Barriers and facilitators to primary healthcare for people with intellectual disabilities / autism: An integrative review

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Title page

Barriers and facilitators to primary healthcare for people with intellectual disabilities / autism: an integrative review.

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Conflict of interest statement

No conflict of interest was declared.

Ethical approval

This was an integrative review and did not require specific ethical approval.
Abstract

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Background

Globally, people with intellectual disabilities, autism, or both, experience health inequalities. Death occurs at a younger age and the prevalence of long-term morbidities is higher than in the general population. Despite this, their primary healthcare access rates are lower, their health needs are often unmet, and their views and experiences are frequently overlooked in research, policy and practice.

Aim

To investigate the barriers and facilitators reported by individuals with intellectual disabilities, autism or both, and / or their carers, to accessing and utilising primary healthcare for their physical and mental health needs.

Design and setting

An integrative review utilising systematic review methodology. (PROSPERO: CRD42018103103).

Method

Electronic databases MEDLINE, Embase, CINAHL and Cochrane were searched for relevant studies (all languages) using a search strategy. Two researchers independently screened the results and assessed the quality of studies.

Results

Sixty-three international studies were identified. Six main themes relating to barriers and facilitators emerged from an analysis of these studies. The themes included training; knowledge and awareness; communication; fear and embarrassment; involvement in healthcare decision-making; and time. All the themes were underpinned by the need for greater care, dignity, respect, collaborative relationships and the need for reasonable adjustments. Opposing barriers and facilitators were identified within each of the main themes.
Conclusions

Adolescents and adults with intellectual disabilities, autism, or both, experience several barriers to accessing and utilising primary healthcare. The findings highlight the reasonable adjustments and facilitators that can be implemented to ensure that these individuals are not excluded from primary healthcare.

How this fits in?

This review synthesises evidence on the barriers and facilitators to accessing and utilising primary healthcare perceived by people with intellectual disabilities, autism or both. The findings highlight important considerations for primary healthcare policy, practice and further research.
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Main text

INTRODUCTION

People with intellectual disabilities and / or autism experience health inequalities (1, 2). Death occurs at a younger age and the prevalence of long-term morbidities is higher than in the general population (2). UK guidance outlines the necessity of tailoring healthcare services to meet their needs (3). Despite this, their access to healthcare services is lower (4,5), their health needs are often unrecognised or unmet (6), and their views and experiences are frequently overlooked (7,8).

Whilst previous reviews have investigated health care access issues for people with intellectual disability (9, 10), a recent mapping of the health and wellbeing needs of adults with both intellectual disability and autism identified an absence of research to determine their needs (11). This lack of understanding represents a significant knowledge gap in efforts to improve their health and wellbeing. There may be overlap between these groups, but their needs may be unique and nuanced (11).

The aim of this study was to identify and synthesise evidence concerning the barriers and facilitators experienced by adolescents and adults with intellectual disabilities (only), autism (only) or both, to accessing and / or utilising primary healthcare for their physical and mental health needs.

METHOD

The integrative review (utilising systematic review methodology) was conducted according to a pre-specified protocol registered on PROSPERO (CRD42018103103) and written in accordance with the PRISMA (2009) checklist and reporting standards (12). The search strategy is contained within the registered protocol. Electronic databases were searched using key terms and MeSH headings combined: Ovid Medline (1946 to 22.08.2018); Embase (1974 to 22.08.2018); CINAHL Complete (Cumulative Index to Nursing & Allied Health Literature) (22.08.2018); and Cochrane (22.08.2018).

Studies were eligible if they were: studies of any design; included people aged 14 years and over (eligible for annual health checks in UK); formally identified, or self-identified, as having an intellectual
disability, and / or an autism condition; their family members, carers, support workers and / or healthcare professionals; studies which explored access to healthcare for any physical and / or mental health conditions involving the target population; any primary healthcare setting e.g. General Practitioners (GPs) and other providers in GP practices, pharmacies, dental surgeries, ophthalmic services, screening and immunisation services; primary healthcare services in the UK and in other countries with similarly structured, funded and resourced primary healthcare services; studies published in all languages; studies published between 2001 and 2018. These publication dates were not stated in the study protocol. Studies conducted before 2001 were excluded due to legislative changes introduced for the target population including the Department of Health & Social Care’s `Valuing People’ Strategy published in 2001. Systematic reviews, book reviews, editorials, commentaries, epidemiological studies, and prevalence studies were excluded.

‘Access to healthcare’ in this review is concerned with helping people to command appropriate healthcare resources to preserve or improve their health, and equity of access is considered in terms of availability, utilisation or service outcomes (13,14).

Titles and abstracts were first screened for relevance independently by two researchers (AJD, JR). Three researchers (AJD, PB, LH) independently read and assessed the full texts of relevant citations, using the pre-specified eligibility criteria. Any citation queries were discussed by the three researchers and a consensus decision reached to resolve any queries.

Study quality was assessed independently by two researchers (AJD, PB) using the Mixed Methods Analysis Tool: MMAT (15,16). The MMAT is designed for use in the appraisal stage of systematic reviews of different types of studies. The critical appraisal of studies considered issues such as the appropriateness of the study’s design to the study’s research objective. This was used to provide context for the findings of the study. The two researchers independently assessed the number of criteria met by each study in each of the MMAT’s domains of assessment (16) and provided each study with a score. Studies were rated as high quality (***** if all of the MMAT criteria were met, good quality (****) if 75% of the criteria were met, satisfactory (*** if 50% of the criteria were met, poor (**) if the study met 25% of the criteria, and very poor (*) if the study met less than 25% of the MMAT scoring criteria.
The researchers’ independent appraisal findings were compared and agreed. Any queries over studies’ appraisal were discussed with two other independent researchers (RL, NW) to reach a consensus decision.

Data were extracted from the included studies using a data extraction tool specifically designed and piloted by the research team. The data from the included studies were analysed using thematic analysis (17). Two researchers (AJD, PB) independently developed the themes. These themes were compared, key themes agreed, and narratively synthesised by the two researchers. A third reviewer (NW) was involved where necessary. NVivo v12 supported the data analysis.

RESULTS

Sixty-three studies met the eligibility criteria for inclusion in the review.

The review identified 53 qualitative studies, five mixed methods studies, four quantitative studies and one randomized controlled trial. Twenty-four studies were conducted between the years 2003 and 2010, and 39 studies were conducted between 2011 and 2018. Most studies were conducted in the United Kingdom (UK) (n=33), the United States of America (USA) (n=13), and Australia (n=8). Forty-nine studies explored the views and experiences of participants with intellectual disabilities, 13 studies explored the views and experiences of participants with autism; and one study explored the views and experiences of participants with both intellectual disabilities and autism.

Quality of studies

Of the 63 eligible studies, 46 (73%) were rated as high quality, two were rated as good quality, seven were rated as being either satisfactory, poor or very poor, and eight were not rated due to their lack of reported information. However, none of the studies were excluded from the review as any appraisal process is potentially only evaluating the reporting of the study rather than its actual conduct and content which may usefully inform the findings and discussion (18).
Themes

Six common themes were identified by the thematic analysis. Participants perceived barriers and facilitators within each of these themes as being integrated in an opposing fashion as narratively discussed below.

1. Training

Healthcare providers (both primary and acute healthcare) may lack specialist training in this field (19-21). Barriers to their training include time constraints, knowledge gaps, and uncertainties over specialist help (22). Despite this, healthcare providers recognise the importance of such specialist training for people with intellectual disabilities and for others with communication challenges (22). Training, knowledge, and awareness-raising for healthcare providers, family members, carers, and support services is essential (23). For example, training for healthcare providers in communicating with people who have intellectual disabilities and/or autism (24). People with intellectual disabilities should be included in the training of healthcare providers as 'experts by experience' (24-26). However, whilst such training may be essential, it may not be routinely undertaken due to resource constraints (21).

2. Knowledge and awareness

Some healthcare providers may lack understanding, knowledge, and awareness about how to support people with autism and intellectual disabilities (27-29), including how to make appropriate reasonable adjustments (28). This lack of knowledge and understanding may lead to poor attitudes such as an abrupt way of speaking or coldness towards people with autism or intellectual disabilities. Poor attitudes held by both healthcare professionals and non-health professionals, across both primary and acute healthcare provider settings, is a recurring theme in the Review’s identified studies (21, 30-32). A warm, friendly and caring attitude from healthcare providers enable service users to access healthcare facilities and discuss sensitive health concerns (21, 33-36).

3. Communication

Communication is a significant barrier for people with autism and/or intellectual disabilities (28, 37-41). It causes problems in primary care as inadequate communication can result in the wrong diagnosis,
inappropriate medication, and it can prevent a person’s access to receiving adequate healthcare (22, 42, 43). Studies found a lack of awareness by healthcare providers about the range of communication issues faced by people with intellectual disabilities and or autism when accessing and attending primary and acute healthcare settings (22, 44 - 46). Healthcare practitioners may rely on communicating with a carer, family member or support worker rather than with the service user directly (22, 24, 35, 44, 46). Carers do not always allow the service user to speak for themselves or carers might try to protect them from perceived harmful communication (27) thereby preventing service users from exerting control over their own healthcare needs (31). Not being listened to created anxiety for some service users with intellectual disabilities and or autism (47).

Good communication between the healthcare provider and the patient with autism and / or an intellectual disability is vital when accessing and utilising healthcare (47, 48). If these patients find healthcare stressful because of poor communication then they may lower their expectations, lower their attendance and feel disaffected, and this may lead to ineffective health care (22).

Some healthcare information may be incomprehensible and or difficult to obtain (24, 33, 49, 50). Accessible healthcare information is perceived to be a high priority by people with intellectual disabilities and or autism (47 - 49). The use of easy-read information, sign language, non-face-to-face communication such as via the telephone, not over-loading the service user with verbal information, and use of virtual reality have been suggested as preferred methods of communication for this population (25, 51).

The ability to see the same healthcare professional is important for people with intellectual disabilities and or autism (33). This notion is also shared by healthcare professionals themselves, with suggestions that this would provide the opportunity to gain a better understanding of the medical history and communication style of the service user (35). Being treated with dignity and respect and being valued by healthcare providers are key facilitators (46, 52) and important in forming good relationships (53). Collaborations between health and other social care providers are also essential (54, 55).
4. Fear and embarrassment

Fear and embarrassment is a barrier to accessing healthcare for individuals with intellectual disabilities and/or autism (56-58). These include fears of being judged over lifestyle choices, of blood tests and vaccinations (37), of medical instruments (30), and fears associated with a lack of understanding about screening procedures (58). Physical examinations can also be a source of embarrassment and or discomfort for individuals with intellectual disabilities (37, 59-62). Some people with intellectual disabilities and or autism find the clinical environment daunting (62, 63) due to unpleasant or alarming noises, odours, and bright lights (30, 64). The waiting room may induce anxiety, especially if the individual is unsure of why they are there (31). Facilitators, in terms of reasonable adjustments in the clinical environment, may include easy-read information, coloured pictures, models, photos, videos, symbols and demonstration dolls (35, 47, 64, 65).

5. Lack of involvement in healthcare decision-making

People with intellectual disabilities and or autism can make choices about their everyday lives, yet decisions about their healthcare may be made by their families, carers and healthcare providers instead (24). Their involvement in the decision-making process is an essential part of their healthcare (33, 65). They feel empowered when they are involved in the decision-making process and gain a better understanding of their treatment and diagnosis (27, 66).

People with intellectual disabilities and or autism value healthcare professionals, their support network and other professionals who work closely with them, and who have specialist knowledge and experience of working with people who have intellectual disabilities (25, 67-72). A joined-up approach, in which the sharing of inter-agency information is key, may help alleviate service users’ healthcare fears (25, 35). Tailored services which are person-centred, flexible and family-centred are highly regarded (25, 72-77).

6. Time

Prolonged times waiting to be seen, and limited time spent with a healthcare professional during an appointment, act as a barrier. Long waiting times are a major cause of anxiety and stress (24, 31, 44,
Additional time is often required for effective communication with people who have intellectual disabilities and / or autism (30, 35, 46, 47). However, despite recognising the importance of spending time getting to know the service user, some healthcare providers can struggle to find extra time to achieve this (21).

**DISCUSSION**

**Summary**

A lack of specialist training in both primary and acute healthcare is an important barrier, which may mean that healthcare providers lack knowledge and awareness of the healthcare needs of people with intellectual disabilities, autism, or both. This may be associated with poor communication between such healthcare providers and service users and a lack of involvement in healthcare decision-making processes for these service users. Effective communication delivered by specialist trained, friendly and caring healthcare providers who treat service users with dignity is essential. Primary healthcare providers need to provide more accessible health-related information for service users (both in terms of availability and format), shorter waiting times and longer consultation times, less daunting clinical environments, improved consistency of care, and greater multi-disciplinary collaborative working to improve the health and well-being of people with intellectual disabilities, autism, or both. Improving these issues could help alleviate some of the fears reported by service users, which is another reported barrier. However, the challenges involved in responding to these identified issues are acknowledged given that primary healthcare services in the UK are currently under intense resourcing pressures (80).

**Strengths and limitations**

A strength of this review is that it provides a timely summary of the recent literature from 2001 to 2018. The review importantly included the views and experiences of people with intellectual disabilities, autism or both, and their families or carers as well as healthcare professionals. It synthesised different types of studies using a rigorous methodology. However, a search of other relevant databases such as PsycINFO, grey literature (including guidance and policy documents), and checking the references and citations of included studies may have yielded additional results. Searches of grey literature and the
checking of reference lists and citations for included studies were not undertaken as originally stated in
the study protocol due to staffing resource constraints.

Most of the identified studies were conducted in High-Income Countries (HICs) and may not reflect the
views and experiences of people with intellectual disabilities, autism or both from Low-and-Middle-
Income countries. Findings are not generalisable to all HICs either as the review included studies from
different HICs such as the USA with different funding and organisational arrangements. The review
was limited to studies involving individuals with intellectual disabilities, autism or both aged 14 years
and over. There may be transferable evidence from studies involving children and younger people from
this population and other cognitive disability populations (e.g. patients with dementia, cerebral palsy,
stroke, or acquired brain injury). The use of convenience samples (22, 55), and self-selection (19) may
also bias the results. Some studies were limited to urban areas as opposed to rural areas which may
pose different healthcare barriers (81, 82).

Comparison with existing literature

This is the first known integrative review of barriers and facilitators to accessing and utilising primary
healthcare experienced by adolescents and adults with intellectual disabilities, autism or both. The
review’s findings are consistent with available UK guidance for general practitioners and other primary
healthcare professionals which outline the necessity of tailoring primary care services for people with
intellectual disabilities (3).

Implications for research and practice

The review’s findings highlight the reasonable adjustments and other modifications that could be
implemented to ensure that people with intellectual disabilities, autism, or both are not excluded from
primary healthcare research and practice. Despite the constraints facing primary healthcare services in
countries such as the UK, their contributions are crucial in addressing the health inequalities
experienced by this population.
References


Figure 1: PRISMA diagram illustrating the literature search strategy

- Records identified through database searching (n = 39979)
- Records after duplicates removed (n = 31582)
- Records screened (n = 31582)
- Records excluded (n = 31325)
- Full-text articles assessed for eligibility (n = 257)
- Full-text articles excluded, with reasons (n = 194)
- Studies included (n = 63) (Of which, 53 qualitative, 5 mixed methods, 4 quantitative studies, 1 RCT)

Not primary care, not views and experiences, not age relevant, epidemiological studies, prevalence studies, commentaries, theses, concerned with diagnosis, providers training, testing evaluation tools, reviews, discussion papers, not about access or utilisation, not target population.