Barriers to Accessing Addiction Treatment for Women at Risk of Homelessness

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Women remain under-represented in addiction treatment, comprising less than a third of clients in treatment services. Shame, stigma, and fear of legal and social repercussions (e.g., child protection involvement) are major barriers impacting on treatment-seeking for women. This is compounded for women at risk of homelessness, with practical and logistical reasons for not engaging in treatment. We conducted a qualitative study with both clinicians and service-providers, and women with lived experience of addiction and at risk of homelessness, to identify barriers to access and help-seeking within this vulnerable population. Adult women with lived experience of homelessness and addiction were invited to participate in an online focus group. Interviews were transcribed and analyzed using framework analysis. Analysis resulted in the identification of barriers to access in three areas. These were system-related, socio-cultural, and emotional barriers. We also present findings from the focus group recorded in real-time, using the novel method of digital illustration. This study highlights key factors impacting on help-seeking and access to treatment for addiction faced by women at risk of homelessness. The findings of this study highlight important areas of consideration for clinicians and service-providers working with women who experience addiction, as well as informing future research directions for this priority population. Findings are discussed in the context of exigent literature.

Keywords: addiction, treatment access, women, lived experience, homelessness

BACKGROUND

The relationship between alcohol and other drug (AOD) use and homelessness is complex. An individual can be defined as homeless if their current dwelling is inadequate, has no tenure, or tenure is short and not extendable, or the dwelling does not allow an individual to have control of, and access to space for social relations (1). There is no clear evidence for a causal link between the two however there is clear evidence that AOD use, and abuse is a significant issue for people who experience homelessness (2–4). Upshur et al. surveyed women across nine health care for homeless clinics in the United States (5). They concluded that when compared to the general population of women, homeless women were four times more likely to be diagnosed with alcohol use disorder and 12 more likely to be diagnosed with a drug use disorder. In Australia, the Australian Institute of Health and Welfare has collected data exploring the intersection of alcohol and other drug (AOD) use and experiences of homelessness over several years (6). In 2019-2020, 1 in 10 clients of Specialist Housing Services (SHS) reported problematic AOD use (6). Clients had significantly more health
issues, were homeless for longer periods, drew upon supports for longer periods of time and were more likely to be homeless at the end of a support period than SHS clients without problematic AOD use (6), highlighting the complex and chronic nature of substance use and homelessness in Australia. Limited access to treatment services, experiences of stigma, person and service level barriers are regularly reported in the literature as barriers to addiction treatment specifically (3, 7), and health care more generally (8, 9) by individuals who are experiencing homelessness.

Pathways to homelessness are wide and varied. From an interactionist perspective, homelessness can be understood as a contextual situation, underscored by a complex combination of human agency and structural factors, including, but not limited to poverty, financial stress, inadequate and unaffordable housing, mental health and/or substance use (10, 11). Pathways to and experiences of homelessness can differ between men and women (12–14). In addition to the aforementioned underlying causes, factors which could affect a women’s homeless status included their own chronic (mental or physical) health conditions (15–17), or their experiences in caring for another with chronic health conditions, harmful and/or violent relationships (18), and substance use issues (19). Women who are homeless report high rates of historical trauma (20, 21). Once homeless, women continue to experience high rates of assault (22) and victimization (23). Homelessness can also adversely impact upon pregnancy and the experience of motherhood. For example, having unstable accommodation can hinder access to reliable contraception and pre-natal health services (24, 25).

Furthermore, navigating motherhood in the context of transition houses or shelters raises specific issues in respect to parent-child relationships (26).

Experiences of AOD use and use disorder also differ across gender (27–29). Not only does the etiology vary (29, 30), but research continues to confirm the differential experiences of AOD use generally and substance use disorder (SUD) specifically across the sexes (31, 32). Experiences of treatment and recovery from SUD can also differ across gender (33, 34). Pregnancy, childbirth, and motherhood are regularly reported as factors preventing women from accessing AOD treatment (35–38). Women are also at an elevated risk of experiencing co-occurring mental health or personality disorders in comparison to men (39–42). Furthermore, shame, stigma, and fear of legal, and social repercussions (e.g., child protection involvement) have also been identified as barriers which can impact upon women seeking AOD treatment (38).

Despite the identification of such barriers, the number of women accessing AOD treatment remains low, and overall, women are underrepresented in treatment. In Australia women made up only one third of recorded public treatment episodes for SUD in 2019–2020 (43). Of those seeking support from Specialist Housing Services (SHS) in Australia, 60% are female (6), in 2019–2020 this equated to ~170,000 clients. Yet, despite the inherent links with problematic AOD use, reports indicate that only 0.8% of women seeking SHS support were referred to AOD treatment services, and only 0.4% or 728 women were provided with AOD counseling. Furthermore, recent economic modeling suggests that even in the presence of a universal healthcare system, out-of-pocket expenses pertaining to health care expenditure inexorably effect lower income earners (44). Taken together, these circumstances suggest that women who experience co-occurring AOD use, and homelessness are at an extraordinarily increased likelihood of not being linked into treatment, compared even with women who use substances but who do not access SHS.

Experiences of AOD use and homelessness interact bidirectionally to prevent women from accessing and engaging effectively in treatment, perpetuating the cycle of disadvantage, and hindering efforts of recovery (5, 45). Despite recognition of the barriers faced by women living with homelessness and SUD, there remains little research into this experience from the perspective of those who live it. Knack et al.’s study exploring women who experienced homelessness engagement with health services in Sweden is one exception (46). They identified three themes which underscored these women’s capacity to access services. Firstly, a demand for a life in order, considered the conditional nature of health care access for this cohort and the requirement for women to have a suitably stable lifestyle. Secondly, the theme of being unwell, unsafe and a woman, explored the multifaceted needs of women as a challenge to the health care system. And finally, the theme of abuse vs. humanity, spoke of the power within health care encounters to reduce or elevate the patient. Despite the importance of these findings, it remains that an exploration of the experiences of women who are homeless and their capacity to access AOD treatment services has not been undertaken.

The aim of this study was to address this gap in lived experience perspectives on the barriers to AOD treatment faced by women with addiction and homelessness.

**METHOD**

This study was a qualitative research project undertaken at a specialist public Australian addiction treatment center, Turning Point, as part of a broader service development initiative aiming to address women’s needs in care. This project was approved by the Monash University Human Research Ethics Committee (MUHREC # 24562).

Inclusion criteria for the qualitative study comprised adult women at risk of homelessness (currently homeless or engaging with SHS due to their risk of homelessness), who also had a history of substance use disorder and a history of seeking treatment for this. Women were recruited through the service’s networks and by approaching service providers and clinicians. Potential participants were asked to speak with the research staff directly to express their interest in participating. Following receipt of an expression of interest, research staff provided potential participants with relevant information and documentation. This process was facilitated by the rapport developed between housing service staff and participants and included addressing practical issues such as access to telephones and internet.

Women were invited to participate in a focus group, which was held online (by Zoom) due to COVID-19 related lockdown.
restrictions. Participants were reimbursed AU$50 (in voucher form) for their participation in the focus group. For women who had challenges accessing suitable technology, they were invited to attend the clinical service on-site and were supported to attend the focus group from a clinical room computer.

The focus group was completed in October 2020. The focus group ran for approximately 90 min and was led by SA, with support from DR and TM. Seven women were recruited; all participants were over the age of 18 years and had experience with addiction treatment services. They, were, or were at risk of becoming, homeless. Three women attended the clinical service on-site, two women were in residential rehabilitation units [accessing devices provided by the service (i.e., tablet)] and two were in private accommodation; using their own devices.

The focus group discussion explored the following questions,

- How do you access addiction services?
- What are the barriers to accessing these services in Victoria?
- How do these barriers affect your capacity to access these same treatment services?
- How do you think services could be adapted/ altered or changed to address the identified barriers?
- How/when do you provide feedback services?
- What are the strengths and limitations of these feedback processes?
- What would you like to see in a feedback program, how would you access and use this?

The focus group was recorded via two processes simultaneously. The first method involved graphic recording, with a professional live scribing artist providing the development of an insitu output; a digital illustration. The document provided a novel way of reporting to stakeholders the findings of the focus group, as well as providing real-time reflection of the key issues being raised in discussion. The artist employed had prior experience working in the context of vulnerable populations and in the addiction space. The artist completed a confidentiality form, attended the focus group, and proceeded to visually summarize the information collected via a digital platform. The second, more traditional method, was the audio-visual recording of the focus group. This was captured, with participants consent, via Zoom functionality. The recording was transcribed verbatim by DR and reviewed by TM and SC for accuracy.

As a study in implementation, framework analysis was used as the primary methodology (47). Developed in the context of healthcare services, with the aim of identifying barriers to services, analysis is undertaken at two levels. Deductive analysis provides for the analysis in the context of a specific question or questions. In the current context, deductive analysis allowed for the exploration of the barriers specific to women at risk of homelessness accessing substance use treatment. Inductive, or “bottom up” analysis aims to identify the thoughts and feelings of the participant regarding the context under consideration. This dual process provided for the identification of practical barriers to treatment, and a deeper exploration of the lived experience of the participants. It is this specific knowledge that can facilitate services to address the barriers in a way that is consumer focused and inclusive.

Analysis occurred iteratively, in which the authors’ experiences of recruitment, planning and implementation of the focus group, the transcript and the digital recording were reviewed both independently and later, simultaneously. For example, coding of the transcript was undertaken, and initial themes were then examined in the context of the digital illustrations and the lived experience of the focus group. This process occurred over several months to provide significant reflection and discussion between authors. Final themes were developed by DR and confirmed by SC and TM.

FINDINGS

Digital Scribe

The digital scribe completed the recording of the focus group in real time, the images below (Figures 1, 2) were derived using the language and experiences of the participants directly. The incorporation of the graphic recording allowed for real-time reflection, within the focus group, of the themes that were raised. Both during, and at the conclusion of the discussion, participants were asked to review the images and provide feedback as to their accuracy.

Framework Analysis

Deductive analysis of the focus group transcript sought to authenticate and develop further the results of the digital scribe. As such, analysis identified the barriers experienced by participants when accessing treatment services. Two themes were identified during this analysis, system-related, and socio-cultural barriers faced by the women in their efforts to access treatment. Inductive analysis provided a space in which we could explore participants’ responses to the identified barriers and how they made sense of their experiences with treatment services. In doing so, we were able to explore more nuanced barriers to addiction treatment. This second level of analysis resulted in identification of the emotional barriers to treatment. These responses mapped directly onto the themes identified in the deductive analysis.

System (Related) Barriers

The women identified several immediate barriers pertaining to accommodation stability, which had, at one time or another, limited their access to AOD treatment services. Participants reported that under a catchment-based allocation of treatment services, having no fixed address meant they could not be waitlisted for a specific service. In the case of frequent changes of address particularly across catchments, eligibility for services was often jeopardized. In these cases, once a change of address was identified, the participant’s place on any single waitlist would be lost. As one participant noted,

“…you don’t know what suburb you are going to be in, do you know what I mean, it is like you are pretty much like wherever you can be and then someone won’t take you because you are not like within that zone or whatever”. The women also spoke of the impact that chronic homelessness had on their capacity to attend different addiction treatment
services. Attendance at long-term residential rehabilitation was not conducive to receiving or maintaining a property in the community. Outpatient treatment was contingent on all other aspects of their circumstances—housing, transport, finances—to be sufficiently secure to permit them to attend an appointment at a specific time and place. Just staying in contact with services could be difficult,

“...not updating your phone number and address, so you just sort of get, yeah, they are obviously trying to contact your old place and you know, your three new places ahead or something, …”

"Someone will steal your phone, or your handbag is gone… you try to reconnect with whoever you made the appointment with, and it is too late“.

Managing eligibility criteria, holding a place on wait list and maintaining contact with services were all impacted by issues related to housing instability. Once admitted to treatment, participants had to navigate appointments and maintain contact with services all within the context of their socio-cultural barriers.

**Socio-Cultural Barriers**

Social and cultural barriers to addiction treatment included multiple levels of illiteracy, socially constructed gender roles, and the chaotic nature of participants’ lifestyles. Barriers were further
reinforced in the face of treatment staff limited understanding of their circumstances.

As a group, participants reported low levels of literacy. In the traditional sense, literacy was expressed in terms of education. However, literacy deficits were further affected by ongoing substance use. Both directly, for example ongoing cognitive difficulties in comprehension and memory because of regular and sustained use, and indirectly, for example one participant reported the impact of an Acquired Brain Injury on her literacy; received when, under the influence of substance, she had a car accident. Participants reported that sometimes they just did not understand the language being used by clinicians or service providers. As this woman noted,

“If there was something that I didn’t understand, I would put it to their attention. I would say listen, I can’t understand, can you please put it in a different context where I can understand”

In the instance that traditional literacy was sufficient to access service information, participants reported that they just did not know where or how to obtain information that could assist in their accessing treatment. Health literacy was poor, participants were unaware of services and treatment types available.

“...not knowing who to go to for what and just trying to sort of find out yeah, what is out there”

Participants felt that they had been adversely affected by the transfer of information to digital platforms. Digital literacy was low and access to devices like smart phones (and associated consumables) was limited by the women’s earning capacity and the insecurity of homelessness.

“Even when I did have a phone cos I am not like very good with computers and stuff, like I would not have had a clue, like I wouldn’t have a clue how to access anything”

“...can’t afford data…. or credit, or whatever yeah you are sort of stumped.”

Low levels of literacy, across multiple domains, limited participants’ ability to access treatment, both in respect to traditional conceptualization, but also in respect to health services and the use of digital devices. Taken together these made it difficult to access the information available, and then to understand what this meant for them specifically.

The women also highlighted the limited suitability of specific programs in respect to their lives. Further to these service level barriers, traditional gender roles inhabited by these women meant that as mothers and carers they had to deal with the day to day needs of their children and other dependants. Inpatient treatment facilities and long-term residential rehabilitation were not viable options in the instance that the women were primary caregivers. Whilst outpatient services continued to operate on a traditional 9-5 structure and provided limited flexibility in appointments. As one woman noted,

“If you miss an appointment like to be assessed for detox or something…you have to start all over again”

Housing, feeding, clothing, and managing the care of the family became a top priority, their own health and wellbeing—including issues of addiction were “the bottom of the list.”

“...everything else being so hectic you tend to yeah, it [your own health and wellbeing] just gets push aside…”

Building upon this list of demands is the biology of addiction itself,

“...you wake up and that is what you think about doing, and then till you get it...you can't function …. everything else stops, and then once you have it you can sort of function…but then the day is nearly over, and appointments are gone”

This statement also alludes to the chaos which characterizes the lives of these women. Neatly summarized by one interaction between two participants, who continue,

“P6: Just also the distress of all of it, like when you are oh great, I have an appointment with ____ at housing or whatever and you will go there and it won't happen...the stress of all that, not getting a result when you are trying, you know what I mean?

P5: Or you get disheartened, yeah,

P6: especially when you have other people to look after and things like that, it gets a lot, you know

P5: …like and with the anxiety you are getting, like trying to keep appointments, and keep the phone and keep the family and the house all operating at the same time so you won’t be homeless...as were about to be...you know what I mean, often that gets, it just gets so much...even getting to this, like even getting to this today was a major event, Okay”

For the women we interviewed managing the practical and emotional aspects of their day-to-day experiences was overwhelming and exhausting. In this context, there is little space to make treatment a priority.

Whilst the socio-cultural milieu of the women who exist in this space between addiction and homelessness not only thwarted efforts to access treatment, during the conversation, it became clear the women believed that generally, service providers did not understand their lives. This was noted by the staff that stood out to the participants,

“____ has been fantastic, but I mean right from the start she knows how hectic and crazy our life can be and she gets [sic] and please can you send me the details the week before, so I don’t stuff it up and all that sort of stuff.”

And in comments such as,

“I know that people like you guys do the best you can”
Whilst acknowledging the support she received from staff in respect to the issue at hand (receiving assistance with computer access), underlying the initial sentiment is a belief that treatment staff cannot really understand the circumstances these women face.

The women interviewed painted a picture of a life that is restricted by multiple socio-cultural barriers. They are unfamiliar with the services available to them and how to access these. They experience discrimination in the face of a digital society, either having limited access to, or being illiterate in the use of such technology. Their roles as mothers are not realized by the services they seek to access. Their efforts to manage these obstacles in the context of active addiction and homelessness are further limited by service providers who are naïve to the realities of their lifestyle. When faced with the multiple complexities of their circumstances, participants felt that services put them in the “too hard basket1." Another felt that the socio-cultural barriers left services saying, "nah, that is too hard, not dealing with this”

**Emotional Barriers**

The emotional experiences of the women interviewed in response to the system-related and social-cultural barriers navigated removed them further from the treatments they sought to access.

Participants expressed frustration with a system that did not sufficiently meet their needs. Eligibility criteria and long waitlists were compounded by their unstable accommodation. They felt that services disregarded their roles as mothers and carers and that they failed to understand the sociocultural circumstances they inhabited. A comment which was noted in respect to the issue at hand (receiving assistance with computer access), underlying the initial sentiment is a belief that treatment staff cannot really understand the circumstances these women face.

But it was not just the mothers who felt vulnerable. As a young female, one participant explained the feelings of increased vulnerability in the detox space.

"...they sit you in _____ with addicts that are a lot older than me, I am only 24, and I was sitting in there with you know, with older people who were much heavier addicts, who were sitting there asking for Valium, Ahhhh, just yeah, it was pretty horrible, I won’t lie"

Finally, the women had to manage the stigmatization they felt when accessing services, they reported embarrassment, shame and distress associated with their lives and substance use.

"P5: you try and think that you are in control of everything and yeah you just don’t want to, I suppose, being honest and open, you know you feel like you are going to be, I don’t know, criticized or stereotyped… P6: it’s embarrassing…"

Frustrated, and vulnerable, they felt abandoned by the system that was advertised to support and assist them.

"they are just druggo’s do you know what I mean, don’t give a shit, you know it feels like that sometimes”

Attempts to access and engage in treatment, increasing frustration, and emotional distress enhanced vulnerabilities, and reinforced stigma in the lives of these women, resulting in entrenched feelings of hopelessness and helplessness. The emotional experiences of these women not only reduced any capacity to engage in the process of treatment but in many cases resulted in the opposite—a return to substance use.

"You think kind of like ‘fuck it’ like why bother and like___ said, you go and get high again”

"You just go what you have been doing [using substances] because they are not going to help, you know it becomes a comfort, so you know it becomes easier I think”

**DISCUSSION**

The wider literature suggests women access services at reduced levels compared to their male counterparts and those who experience homelessness are further restricted in their capacity to access treatment services. To explore the intersection of gender and homelessness as a barrier to addiction treatment, we recruited seven women with lived experience of addiction treatment who were either homeless or at risk of homelessness to participate in a focus group. Our findings are presented in two parts, the output of the digital scribe and the themes identified via framework analysis. Whilst these were separated for presentation, it is of note that our discussion is reflective of the interconnection between the practical barriers identified, the emotional experiences of the women involved and the experiences of the focus group itself. Our findings support the

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1Participant 7.
2Department of Health and Human Services is the state service responsible for child welfare in Victoria Australia.
small but growing body of research exploring women’s access to AOD treatment in the context of homelessness.

Our findings are consistent with the limited extant literature. In one of the few studies identified exploring barriers to AOD treatment experienced by this cohort, women attending homeless healthcare services across nine US sites were surveyed in respect to the type of AOD treatment they had accessed and the barriers they experienced when accessing this treatment (5). Women with three different experiences of addiction were surveyed, those who reported alcohol use only, those who reported drug use only and those who reported use of both drugs and alcohol. The authors identified barriers to treatment from the relevant literature and asked women to identify which they had experienced. Barriers heavily endorsed by those surveyed were reflected in the current findings, including limited program availability, low levels of health literacy and extended wait times for services to become available. Across all three groups of women, the most frequently endorsed barriers were “feeling depressed/not up to going to treatment” and “too busy.” While methodological differences preclude any direct comparison between studies, our findings may provide insight into these highly endorsed barriers. The women we spoke to felt frustrated with and ignored by AOD treatment services. In the context of these experiences, it is not unexpected that motivation to attend AOD treatment services is reduced. This could also reflect a practical issue in which they do not want to, or are unable to, manage the process of attending AOD treatment services in the context of ongoing substance use and homelessness reported by some of our participants. In these instances, participants felt they were at the “bottom” of the to-do list; this may potentially reflect the circumstances of other women in similar situation being “too busy” to attend treatment.

We highlight these comments for discussion, for as clinical staff, it is not uncommon to hear “I am too busy,” or to have the experience of not wanting to engage with an “unmotivated” client. However, taking statements like these at face value perpetuate cultural misunderstanding and reinforce the discrimination these women experience. The women we interviewed were keen to have the research team understand the chaotic nature of their circumstances and the barrier(s) this raised when attempting to access treatment services. Like an entangled ball of twine, to pull one thread, just causes another knot to tighten. A common scenario amongst the women we spoke with is laid out as an example. Their roles as mothers and carers meant that impatient or residential options were not practical. They may receive an opportunity to engage in outpatient sessions, however this can be reliant on their locale and the need for a fixed address to be waitlisted. In the instance that they secured an appointment, they had to be able to record this securely, attend this appointment—finances permitting—on time, with all relevant documentation. They may have to attend this appointment with children, or alternatively, rely on family or friends to provide free childcare.

The existing structure of treatment systems demand “a life in order,” and this is unrealistic for women living with homelessness. Intersectional stigma posits that individuals may experience stigma resulting from the dynamic interaction of multiple marginalized social identities (48, 49). Recently, intersectional stigma has been used to explore the experiences of women in this space (50). Our findings build upon this work by providing specific examples of how intersecting stigmas can present themselves as barriers to treatment. In one example a participant describes feeling shame and embarrassment regarding her circumstances (internalized stigma). She reflects that between her own feelings and the belief (anticipated stigma) that family services would be employed (enacted stigma) should she engage with services, her motivation to access treatment services was reduced. Development of a comprehensive understanding of the sociocultural context of women managing both homelessness and addiction will be an important step in developing treatment services that facilitate, not hinder access.

Knowledge of systems and services provides people with the capacity to make informed choices about their own circumstances, this includes when, where and how to access treatment and health services. Poor literacy and socio-cultural context intersect with addiction to prohibit this group of women from being active participants in their own treatment and recovery. One participant noted that sometimes she just did not understand the language that was used by services and clinicians, another noted that she did not even know “what to search for” when she had wanted to engage with treatment services. Another lamented the wholesale move of information and services, for example, counseling services and peer-based recovery groups, onto digital platforms (which was only expedited in the context of the COVID-19 Pandemic), noting not only limited capacity to navigate digital platforms but limited access to technology.

A lack of information or the capacity to access information, not only left the women interviewed feeling helpless and hopeless in their efforts to access treatment services, low levels of literacy, traditionally, and in respect to health and digital technology, has rendered these women almost powerless. Research highlights the importance of self-efficacy in achieving and sustaining good health, both physically and mentally (51, 52). In the context of poor literacy and the limitations of their sociocultural circumstances they felt shamed, embarrassed, and stigmatized, none of which provide for the growth of self-efficacy, a vital component of successful addiction treatment (53–55).

Taken together similar findings across time and place, suggest barriers preventing women who experience homelessness from accessing AOD treatment involve a complex interaction of individual, service, and wider social characteristics. Due to the limited research in this specific context, strategies of potential redress must be sequestered by exploring the wider literature. Existent research explore the importance of developing a genuine understanding of individuals’ complex lives (25, 56), as well as the importance of prioritizing good relationships between staff and those using services (7, 56, 57). The women we spoke with reported feeling uncomfortable exploring their circumstances with services providers none the less, they spoke openly about their experiences during our focus group. This discrepancy suggests that when provided with a space in which they do not experience fear of punitive state systems (child protection or legal systems) or ongoing stigmatization, they feel comfortable
discussing their experiences and the potential issues they are facing.

Our experiences with the focus group also reflect suggestions that a person-centered approach to care (56, 58) and the engagement of the user in treatment services at every level is crucial. Participants reported that they had never given feedback to the services they accessed (see Figure 2), engaging the experiences of this group can provide services with a better understanding of their experiences and circumstances, but also provide these women with an increased opportunity and competency to articulate their experiences.

Using traditional and novel methods of data collection we were able to contribute to a comprehensive understanding of women’s experience and, importantly, scope to addressing these issues. The current study was undertaken with the aim to explore the lived experience perspectives as to the barriers to AOD treatment faced by women with addiction and homelessness. This study contributes to the expanding body of lived experience perspective literature and highlights the value of multi-dimensional perspectives in informing service delivery. Whilst the women we spoke with identified multiple barriers to accessing AOD treatment, when efforts were made to address these issues, for example supporting access to technology, they were motivated to engage, and open up about their experiences. Incorporating lived experience perspectives into service development offers an opportunity to design services that take into account the real-world barriers that exist, and to adapt interventions using a gendered lens to support more effective engagement.

**DATA AVAILABILITY STATEMENT**

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by Monash University Human Research Ethics Committee. The patients/participants provided their written informed consent to participate in this study.

**AUTHOR CONTRIBUTIONS**

SA conceptualized and designed the research project. SA, DR, and TM undertook data collection. DR, TM, and SC analyzed the data. DR wrote the first draft of the manuscript. All authors contributed to manuscript revision, read, and approved the submitted version.

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