Factors shaping the mental health and well-being of people experiencing persistent COVID-19 symptoms or ‘long COVID’: qualitative study

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Background

Around one in ten people who contract COVID-19 report persistent symptoms or ‘long COVID’. Impaired mental health and well-being is commonly reported, including anxiety, depression and reduced quality of life. However, there is limited in-depth research exploring why mental health and well-being are affected in people experiencing long COVID.

Aims

To explore factors affecting mental health and well-being from the perspective of people with long COVID.

Method

Semi-structured qualitative interviews were audio-recorded and transcribed. Data were analysed using reflexive thematic analysis. Twenty-one people with long COVID participated in the study. Participants were eligible if they self-reported a positive swab test/antibody test or one or more commonly reported COVID-19 symptoms at illness onset, and experiences of one or more long COVID symptoms ≥3 weeks following illness onset.

Results

Five themes were identified across participant accounts regarding factors affecting mental health and well-being, including symptoms causing severe disruption to daily life, lack of service and treatment options, uncertainty of illness trajectories, experiences of care and understanding from others and changes to identity.

Conclusions

People with long COVID experience a range of factors that negatively affect their mental health and well-being. Providing patient-centred health services that integrate rapidly evolving research in this area is important, as are peer support groups and supported approaches to self-management.

Keywords

Long COVID; qualitative research; mental health; COVID-19; phenomenology.

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SARS-CoV-2 infection and long COVID

The COVID-19 pandemic, caused by SARS-CoV-2, has had major consequences for health and well-being globally. The effects of SARS-CoV-2 infection vary widely between individuals, from being asymptomatic to multi-organ failure and death. For many people, symptoms persist far beyond the acute phase of the disease. Various names have been given to this condition, including ongoing symptomatic COVID-19, post-acute COVID-19 syndrome, ‘long-haulers’ and ‘long COVID’. Here we use the term long COVID, as it has become popular among many people with the condition and is widely used in the medical literature; however, we appreciate that discussions continue.

Long COVID prevalence estimates are inconsistent because of the rapidly evolving research base, non-standardised diagnostic criteria and different populations studied. Current research suggests that approximately 10% of people in the UK will have ongoing symptoms 12 weeks after initial infection, and substantial numbers have symptoms lasting over 6 months. There is a growing burden of ill health from long COVID in the UK, alongside patient reports that their needs are not currently being met by existing health services. Long COVID symptoms can be variable and wide-ranging, relating to multiple organ systems, including respiratory, cardiovascular, neurological, psychological/psychiatric, gastrointestinal, musculoskeletal and dermatological; or they can be general and non-specific. Symptoms are likely to result from multiple potentially overlapping disease mechanisms. The most commonly reported symptoms include fatigue, breathlessness, cardiovascular abnormalities, decreased cognitive function, difficulties sleeping and abnormalities in taste and smell. Comparisons have been made between long COVID and other conditions, including post-intensive care syndrome, myalgic encephalomyelitis/chronic fatigue syndrome and the sequalae of infection with coronaviruses like SARS and MERS, which can cause protracted multisystem disease. However, although certain shared features exist, caution should be applied when drawing parallels with symptoms experienced in other conditions, as doing so may detract from appropriate investigation and management.

An evolving research base has sought to understand the types of services and support available to people experiencing long COVID, whereas UK clinical guidelines recommend supported self-management and a multidisciplinary approach to rehabilitation following investigative tests, as well as active monitoring and follow-up. Reports suggest multiple difficulties accessing appropriate care, including uncertainty about treatment options and a lack of coordinated care. These experiences have resulted in patients resorting to self-management of symptoms through trial and error and via information and advice taken from informal peer support networks.

Mental health and wellbeing

Well-being can be defined as functioning well at both a personal and social level. The PERMA (Positive emotion, Engagement, Relationships, Meaning and Accomplishment) theory of well-being represents five elements required for individuals to flourish, including positive emotion, engagement in tasks and activities, connections with others, having meaning and purpose, and a sense of achievement. Adapting to living with a chronic illness has the potential to profoundly affect multiple components of well-being.
as described by the PERMA theory. For example, reduced function can lead to difficulties engaging in daily activities, a reduced sense of purpose and achievement, and increased social isolation, which influences mental health; 18 having a long-term physical health condition is associated with a twofold increased risk of depression or anxiety compared with the general population. 19 Social identity theory also posits that having meaning and purpose in the roles that individuals play within society is crucial for health and well-being, and that any disruption in ability to fulfill these roles, for example as a result of chronic illness, can negatively affect mental health. 20

Effects on mental health and well-being are commonly reported by people with long COVID, in particular features of anxiety, depression and post-traumatic stress disorder, 23 , 25 with cognitive processing disorders (or ‘brain fog’) also frequently reported. 5 , 9 Mental health effects are likely multifactorial in origin, including the psychosocial context of the pandemic, 23 , 24 pre-existing individual susceptibilities 23 , 25 and physical biological processes related to SARS-CoV-2 infection. 24 , 25 Although mental health effects are widely reported, there is limited in-depth research exploring the perspectives of people with long COVID regarding factors shaping their mental health and well-being. Addressing this gap has been identified as a key priority by an international, multi-stakeholder forum 26 to improve understanding, promote empathic healthcare interactions and facilitate development of patient-centred and appropriate healthcare provision for people with long COVID. Therefore, this study aimed to explore the perspectives of people with long COVID with regards to how their mental health and well-being has been affected.

Method

Sample and recruitment

We conducted semi-structured qualitative interviews with people who self-reported long COVID. Participants were recruited via an online long COVID support group, an advertisement in the University College London COVID Social Study 27 newsletter and through social media. Seventy-four people responded to the invitation to take part, and from this participant pool, 53 people were purposively sampled to ensure that people of different ages, ethnicity and genders were represented in the study. All 53 potential participants were sent a study information sheet and a screening form to determine eligibility. Twenty-nine screening forms were returned, we received no further response from 23 people and one person declined further participation. Twenty-one people went on to take part in the study.

To assess participant eligibility for the study, we used the post-COVID-19 symptom criteria proposed by Greenhalgh et al, which was broadly representative of the consensus at the time. 28 Eligibility criteria consisted of a positive swab test/antibody test or one or more commonly reported COVID-19 symptoms at illness onset (persistent cough, loss or change in taste/smell, high temperature); and experiencing one or more broader symptoms ≥3 weeks following the onset of their first symptoms 29 (Supplementary Appendix 1 available at https://doi.org/10.1192/bjo.2022.38). Eligible participants were encouraged to ask questions about the study before providing written informed consent to take part and completing a demographics form. All participants gave their written informed consent to participate in the study. The authors assert that all procedures contributing to this work comply with the ethical standards of the relevant national and institutional committees on human experimentation and with the Helsinki Declaration of 1975, as revised in 2008. All procedures involving human patients were approved by the University College London Ethics Committee (project identifier 14895/005).

Data collection

All interviews followed a semi-structured topic guide and were audio recorded. Questions explored onset and initial impact of COVID-19 symptoms, ongoing impact and development of longer-term health problems, impact of long COVID on social lives and mental health, and worries about the future. Example topic guide questions are listed in Fig. 1 and the full topic guide is provided in Supplementary Appendix 2. Interviews were conducted by a female postdoctoral applied mental health services researcher (A.B.) via video (14 interviews) or telephone call (seven interviews), depending on participant preference. Interviews lasted on average 67 min (range 41–99 min), and participants received a £10 shopping voucher for their time. Data collection continued until no new concepts related to mental health and well-being were discussed by interviewees. 29

Data analysis

Audio files were transcribed verbatim by an external transcription company. Personal and identifiable data were removed from transcripts to maintain confidentiality. Transcripts were then imported into NVivo 12 software for Windows (QSR International Pty Ltd, Doncaster, Australia; see https://www.qsrinternational.com/nvivo-qualitative-data-analysis-software/support-services/nvivo-downloads) and analysed using reflexive thematic analysis. 30 An initial coding framework was built based on concepts taken from the topic guide. Initial codes were organised and labelled under the following headings: impact of long COVID symptoms on normal life, social activities, health and well-being; ability to access services and support; and activities that protect mental health. Codes were applied to the text with line-by-line coding, and when new concepts were identified within the text, additional codes were added to the framework. Three transcripts (14%) were independently double-coded by three researchers (A.B., H.A. (a male medical trainee and doctoral student in behavioural science and health) and K.E.J.P. (a male physician and clinical scientist)).

Fig. 1 Topic guide question examples.
research fellow with training in qualitative research)) who met to discuss emerging codes and themes and generate the coding framework. H.A. then applied the coding framework to all remaining transcripts, adding new codes as transcripts became available and until no new codes were identified. Codes were then arranged into themes pertaining to the research questions. The research team met to finalise themes and approve the final report.

Results

We interviewed 21 participants between November 2020 and September 2021. All but one interview was conducted within a 4-month period (between November 2020 and March 2021). Participants were aged between 26 and 70 years (mean 47 years) and were predominantly female (67%) and White British (67%). All participants reported a case of suspected or confirmed COVID-19 between 28 February 2020 and 11 January 2021, and experienced on average 12 symptoms (range 4–18 symptoms) for 29 weeks (range 8–52 weeks) at the time of interview, with the most common symptoms being fatigue (95%), muscle pain/weakness (86%), shortness of breath (81%), difficulties concentrating (76%) and memory lapses (71%). A total of 52% of participants reported a confirmed positive swab test or had tested positive for antibodies. Four participants were hospitalised during the initial phase of their illness. All participants were still experiencing symptoms at the time of the interview. See Table 1 for participant characteristics and Table 2 for symptoms and self-reported laboratory confirmation of SARS-CoV-2 infection.

Themes

We identified five themes regarding factors shaping mental health and well-being for people living with long COVID: symptoms causing severe disruption to daily life, and well-being (theme 1) and uncertainty of illness trajectories, care and understanding from others, and changes to identity. Fig. 2 demonstrates the relationships between themes.

| Table 1 | Participant characteristics |
|---------|-----------------------------|
| Characteristics | N = 21 (%) |
| Age, range (mean) | 26–70 (47) |
| Female | 14 (67%) |
| Ethnicity (self-reported) | |
| White British | 14 (67%) |
| Asian (Indian/Pakistani) | 5 (24%) |
| White Irish | 1 (5%) |
| Other Asian background | 1 (5%) |
| Marital status | |
| Married/civil partnership/living with partner | 13 (62%) |
| Single | 5 (24%) |
| Divorced/separated | 2 (10%) |
| With partner but not living together | 1 (5%) |
| Living situation | |
| With others | 16 (76%) |
| Alone | 5 (24%) |
| Employment | |
| Employed | 9 (43%) |
| Currently unable to work because of long COVID | 8 (39%) |
| Reduced working hours because of long COVID | 2 (10%) |
| Retired | 2 (10%) |
| Pre-existing physical health problem | 10 (48%) |
| Pre-existing mental health problem | 2 (10%) |

a. Data are self-reported via a demographics form.

Long COVID symptoms led to severe disruption to daily life and activities (theme 1) and prevented participation in enjoyable activities that might normally protect participant well-being. This disruption was exacerbated by a lack of service and treatment options (theme 2) and by uncertainty of illness trajectories (theme 3). This, in turn, made managing uncertainties regarding trajectories, and pushing for service and treatment options even more challenging.

A lack of service and treatment options (theme 2) led to worries of exacerbation of symptoms and continued uncertainty around illness trajectories (theme 3); uncertainty of illness trajectories (theme 3) led to difficulties planning daily activities (theme 1), and subsequently affected ability to engage in enjoyable activities.

Positive experiences of care and understanding from others (theme 4) protected mental health, whereas a lack of care negatively affected well-being.

Ongoing long COVID symptoms that led to severe disruption to daily activities (theme 1) and uncertain illness trajectories (theme 3) led to changes to identity (theme 5), which negatively affected mental health because people were no longer able to fulfil their usual social roles.

Theme 1: symptoms causing severe disruption to daily life

Most participants described leading active lives before COVID-19 infection, working in full-time employment, participating in regular exercise and having busy social lives; however, ongoing symptoms had significantly curtailed activities of daily living.

Daily activities. Ongoing fatigue and ‘brain fog’ were commonly described as having a huge impact on people’s working lives, with almost half of the sample not able to fully return to work: ‘I couldn’t have even picked the laptop up and opened it, to be honest, you’re completely just wiped out’. The impact on daily life for some participants was profound, resulting in difficulties ‘even being able to manage basic household chores’ or struggling to maintain personal care:
'I haven’t showered for 3 days. I have to think about showering, I have to sit down and spend like 2 hours thinking about will I be able to manage showering today? Because even taking a bath makes me tired’ (550_female_aged_25-29).

Some participants described experiencing a prolonged loss or change in taste and smell that resulted in no longer being able to enjoy food. For those who had previously used cooking as a way to de-stress, this was experienced as particularly detrimental to well-being:

'That’s kind of disheartening … I could make a really nice meal which before lockdown would’ve been the kind of thing that I would like to do, but also for me personally, I see it as more of if you’ve had a bad day I’m going to do a nice thing for myself and invest in making a nice dinner for maybe me and someone else’ (565_male_aged_25-29).

Social life. Many participants were no longer able to participate in social activities, which had a profound impact on their mental health and well-being:

'I want to be able to have laughter, I want to be able to go out and be the life and soul of the party, which I’m just not anymore, so I do grieve’ (557_female_aged_35-39).

For some, the severity of their ongoing symptoms meant they were unable to socialise, mainly because of fatigue and the need to conserve energy by pacing oneself: ‘I haven’t got the energy to interact.’ For one participant, socialising became difficult because of breathlessness that made holding a conversation difficult:

'I couldn’t really talk to people … I just limited my [family member] to twice a week calls and I had to have short, 5-minute calls because more than that then I’d have difficulty breathing, So that was a bit frustrating that I couldn’t actually talk to people’ (569_male_aged_55-59).

Participants also described withdrawing from their social lives because of ‘not wanting to worry people’ and feelings of ‘guilt because I feel like I’m pouring more pressure on [my family]’ by sharing their experiences with others. Others withdrew because they felt that they had little to contribute to friendships because they were unable to focus on anything other than their health: ‘I was reflecting and thinking maybe I was talking about it all the time’.

'Well, early days, I’d get emails and stuff and then it’s just drifted off. It’s my own fault because I could phone up and chat to people, but I just feel it’s a bit boring. How are you? You tell them how you are … How’s your heart? You tell them, so, I’d rather not dwell on it. And I’ve got nothing else to talk to them about now’ (564_female_aged_70-74).

Enjoyable activities. Participants described the importance of engaging in new hobbies or activities or adapting previous activities to give them reprieve from their health problems, to support their well-being and to create a sense of achievement. Just under half of participants described engaging in low-grade physical activity such as walking: ‘I’m just doing some very gentle, gentle stuff. I’m walking’. Playing or listening to music or podcasts were important for distraction and relaxation:

'When my relapse started and into 3 weeks of it I think it was when I was in bed most of the time, I had these awful, awful feelings of almost … I felt like crying but I wouldn’t, but I’d change my mindset to, oh I’ll do something more positive than lying in bed. I had some pastels and I was doing a little bit of drawing, and I’d listen to the radio and put music on and stuff. I’d just alter it just by changing my mind from focusing on, oh I feel ill and will I ever work again, will I ever be able to do anything again, to oh well I’ll do this’ (559_female_aged_45-49).

However, for many participants, participating in enjoyable activities that could improve or protect mental health was not possible, such as physical activity: ‘I can’t do my exercises, and I can’t keep my body fit’, getting out of the house: ‘I’ve been stuck at home for the whole year’, or engaging in arts and culture:

'I’ve been trying to do some interesting things, so I watched a ballet online from the Royal Opera House, I went to a book launch of a book. I thought, I’ve never done that before, I’ll do that online. All sorts of opportunities. But even the things that are joyful and enriching, and would normally be lovely and powerful, if I get really excited about it, it just makes me tired’ (563_female_aged_60-64).
Some participants described replacing their usual physical activities with competitive activities that could be undertaken at home, such as online gaming:

‘That (online game) has been an absolute lifesaver for me because it’s kept my brain ticking over a bit, and given me, just a challenge, so I’m very fortunate with that. I’m sure other people have other things, but because I’ve got, naturally a competitive nature, with my golf, and I used to play tennis’ (564_female_aged_70-74).

Theme 2: access to services and treatments

Although just over half of participants had received investigative tests, many participants described a lack of follow-up: ‘we’re now on week 15 and I’ve still not had any results’ and ongoing stress and worry that the treatments or services they needed to manage their symptoms were unavailable, inaccessible or indeed were available, but not being offered to them:

‘What’s bothering me at the moment is I’m not in any plan or regime. So, I’m not on any pharmacological medicine. I’m not on any physical rehabilitation. So, I’m not on any programme. So, I just feel like I’m ambling. I’m just in the ether ... A year of my life has gone. I can’t carry on’ (553_female_aged_50-54).

Several participants described their frustration at being unable to access treatments or services because diagnostic tests were inconclusive or produced negative results despite ongoing symptoms:

‘I had several chest x-rays, and things, so I was looked after. But there was nothing wrong as far as they could see, but obviously, I was still having symptoms. But I was a bit frustrated. It was several months by then, and I was thinking why has nothing been set up yet for people who are suffering?’ (552_female_aged_50-54).

Although many participants described being aware of long COVID clinics, there was confusion around the referral process: ‘we’d been rejected because they’ve changed the criteria’, or there was a lack of available services in different areas of the UK:

‘There are no COVID clinics or anything in this area, so no. I have spoken to the doctors quite a lot since I’ve been ill, and they’ve been marvellous, but they’re still, we don’t know, we’re still waiting for the clinic to open’ (561_female_aged_50-54).

Just under a third of participants who described being unable to access National Health Service (NHS) care had gone on to access private healthcare, either by paying for care themselves: ‘the only door that is half-open, is if I decide to pay and go privately’, or through insurance provided by their employer:

‘So, without that private healthcare, I don’t think I would maybe have got over some of the symptoms that I had because, as I say, the GP [general practitioner] was just obviously it’s new for them as well, but they just weren’t offering any kind of help at all. And I remember when I went through them and says, well, I can’t actually refer you to the specialists with what you’re wanting under the NHS’. (564_female_aged_70-74).

Theme 3: uncertainty of illness trajectories

Most participants described feelings of anxiety around the uncertain symptom trajectories they were experiencing, with fluctuating periods of having ‘good days’ of feeling well and even, at times, ‘euphoric’ followed by ‘bad days’, periods of intense exhaustion and being unable to get out of bed. One participant described ‘the ebb and flow of it all’ as the most difficult thing to deal with:

‘There’s a lot of relapses involved ... which, initially, was very, very demoralising, because you think, or I would think, I’ve turned a corner, I’m going to get better, and then it would just suddenly get, like take two steps forward and three or four steps back. And that’s characterised the entire journey I’ve had with COVID, up until now really’ (560_female_aged_50-54).

For others, uncertainty about the future and how long their symptoms might continue led to increased anxiety about employment, finances and family life. Participants described feeling less in control of their lives as time went on:

‘I can’t live like this, how am I going to go on in the future, am I going to get worse, am I going to get better? Obviously, you think of everything. How are you going to look after your kids, the family, your work situation, how are you going to pay your bills’ (555_male_aged_45-49).

Participants also described having to weigh up the potential consequences of participating in activities that protected their well-being but that ultimately would lead to a relapse, against the immediate short-term benefits to their mental health:

‘I take the opportunities I can. So, a friend’s been off work and she messaged me yesterday and said, weather’s nice ... do you fancy a walk in the park? And I said oh great, yes, lovely, I’m free from here and here. Technically, that was too much in a day, and I knew it was too much and I paid for it later in the evening and this morning, but it was so nice to see her’ (563_female_aged_60-64).

Theme 4: care and understanding from others

Having a supportive and understanding social network, including friends, family and health professionals, was described as important for participant well-being, particularly when faced with fluctuating symptoms and uncertainty (theme 3). When this support was not available, participants described negative effects on their well-being and feelings of anxiety and loneliness.

Health professionals. Participants frequently reported feelings of frustration or anxiety because of a lack of understanding and support from health professionals. Experiences that exacerbated poor mental health included feeling ‘ignored’, ‘abandoned’ or ‘being brushed aside’, feeling that health professionals were unwilling to help or listen, a lack of continuity of care, and instances of diagnostic overshadowing whereby symptoms of long COVID were attributed to the participant’s mental health:

‘I have come across a few doctors where I’ve felt they’ve made me feel a bit stupid when I’m worrying about certain things ... So yes, there’s that worrying that people just think you’re making it up. I think it’s just that anxiety thing, for me that I don’t want to be an anxious person. And I think if I sense that I’m talking to a doctor, or sometimes they’ve said it outright, you’re an anxious person, and I think oh great they just think I’m anxious!’ (558_female_aged_35-39).

Even when health professionals were unable to offer appropriate medical treatments or answers, it was still important for participants to feel heard and for their stories to be believed and thus validated:

‘In the early days, my GP was fantastic ... I sent him a letter to tell him what was going on in my life, all my symptoms and everything. And I said, I am so sorry about harassing you. And he phoned me up, and he said, keep harassing me, he said, if you don’t tell me, I don’t know’ (564_female_aged_70-74).
Support from family and friends. Participants described the importance of receiving support and empathy from their social networks to protect their well-being during difficult times:

‘I’ve had a couple of other friends who said, look, don’t sit alone at home being miserable. If you need anything, if you just need to talk to someone, just give me a bell, or message me or whatever. So, that’s been nice, I think a lot of people didn’t realise how ill I was, and I think that when I went back to work, they were quite shocked, and were keen to reach out’ (560_female_aged_50-54).

In the initial stages of illness, participants described receiving emotional and practical support from family and friends: ‘[My sibling] used to do the shopping, and she’d sometimes do the cooking and bring it over’. For some participants, however, as their illness progressed and symptoms remained, emotional support became less available because of a lack of understanding or unwillingness to acknowledge that participants could still be unwell:

‘A lot of people don’t understand long COVID, so when you explain to them, I’m still not feeling right 6 months down the line, a lot of people have said, I think you’re just worrying too much. That’s what I think my parents come back with. They keep saying to me, you worry too much, there’s nothing wrong with you’ (555_male_aged_45-49).

Although most participants had not felt stigmatised because of their long COVID, a small number of participants reported experiencing external stigma directed at them from others. This included examples of family members ‘avoiding us like the plague’ for almost a year after initial symptoms or people behaving differently toward participants once they disclosed that they still had not recovered:

I’d walk up to the shops, buy one thing from the chemist or something and walk back. And I’d meet someone I knew, and I’d say, I’ve had COVID, they would step back, and so I stopped doing it because I realised that for a lot of people, I was still contagious, and I understand that. And I think that’s why I isolated, because it was quite difficult being around people that knew, who were not comfortable around me. And that’s even after 3 or 4 months afterwards, and even now, if I mention to somebody that I had COVID in March, they step back, and it is quite a difficult thing to deal with’ (564_female_aged_70-74).

For one participant, feelings of stigma arose from the impact of long COVID on mental health and worries that she might be judged by others for not being able to cope:

‘I think the stigma that was I mean around being depressed and having any mental conditions. I think people kind of judging for that. I’ve seen people do that. Especially in my friends’ circle’ (550_female_aged_25-29).

Discussion

Our study found that people with long COVID experience unique challenges to their mental health and well-being related to the impact of the symptoms on their lives, as well as interactions with health services, family and friends. Limited care and understanding from others, a lack of service and treatment options, disruptions to daily life and activities brought about by symptoms, the uncertainty of illness trajectories and changes to identity all contributed to a deterioration in mental health. Participants often described barriers to engaging in activities that might protect their mental health; however, being listened to and feeling validated by health professionals, family and friends, accessing peer support and engaging in enjoyable activities within the limitations of their condition, were identified as important for psychological well-being. The relationships between the factors identified in our themes and poorer mental health and well-being are likely to be mutually reinforcing. As people experience worsening mental health and well-being, their ability to search out and fight for the limited services that exist is also likely to decrease. Similarly, the mental health and well-being symptoms experienced may compound other symptoms such as fatigue and breathlessness. As such, the bidirectional...
relationships between many of the drivers of poorer mental health, and worse mental health itself, have the potential to create detrimental cycles of reinforcement.

Our findings align with, and extend, existing research into experiences of long COVID. They echo work that has previously described symptoms, impact on daily life and interaction with healthcare providers.\(^{14,22,31}\) However, our work extends previous findings by highlighting which features of long COVID are seen as most significant in shaping mental health and well-being from the perspective of people with lived experience. Our findings also improve understanding of how these features are experienced by people with long COVID, and as such, present potentially useful considerations for healthcare, self-management and wider society.

In keeping with the growing body of research, our participants reported multiple, varied, relapsing-remitting symptoms often severely affecting their ability to engage in daily life, resulting in a reduced sense of achievement, meaning and purpose.\(^{14,22,31}\) The PERMA theory of well-being posits that engagement in tasks and activities is vital to maintaining positive well-being.\(^{32}\) Therefore, these limitations have strong negative effects on mental health through limiting participation in enjoyable and meaningful aspects of life and those that gave individuals their sense of identity, including work, leisure and social activities. Social identity theory suggests that we derive meaning and purpose from the roles we play within communities and groups, and that this positively affects individual health and well-being.\(^{33}\) When participants in our study were no longer able to connect with their previous positive identities and these were replaced with illness-related identities, this posed a threat to well-being. Experiences of loss and threatened identity were also found in a qualitative study involving 114 people with long COVID,\(^ {14}\) and research regarding other chronic diseases such as chronic fatigue syndrome.\(^ {34}\) Relatedly, a small number of participants in our study described a lack of understanding from and feeling stigmatised by others as detrimental to well-being. Previous work on chronic illness has theorised that these experiences of illness-related stigma may reduce well-being via a reconstruction of an individual’s identity with both illness and stigma incorporated.\(^ {35}\)

Our findings identified the relevance of relapsing-remitting disease trajectories as important drivers of negative mental states because of related uncertainty about the future, which is well-established as a driver of anxiety and depression for people with various chronic diseases.\(^ {36,37}\) Other research, particularly from earlier on in the pandemic, also identified various barriers to healthcare,\(^ {1,14,22,31}\) including a lack of specialist service provision and limited understanding of the illness from healthcare professionals. Our study also shows that some people resorted to accessing private healthcare and therefore felt better supported to manage their symptoms. This finding raises concerns around health inequalities for people with long COVID who are unable to afford private healthcare. Potentially, with the creation of specialist long COVID services, increasing research and clinical guidelines, such barriers should become less of an issue over time, although given other competing needs faced by healthcare services, access could remain an issue. Feeling ignored or not believed by healthcare professionals, family members and friends has been commonly reported.\(^ {1,14,22,31}\) These experiences are similar to those described by people with myalgic encephalomyelitis/chronic fatigue syndrome.\(^ {38}\) Our findings suggest such experiences have particularly powerful negative effects on mental health.

Factors positively affecting mental health and well-being were infrequently mentioned. As in previous work, we did find that interactions where people felt listened to and believed were greatly valued, and empathy from health professionals, even when faced with limited treatment and referral options,\(^ {13}\) was important. Additionally, long COVID peer support groups appear valued by some individuals, although the potential for overwhelming and inaccurate information, as reported by participants in our study, is an important consideration.

**Strengths and limitations**

Certain limitations should be discussed. First, our sample included people who met clinical diagnostic criteria but without laboratory confirmation of COVID-19. This is in keeping with relevant guidance and excluding people without laboratory confirmation would have limited our ability to capture experiences of long COVID symptoms lasting up to 12 months, as testing was limited in the first part of the pandemic. Second, a degree of selection bias may have occurred toward those with mental health and well-being effects, and people may have been more likely to share their experiences if they were experiencing extremely positive or negative effects to their well-being. Third, our sample consisted of 21 people living in the UK, therefore assessing the external validity beyond this context is not possible, but likely to be relevant. Also, we did not collect specific data on where people lived within the UK, and we are therefore unable to comment on any potential regional variations in service provision and support. Most interviews were conducted over a 4-month period between November 2020 and March 2021, which may not have captured significant changes in long COVID healthcare provision and an increase in knowledge and awareness of long COVID over time. Finally, online methods of recruitment and data collection may have excluded some participants without access to the internet from taking part, although this method did enable participants to be interviewed in their own homes across different parts of the UK.

**Implications**

Our findings emphasise and support the need for accessible, patient-centred, specialist multidisciplinary healthcare provision. From healthcare professionals to family members and friends, ensuring people with long COVID are listened to, and their experiences validated, is vital. Importantly, optimising these experiences now features in learning modules and guidance for general practitioners.\(^ {36}\) Our findings regarding challenges in communicating long COVID experiences to family and friends suggest that it could be of value to adapt similar education and support materials and guidance for informal support networks. Given participants described specific mental health components of long COVID, these should be acknowledged as part of long COVID treatment pathways, and potential contributory factors identified and addressed where possible. Supporting individuals with long COVID in adapting their work, social and physical activities to meet their health capabilities, and to help individuals maintain or adapt their social roles and identity, could also be valuable as part of tailored occupational health programmes.\(^ {32}\) Long COVID guidelines now include the ‘Pace, Plan and Prioritise’ principles around ‘pausing’ physical activity for self-management,\(^ {37}\) but whether this strategy improves symptom management remains unclear, and participants in previous research describe receiving conflicting advice.\(^ {13}\) Additionally, effects on social participation suggests that there may be a role for tailored community groups and social prescribing, although this should be accompanied by research and evaluation to assess appropriateness and effectiveness.

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Supplementary material

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Data availability

The data are not publicly available due to their containing information that could compromise the privacy of research participants.

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Author contributions

All authors meet the criteria for authorship as recommended by the International Committee of Medical Journal Editors. A.B. and D.F. had the original idea for the study. All authors were involved in designing the study. A.B. provided guidance on qualitative methods, including design and analysis. A.B. conducted the interviews. H.A. led the analysis, with support from A.B. and K.E.J.P. Preliminary findings were discussed with the other authors and refined. The first draft of the manuscript was written by A.B. and K.E.J.P. All authors read, contributed to and agreed on the final manuscript draft.

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Declaration of interest

None.

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