Attitudes toward and experiences of clinical and non-clinical services among individuals who self-harm or attempt suicide: a systematic review

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Abstract

The prevalence of self-harm has increased substantially in recent decades. Despite the development of guidelines for better management and prevention of self-harm, service users report that quality of care remains variable. A previous systematic review of research published to June 2006 documented largely negative experiences of clinical services among patients who self-harm. This systematic review summarized the literature published since then to July 2022 to examine contemporary attitudes toward and experience of clinical and non-clinical services among individuals who self-harm and their relatives. We systematically searched for literature using seven databases. Quality of studies was assessed using the Mixed-Methods Appraisal Tool and findings were summarized using a narrative synthesis. We identified 29 studies that met our inclusion criteria, all of which were from high- or middle-income countries and were generally of high methodological quality. Our narrative synthesis identified negative attitudes toward clinical management and organizational barriers across services. Generally, more positive attitudes were found toward non-clinical services providing therapeutic contact, such as voluntary sector organizations and social services, than clinical services, such as emergency departments and inpatient units. Views suggested that negative experiences of service provision may perpetuate a cycle of self-harm. Our review suggests that in recent years there has been little improvement in attitudes toward and experiences of services for patients who self-harm. These findings should be used to reform clinical guidelines and staff training across clinical services to promote patient-centered and compassionate care and deliver more effective, acceptable and accessible services.

Introduction

Self-harm refers to any act of intentional self-injury or self-poisoning, irrespective of level of motivation or suicidal intent (Hawton et al., 2003a; NICE, 2022). The prevalence of self-harm has increased globally, with evidence of this in countries such as Norway (Tormoen, Myhre, Walby, Groholt, & Rossow, 2020), England (McManus et al., 2019), the United States, China, and India (McManus et al., 2019; Muehlenkamp, Claes, Havertape, & Plener, 2012; Tormoen et al., 2020). Psychologically, self-harm is associated with low self-esteem, interpersonal difficulties, and hopelessness (Fox et al., 2015; Hawton, Saunders, & O’Connor, 2012). Physically, self-harm can result in severe scarring, muscle and nerve damage, infection, and premature death (Hawton et al., 2012; Witt et al., 2021b). Self-harm is the strongest predictor of suicide (Carr et al., 2017; Geulayov et al., 2019; Hawton, Zahi, & Weatherall, 2003b) with approximately 50% of individuals who die by suicide having previous episodes of self-harm (Fazel & Rüsenvon, 2020; Foster, Gillespie, & McClelland, 1997).

Healthcare services have been criticized over their management of self-harm. Studies demonstrate a high degree of variation in self-harm management across general hospital settings (Arensman et al., 2018; Cooper et al., 2013). For example, the proportion of patient presentations for self-harm receiving psychosocial assessments in emergency departments in England was approximately 58% although it ranged by hospital from 28% to 91% (Cooper et al., 2013) despite this being recommended practice for self-harm presentations (NICE, 2022). There is also evidence to support the effectiveness of interventions in preventing repeat self-harm or suicide following a first episode (Witt et al., 2021a, 2021b). Rates of readmission to psychiatric inpatient care for self-harm are highest in the following year, with one third of these occurring.
in the first month after discharge (Gunnell et al., 2008). Despite this, national guidelines for the short-term management of self-harm have been found to be implemented by healthcare professionals in less than half of the encounters they have with patients (Leather et al., 2020). Together, this evidence highlights a need for improved care for people who self-harm, both in relation to psychosocial assessment and aftercare.

Eliciting patients’ attitudes toward services providing interventions for self-harm are essential as they identify barriers to service delivery and influence treatment engagement (Ribeiro Coimbra & Noakes, 2022). The ‘Interpersonal cycle of reinforcement of self-injury’ (Rayner, Allen, & Johnson, 2005) posits that patients’ experiences of stigmatizing attitudes from staff and negative therapeutic relationships can feed into negative cognitions about themselves, which can lead to treatment disengagement. Understanding patients’ experiences of services therefore enables identification of key areas of improvement to enhance treatment adherence and improve outcomes (Kapur et al., 2013b; Rayner et al., 2005; Ribeiro Coimbra & Noakes, 2022).

A systematic review of patients’ attitudes toward clinical services following self-harm published in 2009 identified predominantly negative perceptions, including poor communication between patients and staff; limited staff knowledge of self-harm and poor therapeutic relationships (Taylor, Hawton, Fortune, & Kapur, 2009). Many patients suggested a need for improvements in psychosocial assessment, referral pathways and access to aftercare. As that review was completed over a decade ago and focused only on clinical services, an update of the literature is needed to reflect contemporary practice, widening the scope to the full range of services currently available to people who self-harm.

The present systematic review aimed to examine attitudes of patients and their families toward clinical and non-clinical self-harm services from research published since the final search date of the previous review (July 2006). We also aimed to compare patients’ experiences of clinical and non-clinical services, defining clinical services as those provided by public or private healthcare providers (primarily consisting of clinicians), and non-clinical services as charitable and voluntary sector organizations, social services, and faith-based organizations.

**Method**

Our review followed the Preferred Reporting Items for Systematic Reviews and Meta-analyses (PRISMA) guidelines (Moher, Liberati, Tetzlaff, Altman, & Prisma Group, 2009). We pre-registered the review protocol with PROSPERO (CRD42021264789).

**Search strategy**

As our review represented an update of a previous systematic review (Taylor et al., 2009), we replicated their methodology but expanded our search terms to include clinical and non-clinical services, and updated terminology (supplementary materials: S1).

We searched seven electronic databases (EMBASE, MEDLINE, PsycINFO, Global Health, AMED, HMIC and CINAHL). We also searched Google Scholar and OpenGrey for gray literature. Eligible studies were limited to those in English language and published from July 2006 as the previous review included studies published up until June 2006 (Taylor et al., 2009). The initial search was conducted in July 2021 and the final search was conducted on 1 July 2022. The reference lists of included studies were hand-searched to identify further eligible studies.

**Inclusion and exclusion criteria**

We included published and unpublished primary research studies capturing the experiences or attitudes toward services of people who self-harm. Eligible studies were those that included participants with at least one episode of self-harm, irrespective of suicidal intent. Studies were excluded if participants experienced attempts of assisted suicide, euthanasia attempts or experienced harm without explicit intent (e.g. accidental overdose). We also included studies capturing the attitudes of carers and relatives of individuals who self-harmed. Studies were included if participants received any medical or psychosocial intervention for their self-harm episode from clinical services (primary or secondary healthcare) or non-clinical services (services outside of healthcare settings including but not limited to social, voluntary sector or faith-based services). In order to maximize the evidence, qualitative, quantitative and mixed methods studies were included, as was the case in the previous review (Taylor et al., 2009). Secondary analyses of data and systematic reviews were excluded.

**Study selection**

Search results were exported into Covidence (Covidence Systematic Review Software, 2021) and de-duplicated. All titles and abstracts were first screened by one reviewer (TU). Full text articles of eligible studies were then screened by a second independent reviewer (ZK or GB) using the predetermined inclusion and exclusion criteria. Any disagreements were resolved through discussions with a third reviewer (SR).

**Data extraction**

A data extraction table was used to extract information on authors, publication year, country of origin, sample size, sample characteristics (i.e. demographic information), type of self-harm behaviors, type of services and interventions, methodology, measures of attitudes and relevant quantitative and/or qualitative findings. All data were extracted by one reviewer (TU) and a second reviewer (ZK or GB) independently completed data extraction for 25% of articles to compare level of agreement.

**Quality assessment**

The quality of included studies was assessed using the Mixed Methods Appraisal Tool (MMAT) (Hong et al., 2018) by one reviewer (TU). A second reviewer (ZK or GB) independently conducted quality assessment of 25% of the papers to compare level of agreement. The MMAT has previously been validated for use in systematic reviews and was selected as it is designed to appraise a variety of study designs (Hong et al., 2018). Calculating an overall quality score is discouraged when using the MMAT, therefore, we reported scores for each criterion. There were high levels of agreement between the reviewers, with only one paper requiring discussion. All studies were given equal value in terms of contributing to the summary findings.

**Data synthesis**

We summarized quantitative and qualitative findings using a narrative synthesis approach as we anticipated a wide variety of study
designs, sample populations and measures and therefore substan-
tial heterogeneity of findings. We used validated guidelines for
narrative syntheses from the Economic and Social Research
Council framework to follow established practice (Popay et al.,
2006).

One researcher (TU) first grouped studies by methodology,
setting and population, tabulating key findings relevant to atti-
tudes toward services using these categories. Team discussions
were used to agree these categories. Findings were then compared
across studies to categorize similarities and differences in attitudes
by setting and population, and to identify meaningful higher-level
constructs (Popay et al., 2006). The final constructs were synthe-
sized following critical discussion with the wider team until com-
plete agreement on structure and content was reached. We have
reported findings by age group, highlighting similarities or differ-
ences in experiences or attitudes between young people and
adults. We have defined ‘young people’ as below 25 years old,
as it has been recommended that adolescence should be regarded
as continuing to age 24 (Sawyer, Azzopardi, Wickremarathne, &
Patton, 2018).

Finally, we sought the perspective of an individual with lived
experience of accessing self-harm services to help us interpret
findings.

Results

Study selection

The initial search identified 9443 studies, which was reduced to
6028 studies following deduplication. Full text screening was com-
pleted on 142 studies, with 26 studies deemed eligible and
included in the review. Three further studies were identified
from hand-searching the reference list of these included articles.
A total of 29 studies were included (Fig. 1).

Study characteristics

Characteristics of the included studies are summarized in Table 1. Studies were published between 2007 and 2022. However, 27 out
of 29 studies were published from 2015 onward. All in high- and
middle-income countries. These included 11 studies from the UK,
four from Sweden, two from Canada, two from China, two from
Norway, two from the USA, one from Australia, one from
Belgium, one from Finland, one from Ireland, one from
Portugal, and one from South Africa.

The gender profiles of participants were reported in 24 studies.
While one study included only female participants (Lindkvist
et al., 2021) and one included only male participants (Hassett
& Ibister, 2017), all other studies included a mix of female and
male participants. Five studies included participants who identi-
fied as trans, non-binary and/or gender diverse (Byrne et al.,
2021; Cliffe & Stallard, 2023; Mitten, Preyde, Lewis,
Vanderkooy, & Heintzman, 2016; Mughal, Dikomitis,
Babatunde, & Chew-Graham, 2021; Worsley, Barrios, Shuter,
Pettit, & Doupunik, 2019). Only three studies reported on partici-
pants’ ethnicity (Cross & Clarke, 2022; Johnson, Ferguson, &
Copley, 2017; Xanthopoulou, Ryan, Lomas, & McCabe, 2022),
all of which included exclusively or majority White participants.
The age range of participants was not reported in six studies
(Bantjes et al., 2017; Ejneborn Looi, Engström, & Sävenstedt,
2015; Fu et al., 2021; Michaud, Dorogi, Gilbert, & Bourquin,
2021; Miettinen, Kaunonen, Kylma, Rissanen, & Aho, 2021;
Mitten et al., 2016; Williams, Nielsen, & Coulson, 2020). Eight
studies included children and young people only (19 years and
under) (Cross & Clarke, 2022; Hassett & Ibister, 2017;
Holliday & Vandermause, 2015; Johnson et al., 2017; Lindkvist
et al., 2021; Mitten et al., 2016; Simoes, Dos Santos, &
Martinho, 2021; Worsley et al., 2019), three studies included
young adults only (18–24 years) (Byrne et al., 2021; Idenfors,
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<tr>
<th>Study (country)</th>
<th>Sample characteristics</th>
<th>Phenomena of interest</th>
<th>Type of service &amp; intervention</th>
<th>Study Methods</th>
<th>Measures of attitude</th>
<th>Data analysis</th>
<th>Summary of findings</th>
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<tbody>
<tr>
<td><strong>Qualitative studies</strong></td>
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<tr>
<td>Bantjes et al. (2017) South Africa</td>
<td>N = 80 Demographics N/S</td>
<td>Attempted suicide</td>
<td>Emergency psychiatric unit Generic short-term provision</td>
<td>In-depth interviews</td>
<td>Two interview topics: • Experience of hospital care • Ideas of how hospitals can reduce risk of future harm</td>
<td>Narrative synthesis using ethnological approach</td>
<td>• Positive experience of receiving care e.g. empathy and understanding but disruption of therapeutic relationships • Poor staff communication • Busy service experienced as unsettling and frightening</td>
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<tr>
<td>Berg et al. (2020) Norway</td>
<td>N = 18 Patients’ age: 18–57 years (M = 40.0) Patients’ gender: 11 female, 7 male Ethnicity N/S</td>
<td>Attempted suicide</td>
<td>Specialist service, acute medical wards, short term crisis ward Intervention N/S</td>
<td>Semi-structured interviews</td>
<td>Interview guide including experience of hospitalization, safety, treatment from staff, communication, and impact of care on subsequent suicidal ideation</td>
<td>Inductive thematic analysis using phenomenological approach</td>
<td>Three themes: • Positive experience of safe and sensitive care to suicidal ideation • Mixed experience of tailor-made, collaborative care • Mixed experience of protected adaptive practice</td>
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<tr>
<td>Byrne et al. (2021) Australia</td>
<td>N = 13 Age: 17–25 years (M = 21.2) Gender: 11 female, 1 male, 1 non-binary Ethnicity N/S</td>
<td>Self-harm or attempted suicide</td>
<td>ED Generic short-term provision</td>
<td>Semi-structured interviews</td>
<td>Three interview topics: • What happened during time at ED • Positive aspects of the experience • Negative aspects of the experience • Recommendations for service improvement</td>
<td>Thematic analysis</td>
<td>Three interrelated themes: • ED was experienced as distressing • ED environment and care was often counter-therapeutic • Hospital staff perceived as disinterested, dismissive, and lacked knowledge</td>
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<tr>
<td>Cliffe and Stallard (2023) UK</td>
<td>N = 25 Age: 18–31 Years Gender: 20 Female, 4 Male, 1 Non-Binary Ethnicity: N/S</td>
<td>Self-harm</td>
<td>Mixed interventions – online and face to face professional support</td>
<td>Semi structured interviews</td>
<td>Interview questions focused on student experiences and preferences of self-harm interventions</td>
<td>Thematic analysis</td>
<td>• Some students felt that they were made to feel self-conscious when accessing face to face support • Online support was helpful as it could be accessed at any time. This was seen as better in managing impulses to self-harm than waiting for face-to-face help. • Human connection was seen as a valuable tool in making patients feel safe, secure, and heard. • Most interventions are too short term which is not helpful in addressing the reasons behind self-harming.</td>
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<td>Study</td>
<td>Country</td>
<td>N</td>
<td>Age Range</td>
<td>Gender</td>
<td>Ethnicity</td>
<td>Study Design</td>
<td>Data Collection</td>
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<td>Enoksson et al. (2022)</td>
<td>Sweden</td>
<td>16</td>
<td>21–44 Y</td>
<td>14</td>
<td>N/S</td>
<td>Self-harm</td>
<td>Brief self-admission</td>
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<td>(M: 32.5)</td>
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<td>2 male</td>
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<td>Fu et al. (2021)</td>
<td>China</td>
<td>15</td>
<td>N/S</td>
<td>11</td>
<td>N/S</td>
<td>Self-harm or attempted suicide</td>
<td>Psychiatric inpatient unit intervention N/S</td>
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<td>female (mothers), 4 male (fathers)</td>
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<td>Hagen et al. (2018)</td>
<td>Norway</td>
<td>5</td>
<td>33–54 Y</td>
<td>4 female, 1 male</td>
<td>N/S</td>
<td>Attempted suicide (self-poisoning or hanging)</td>
<td>Psychiatric hospital unit</td>
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<td>Holliday and Vandermause (2015)</td>
<td>USA</td>
<td>6</td>
<td>15–19 Y</td>
<td>5 female, 1 male</td>
<td>N/S</td>
<td>Attempted suicide</td>
<td>ED</td>
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<td>Hassett and Isbister (2017)</td>
<td>UK</td>
<td>8</td>
<td>16–18 Y</td>
<td>all male</td>
<td>N/S</td>
<td>Self-harm: cutting, overdose, scratching, burning, strangulation, head banging, punching walls</td>
<td>Engagement with CAMHS</td>
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<th>Study (country)</th>
<th>Sample characteristics</th>
<th>Phenomena of interest</th>
<th>Type of service &amp; intervention</th>
<th>Study Methods</th>
<th>Measures of attitude</th>
<th>Data analysis</th>
<th>Summary of findings</th>
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</thead>
</table>
| Hume and Platt (2007) UK | N = 14 | Self-harm | Inpatient units, EDs, GPs, community psychiatric teams, social care, Samaritans | Semi-structured interviews | Seed categories used to guide interview including questions on experiences, attitudes and feelings about treatments and interventions | Grounded theory | Various experiences of care:  
• Dissatisfaction with after-care  
• Need for mutual understanding and support from healthcare staff  
• Preference for community care over hospital-based settings |
| Hunter et al. (2013) UK | N = 13 (7 at follow-up) | Self-harm (cutting and/or poisoning) | Specialist self-harm unit Psychosocial assessment | Semi-structured interviews | Initial interview topics:  
• Experience service  
• Experience of psychosocial assessment  
• Outcome of assessment  
• Improving suggestions  
• Follow-up interview topics:  
• Impact/outcome of attendance  
• Further self-harm behavior | Interpretative phenomenological analysis | • Unclear purpose of psychosocial assessment  
• Positive experience of validation and emotional support  
• Negative experience of shame and feeling ignored, particularly for those with a personality disorder  
• Mixed experience of transitioning out of care and of aftercare |
| Idenfors et al. (2015) Sweden | N = 9 | Self-harm (cutting and/or poisoning) | ED, child and adolescent psychiatric unit, psychiatric ward | Semi-structured interviews | Open-ended questions based on experiences of professional care, highlighting shortcoming, positive aspects, and barriers | Content analysis | Three themes:  
• A need to be in good hands by professionals speaking the same language and showing trust  
• Help not matching life circumstances e.g. practical support and individual needs  
• A lack of autonomy in care |
| Johnson et al. (2017) UK | N = 7 | Self-harm | Residential care or Secure unit | Semi-structured interviews | Interview questions based on helpful and unhelpful staff responses, how to improve practice and support from staff | Content analysis | Global themes of safety and care:  
• Good levels of safety but often intrusive  
• Staff had adequate level of care but need for collaboration and empathy |
| Leung et al. (2019) China | N = 11 | Self-harm or attempted suicide (overdose, burning, cutting) | Community social services | Semi-structured interviews | Interview questions based on access to services, staff involvement, helpful and unhelpful aspects, and suggestions for improvement | Content analysis | Four main themes:  
• Mixed experience of service availability  
• Negative experience of accessibility |
<table>
<thead>
<tr>
<th>Study</th>
<th>Country</th>
<th>N</th>
<th>Demographics</th>
<th>Intervention</th>
<th>Data Collection</th>
<th>Analysis Method</th>
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<tr>
<td>Lindkvist et al. (2021)</td>
<td>Sweden</td>
<td>19</td>
<td>Gender: all female Age: 14–19 Years Ethnicity: N/S</td>
<td>Recurrent self-harm and suicidal behavior</td>
<td>Brief admission Semi structured interviews</td>
<td>Two interview topics: Experience of care Long term effects</td>
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<td>Thematic analysis</td>
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<td>Feeling safe and relieved Welcoming atmosphere by professionals Feeling independent &amp; less of a burden on loved ones Growing from self-reflection Receiving insufficient attention Feeling less prioritized than others Reports of unprofessional behavior from staff</td>
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<tr>
<td>Ejneborn Looi et al. (2015)</td>
<td>Sweden</td>
<td>19</td>
<td>Demographics N/S</td>
<td>Self-harm</td>
<td>Psychiatric inpatient unit intervention N/S</td>
<td>Online self-report survey with open-text responses</td>
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<td>Content analysis</td>
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<td>Three themes: Negative experiences of neglect and a desire to be understood and heard Experience of distrust between staff and patients and a need for mutual relation Experience of counterproductive and superficial care</td>
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<td>Michaud et al. (2021)</td>
<td>Canada</td>
<td>41</td>
<td>Demographics N/S</td>
<td>Suicide attempts</td>
<td>Specialized intervention group following a suicide attempt (Having a case manager, a crisis plan, meetings and follow up calls)</td>
<td>Semi structured interviews</td>
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<td>Thematic analysis</td>
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<td>Valued the human and professional qualities of nurses Appreciated follow-up calls and meetings Negative perceptions of joint planning with poor therapeutic relationship and understanding of difficulties</td>
</tr>
<tr>
<td>Miettinen et al. (2021)</td>
<td>Finland</td>
<td>27</td>
<td>Demographics N/S</td>
<td>Self-harm</td>
<td>A range of professional services.</td>
<td>Essays and interviews</td>
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<td>Content analysis</td>
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<td>Feelings of not being taken seriously in their distress. Feelings that professionals were ill equipped to address matters of self-harm upon seeing the injuries. Lack of information regarding the program of treatment and then a lack of monitoring led to feelings of abandonment. Frequent change of professionals led to</td>
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<td>Study (country)</td>
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</table>
| Mitten et al. (2016) Canada | $N = 12$
Age: 15–19 years ($M = 15.74$)
Gender: 11 female, 1 male, 1 non-binary
Ethnicity N/S | Self-harm | Inpatient psychiatric unit
Generic short-term provision, crisis intervention | Semi-structured interviews | Interview questions on stigma and experience of care | Content analysis | • Experience of stigma from staff a
• Positive experiences of physical and mental healthcare e.g. procedural knowledge from staff |
| Mughal et al. (2021) UK | $N = 13$
Age: 19–25 years ($M = 22.08$)
Gender: 12 females, 1 transgender male
Ethnicity N/S | Self-harm | GP
Generic short-term provision | Semi-structured interviews | Interview topic guide including:
• Experiences of GP care
• Access to GP care | Reflexive thematic analysis | Three themes:
• Difficulties in help-seeking avenues
• Barriers to support from GPs e.g. superficial care, dismissal
• Facilitators to care e.g. listening, understanding, and relationship-based care |
| O’Keeffe et al. (2021) UK | $N = 27$ (19 patients and 8 carers)
Patient gender: 16 female, 3 male
Carer gender: all female
Patient age: 17–77 years ($M = 39$)
Carer age: 48–77 years ($M = 59$)
Ethnicity: N/S | Self-harm | ED
Semi structured interviews. | Experiences of receiving care | Thematic analysis | • Patients felt judged for seeking help which exacerbates feelings of distress
• Patients felt accessing emergency services lacked human connection which they need to feel like their life is not hopeless.
• Patients felt that the standard questions asked are not an adequate way of addressing individual needs |
| Quinlivan et al. (2021) UK | $N = 102$
(88 patients and 14 carers)
Patient age: 18–75 years ($M = 34$)
Carer age: 41–73 years ($M = 56$)
Patient gender:
72 female, 16 male
Carer gender:
13 female, 1 male
Ethnicity N/S | Self-harm | ED
Psychosocial assessment | Online free-text survey responses | Interview questions designed to explore patient and carer experiences of assessment following self-harm | Thematic analysis | Themes based on before, during and after assessment:
• Compassionate, collaborative, and supportive care reduced likelihood of repeat self-harm
• Greater distress experienced due to stigma, overly standardized assessments, and invasive questions
• Poor after-care leading to despondency |
<table>
<thead>
<tr>
<th>Study</th>
<th>Country</th>
<th>N</th>
<th>Demographics</th>
<th>Intervention</th>
<th>Data extraction</th>
<th>Analysis</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Simoes et al. (2021)</td>
<td>Portugal</td>
<td>33</td>
<td>Age: 10–19 years Gender: 24 female, 9 male Ethnicity: N/S</td>
<td>Semi-structured interviews.</td>
<td>Four interview topics: Most important aspects of hospitalization Experience of care post discharge Family involvement Protective factors</td>
<td>Thematic analysis</td>
<td>Positive feelings of being understood and acknowledged Being able to be away from everything to be taken care of Negative experiences around feeling locked away and isolated Not enough one on one activities Lack of personalized care Medication changes without patient consultation.</td>
</tr>
<tr>
<td>Vandewalle et al. (2021)</td>
<td>Belgium</td>
<td>14</td>
<td>Age: 23–66 years (M = 42) Gender: 10 females, 4 males Ethnicity: N/S</td>
<td>Semi-structured interviews.</td>
<td>Two interview topics: Expectations v. experience of care for family member Own experience of care and support</td>
<td>Thematic analysis</td>
<td>Carers felt was listened to by professionals Concerns over lack of personalized care Feeling uninvolved and uninformed in care Concerns that carers were left unsupervised for long periods of time following a patient’s suicide attempt Not enough continuity or handover between professionals which led to patient/carers being asked the same emotionally charged questions repeatedly Insufficient support following discharge.</td>
</tr>
<tr>
<td>Williams et al. (2020)</td>
<td>UK</td>
<td>209</td>
<td>Demographics N/S</td>
<td>Online messages forum from self-harm support communities</td>
<td>Data extraction of information relating to service experience from online message forum</td>
<td>Thematic analysis</td>
<td>Difficulty in accessing services Positive experiences of medical support as a gateway to psychological therapy Fears over stigma, being misunderstood and confidentiality.</td>
</tr>
<tr>
<td>Worsley et al. (2019)</td>
<td>USA</td>
<td>27</td>
<td>Age: 9–18 years Gender: 11 female, 12 male and 4 transgender or non-binary Ethnicity N/S</td>
<td>Semi-structured interviews.</td>
<td>Interview topics: Perceptions of interactions with the clinical team What practices were beneficial Positive or negative experiences about care and hospital stay What should be changed about hospital stay</td>
<td>Thematic analysis</td>
<td>Eight themes: Positive experiences of clinical interactions Unmet information needs Distressing experiences of repetitive inquiries Safety concerns Bringing up fears related to previous hospital treatment Need to be engaged in activities to avoid boredom Positive physical comfort Mixed emotions following hospital care.</td>
</tr>
</tbody>
</table>
### Table 1. (Continued)

<table>
<thead>
<tr>
<th>Study (country)</th>
<th>Sample characteristics</th>
<th>Phenomena of interest</th>
<th>Type of service &amp; intervention</th>
<th>Study Methods</th>
<th>Measures of attitude</th>
<th>Data analysis</th>
<th>Summary of findings</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Xanthopoulou Et Al. (2022)</strong>&lt;br&gt;UK</td>
<td>$N = 28$&lt;br&gt;Age: 18–78 ($M = 35.36$)&lt;br&gt;Gender: 19 female, 9 male&lt;br&gt;Ethnicity: 27 White Caucasian, 1 Indian</td>
<td>Self-harm or attempted suicide (Overdose, hanging, drowning)</td>
<td>ED/Liaison Psychiatry Psychosocial assessment and referral</td>
<td>Semi-structured interview</td>
<td>Interview questions regarding professional conduct, communication, treatment planning, feeling after assessment and areas for improvement</td>
<td>Inductive thematic analysis</td>
<td>Two themes: • Positive experiences of therapeutic conversations • Negative experiences of formulaic assessment</td>
</tr>
<tr>
<td><strong>Mixed-methods studies</strong></td>
<td></td>
<td>Self-harm (Cutting, scratching, hitting, pinching, reducing intake, poisoning)</td>
<td>Community based therapeutic program.</td>
<td>Semi structured questionnaire</td>
<td>Qualitative: Semi structured questionnaire. Quantitative: Satisfaction questionnaire.</td>
<td>Qualitative data: Thematic analysis. Quantitative data: Wilcoxon Signed-Ranks test</td>
<td>Qualitative: • Friendly, responsive, and non-judgmental practitioners. • Quick speed at which they were contacted and seen. • An appreciation for individually tailored care; useful tools and strategies provided. Quantitative: statistically significant reduction in: • depression scores ($p &lt; 0.001$); • total anxiety scores ($p &lt; 0.001$); • total anxiety and depression scores ($p &lt; 0.001$), all with moderate effect size ($r = 0.47, 4 = 0.45$ and $r = 0.49$; respectively)</td>
</tr>
<tr>
<td><strong>Cross and Clarke (2022)</strong>&lt;br&gt;UK</td>
<td>$N = 61$&lt;br&gt;Age: 12–17 Years&lt;br&gt;Gender: 56 Female 5 Male&lt;br&gt;Ethnicity: 64.5% White, 27.4% Not disclosed</td>
<td></td>
<td>Community based therapeutic program.</td>
<td>Semi structured questionnaire</td>
<td>Qualitative: Semi structured questionnaire. Quantitative: Satisfaction questionnaire.</td>
<td>Qualitative data: Thematic analysis. Quantitative data: Wilcoxon Signed-Ranks test</td>
<td>Qualitative: • Friendly, responsive, and non-judgmental practitioners. • Quick speed at which they were contacted and seen. • An appreciation for individually tailored care; useful tools and strategies provided. Quantitative: statistically significant reduction in: • depression scores ($p &lt; 0.001$); • total anxiety scores ($p &lt; 0.001$); • total anxiety and depression scores ($p &lt; 0.001$), all with moderate effect size ($r = 0.47, 4 = 0.45$ and $r = 0.49$; respectively)</td>
</tr>
<tr>
<td><strong>Cully et al. (2022)</strong>&lt;br&gt;Ireland</td>
<td>$N = 32$&lt;br&gt;Age: 18–68 years ($M = 42$)&lt;br&gt;Gender: 18 female, 14 male&lt;br&gt;Ethnicity N/S</td>
<td>Self-harm</td>
<td>EDs, inpatient unit Psychosocial assessment, medical treatment, generic short-term provision</td>
<td>Qualitative: semi-structured interviews Quantitative: closed questionnaire</td>
<td>Interview topics: • Contact with mental health service • Role of services following self-harm presentation • Challenges with appointments • Benefits experienced from services</td>
<td>Qualitative data: thematic analysis Quantitative data: statistical analysis using Chi Squared or t test</td>
<td>Qualitative: • Mixed experience of care and support • Care described as comprehensive but lacked continuity • Experience of safety during crisis Quantitative: *Significant relationship between unsupportive care and: • feelings of hopelessness ($p = .008$) • repetition of self-harm ($p = 0.037$) • lower self-efficacy ($p = 0.038$)</td>
</tr>
</tbody>
</table>

N/S, not specified by authors; N/A, not applicable; M, mean; ED, emergency department; GP, general practitioner.
Kullgren, & Salander Renberg, 2015; Mughal et al., 2021) and twelve studies included participants across adulthood (18 years and over) (Berg, Rortveit, Walby, & Aase, 2020; Cliffe & Stallard, 2023; Cully, Leahy, Shiely, & Arensman, 2022; Enoksson, Hultsjö, Wardig, & Stromberg, 2022; Hagen, Knizek, & Hjelmeland, 2018; Hume & Platt, 2007; Hunter, Chantler, Kapur, & Cooper, 2015; Leung, Chow, Ip, & Yip, 2019; O’Keeffe, Suzuki, Ryan, Hunter, & McCabe, 2021; Quinlivan et al., 2021; Vandewalle et al., 2021; Xanthopoulou et al., 2022). We have reported findings by age group, highlighting similarities or differences in experiences or attitudes between young people and adults. We have defined ‘young people’ as below 25 years old, as it has been recommended that adolescence should be regarded as continuing to age 24.

Overall, the studies examined attitudes of patients/carers following a patient’s presentation for self-harm (n = 16), attempted suicide (n = 8) or a mixture of self-harm and attempted suicide (n = 5). Studies examined patients’ attitudes or experiences solely (n = 24), relatives’ attitudes or experiences solely (n = 2) or both patients’ and relatives’ attitudes and experiences (n = 3). Studies exclusively examined one type of service (n = 18) or a combination of services (n = 11). The clinical services included in studies were psychiatric/inpatient units (n = 12), emergency departments (EDs; n = 10), primary care (n = 4), secure units (n = 1), crisis wards/brief admission units (n = 3), community-based psychiatric teams (n = 3), community-based crisis care (n = 2), specialist psychiatric wards (n = 1), acute medical wards (n = 1) and Child and Adolescent Mental Health Services (n = 1). The non-clinical services included in studies were voluntary sector community-based programs (n = 1), social services (n = 2) or a voluntary sector helpline (Samaritans; n = 1). Based on these categories we made a team decision to group findings by clinical v. non-clinical services.

Quality assessment

Quality assessment ratings for the studies are presented in Tables 2 and 3. We judged 25 of the 27 qualitative studies to be of high methodological quality. Both the mixed-methods studies were assessed to be of moderate risk of bias.

Attitudes toward services from individuals who self-harm and their relatives

Our narrative synthesis of studies resulted in the development of four overarching constructs: staff attitudes, therapeutic contact, clinical management, and organizational barriers.

Staff attitudes

Professional stigma

The stigmatizing attitudes of professionals were reported in nine studies that examined clinical services. Across EDs and inpatient units, patients experienced negative judgements, service gatekeeping or belittling comments regarding their injuries (Mitten et al., 2016; Quinlivan et al., 2021; Williams et al., 2020).

Five studies reported a perception that professional stigma acted as a barrier to disclosure, with shame and fear impairing disclosure within psychosocial assessments and when help-seeking (Byrne et al., 2021; Hunter et al., 2013; Mitten et al., 2016; O’Keeffe et al., 2021; Xanthopoulou et al., 2022). Patients reported how their own low self-esteem and self-blame were reinforced by professionals’ stigmatizing attitudes (Byrne et al., 2021; Quinlivan et al., 2021; Vandewalle et al., 2021).

Experiences of professionals’ stigmatizing attitudes varied between clinical and non-clinical services, with the latter preferred for being more accepting. In one study, patients showed preferences for social services and voluntary sector organizations over hospital services, with the former described as more supportive and having the potential to build long-term relationships with patients (Hume & Platt, 2007). In one community-based program, staff (voluntary sector youth workers) were described as non-judgemental and friendly, reducing any shame felt by clients (Cross & Clarke, 2022).

Two studies set in clinical services described perceptions of stigma surrounding mental health diagnoses. Patients highlighted how professionals’ interest and compassion diminished after disclosure of a diagnosis of a ‘personality disorder’, with labels of ‘time-waster’ and ‘attention-seeker’ applied (Quinlivan et al., 2021). Whilst one UK-based qualitative study reported experiences of staff withdrawal and rushed assessments (Hunter et al., 2013), another UK-based qualitative study reported perceptions of psychiatric diagnoses being wrongfully used by professionals to minimize the severity of a patient’s self-harm on the basis it was expected or normalized (Quinlivan et al., 2021).

Young people and adults reported similar experiences of professional stigma, particularly in the ED setting (Mitten et al., 2016; O’Keeffe et al., 2021).

Minimization of distress

A tendency to minimize patients’ distress was reported in nine studies, in samples of young people and adults. Across EDs, GPs and inpatient units, staff were described as uninterested and dismissive of physical and psychosocial distress (Ejneborn Looi et al., 2015; Hagen et al., 2018; Lindkvist et al., 2021; Mughal et al., 2021; Xanthopoulou et al., 2022). Three studies set in clinical services reported experiences of staff prioritizing cases that they perceived as more ‘serious’ and patients whose injuries were not self-inflicted, further demonstrating professional discrimination (Ejneborn Looi et al., 2015; Fu et al., 2021; Hagen et al., 2018). Minimization also resulted in care being withheld; patients were told that pain medication and medical treatments were unnecessary, with staff making comments about a ‘waste’ of beds and resources (Byrne et al., 2021; Hagen et al., 2018; Quinlivan et al., 2021). Minimization led to patients viewing services as ‘cold’ and ‘robotic’, only responding if a ‘threshold’ of seriousness was met (Byrne et al., 2021).

We noted apparent age differences in findings, in that minimization of distress was more often mentioned in studies of young people (n = 6) than adults (n = 2). It was reported that some GPs treated young people’s disclosure of self-harm casually or were dismissive (Mitten et al., 2016; Mughal et al., 2021), and young people reported being told ‘it’s just a phase’, ‘heaps of young people your age do this, it’s normal, you’ll get over it when you’re older’ (Byrne et al., 2021). In one study, it was said that presentations were taken more seriously when a young person was accompanied by a family member (Byrne et al., 2021).

Therapeutic contact

Staff-patient relationship

Twenty-one studies presented data describing relationships with staff. Within non-clinical services (social services and voluntary sector services), clients generally described a strong rapport...
Table 2. Quality assessment ratings for qualitative studies using the MMAT

<table>
<thead>
<tr>
<th>Study</th>
<th>Is the qualitative approach appropriate to answer the research question?</th>
<th>Are the qualitative data collection methods adequate to address the research question?</th>
<th>Are the findings adequately derived from the data?</th>
<th>Is the interpretation of results sufficiently substantiated by data?</th>
<th>Is there coherence between qualitative data sources, collection, analysis and interpretation?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bantjes et al. (2017)</td>
<td>✓</td>
<td>✓</td>
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<td>Cliffe and Stallard (2023)</td>
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<td>Enoksson et al. (2022)</td>
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<td>Fu et al. (2021)</td>
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<td>Hagen et al. (2018)</td>
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<td>Hassett and Isbister (2017)</td>
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<td>Hume and Platt (2007)</td>
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</table>

✓ = yes, x = no, ? = can’t tell.
between themselves and staff, based on mutual understanding, non-judgemental care, and trust (Cross & Clarke, 2022; Hume & Platt, 2007; Leung et al., 2019). However, experiences within clinical services were variable. Studies reporting positive experiences highlighted genuine and sensitive contact as well as mutual understanding to empower patients and encourage them to collaboratively explore their distress (Cliffe & Stallard, 2023; Enoksson et al., 2022; Hagen et al., 2018; Hassett & Isbister, 2017; Lindkvist et al., 2021; Michaud et al., 2021; Xanthopoulou et al., 2022). This rapport allowed staff to respond to patients’ needs for more effective care, such as reacting to fluctuations in suicidality, distress, and instability (Berg et al., 2020; Byrne et al., 2021; Miettinen et al., 2021; O’Keeffe et al., 2021; Quinlivan et al., 2021). Two studies described similar experiences (Bantjes et al., 2017; Cully et al., 2022; Idenfors et al., 2021; Quinlivan et al., 2021). This led to a lack of confidence in staff and doubts over the quality of care. Relatives reported superficial, rushed, and formulaic, where generic tick-box questions were reported in eleven studies. Assessments were described as superficial and judgemental staff contact, particularly during sensitive discussions about the patients’ care and self-harm. This led to a lack of confidence in staff and doubts over the quality of care (O’Keeffe et al., 2021; Quinlivan et al., 2021). Two studies described similar experiences by young people and adults, whereby carers felt under-involved in decision-making but were overly depended on to keep the person safe (O’Keeffe et al., 2021; Vandewalle et al., 2021).

### Relationships with relatives
Relatives of patients also reported negative experiences within EDs and inpatient units, with four studies highlighting their observations of poor communication from staff. Relatives were often excluded from discussions about patients’ care, felt inadequately informed about prognosis and had their concerns dismissed (Fu et al., 2021; Quinlivan et al., 2021; Vandewalle et al., 2021). Relatives experienced superficial and judgemental staff contact, particularly during sensitive discussions about the patients’ care and self-harm. This led to a lack of confidence in staff and doubts over the quality of care (O’Keeffe et al., 2021; Quinlivan et al., 2021). Two studies described similar experiences by young people and adults, whereby carers felt under-involved in decision-making but were overly depended on to keep the person safe (O’Keeffe et al., 2021; Vandewalle et al., 2021).

### Clinical management

#### Psychosocial assessment
Attitudes toward psychosocial assessments within clinical settings were reported in eleven studies. Assessments were described as superficial, rushed, and formulaic, where generic tick-box questions denied opportunities to explore individual experiences and psychosocial difficulties (Berg et al., 2020; Byrne et al., 2021; Quinlivan et al., 2021; Simoes et al., 2021). While good staff knowledge of psychosocial assessment protocols was reported in EDs and psychiatric wards, knowledge about mental health in those settings was seen as insufficient, with patients recommending staff training to help them better assess the context for and
severity of a patient’s suicidality (Hagen et al., 2018; Holliday & Vandermause, 2015).

Across clinical services, patients and relatives reported a lack of involvement in treatment planning, with unnecessary repetition of questions leading them to believe that staff did not listen or understand their individual experiences (Fu et al., 2021; Quinlivan et al., 2021). However, care was positively experienced when staff were sensitive to patients’ emotional distress when completing an assessment, collaboratively explored the factors leading to self-harm and involved patients in treatment decisions (Johnson et al., 2017; Michaud et al., 2021; Worsley et al., 2019; Xanthopoulou et al., 2022). We could not compare young people and adults’ experiences or attitudes on psychosocial assessment, as we could not separate findings by age group.

Use of restrictions and coercive care

Eleven studies reported variable attitudes toward coercive care in clinical services. In five studies, patients and relatives described the benefits of restrictions and removal of potentially lethal objects to protect against further self-harm (Berg et al., 2020; Cully et al., 2022; Hassett & Isbister, 2017; Idenfors et al., 2015; Vandewalle et al., 2021). Many patients experienced EDs and inpatient wards as ‘safe havens’ that removed them from distressing environments (e.g. difficult home dynamics) meaning patients could effectively shift focus toward recovery (Cully et al., 2022; Worsley et al., 2019). Brief admissions were felt to empower some patients as they felt they were given more control over care through joint decision making (Enoksson et al., 2022; Lindkvist et al., 2021). The authors defined these as specialist units where patients had the autonomy to self-refer for brief periods (for example three-day admissions) to manage escalating risk. However, other clinical services such as EDs and more traditional psychiatric inpatient care were experienced more negatively as patients reported feeling disempowered by restrictions (Quinlivan et al., 2021; Simoes et al., 2021). In light of this, patients and relatives expressed the importance of communicative practice when imposing restrictions: where staff in EDs explained the rationale behind restrictions and used collaborative assessments, these mitigated feelings of anxiety and disempowerment (Quinlivan et al., 2021). Similar mixed feelings and experiences toward restrictions and coercive care were described by both young people and adults.

Discharge and aftercare

Negative experiences of discharge following an assessment for self-harm in clinical services were reported across 12 studies. Studies reported how patients felt ill-prepared and unsafe at discharge where feelings of abandonment diminished their trust in clinical services and triggered repeat self-harm (Berg et al., 2020; Byrne et al., 2021; Hume & Platt, 2007; Idenfors et al., 2015; Xanthopoulou et al., 2022).

Regarding discharge, some patients were not contacted by services at all, whilst other patients faced long waiting times (Hunter et al., 2013; Quinlivan et al., 2021). Those who did receive follow-up care were often disappointed due to its brief length, low number of appointments given, and prioritization of discussions about medication over psychology (Cully et al., 2022; Holliday & Vandermause, 2015; Miettinen et al., 2021; Quinlivan et al., 2021). However, two studies of clinical services investigating experiences of patients on brief admission units described positive accounts of detailed discharge plans and safety planning which provided patients with a sense of security (Enoksson et al., 2022; Lindkvist et al., 2021). Greater control over their care meant patients could readjust back into society comfortably (Enoksson et al., 2022; Lindkvist et al., 2021). Although the attitudes and experiences of discharge and aftercare were similar between young people and adults, we noted that findings of young people were more focused on concerns about being discharged too early or premature endings in treatment (n = 4) compared to adults (n = 1). Findings with mixed samples of adults and young people, were more focused on dissatisfaction of aftercare (n = 6).

Psychotropic medication

Seven studies reported on attitudes toward medication administration after self-harm, all of which were within clinical services: EDs, inpatient units and community-based psychiatric care. While medication was seen as helpful, staff were perceived to focus more on describing benefits whilst tending to minimize information on side-effects and risks (Ejneborn Looi et al., 2015; Idenfors et al., 2015). Changes in medication without follow-up consultations from staff led patients to view services as negligent (Hagen et al., 2018; Simoes et al., 2021). Patients and relatives reported that medication was often administered without adjunctive psychological interventions, which they experienced as avoiding problems rather than an effective resolution (Fu et al., 2021; Hunter et al., 2013; Vandewalle et al., 2021). Similar experiences and attitudes about psychotropic medication were described by young people and adults. Both groups had a desire for more information about medication side effects (Hagen et al., 2018; Idenfors et al., 2015).

Organizational barriers

Waiting times

Nine studies described negative experiences in clinical services of long waiting times across services for young people and adults. For EDs, inpatient and crisis management teams, lengthy waiting times for a psychosocial assessment led to feelings of anxiety, particularly when in busy and loud environments (Bantjes et al., 2017; Byrne et al., 2021; Miettinen et al., 2021; Quinlivan et al., 2021; Williams et al., 2020). Patients and relatives also received little communication regarding the purpose of the wait, reasons for delays and progress (Cully et al., 2022; Vandewalle et al., 2021). Beyond the ED, there were also experiences of long waiting times for aftercare following an initial assessment (Byrne et al., 2021; Miettinen et al., 2021).

In non-clinical settings, experiences were variable. One community-based program had an average waiting time of 1.7 days between assessment and referral contact, which clients cited as a key reason for high satisfaction (Cross & Clarke, 2022). However, long waiting times within social services were found to heighten client anxiety (Leung et al., 2019).

Access to care

Nine studies reported on access to care across clinical services. Young people, adults, patients, and carers, perceived that the broader system was failing individuals who self-harm. They often found themselves limited to crisis support because they face exclusion from services or endure lengthy waiting lists, resulting in a recurring cycle of ED attendance (Byrne et al., 2021; O’Keeffe et al., 2021; Quinlivan et al., 2021, p. 52). EDs, inpatient units and brief admission units were reported as having a lack of beds and staff, which patients felt contributed to excessive waiting
times, inappropriate transfers, and premature discharges (Byrne et al., 2021; Enoksson et al., 2022; Johnson et al., 2017; Miettinen et al., 2021). For brief admission, some patients felt the care was less specialized compared to what they would receive in EDs and wanted more options for psychological support (Lindkvist et al., 2021). However, others felt that they could call on staff freely within brief admission wards and also a sense of predictability and safety, unlike in busy and intense EDs (Lindkvist et al., 2021).

Many patients were unaware which non-clinical services were available to them and felt that they should be better integrated with clinical services for more accessible care following discharge (Cross & Clarke, 2022; Leung et al., 2019). For social and voluntary services, they suggested extended services hours, telephone/digital appointments, and better staffing to improve accessibility (Idenfors et al., 2015; Leung et al., 2019; Williams et al., 2020).

Discussion

Main findings

This systematic review of 29 studies examined attitudes toward and experiences of clinical and non-clinical services of individuals who self-harm, as well as the views of their relatives. Our findings relating to clinical services are comparable to those of the previous systematic review (Taylor et al., 2009) describing negative attitudes toward organizational barriers and clinical management. This suggests little systemic change in clinical service provision for self-harm in the last 16 years. However, our review also included views on non-clinical services, where staff attitudes and therapeutic contact were experienced more positively than in clinical settings.

Patients and relatives reported a lack of individualized and collaborative care within clinical services. This was characterized by superficial and formulaic contact that failed to recognize the complexity of self-harm presentations. These findings may be underpinned by the use of increasingly manualized approach within clinical settings as a means of managing high service demands (Hawton, Lascelles, Pitman, Gilbert, & Silverman, 2022). Clinical staff themselves have previously reported conflicts between meeting professional regulations and providing holistic care (Bhui, 2016). The only age patterning of constructs we believed in busy and intense EDs is that staff freely within brief admission wards and also a sense of predictability and safety, unlike in busy and intense EDs (Lindkvist et al., 2021).

Our review highlighted that genuine and sensitive therapeutic contact in clinical and non-clinical services was viewed as a positive experience that patients linked to promoting recovery, a finding which comes as no surprise. Previous research has shown how strong therapeutic rapport enables patients to feel valued and listened to and enables them to produce training for mental health professionals that is trauma-informed and reduces stigma, particularly for those with personality disorders.

Limitations

Our quality assessment highlighted four studies of low to moderate quality (Bantjes et al., 2017; Cross & Clarke, 2022; Cully et al., 2022; Mughal et al., 2021), but we included these with equal weighting to other studies in our synthesis for comprehensiveness. However, we acknowledge that these lower quality studies may potentially have introduced bias. We limited our initial search to studies published in English, which may explain why all included studies were published in high-and middle-income countries. Moreover, only three of the included studies provided information on participant ethnicity, having either a majority or only white-Caucasian participants. Research has demonstrated that Black, Asian, and Minority Ethnic (BAME) groups experience poor access and quality of care from services due to poor cultural sensitivity and discrimination (Al-Sharifi, Krynicki, & Uphogrove, 2015; Memon et al., 2016). Important attitudes from BAME groups may not have been captured in this review. We differentiated findings by age group where possible. However, we could only do so for 14 out of 29 papers, as the remaining 15 papers had samples of mixed ages that spanned both adolescence and adulthood. This is important considering that self-harm is most prevalent in young people, with both young people and older adults demonstrating high levels of undisclosed self-harm and reduced help-seeking (Gillies et al., 2018; Memon et al., 2016; Troya et al., 2019). The study by Worsley et al., included individuals as young as nine and isolating their experiences to see how they differ from older adolescents, may
have provided useful insights (Worsley et al., 2019). Different services are also available for different age groups (e.g. child and adolescent services or adult services), leading to potentially different attitudes. Finally, it was not possible to compare findings by gender. This is important as males have typically been underrepresented in studies focusing on self-harm mental health (Hassett & Isbister, 2017), and people identifying as trans, non-binary, and/or gender-diverse are at greater risk of self-harm (Marshall, Claes, Bouman, Witcomb, & Arcelus, 2016). Overall, there is a clear research need to explore attitudes toward services by different demographic groups.

Included studies inconsistently reported on patients’ histories of self-harm and clinical management. Therefore, we could not interpret findings in the wider context of patients’ previous experiences of services. Similarly, none of the included studies explicitly examined level of suicidal ideation, and the studies examining both attempted suicide and self-harm presentations did not differentiate findings between the two. While in the UK it is customary not to distinguish between episodes on the basis of intent (Kapur, Cooper, O’Connor, & Hawton, 2013a), it is possible that one-off or frequent attendance for recurrent non-suicidal self-harm elicits a less intense service response than presentations where suicidal intent is expressed, creating different experiences of care. As included studies did not permit us to examine this, there is a need for further research examining how experiences differ by suicidal intent.

Implications

Our findings show that attitudes toward clinical services have shown little improvement in the 16 years since the previous review by a UK-based team (Taylor et al., 2009). This suggests that the range of UK-based (Department of Health, 2017; NICE, 2013) and international (World Health Organization, 2014) guidelines and policies designed to support service provision have had limited impact. To drive real progress in service provision it may be useful to review guidelines based on these findings. Furthermore, the problems commonly identified by patients (long waiting times, understaffing and limited access to services) have clear implications for the expansion of services, which should be a priority for governments internationally.

With negative staff interactions having a major impact on patient attitudes (Ejneborn Looi et al., 2015; Holliday & Vandermulse, 2015; Hume & Platt, 2007), policymakers should consider recommendations previously made regarding effective staff training and clinical supervision within clinical services (Taylor et al., 2009). Widespread implementation of training, based on the Self-harm and Suicide Prevention Competence Framework (Leather et al., 2020; National Collaborative Centre for Mental Health, 2018) would provide mental health professionals, clinical managers and service commissioners with guidance for best practice. However, this framework does not seek to prescribe what should be done, but instead there is flexibility in its application that allows for person-centered care (National Collaborative Centre for Mental Health, 2018). Improving staff attitudes and knowledge has been shown to have a wide-scale impact on service quality (Ferguson et al., 2019). This, in turn, has the potential to improve the therapeutic value of psychosocial assessment and improve outcomes (Hawton et al., 2022). It may also reduce costs and pressure on services (Kapur et al., 2013b).

Our review also highlighted problems with staff interactions viewed as too standardized and superficial. This demonstrates the importance of the therapeutic relationship, whereby staff should build strong rapport with patients and relatives, involve them in treatment decisions and encompass sufficient flexibility in treatments to ensure that practice is person-centered.

This review substantiates the need for integrated services to maintain quality of care during therapeutic contact, discharge, and transitions in treatment. This is of particular importance during repeated service redesign, especially throughout periods in which the COVID-19 pandemic has impacted service provision. With transformations in services and diversions away from EDs toward other primary, community-based, and remote treatments, including mental health crisis hubs, better collaboration between services can promote effective care while reducing service pressure.

Future research

With findings demonstrating little improvement in clinical services in the last 16 years, health service researchers and policymakers should monitor the implementation of service guidelines. Research should also address the large gap in the literature pertaining to the attitudes of under-represented groups including older adults, BAME communities, LGBTQI+ communities and those from low-and middle-income countries. Such groups can offer vital insights that may have not yet been uncovered to broaden our understanding of the quality-of-service provision. Finally, research should evaluate the impact of training and specific service changes on patients and carers’ perceptions of services.

Conclusions

The findings of this review provide insights into attitudes of individuals who self-harm and their relatives toward clinical and non-clinical services, which remain largely unchanged since a previous review 16 years ago. Across services, experiences of organizational and clinical management were largely negative, while staff attitudes and therapeutic contact were more positively experienced in non-clinical services compared to clinical services. Our findings have important implications for staff training and practice and should be used to reform existing healthcare guidelines for acceptable care for patients who self-harm.

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