The lockdown and its consequences—Perspectives and needs of people at increased risk of severe illness from COVID-19

Results from a qualitative longitudinal study

Erika Mosor · Valentin Ritschl · Margaret R. Andrews · Maisa Omara · Paul Studenic · Gertraud Schaffer · Ernst Leitgeb · Claudia Oppenauer · Linda C. Li · Tanja Stamm

Received: 17 August 2021 / Accepted: 27 October 2021 / Published online: 24 November 2021
© The Author(s) 2021

Summary

Background There is a lack of knowledge on how people at increased risk of severe illness from Coronavirus disease 2019 (COVID-19) experienced the infection control measures. This study aimed to explore their perspectives and needs during the coronavirus outbreak.

Methods A qualitative longitudinal interview study was conducted in Austria during lockdown due to COVID-19 containment and afterwards. People older than 65 years of age and/or affected by a chronic medical condition participated in individual telephone interviews at two time points. Thematic analysis was used to analyze the data and saturation was defined as no new emerging concepts in at least 10 subsequent interviews.

Data sharing statement We used a qualitative research design, and therefore the data generated are not suitable for sharing beyond that contained within the report. Further information can be obtained from the corresponding author. The study protocol was published on Research Gate (https://doi.org/10.13140/RG.2.2.12018.45762) and is freely accessible to everyone.

Supplementary Information The online version of this article (https://doi.org/10.1007/s00508-021-01979-9) contains supplementary material, which is available to authorized users.

E. Mosor, PhD, MSc · V. Ritschl, MSc · M. R. Andrews, MPH · M. Omara, MSc · C. Oppenauer, PhD · Univ.-Prof. Mag. Dr. T. Stamm, PhD, MSc, MBA (✉) Section for Outcomes Research, Center for Medical Statistics, Informatics, and Intelligent Systems, Medical University of Vienna, Spitalgasse 23, 1090 Vienna, Austria
tanja.stamm@meduniwien.ac.at

Results Thematic saturation was reached when 33 individuals (75.8% female, mean age ± standard deviation [SD] 73.7±10.9 years) were included. A total of 44 lower level concepts were extracted and summarized into 6 higher level concepts. They included (i) a general positive attitude toward COVID-19 measures, (ii) challenges of being isolated from the community, (iii) deterioration of health status, (iv) difficulties with measures due to their health condition, (v) lack of physical contact and (vi) lack of information versus overload. Participants suggested environmental adaptations for strengthening resilience in people at increased risk of severe illness from COVID-19.

Conclusion Strategies and interventions are needed to support people at risk under pandemic conditions. Their perceptions and needs should be addressed to reduce the potential deterioration of health conditions...
and ensure well-being even during prolonged periods of crisis.

**Keywords** Coronavirus · Aged · Chronic disease · Qualitative research · Patient preference

**Introduction**

The novel Coronavirus disease 2019 (COVID-19) spread rapidly worldwide, and the number of cases increased at an accelerated pace [1]. While unexpected changes in daily life affect all people, some persons are more likely to become seriously ill from COVID-19 than others. The Centers for Disease Control and Prevention (CDC) define these people like older adults and people of all ages with certain (chronic) health conditions and severe illness from COVID-19 as an increased risk for hospitalization, intensive care, need for a ventilator and/or death [2]. While the governments of many countries implemented measures to mitigate the pandemic [3], there is a lack of knowledge of how people with an increased risk of severe illness from COVID-19 specifically experienced these measures and under which conditions they were able to best adhere to them.

Recent work identified increased anxiety, depression, insomnia, and stress during the COVID-19 outbreak in all societal groups [4]; however, people who are more likely to be seriously ill from COVID-19 could be even more affected by collateral health damage and negative psychosocial consequences [5]. People from vulnerable groups might be more challenged in daily life and face more severe quarantine consequences, including deteriorated health and well-being, than people from other population groups [6]. Physical distancing measures, such as strict controls of any outdoor activities in response to the COVID-19 pandemic, aim to cut transmission by reducing close social contacts and have been recommended globally to control the community spread of the virus [7]. In Austria, people living in geriatric facilities were particularly protected by the government. Moreover, providers and managers of long-term inpatient care facilities had taken drastic measures, such as instituting visiting bans, curfews, and isolation of residents beyond governmental regulations.

Understanding people's living environments and preferences at risk of severe illness from COVID-19 in greater depth and incorporating their perspectives into the measures to fight the current or future pandemics are essential aspects of effective, holistic crisis management. Including these individuals' perspectives and health needs in planning measures to mitigate the pandemic [3], there is a lack of knowledge, few studies have investigated these people specifically experienced these measures and under which conditions they were able to best adhere to them.

Therefore, this study aimed to explore the perspectives and needs of people who are more likely to become seriously ill from COVID-19 during different phases of the coronavirus outbreak.

**Subjects and methods**

**Study design and participants**

A qualitative longitudinal interview study was conducted with individuals from different parts of Austria. Qualitative research is used to explore patients’ perspectives and understand reasons and motivations for behavior as well as preferences and values without imposing a preunderstanding [9]. After the first confirmed cases of COVID-19 in Austria on 25 February 2020, the Austrian government ordered nationwide restriction measures from 16 March 2020 onwards. Until mid-April, public life in Austria remained severely restricted by these measures before the first easing measures were implemented [10].

People older than 65 years and those affected by a chronic health condition, such as heart disease, diabetes, cancer, chronic lung disease and/or any immunodeficiency, were eligible for the current study. Exclusion criteria were insufficient language skills to participate in the interviews (either German or English language) and severe hearing and/or cognitive impairments that would make telephone interviews unfeasible. A maximum variation strategy regarding gender, age, educational level, comorbidity, living situation, and other socio-demographic characteristics was applied [11]. Qualitative research typically uses small sample sizes with a diverse range of participants to explore people’s personal experiences and views on a specific topic in depth [12–14]. Purposive sampling was used as it focuses on particular characteristics of a population of interest and therefore allows identification and selection of information-rich participants [15, 16].

Eligible individuals were identified by patient organizations, geriatric institutions, health professionals and those who participated in the interviews themselves. Appointments were made for remote interviews with participants. The most widely used principle for determining the appropriate sample size and evaluating its sufficiency in a qualitative study is data saturation [17], defined as “the point in data collection when new data no longer bring additional insights to the research questions” [18]. Data analysis started when the first transcripts were available and proceeded parallel to data collection to determine the point of saturation [19]. Despite different approaches in defining data saturation, researchers agreed on some general principles and concepts: no new data, no new themes, no new coding and the ability to replicate the study [20]. In our study, recruitment continued until data saturation was reached,
which was defined as no new concepts coming up in at least 10 subsequent interviews [13, 20, 21].

The study was approved by the Ethics Committee of the Medical University of Vienna (EK Number 1388/2020). Participants were informed about the study’s purpose and procedures and provided written informed consent by postal service or email in accordance with the Declaration of Helsinki [22].

Data collection

Two remote, semi-structured interviews were held with every participant. The research team, including patient research partners (GS, EL), co-developed, piloted and finalized the semi-structured interview guide for the remote interviews [23]. The interview questions focused on people’s perspectives, needs and preferences regarding the impact of COVID-19 measures on their mental and physical health, autonomy, social connectedness, activities, and work/productivity in daily life (Table 1).

The interviews started during the first lockdown, while the follow-up interviews were conducted in early May 2020 after the government eased measures for the first time. Therefore, the follow-up interview questions were adjusted for the different situation and the initial findings from the first interviews. The first author (EM) experienced in qualitative research data acquisition and analysis, performed the interviews. All interviews were conducted in German, audio recorded, transcribed verbatim, and analyzed centrally in Vienna, Austria, by EM with input from the research team.

Data analysis

Thematic analysis of qualitative data followed a modified form of “meaning condensation” [23], facilitated by using ATLAS.ti software developed by ATLAS.ti Scientific Software Development GmbH, Berlin, Germany [24] to organize the data. The analysis comprised the following procedures: in a first step, all transcripts were checked against the audio recordings for accuracy and read several times to gain a broad understanding. After familiarization, data were divided into meaning units (defined as specific parts of the text, a few words or a few sentences with a common meaning). Subsequently, initial codes were assigned to these meaning units. Associated codes were then grouped into lower level concepts. In a final step, the lower level concepts were summarized into higher level concepts. Based on the biopsychosocial model proposed in the International Classification of Functioning, Disability and Health (ICF) and used as a frame of reference in this study [25], the concepts that emerged from the qualitative analysis were used to derive proposals for strengthening resilience that might lead to good individual and community health outcomes [26].

Descriptive statistics were calculated to summarize the characteristics of participants by using R (www.r-project.org). Metric variables were tested for normal distribution. In the case of non-normally distributed data, we depicted medians and interquartile ranges (IQR) in addition to mean values and standard deviations (SD).

Rigour and accuracy of the study

Several strategies were used to enhance the trustworthiness of the qualitative data [27]. Data triangulation was achieved by interviewing persons of different ages, various diseases and disabilities, rural and urban areas and comparing findings to scientific literature and policy documents throughout the project duration [28]. Reflective memos and debriefing notes were recorded after each interview. Moreover, the first four interviews were independently coded by another experienced investigator (VR). After analyzing all interviews, the results were reviewed and discussed with all researchers and patient research partners (GS, EL), who were not involved in the analysis of the transcripts, until consensus was achieved. Original quotes used for publication were translated from German into English by a bilingual native speaker (MA). Following the translation, data were checked to avoid any translation mistakes by the first author (EM). Finally, we reported the results according to the consolidated criteria for reporting qualitative research (COREQ) checklist ([29]; supplemental table A).

Results

Descriptive characteristics of the participants and the data

Thematic saturation (supplemental table B) was reached after including 33 individuals with a mean age ± standard deviation (SD) of 73.7 (±10.9) years.
Table 2  Characteristics of participants

| Characteristics                                  | Total          |
|-------------------------------------------------|----------------|
| **n (%)**                                        | **33 (100)**   |
| **Women**                                        | **25 (75.8)**  |
| **Ø Age (years ±SD) participants**               | **73. (±10.9)**|
| **Median (range of age in years; IQR)**          | **76 (46–92;12)**|
| **Personal living situation n (%)**              |                |
| Living alone                                     | **7 (21.2)**   |
| Living with others                               | **18 (54.5)**  |
| Living in a care facility                        | **8 (24.2)**   |
| **Diagnosed with the following health condition(s) n (%)** |            |
| Diseases of the cardiovascular system            | **13 (39.4)**  |
| Diseases of the digestive system                 | **1 (3.0)**    |
| Endocrine, nutritional, and metabolic diseases   | **6 (18.2)**   |
| Diseases of the eyes, ears and related structures| **4 (12.1)**   |
| Diseases of the musculoskeletal system and connective tissue | **15 (45.5)** |
| Diseases of the nervous system                   | **6 (18.2)**   |
| Diseases of the respiratory system               | **7 (21.2)**   |
| Diseases of the urogenital system                | **1 (3.0)**    |
| Malignancies                                     | **6 (18.2)**   |
| None                                            | **2 (6.1)**    |
| **Employment status n (%)**                      |                |
| Full-time (38.5 h or more per week)              | **1 (3.0)**    |
| Part-time (less than 38.5 h per week)            | **2 (6.1)**    |
| Unemployed                                       | **1 (3.0)**    |
| Retired                                          | **29 (87.9)**  |
| **N (%) total number of participants, n (%) number of participants, Ø Age (±SD) mean age (standard deviation), IQR interquartile range** | |

Two telephone interviews with each participant were conducted between 8 April and 15 May 2020. The typical participant was female (75%), had a cardiovascular (39%) and/or musculoskeletal (45%) disease, and was retired (88%), see Table 2. Key characteristics of each participant are depicted in supplemental table C. In total, 27 h and 30 min of interview time were collected (mean duration was 25 min), resulting in 220 pages of transcript. We extracted 44 lower level concepts and summarized them into six higher level concepts (Table 3).

A general positive attitude

The participants expressed a general positive attitude towards measures that have been taken to mitigate the spread of COVID-19. Some people felt particularly well protected under strict home isolation and appreciated positive aspects of the crisis that helped them better cope with this exceptional situation. They had been offered support from their families, carers, neighbours, and other people they had not known before. Several participants even experienced increased connectedness and cohesion in times of the pandemic, like a 73-year-old woman (No 9) who said:

And still, I feel better now, because I have much more contact with people, even if it is only by phone. They ask me how I am, what I'm doing. It's good for my well-being.

Moreover, some people found more time to relax, tried out new activities during home isolation, and supported others to better deal with this exceptional situation. In the absence of regular treatment and therapy, several study participants (8,24%) reported increased self-management regarding their health.

Challenges of being isolated from the community

Strict home or community isolation was described as an experience of being “captured and imprisoned”. A 90-year-old man (No 19) living in a geriatric facility experienced this situation as follows:

We all yearn to sit at a table at noon and talk. To be locked up in a room and eating alone is kind of torture.

A woman (No 11) at the age of 79 years who lived independently in an assisted-living facility described a situation in which she felt at the mercy of others as follows:

They lock us up in here and won't let us out. Security is down there around the clock. We have to stay in our apartments and can't have any contact with the outside world, except for by phone. That's no kind of life!

Deterioration of health status

Almost one third of the participants (10, 30.3%) reported a significant deterioration in their health status caused by reduced routine care and therapy during the lockdown. In their view, health care changed dramatically from one day to the next, and there were only a few telehealth offers. A 64-year-old woman (No 5) with a physical disability said:

I notice that the longer this goes on, the more difficult it becomes, physically and mentally. Especially physically. My body doesn't work that way anymore. I stumble through the cramps more often, although I really do my exercises umpteen times a day at home and I am really active, but I can't compensate for this alone, I lack the therapy.

People with already existing health problems lacked medical care, treatment, and supply of assistive devices and regular personal contact with their physicians and therapists during this period; however, some people cancelled therapeutic interventions and home care services themselves out of fear of infection with SARS-CoV-2, which led to an enormous additional burden in everyday life. One woman (No 28) at the age of 81 years, suffering from multiple health
Table 3  Overview of 6 higher level and 44 lower level concepts including original quotes from the study participants

| Higher level concepts | Lower level concepts | Example quotes (including sex and age) |
|-----------------------|----------------------|----------------------------------------|
| 1. A general positive attitude | Being a person who can be well alone | I don’t dare to say it out loud, but this crisis has now put me at peace. I can have breakfast with my family again. It’s nice to sit at the breakfast table and discuss all kinds of things, and then everyone does something on their own again, but we’re actually happy. (No 26, female, 66) |
| | Being able to relax at last | Being offered support | I really liked the way the government acted at the beginning of the crisis. It was clear, and one felt that they were saying the same thing and wanted to achieve the same thing. This was great! (No 4, female, 62) |
| | Coping well with the current situation | Feeling safe and protected | Very calm and relaxed . . . we have a lot to tell, we have the garden, and sometimes we do nothing at all and just watch the birds or whatever. |
| | Experiencing increased connectedness and cohesion | | Very comfortable. (No 33, female, 71) |
| | Increasing self-responsibility regarding one’s own health | Ongoing communication in various ways | We experienced an excellent neighbourhood. To this day, they do the shopping for us and do smaller things, and we get homemade bread, very touching how we are taken care of. (No 16, female, 78) |
| | | Opportunity for new activities | You’re alone all the time, and suddenly there’s a knocking, and they [caregivers] are standing out there asking if you want a coffee . . . and that’s soothing. Very soothing. (No 9, female, 73, living in a geriatric facility) |
| | | Supporting others | |
| 2. Challenges of being isolated from the community | Being distressed | At the beginning of the corona crisis, I ordered a toilet seat, and it was so difficult with the company. Finally, they sent the wrong one, and I haven’t been able to exchange this toilet seat yet because I’m afraid to go there and get infected. I haven’t been able to reach anybody either. The phone did not work. I called them a dozen times, and no one answered. (No 16, female, 78) |
| | Changes in the living environment | | What I was afraid of was that I might infect my husband. That was and still is a great fear, I must say. (No, 31, female, 70) |
| | Criticism of the behavior of others | | My children and my husband have been very strict about following government guidelines or measures to not endanger me in any way as a risk patient. And that just creates a certain amount of tension. (No 23, female, 46) |
| | Dealing with risk of infection | Increased conflict potential | My husband worked from home; I worked from home, the big ones did their studies over and over again—I just wanted to avoid any risk of contagion. (No 5, female, 64) |
| | Desire for easing COVID-19 measures | Lack of basic digital literacy skills as an additional obstacle | |
| | Increased conflict potential | Loss of autonomy | |
| | Lack of basic digital literacy skills as an additional obstacle | Necessary changes in future plans | |
| | Loss of autonomy | Regular leisure activities no longer possible | |
| | Necessary changes in future plans | Worries about a second wave | |
| 3. Deterioration of health status | Cancellation of health care services | Not having therapy was the hardest thing for me during this time. I had to take pills because I couldn’t stand the pain anymore. And I usually take pills very, very rarely. (No 5, female, 64) |
| | Increased health problems due to lack of treatment and therapy | | |
| | Reduced availability of doctors and therapists | | |
| 4. Adaptations to improve the implementation of COVID-19 measures | Difficulties to follow pandemic measures due to disability | It’s been 2 or almost 3 weeks now since. I’ve tried to organize masks for myself that I can get along with. And it is not easy when you live alone with a disability and should have no contact at all! I have still found no suitable masks or adaptations. It is so difficult to put the “normal” mask on. It takes me 10 minutes or more. As a healthy person, you can’t even imagine that. I can hardly stand it anymore, physically and mentally. (No 5, female, 64) |
| | Facing alienation during hospital care | Risk to carers due to long waiting time for test results | |
| | Risk to carers due to long waiting time for test results | | |
| 5. Lack of physical contact | Being concerned about others | This is an assisted living facility, but I feel like I’m living alone in the house. No life. Nothing at all. People pull away. You just don’t meet anybody. On Saturday I saw some people, and it was the first time after 6 weeks, so I stopped and watched them, mainly because they were talking and I hadn’t heard that for a long time. I just stood there and was happy about the kids playing, laughing and being happy. (No 9, female, 73) |
| | Implementation of social/physical distancing measures | Importance of being close to each other | |
| | Importance of being close to each other | Increasing loneliness and depression | |
| | Increasing loneliness and depression | People have been abandoned | So if I had gotten sick, who would have taken care of my husband? I honestly don’t know what to do. I don’t want my husband in the hospital either. That’s a difficult thing. That’s why I’m so overprotective. (No 16, female, 78) |
| | People have been abandoned | Restrictions despite palliative care | |
| | Restrictions despite palliative care | Use of protective measures like masks | |
| | Use of protective measures like masks | | |

Example quotes (including sex and age)
Table 3 (Continued)

| Higher level concepts | Lower level concepts | Example quotes (including sex and age) |
|-----------------------|----------------------|----------------------------------------|
| 6                     | 44                   | —                                      |
| 6. Lack of information versus overload | Feeling fooled by the government/loss of credibility | Despite the ban, I still had my son brought to us every Monday by the transport service. It was important for me because the therapies did not take place, my sister did not visit him, and he has no one else. But he needs someone to talk to, someone to understand him. He said I was breaking the law. Later, the government said that this was not against the law at all. That was not right! (No 28, female, 81) |
|                        | Information status and channels used | At the end of the day, when the measures were eased, it was said: “You could have always met the family in private spaces; there would have been no objections.” I didn’t like that, because I simply want to be treated as an adult and not like a fool. Their credibility suffers, and I think that is a pity. (Nr 4, female, 62) |
|                        | Measures affecting working conditions | It is such a strange situation, on the one hand, you are constantly told how to behave, and on the other hand, you realise that it doesn’t work that way in everyday life, so we don’t work that way, our life doesn’t work that way. (No 23, female, 46) |
|                        | Need to justify own decisions regarding COVID-19 measures to others | |
|                        | Perspective regarding government work | |
|                        | Reasons for accepting the measures | |
|                        | Relief through the easing of measures | |
|                        | Trust in information provided | |
|                        | Weighing up the amount of news so as not to be misinformed or overwhelmed, bored or frightened | |
|                        | Worry about the economic situation | |

conditions herself, described the reasons for cancelling her husband’s care:

_We were afraid; the carers have children or go to people who are even worse off than my husband. […] Now I just have to find a way to wash him._

Adaptations to improve the implementation of COVID-19 measures

Some people reported that particular adaptations of COVID-19 measures would have made it easier to implement them. More than half of the participants (18, 55%) in this study had problems following basic COVID-19 protection measures due to their functional limitations in daily life. They could not wear masks because of physical impairment, had panic due to past traumatic experiences when wearing masks or experienced breathing difficulties. Moreover, participants described barriers to implementing basic hygiene measures, such as disinfection stands which are often not accessible for people using wheelchairs or persons of small stature. Like a woman (No 5) aged 64 years with restrictions in her mobility, some people also experienced difficulties in enacting physical distancing because of additional support needs.

_ I need some personal assistance while walking because I feel much safer then, but this is not allowed. So, my only option is the wheelchair or not playing by the rules._

Others also experienced difficulties in communication due to the use of face masks and Plexiglas barriers. One female participant (No 8), who had to stay in hospital for a few days, talked about washbasins that she could not reach:

_ I am 106cm tall and have difficulty reaching the sink in the hospital. And then they always tell you to wash your hands. I asked if they could please bring me a bottle of disinfectant, but they said, “No, we can’t do that—no distribution to patients.”_

Lack of physical contact

Although most participants in this study (25, 76%) have stayed connected and maintained their social networks in a non-face-to-face way, they still suffered from the ban on meeting others in person. They emphasized the importance of personal contact and closeness. From their perspective, prohibiting physical contact and closeness led to social isolation and deprivation, which constituted a massive threat to people’s physical and mental health. They reported increased confusion, malnutrition, sadness, increasing desperation and lack of physical activity among relatives and friends they were no longer allowed to visit. They felt that especially older adults, people with cognitive impairment, hearing problems, and those who were seriously ill or dying, have been abandoned during the first weeks of the pandemic, and nobody cared. A woman (No 3), 65 years old, gave the following example:

_… and then people like my aunt at 103 years die due to isolation or loneliness, or depression because they are so alone. This just can’t go on! Something must be done quickly._

Lack of information versus overload

People reported that they repeatedly had to weigh up how much information they needed not to be misinformed or overloaded or become highly concerned about their own health or bored. A 75-year-old man (No 29) expressed his experience in the following statement:
Table 4  Higher level concepts derived from the qualitative analysis, mapped ICF categories [33], and corresponding environmental adaptations and personal support needs

| Higher level concepts to improve the implementation of COVID-19 measures | Mapped ICF categories | Environmental adaptations and personal support needs from the perspective of the participants |
|---|---|---|
| A general positive attitude | d230 Carrying out daily routine d240 Handling stress and other psychological demands d2402 Handling crisis d350 Conversation d360 Using communication devices and techniques e310 Immediate family e320 Friends e325 Acquaintances, peers, colleagues, neighbours, and community members e355 Health professionals | Promotion of healthy behavior and education in self-management and the use of digital technology for communication Educate people on how best to stay physically active; connect with others by using new technologies such as telephone consultations, text messaging and video conferencing; and manage the stress of daily life through the challenging times of a crisis |
| Challenges of being isolated from the community | d400 Moving around in different locations d470 Using transportation d485 Acquiring, keeping and terminating a job d910 Community life d920 Recreation and leisure d930 Religion and spirituality d940 Human rights | Information to avoid feelings of helplessness and outside control |
| Deterioration of health status | b280 Pain b710 Mobility of joint functions b735 Muscle tone functions b770 Gait pattern functions s 750 Structure of lower extremity e580 Health services, systems and policies | Delivery of routine health service at any time Undertake immediate action to improve care and support to prevent any deterioration in health status resulting from isolation measures for people at risk (such as chronically ill, disabled, old, frail, or cognitively impaired). Therefore, routine health service delivery may need to be adapted to be reliably continued for people at higher risk of serious illness from COVID-19, even during a future crisis |
| Adapations to improve the implementation of COVID-19 measures | e1101 Drugs e1151 Assistive products and technology for personal use in daily living | Identification and elimination of obstacles and barriers to best follow the infection control measures Implement points of contact to best support people and address their specific needs to maintain their health, safety, and independence in the community throughout the COVID-19 outbreak and related health emergencies |
| Lack of physical contact | d710 Basic interpersonal interactions d720 Complex interpersonal interactions | Assurance of personal contact if needed Raise awareness that especially older people and people with underlying medical conditions should never be left alone for long periods. This might have negative consequences for their health and well-being. The involvement of family members and other support networks should be allowed even in times of crisis with care and consideration; without preventing an encounter per se |
| Lack of information versus overload | d310 Communicating with—receiving—spoken messages d320 Communicating with—receiving—formal sign language messages d325 Communicating with—receiving—written messages | Incorporation of people’s perspective in decision-making Involve representatives of people at increased risk of severe illness from COVID-19 in expert groups to identify context-relevant essential information during an outbreak |

…”it gets on my nerves, this constant information about things I already know. I think there are already more experts than corona patients. So I’m switching off immediately.

Some participants felt that the information they received was not sufficiently concise and relevant to their underlying conditions. This was perceived as a potential risk to their personal health. For example, a woman (No 23) at the age of 46 years with severe lung disease, the information provided by the government did not meet her needs.

I think this information about the course of the disease was somehow minimal. I would like to have a more informed discussion of the facts. Instead, we were treated as minors.

Only one third of the participants (11 out of 33) were concerned about possible economic consequences that might affect them personally.

Longitudinal perspective

Almost all participants welcomed the easing of measures that took place between the first and subsequent telephone interviews. Over time, however, they experienced (i) an increasing concern about misconduct by others regarding compliance with infection control measures, (ii) increased levels of loneliness and depression, (iii) a feeling of being overwhelmed or bored by the amount of information provided about the pandemic and the measures taken to mitigate its effects and (iv) difficulties in adhering to specific measures due to their functional limitations/disabilities when they started to leave home again.

ICF mapping and environmental adaptations from the perspective of the participants

The higher level concepts were mapped to 29 ICF categories in total; 4 (14%) related to body functions and
Discussion

The present study comprehensively explored the experiences and needs of people at increased risk of severe illness from COVID-19 in response to the early phases of the pandemic. The actions that were implemented during the lockdown had significant positive and negative impacts on the functioning and health of the selected population. Although the initial infection control measures taken by the government were generally accepted and valued by older adults and people with underlying chronic health conditions, individuals also expressed challenges in daily life and potential negative consequences of the lockdown that may have been given too little consideration. These collateral damages to the population at risk have not been well understood thus far. Studies have mainly focused on health care providers [30–32], often not considering the needs of people at increased risk of severe illness from COVID-19.

Some participants in our study reported additional limitations in their functioning in daily life caused by the infection control measures [33]. They experienced reduced access to health care services, such as hospitals and outpatient clinics, and the suspension of many health care interventions. Moreover, some of them would have needed specific exemptions to follow the infection control measures properly. Especially in high strain on the health care system, it is crucial to ensure that all population groups receive good care. Physical and mental health are essential for resilience throughout the COVID-19 outbreak and related health emergencies [34]. Health in all policies requires equity and includes strategies to achieve the best health outcomes and a sustainable health care system for everyone in our society [35]. Participants in this study were convinced that incorporating their perspectives into the design of health care in the context of the pandemic would make an important contribution to societal health ensuring adequate compliance with the specific measures taken.

People also pointed out that constant social contact with people with cognitive, mental, psychosocial and/or physical disabilities or chronic health conditions is urgently needed to prevent a deterioration in health and promote well-being despite times of crisis. In their work, Cacioppo et al. pointed out that social isolation impairs older people's mental and cognitive health and is also accompanied by a significantly higher risk of mortality [36], underpinned by a meta-analytic review done by Holt-Lunstad et al. [37]. Using existing support networks with care and considering the risk of infection might help overcome the disadvantage of this vulnerable group even in times of a pandemic.

Interestingly, and in contrast to population-based surveys on the population's health status during pandemics in other countries [38], participants in this study were less concerned about possible financial uncertainties caused by the crisis; however, most of them were retired and trusted the existing social and health system, which might be different for people living in countries with a less established or even not existing safety net.

In their work, Teti et al. concluded that the needs and preferences of people at risk must be studied in depth to plan health care efficiently [8]. With this study we provided comprehensive longitudinal data but remotely collected, which might be regarded as a limitation of our work. Nevertheless, our findings highlight the need for further research to understand the role of resilience in managing the extraordinary challenges due to COVID-19 among people at risk of severe illness and to explore interesting changes in the experience of the applied measures over time in more depth. The evidence provided will support the formulation of preparedness and response strategies targeted and based on the preferences of this vulnerable group of individuals. Targeted strategies and interventions such as maintaining access to health care providers, facilitating the use of digital technology to overcome social isolation, and adapting the environment are needed to increase compliance with risk reduction measures, prevent collateral health deterioration in this group, and ensure well-being even during prolonged periods of crisis.

Conclusion

Participants in this study have identified several opportunities to promote and protect the health and well-being of people at higher risk of serious illness from COVID-19 and reduce health inequity. The study results might guide policy and health care providers to minimize direct and indirect harm and effectively
support people living under demanding conditions in such a pandemic. In addition, the knowledge gained could be used to revise outbreak response activities and develop action plans and interventions for future waves of infection or pandemics to promote best and protect population health and well-being in different population groups.

Acknowledgements We want to thank all people who participated in this study to share their valuable perspectives and Emilie Mosor for transcribing all the interviews.

Author Contribution Erika Mosor, Valentin Ritschl, Margaret R. Andrews, Maisa Omara, Paul Studenic, Gertrude Schaffer, Ernst Leitgeb, Claudia Oppenauer, Linda C. Li and Tanja Stamm were responsible for the study concept and design. Erika Mosor conducted the interviews. Erika Mosor and Valentins Ritschl verified the underlying data, extracted and analyzed them. Erika Mosor, Valentin Ritschl, Margaret R. Andrews, Maisa Omara, Paul Studenic, Gertrude Schaffer, Ernst Leitgeb, Claudia Oppenauer, Linda C. Li and Tanja Stamm were involved in interpreting the data, writing and reviewing the manuscript. Erika Mosor and Valentin Ritschl were responsible for the tables. Gertrude Schaffer and Ernst Leitgeb are patient representatives/advocates.

Funding Open access funding provided by Medical University of Vienna.

Declarations

Conflict of interest E. Mosor, V. Ritschl, M.R. Andrews, M. Omara, P. Studenic, G. Schaffer, E. Leitgeb, C. Oppenauer, L.C. Li and T. Stamm declare that they have no competing interests. There was no financial support from any funding organization for the submitted work.

Ethical standards All procedures performed in studies involving human participants or on human tissue were in accordance with the ethical standards of the institutional and/or national research committee (Ethics Committee of the Medical University of Vienna (EK Number 1388/2020)) and with the 1975 Helsinki declaration and its later amendments or comparable ethical standards. Informed consent was obtained from all patients included in the study.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

1. European Centre for Disease Prevention and Control. Outbreak of novel coronavirus disease 2019. 2020. https://www.ecdc.europa.eu/sites/default/files/documents/RRA-sixth-update-Outbreak-of-novel-coronavirus-disease-2019-COVID-19.pdf. Accessed 15 June 2020.
2. Centers for Disease Control and Prevention. Evidence used to update the list of underlying medical conditions that increase a person’s risk of severe illness from COVID-19. 2020. https://www.cdc.gov/coronavirus/2019-ncov/need-extra-precautions/evidence-table.html. Accessed 25 July 2020.
3. Bedford J, et al. COVID-19: towards controlling of a pandemic. Lancet. 2020;395(10229):1015–8.
4. Huang Y, Zhao N. Generalized anxiety disorder, depressive symptoms and sleep quality during COVID-19 outbreak in China: a web-based cross-sectional survey. Psychiatry Res. 2020;288:112954.
5. World Health Organisation. Disability considerations during the COVID-19 outbreak. 2020. https://www.who.int/publications/i/item/WHO-2019-nCoV-Disability-2020-1. Accessed 24 June 2020.
6. Brooks SK, et al. The psychological impact of quarantine and how to reduce it: rapid review of the evidence. Lancet. 2020;395(10227):912–20.
7. World Health Organization. COVID-19: operational guidance for maintaining essential health services during an outbreak: interim guidance, 25 March 2020. Geneva: World Health Organization; 2020.
8. Tiet M, Schatz E, Liebenberg L. Methods in the Time of COVID-19: The Vital Role of Qualitative Inquiries. Los Angeles: SAGE; 2020.
9. Curry LA, Nembhard IM, Bradley EH. Qualitative and mixed methods provide unique contributions to outcomes research. Circulation. 2009;119(10):1442–52.
10. Pollak M, Kowarz N, Partheymüller J. Chronology of the corona crisis in Austria—part 1: background, the way to the lockdown, the acute phase and economic consequences. 2020. https://viecer.univie.ac.at/en/projects-and-cooperations/austrian-corona-panel-project/corona-blog/corona-blog-beitraege/blog51/. Accessed 30 Aug 2020.
11. Guetterman TC. Descriptions of sampling practices within five approaches to qualitative research in education and the health sciences. Forum Qual Soc Res. 2015;16(2), Art. 25.
12. Stamm TA, et al. Concepts of functioning and health important to people with systemic sclerosis: a qualitative study in four European countries. Ann Rheum Dis. 2011;70(6):1074–9.
13. Moser E, et al. I would never take preventive medication! perspectives and information needs of people who underwent predictive tests for rheumatoid arthritis. Arthritis Care Res. 2020;72(3):360–8.
14. Francis JJ, et al. What is an adequate sample size? Operationalising data saturation for theory-based interview studies. Psychol Health. 2010;25(10):1229–45.
15. Patton MQ. Qualitative research and evaluation methods. Thousand Oaks: SAGE; 2002.
16. Palinkas LA, et al. Purposeful sampling for qualitative data collection and analysis in mixed method implementation research. Admin Policy Ment Health. 2015;42(5):533–44.
17. Vasileiou K, et al. Characterising and justifying sample size went predictive tests for rheumatoid arthritis. Arthritis Care Res. 2020;72(3):360–8.
19. Pope C, Ziebland S, Mays N. Qualitative research in health care. Analysing qualitative data. BMJ. 2000;320(7227):114–6.
20. Guest G, Bunce A, Johnson L. How many interviews are enough? An experiment with data saturation and variability. Field Methods. 2006;18(1):59–82.
21. Saunders B, et al. Saturation in qualitative research: exploring its conceptualization and operationalization. Qual Quant. 2018;52(4):1893–907.
22. World Medical Association. World medical association declaration of Helsinki: ethical principles for medical research involving human subjects. JAMA. 2013;310(20):2191–4.
23. Kvale S. Interviews: An introduction to qualitative research interviewing. Thousand Oaks: SAGE; 1996.
24. ATLAS.ti, Atlas.ti [8.0]. 2018, Atlas.ti Scientific Software Development GmbH: Berlin.
25. Organization WH. International classification of functioning, disability and health: ICF. Geneva: World Health Organization; 2001.
26. Ziglio E. Strengthening resilience: a priority shared by health 2020 and the sustainable development goals. Copenhagen: WHO Regional Office for Europe; 2017.
27. Mays N, Pope C. Qualitative research in health care. Assessing quality in qualitative research. BMJ. 2000;320(7226):50–2.
28. Denzin NK, Lincoln YS. The Sage handbook of qualitative research. Thousand Oaks: SAGE; 2011.
29. Tong A, Sainsbury P, Craig J. Consolidated criteria for reporting qualitative research (COREQ): a 32-item checklist for interviews and focus groups. Int J Qual Health Care. 2007;19(6):349–57.
30. Sun N, et al. A qualitative study on the psychological experience of caregivers of COVID-19 patients. Am J Infect Control. 2020;48(6):592–6.
31. Greenberg N, et al. Managing mental health challenges faced by healthcare workers during covid-19 pandemic. BMJ. 2020;368:m1211.
32. Chen Q, et al. Mental health care for medical staff in China during the COVID-19 outbreak. Lancet Psychiatry. 2020;7(4):e15–e6.
33. WHO. ICF—international classification of functioning, disability and health. 2005. http://www.dimdi.de/static/de/klassi/icf/. Accessed 10 Jan 2016.
34. United Nations. COVID-19 outbreak and persons with disabilities. 2020. https://www.un.org/development/desa/disabilities/covid-19.html. Accessed 8 June 2020.
35. de Wit M, et al. Practical guidance for engaging patients in health research, treatment guidelines and regulatory processes: results of an expert group meeting organized by the World Health Organization (WHO) and the European Society for Clinical and Economic Aspects of Osteoporosis, Osteoarthritis and Musculoskeletal Diseases (ESCEO). Aging Clin Exp Res. 2019;31(7):905–15.
36. Cacioppo JT, et al. Loneliness as a specific risk factor for depressive symptoms: cross-sectional and longitudinal analyses. Psychol Aging. 2006;21(1):140.
37. Holt-Lunstad J, Smith TB, Layton JB. Social relationships and mortality risk: a meta-analytic review. PLoS Med. 2010;7(7):e1000316.
38. Wozniak A, et al. COVID impact survey. 2020. https://www.covid-impact.org. Accessed 27 July 2020.

Publisher’s Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.