & Goes, 2013). The interventions were categorised as behavioural, structured, and personalised, though individual differences create uncertainty on their external validity. While some adaptations provided a rationale and were overseen by consultation, others were not justified by an evidence-base, thus highlighting the need for further research in this area.
and dementia care, aimed at strengthening person-centred care.

Disclosure statement
There are no relevant financial or non-financial competing interests to report.

Funding
This project is funded by the National Institute for Health and Care Research (NIHR) under its Research for Patient Benefit (RfPB) Programme (Grant Reference Number NIHR201934). The views expressed are those of the author(s) and not necessarily those of the NIHR or the Department of Health and Social Care.

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