COVID-19 wellbeing study: a protocol examining perceived coercion and psychological well-being during the COVID-19 pandemic by means of an online survey, asynchronous virtual focus groups and individual interviews

Veronica Ranieri, Andrea Sem Stoltenberg, Elena Pizzo, Chiara Montaldo, Emanuele Bazzi, Sarah Edwards, Sunjeev Kamboj

ABSTRACT

Introduction The COVID-19 pandemic has resulted in many countries applying restrictive measures, such as lockdown, to contain and prevent further spread. The psychological impact of lockdown and working as a healthcare worker on the frontline has been chronicled in studies pertaining to previous infectious disease pandemics that have reported the presence of depressive symptoms, anxiety, insomnia, and post-traumatic stress symptoms. Potentially linked to psychological well-being and not yet studied is the possibility that lockdown and working on the frontline of the pandemic are associated with perceptions of coercion.

Methods and analysis The present study aimed to examine perceived coercion in those who have experienced COVID-19-related lockdown and/or worked as a frontline healthcare worker across three European countries. It aimed to describe how such perceptions may impact on psychological well-being, coping and post-traumatic growth. It will employ an explanatory mixed-methods research methodology consisting of an online survey and online asynchronous virtual focus groups (AVFGs) and individual interviews. χ² tests and analyses of variance will be used to examine whether participants from different countries differ according to demographic factors, whether there are differences between cohorts on perceived coercion, depression, anxiety and post-traumatic growth scores. The relationship between coercion and symptoms of distress will be assessed using multiple regression. Both the AVFGs and the narrative interviews will be analysed using thematic narrative analysis.

INTRODUCTION

COVID-19 is an infectious disease caused by a novel betacoronavirus believed to originate in Wuhan, China. Early indications of the presence of this virus emerged in December 2019, when several individuals displayed clinical presentations akin to viral pneumonia. Although our knowledge of the virus is still accumulating, severe cases of infection may lead to serious health complications and death. As of 11 March 2020, the WHO declared the disease a pandemic. In addition to this declaration, individuals and governments were advised to take precautionary and restrictive measures to reduce virus transmission, such as social distancing, self-isolation, quarantine and lockdown, depending on prevalence and health service capacity within each country.

The psychological impact of restrictive measures and epidemics has received some attention in the mental health literature, with studies reporting the presence of depressive symptoms, anxiety and insomnia in those who...
have experienced quarantine and those who have worked as healthcare workers during epidemics.\textsuperscript{5–13} Such distress, as well as its possible impact on suicidality, is expected as a result of restrictive practices linked to the COVID-19 pandemic, with researchers and clinicians worldwide commenting on the effect of secondary stressors associated with both the lockdown and the threat of infection on the general population’s mental health.\textsuperscript{14–19} In healthcare workers, it has been linked to a number of COVID-related work stressors, including increased workloads, unfamiliar tasks, being at increased risk of infection and fears of infecting others.\textsuperscript{12,20–23} Linked to psychological well-being and not yet studied is the potential for these experiences to give rise to perceptions of coercion, both in those living under lockdown and those providing frontline healthcare. Such perceptions arise from experiencing a dearth of choice, freedom, influence or control with regard to one’s situation.\textsuperscript{24,25} These are most commonly reported when individuals experience a situation that they feel is forced on them without justification, and where they feel excluded from the decision-making process or do not have an opportunity to express their viewpoint.\textsuperscript{26} Such perceptions have been commonly reported in the mental health literature, particularly in relation to restrictive measures such as involuntary admissions to hospital, which, in ways, may be comparable in populations where restrictions were severe and individuals were legally enforced to stay at home.\textsuperscript{27} In mental health settings, higher levels of perceived coercion are indicative of a poorer prognosis.\textsuperscript{28} It is, however, unclear as to whether a pandemic-related lockdown will give rise to similar perceptions of coercion. Understanding such perceptions is critical if governments are to secure the cooperation of their citizens in enacting mitigation (lockdown) efforts, and the cooperation and mental well-being of frontline workers, in future scenarios.

In light of the aforementioned research, the present study primarily examines perceived coercion in those who have experienced COVID-19-related lockdown and/or frontline roles across three European countries with different demographic characteristics and healthcare systems, and which enacted differentially stringent mitigation approaches. In healthcare workers, it examines how perceptions of coercion and pressures may be related to perceived risk of infection and COVID-19-related stressors in the workplace. Using the transactional theory of stress and coping, it aimed to describe how such perceptions may impact on psychological well-being, coping and post-traumatic growth,\textsuperscript{29} and, in healthcare workers, on professional quality of life. Finally, we will also preliminarily investigate whether there is an increase in healthcare resource usage (in the general population sample), as a coping strategy, and whether this is linked to the aforementioned constructs.

**Key research questions**

Our research questions are the following:

1. What are the experiences of those who have been under lockdown or have been working as healthcare workers during the COVID-19 pandemic? How have these experiences impacted on their psychological well-being?
2. To what extent does the general population perceive the lockdown as coercive, pressured and procedurally just? To what extent do healthcare workers perceive having to work on the frontline with patients with COVID-19 as coercive? Do these perceptions change over time?
3. Are perceptions of coercion associated with psychological distress, after controlling for demographic and background factors?
4. Do perceptions of coercion and psychological distress in the general population and among key workers vary across affected countries after controlling for demographic and background factors?
5. What practical recommendations can we highlight to policy-makers and healthcare management about how to improve the psychological support provided to both the general population and healthcare workers across affected countries?

**METHODS AND ANALYSIS**

This study will employ an explanatory mixed-methods research methodology consisting of (1) an online survey and (2) online asynchronous virtual focus groups (AVFGs) or individual interviews.

**Sample**

**Online survey**

Participants will consist of individuals who have experienced governmental lockdown and/or who are key workers during the coronavirus pandemic. Participants will be recruited from the UK, Italy and Norway with the potential to be extended to other countries. Individuals aged ≥18 years and who have experienced governmental lockdown and/or are key workers working on the frontline in the UK, Italy or Norway will be invited to participate. We aimed to recruit 2000 individuals who experienced lockdown per country, as per Martínez-Mesa et al and Maxwell.\textsuperscript{30,31} Participants will be recruited using social media platforms such as Twitter, Facebook and Instagram. Advertisements for the study will also be cascaded via email and other social media messaging technology such as WhatsApp.

**AVFGs and individual interviews**

A sample of individuals from the quantitative study will be invited to participate in either an AVFG or individual interview, depending on preference and availability. A total of six to nine AVFGs will be conducted simultaneously, with two to three AVFGs/country. AVFGs will be limited to 6–10 participants/group. Purposive sampling will be used to select individuals according to their age, gender and geographical location, or other distinctive
factors where possible, so that each group consists of participants from a diverse range of backgrounds.

**Setting**

The study will be completed entirely online using a general data protection regulation (GDPR)-compliant data collection tool (ISO 27001 certified). AVFG participants will be asked to log onto a virtual learning environment\(^{22-33}\) that provides a secure and confidential space for research participants to write and interact with each other. Individual interviews will take place by telephone or using online conferencing software.

**Procedure**

**Online survey**

Advertisements will be posted and shared on social media. Clicking on the study’s link will direct participants to the survey’s home page from which they can be directed to the study information page. Should they wish to participate after reading the study information, they will provide informed consent online and proceed to the online survey. After completing the survey, participants will be invited to take part in a shorter follow-up survey within a period of 3 months and/or further qualitative research. Those who wish to take part will be asked to enter their email address.

**AVFGs or individual interviews**

AVFG participants will be registered onto a virtual learning environment\(^{22,33}\) under a chosen alias (to preserve anonymity). Then they will familiarise themselves with the platform and documents listed as ‘essential information’, which include study information, addition consent form, researcher contact details and support, and netiquette guidelines. Once all participants provide consent, the first focus group question will appear as a discussion topic. Participants will be asked to engage in discussion about a different question (outlined as follows) each week for 3 weeks. Participants will be able to post as often as they wish each week, both in response to the question and in reaction to other participants’ responses. Participants will receive an email notification for each new question posted. Discussion boards will be moderated two times per day by the researchers to monitor the content of what is posted and to delete if offensive, and to probe and clarify participants’ responses.

Should participants prefer to take part in an interview rather than an AVFG, the researcher will email them a copy of the information sheet and consent form and arrange a time for interview. All interviews will be audio-recorded and then transcribed. The interviews are expected to last up to 1 hour. All identifiable data will be stored on Data Safe Haven, in accordance with University College London (UCL) policy.

**Instruments**

**Online survey**

All participants will complete a questionnaire constructed by the researchers that asks individuals about general demographic details and their clinical background at baseline. Participants will also complete the following measures at baseline and within a 3-month follow-up period: (1) an adapted version of the MacArthur Admission Experience Survey\(^{24}\) to measure the extent to which individuals perceive their circumstances as coerced, pressured and procedurally just in the general population and a Perceived Coercion Scale for Healthcare Workers constructed by researcher AS, (2) the Depression, Anxiety and Stress Scale\(^{35}\) to indicate the presence of depression, anxiety, stress and post-traumatic growth; (3) the Post-traumatic Growth Inventory-Short Form\(^{36}\) to assess whether individuals experience positive outcomes relating to the COVID-19 pandemic; and (4) the brief COPE\(^{37}\) to observe individuals’ methods of coping within the context of the COVID-19 pandemic. Healthcare workers will also be asked to complete an adapted version of the Perceived SARS Related Risk Scale\(^{38}\) and the Professional Quality of Life measure\(^{39}\). The adapted versions of the scales may undergo minor modification after piloting. A copy of the survey can be accessed online (www.ovid19wellbeingstudy.org). A copy of the survey questions is attached as an online supplemental file with this article.

**AVFGs**

Members of the general population taking part in the AVFGs will be asked to answer the following questions and to engage with other participants’ responses:

Week 1: Can you tell us about how you felt when you were first told to stay home during the coronavirus pandemic? Looking back on the lockdown, have your feelings changed towards it? If yes, in what way? Did you agree or disagree with the governmental lockdown? In what way? How do you feel about the lockdown being lifted? What are your feelings regarding a future potential lockdown?

Week 2: Did you self-isolate prior to the lockdown or did you remain home solely because the lockdown came in? For example, did you first experience forced isolation due to confirmation of illness? Did you first practice self-isolation due to you or someone in your household being symptomatic, or having a condition that places you at a greater health risk? Or was your first experience of isolation a result of governmental lockdown? Did you feel you had any control over your isolation? Was it something you chose initially or something you felt was forced on you? How did this make you feel?

Week 3: What has been the impact of the lockdown on your psychological well-being? Have you, someone in your household or someone you know experienced an onset of coronavirus-type symptoms? How did this impact you emotionally? What support do you feel you need for your emotional well-being? What have you done so far to try to stay well?

Healthcare workers will be asked to answer and engage with other participants’ responses for the following questions:
Week 1: Can you tell us about how you felt when you first started working with patients confirmed with or suspected of having COVID-19? Were you asked or told to work with this patient group? Did you volunteer to work with this patient group? Did you feel that working with this patient group is part of your work role, in spite of potential risk? How have your feelings changed about working with this patient group between then and now, if at all? Week 2: Under what circumstances did you work with patients confirmed with or suspected of having COVID-19? Are you redeployed or did you remain in your workplace? Did you feel that you were adequately physically protected? Did you feel adequately supported within your workplace? Did you feel your needs were considered? Did you encounter barriers to accessing protection or support within your workplace? In what way? Week 3: What has been the impact of working as a key worker with individuals with COVID-19 on your psychological well-being? What supports do you feel you need for your emotional well-being? Did you seek psychological support, and from whom? What have you done so far to try to stay psychologically well?

Individual interviews
Semistructured interviews will be adopted to enable the researchers to capture detailed insights about each individual’s personal experiences and perceptions and the context in which these occur. These interviews will focus on individuals’ perceptions regarding lockdown (inclusive of perceived coercion, pressures, and procedural justice), their experiences of isolation and their psychological well-being. Individuals who identify themselves as healthcare workers in the online survey may be invited to interview depending on their availability. These interviews will be slightly tailored to focus on clinicians’ experiences and perceptions of working during the coronavirus pandemic (inclusive of perceived coercion, pressures and procedural justice), and their psychological well-being.

Analysis
Online survey
All numerical data will be inputted into a statistical analysis software programme (SPSS) version 26. The data will be tested for normality by conducting quartile–quartile plots and Kolmogorov–Smirnov tests for each measure (and subscale, where applicable). Following the aforementioned initial management of the data, we will describe demographic and background characteristics according to country and respondent type. A multiple regression model will be used with perceived coercion as the main predictor and measures of psychological distress (ie, depression, anxiety and post-traumatic stress) as the main outcome variables. The secondary outcome variables will include measures of adaptability (ie, coping and post-traumatic growth) and demographic (ie, age, gender, ethnicity and education level) and background variables (ie, job role and work environment) as covariates. In healthcare workers, COVID-19-related stressors and perceptions of COVID-related risk will also be analysed as possible predictors, mediators and/or moderators. Further exploratory analysis may be conducted to assess for interaction between variables. Analyses of variance will be used to examine whether there are differences between UK, Italian and Norwegian cohorts on perceived coercion, stress, depression, anxiety, trauma scores and professional quality of life.

AVFGs and individual interviews
The qualitative substudy will employ a phenomenological approach as its focus is on understanding the subjective experiences of individual participants and the meaning that participants attribute to these experiences. It will also employ an interpretative framework, using the stress-coping paradigm, whereby the focus of our questions and analysis will be on creating a picture of what influences individuals’ appraisals of perceived coercion, pressures and procedural justice and how these appraisals influence and are influenced by coping. Focus group data will be downloaded directly as text from the virtual learning environment platform. Interviews will be audio-recorded and transcribed. Computer Assisted Qualitative Data Analysis Software, such as NVivo, will be used to code the transcripts.40 Text scripts and transcripts will be analysed, adapting Braun and Clarke’s thematic analysis.41 This will involve both sequencing events and experiences, and grouping commonalities and experiences within chronological sequences, and between the two types of participants. Themes and narratives will be explored by two researchers with one principally analysing the data and the second overseeing the emerging themes. Any disagreements will be discussed and resolved within the wider research team. Concepts from the narratives will be both derived inductively from the data and applied deductively to our theoretical framework.

Patient and public involvement
No patient involvement. The background questionnaire was co-constructed with the authors who are healthcare workers. The questionnaire was also piloted with healthcare professionals who gave feedback on it.

ETHICS AND DISSEMINATION
This study received approval by UCL’s Research Ethics Committee as application 7335/004.

Informed consent
Individuals who click on the study’s advertisement on social media will be brought to the survey’s home page. Should they wish to proceed, they will be brought to an information page detailing the purpose of the study, how their confidentiality and anonymity will be preserved and how their data will be treated. Should individuals wish to participate after reading the information page, they will
be asked to provide informed consent online. At the end of the survey, participants will be asked if they would like to take part in a follow-up assessment and/or an online focus group or interview. Those who wish to take part will be asked to enter their email address.

Participants who provide their email address and show a preference to take part in an online group will be registered onto a virtual learning environment (i.e., Blackboard) under an alias chosen by them in order to preserve their anonymity. Once registered, they will be asked to log in and familiarise themselves with both the platform and all documents listed as ‘essential information’ within it. Essential information will include the study’s information sheet and separate consent form, contact details for the researchers and support, and netiquette guidelines. Once all participants provide consent online, the first focus group question will appear as a discussion topic. Should participants prefer to take part in an interview, the researcher will email them a copy of the information sheet and consent form and arrange a time for interview.

Data protection and confidentiality
An online data collection tool that is GDPR compliant and certified to ISO 27001 standard will be used for the online survey. Participants will be given the opportunity to choose their own pseudonym for the AVFGs to help preserve their anonymity. All data collected during the course of the research will be kept strictly confidential. All identifiable information will be stored on UCL’s Data Safe Haven, a GDPR-compliant, encrypted system for the duration of the study. Audio recordings will be kept on Data Safe Haven until these are transcribed into an anonymous format. Non-identifiable data will be stored on a password-protected, encrypted drive on a UCL desktop for a period of 5 years.

Dissemination
Participants will not be identifiable in any ensuing reports or publications. Results will be disseminated by means of peer-reviewed publications and at national and/or international conferences. We will aim to publish the findings in an open-access journal to make the study’s findings accessible to the general population. A summary of the findings will be shared on our website (thecovid19wellbeingstudy.org)

REFERENCES
37 Carver CS. You want to measure coping but your protocol’s too long: consider the brief cope. Int J Behav Med 1997;4:92–100.