Acute impact of a national lockdown during the COVID-19 pandemic on wellbeing outcomes among individuals with chronic pain

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Abstract
Changes to wellbeing in a community-based sample of 638 adults with non-malignant chronic pain were assessed during a period of mandated lockdown measures in the UK to control the COVID-19 outbreak. Participants completed an online survey pre-lockdown and were followed up during lockdown. Multivariate analysis demonstrated that decreased ability to self-manage pain, restricted access to healthcare and increased dependence on others were associated with negative wellbeing outcomes related to sleep, anxiety and depression. Essential but non-urgent services are required during periods of lockdown to maintain independence and self-management in order to preserve wellbeing in this population.

Keywords
anxiety, chronic pain, Covid-19, depression, sleep

Introduction
Throughout 2020, the COVID-19 pandemic prompted governing bodies across the globe to impose combative public health measures to contain the virus and reduce infection rates and mortality (Chakraborty and Maity, 2020). In the United Kingdom (UK), measures included social distancing, closure of schools and public spaces, and a ‘stay at home’ order except in essential circumstances (Jarvis et al., 2020). These restrictions, described as ‘lockdown’, brought considerable disruption to social and economic activities, and reduced access to many essential but non-urgent services (Gupta et al., 2020; Iacobucci, 2020). Research during the pandemic has found evidence of high psychological distress in the general population across different cultures and regions (Liang et al., 2020; Odriozola-González et al., 2020; Xiong et al., 2020). Reasons for this may stem from a combination of factors which initiate and perpetuate psychological distress such as: health-related fear, isolation, confinement, misinformation, changes in daily routine and economic uncertainty (Giordani
et al., 2020; Odriozola-González et al., 2020; Ornell et al., 2020).

**Impact on wellbeing among chronic pain populations**

It is estimated that around one third of the UK population lives with a form of chronic pain and one in four individuals experience poor mental health at any one time (Fayaz et al., 2016b). The high prevalence of chronic pain and poor mental health negatively impacts individuals in terms of their wellbeing, and accounts for large losses of economic productivity at societal level. The biopsychosocial model of health implicates an interaction between biological, psychological, and social factors which affect an individual’s illness perception, pain-management and overall wellbeing (Bevers et al., 2016). Chronic pain populations report higher levels of anxiety, depression, post-traumatic stress disorder and poor sleep quality than the general population, as well as higher comorbidity of physical illnesses (Fayaz et al., 2016a; Gatchel, 2004; Morin et al., 1998). The relationship between sleep and pain has been shown to be bidirectional and particularly impacts this population. Pain may impact sleep by increasing micro-arousals, which over time cause fragmented sleep patterns. Equally, sleep disturbance has been shown to reduce pain thresholds with severe consequences for chronic pain populations who experience pain in episodic or persistent manners (Smith and Haythornthwaite, 2004).

Furthermore, chronic pain populations may be disproportionately impacted by public health restrictions during a pandemic because of reduced access to healthcare, risk of increased disease severity from contracting COVID-19, and closure of social and community support services (Kang et al., 2020; Rhodes et al., 2020). There is clear evidence that self-management strategies play an important role for psychological coping and management of chronic pain (Nicholas and Blyth, 2016). These strategies include adhering to prescribed medication or physical activity regimen, identifying treatments jointly with a health practitioner, in addition to managing the impact on mood and relationships due to pain interference. The ability to adopt self-management strategies has been shown to improve mental health in this population, however, evidence also suggests that socioeconomic status (SES) may influence self-management outcomes as those with low SES may have access to fewer resources than those with high SES (Hardman et al., 2020). Additionally, access to healthcare greatly impacts an individual’s health journey as those with chronic pain rely on a combination of assessments, diagnostics, and interventions, involving frequent interaction with the health system (Reid et al., 2015). Losing the ability to self-manage or restricting healthcare access for prolonged periods may have a significant impact on wellbeing including sleep behaviours and mental health. The pandemic brings risk and burden to chronic pain populations regarding disease management, as well as the potential to impact social and health behaviours. It is therefore important to study this population during periods of lockdown to ensure their wellbeing is not severely affected and mitigate risk in future waves where possible (Arora and Grey, 2020).

The objective of this study was to explore changes in wellbeing outcomes as related to sleep, anxiety and depression within a community sample of adults living with chronic pain between the start of the COVID-19 outbreak, pre-lockdown and during a period of lockdown in the UK. Two aims were identified to pursue the study:

1. To measure changes in wellbeing outcomes (sleep, anxiety and depression) observed between the pre-lockdown period (Time 1) and during lockdown (Time 2).
2. To identify if COVID-19 specific changes related to (a) dependence on others for support, (b) ability to self-manage pain condition and (c) access to healthcare were associated with wellbeing outcomes.
Given current global and national estimates of COVID-19, it is likely physical distancing measures and reduced service provision will continue for some time. Identifying factors which could impact wellbeing in this population may help health and social care services dealing with the pandemic’s response and recovery process.

Methods

Design

A cross-sectional survey was conducted related to a separate protocol pre-lockdown (Time 1: February and March 2020). To evaluate the impact of COVID-19 and the lockdown measures, a subsequent follow-up survey (Time 2: April and May 2020) was released to the same participants whilst lockdown measures were in place. Both surveys were hosted on Qualtrics (www.qualtrics.com). This research was granted ethical approval by UCL Institute of Education’s Ethics Committee. All participants were asked for explicit and voluntary consent at the start of each survey.

Participants

A total of 1,234 adults with existing non-malignant chronic pain, who completed a cross-sectional survey at Time 1, were contacted to complete a follow-up study via online means at Time 2. A sample of 638 participants, living in the UK during the pandemic, responded and were included in this study (a response rate of 52%). During the analysis stage, participants with missing values for independent or dependent variables were excluded from individual analyses.

Participant characteristics are described in Table 1. Eligibility was based on participants being of adult age (18+ years), with an existing chronic pain condition unrelated to cancer. Participants were screened for mental and physical health conditions co-existing with their chronic pain condition. The sample was recruited via social media advertisements as well as online news bulletins and chronic pain charity websites. Participants did not receive compensation for taking part in the study.

Measures

Background measures and independent variables related to the pandemic. At Time 1, background measures collected demographic indicators: age, sex, ethnicity, education status and a brief history in relation to participants’ chronic pain condition.

At Time 2, participants were asked questions relating to COVID-19 and personal circumstances during the lockdown period. These were converted into categorical variables and were used as independent variables during the analyses. Significant independent variables related to the COVID-19 lockdown were:

1. ‘Has your dependence on others such as family/friends/local community changed during the COVID-19 pandemic for practical and emotional support?’
2. ‘Since the lockdown measures were put into place, have you felt any change in your ability to self-manage your pain?’
3. ‘Has your access to healthcare (for non-COVID-19 related care) been affected by the COVID-19 pandemic?’

Wellbeing measures related to anxiety, depression and sleep were collected during Time 1 and Time 2.

Wellbeing measure: Anxiety and depression. The Hospital Anxiety and Depression Scale (HADS) is a 14-item validated measure designed to measure anxiety and depression symptoms during the past week (Zigmond and Snaith, 1983). It comprises of two subscales assessing anxiety (HADS–A) and depression (HADS–D). Items are rated on a four-point Likert scale (e.g. 0 = not at all to 3 = most of time). Five items require reverse scoring. Scores for each item are summed for each subscale, scores above eight suggest anxiety and depression (Zigmond and Snaith, 1994). Cronbach’s alpha
coefficients were found to be 0.83 for the anxiety subscale and 0.82 for the depression subscale (Bjelland et al., 2002).

**Wellbeing measure: Sleep quality.** The Pittsburgh Sleep Quality Index (PSQI) is a validated measure for sleep research and consists of 24 items (Buysse, 1989). The scale comprises seven components which measure (1) subjective sleep quality, (2) sleep latency, (3) sleep duration, (4) sleep efficiency, (5) sleep disturbance, (6) daytime dysfunction and (7) sleep medication over the past month. Each component generates a score from 0–3 where higher scores indicate poorer sleep outcomes. A sum of the seven component scores can be used to generate a global PSQI score ranging from 0 to 21. A global score above five indicates poor sleep

| Variable                     | Category                              | N   | %   | Mean + SD |
|------------------------------|---------------------------------------|-----|-----|-----------|
| Age                          |                                       | 637 | 42.9| (13.4)    |
| Gender                       | Male                                  | 70  | 11  |           |
|                              | Female                                | 557 | 87  |           |
|                              | Prefer not to say                     | 11  | 2   |           |
| Ethnicity                    | White – any background                | 601 | 94  |           |
|                              | Black/Black British                   | 5   | 1   |           |
|                              | Asian/Asian British                   | 7   | 1   |           |
|                              | Mixed                                 | 17  | 3   |           |
|                              | Other                                 | 6   | 1   |           |
| Highest level of education   | Secondary school                      | 70  | 11  |           |
|                              | Higher secondary or further Ed        | 185 | 29  |           |
|                              | Undergraduate degree                  | 221 | 35  |           |
|                              | Postgraduate degree                   | 154 | 25  |           |
| Cause of chronic pain        | Chronic widespread pain               | 218 | 34  |           |
|                              | Musculoskeletal                       | 234 | 37  |           |
|                              | Headaches                             | 66  | 10  |           |
|                              | Neuropathic                           | 93  | 15  |           |
|                              | Visceral                              | 19  | 3   |           |
|                              | Other                                 | 8   | 1   |           |
| Time since pain inception    | Up to 1 year                          | 15  | 2   |           |
|                              | 1–2 years                             | 32  | 5   |           |
|                              | 2–3 years                             | 26  | 4   |           |
|                              | 3–5 years                             | 58  | 9   |           |
|                              | 5–10 years                            | 138 | 22  |           |
|                              | Over 10 years                         | 365 | 57  |           |
|                              | Unsure                                | 4   | 1   |           |
| Pain medication              | Yes                                   | 574 | 90  |           |
|                              | No                                    | 64  | 10  |           |
| Co-morbid physical health conditiona | Yes                                      | 605 | 95  |           |
|                              | No                                    | 33  | 5   |           |
| Co-morbid mental health conditionb | Yes                                     | 353 | 55  |           |
|                              | No                                    | 285 | 45  |           |

*a Co-morbid physical health conditions such as diabetes, cardiovascular and respiratory conditions. *b Co-morbid mental health conditions such as depression and anxiety disorders.
quality. The scale has good reliability with Cronbach’s alpha scores above 0.8 (Carpenter and Andrykowski, 1998).

**Pain.** The Brief Pain Inventory (BPI) short form is a widely used self-report measure for clinical pain (Cleeland and Ryan, 1994). The BPI is composed of two subscales which rate severity of pain and the degree to which pain interferes with common dimensions of feeling and function in the past 24 hours. These are referred as (1) pain severity and (2) pain interference. Both subscales range from 0 to 10 with higher scores indicating higher levels of pain severity and interference. These scales have good internal consistency with Cronbach’s alpha of 0.85 and 0.88 for the severity and interference scales respectively (Tan et al., 2004).

**Statistical analyses.** To address the first aim and measure changes in wellbeing (sleep, anxiety and depression) and pain outcomes, paired t-tests were conducted to compare mean scores between Time 1 and Time 2 survey data.

To address the second aim and analyse the relationships between the grouping variables at Time 2; namely levels of dependence on others during lockdown, levels of ability to self-manage pain during lockdown, and access to healthcare during lockdown, and the dependent variables (global sleep quality, anxiety, and depression), three multivariate analysis of variance (MANOVA) were conducted. SPSS version 25 was used to conduct all analyses and significance levels were set at $p \leq 0.05$.

**Data sharing statement.** The current article includes the complete raw dataset collected in the study including the participants’ data set, syntax file and log files for analysis. Pending acceptance for publication, all of the data files will be automatically uploaded to the Figshare repository.

**Results**

Participants were mostly female (87%), of a white ethnic background (94%) and had a mean age of 42 years (SD = 13 years). Respondents were asked their chronic pain condition and responses were later categorised using the International Classification of Diseases (ICD) 11th edition which groups chronic pain into seven types (Treede et al., 2015). The sample included chronic widespread pain (e.g. fibromyalgia; 34%), musculoskeletal (e.g. osteoarthritis; 37%), headache (e.g. chronic migraine; 10%), visceral (e.g. pelvic pain; 3%), neuropathic (e.g. trigeminal neuralgia; 15%) and other (1%). Nearly all participants reported co-existing physical health conditions (95%), and over half reported a mental health condition (55%).

**Addressing Aim 1: Changes in wellbeing and pain outcomes**

Table 2 displays the paired t-test results comparing mean scores across Time 1 (Feb-Mar) and Time 2 (Apr-May) for all eight PSQI components of sleep (global sleep quality, subjective sleep quality, sleep latency, sleep duration, sleep efficiency, sleep disturbance, daytime dysfunction and sleep medication), anxiety, depression, pain interference and pain severity. Across the whole sample, statistically significant improvements were reported at Time 2 in global sleep quality, subjective sleep quality, sleep latency, sleep duration, sleep efficiency, sleep disturbance, daytime dysfunction and sleep medication, anxiety, depression, pain interference and pain severity. These improvements did not reach the minimum importance difference (MID) of 1.75–3 points for global sleep quality, 1.7 points for depression or 2.2 points for pain severity (Lu et al., 2013; Mease et al., 2011; Puhan et al., 2008). There were no significant changes in anxiety levels between Time 1 and Time 2.

**Addressing Aim 2: Differences in wellbeing outcomes at Time 2**

Three MANOVA were conducted using cross-sectional Time 2 data to determine whether there were any differences in wellbeing outcomes (sleep, anxiety and depression) based on changes in dependence on others, ability to
Differences in wellbeing outcomes based on changes of dependence on others during the pandemic

The results of the first MANOVA presented in Table 3 yielded a significant main effect of dependence on others during the pandemic on global sleep quality, anxiety and depression; Pillai’s Trace = 0.02, $F(6,1258) = 2.47$, $p = 0.02$, $n^2 = 0.01$. Between-subject effects showed significant group differences for sleep, anxiety and depression. Scheffe’s pairwise comparisons revealed that individuals whose dependence on others had increased during the pandemic reported significantly poorer sleep ($M = 13.50$) than individuals who reported less dependence on others during the pandemic ($M = 12.60$). A similar trend was observed with regards to anxiety scores. Individuals with increased dependence on others reported significantly higher anxiety scores ($M = 10.37$) than those who had become less dependent during the pandemic ($M = 9.01$). Finally, those who reported increased dependence on others, had significantly higher depression scores ($M = 8.75$) than those individuals who reported less dependence on others ($M = 7.67$).

Table 2. Paired t-test results comparing averages for sleep, anxiety, depression, pain interference and pain severity at Time 1 and Time 2.

| Variable                      | Time point     | N   | Mean ± SD | t    | p Value |
|-------------------------------|----------------|-----|-----------|------|---------|
| Global sleep quality (PSQI)   | Pre-lockdown   | 635 | 13.83 ± 3.67 | 3.344 | 0.001   |
|                               | During-lockdown|     | 13.19 ± 3.74 |      |         |
| Subjective sleep quality (PSQI)| Pre-lockdown   | 638 | 2.07 ± 0.7 | 3.235 | 0.001   |
|                               | During-lockdown|     | 1.94 ± 0.75 |      |         |
| Sleep latency (PSQI)          | Pre-lockdown   | 636 | 2.3 ± 0.96 | −0.23 | 0.818   |
|                               | During-lockdown|     | 2.31 ± 0.92 |      |         |
| Sleep duration (PSQI)         | Pre-lockdown   | 637 | 2.08 ± 1.05 | 6.648 | <0.001  |
|                               | During-lockdown|     | 1.71 ± 1.06 |      |         |
| Sleep efficiency (PSQI)       | Pre-lockdown   | 638 | 2.15 ± 1.1 | 0.473 | 0.636   |
|                               | During-lockdown|     | 2.12 ± 1.05 |      |         |
| Sleep disturbance (PSQI)      | Pre-lockdown   | 638 | 2.06 ± 0.6 | 0.154 | 0.878   |
|                               | During-lockdown|     | 2.05 ± 0.58 |      |         |
| Daytime dysfunction (PSQI)    | Pre-lockdown   | 638 | 2.07 ± 0.79 | 5.472 | <0.001  |
|                               | During-lockdown|     | 1.85 ± 0.79 |      |         |
| Sleep medication (PSQI)       | Pre-lockdown   | 638 | 1.11 ± 0.53 | −1.507 | 0.132   |
|                               | During-lockdown|     | 1.22 ± 0.53 |      |         |
| Anxiety (HADS-A)              | Pre-lockdown   | 636 | 9.98 ± 4.69 | 0.368 | 0.713   |
|                               | During-lockdown|     | 9.89 ± 4.76 |      |         |
| Depression (HADS-D)           | Pre-lockdown   | 636 | 9.09 ± 4.40 | 3.233 | 0.001   |
|                               | During-lockdown|     | 8.35 ± 4.44 |      |         |
| Pain interference (BPI)       | Pre-lockdown   | 638 | 6.36 ± 2.19 | 3.552 | <0.001  |
|                               | During-lockdown|     | 5.96 ± 2.32 |      |         |
| Pain severity (BPI)           | Pre-lockdown   | 638 | 5.43 ± 1.8 | 2.879 | 0.004   |
|                               | During-lockdown|     | 5.16 ± 1.86 |      |         |

SD: standard deviation; PSQI: Pittsburgh sleep quality index; HADS-A: Hospital Anxiety and Depression Scale (Anxiety); HADS-D: Hospital Anxiety and Depression Scale (Depression); BPI: brief pain inventory. Statistically significant differences are in bold.
The results of the second MANOVA presented in Table 4 yielded a significant main effect of ability to manage pain during the pandemic on global sleep quality, anxiety and depression; Pillai’s Trace = 0.08, \( F(6,1258) = 8.40, p < 0.001, n^2 = 0.04 \). Between-subject results yielded significant group differences for sleep, anxiety and depression. Scheffe’s pairwise comparisons revealed that individuals who found it harder to self-manage their pain during the pandemic had significantly poorer sleep (\( M = 13.79 \)) than both individuals who reported no change (\( M = 12.39 \)) and those who reported easier self-management of their pain during the pandemic (\( M = 10.92 \)). Similarly, individuals who found it harder to self-manage their pain during the pandemic reported significantly higher levels of anxiety (\( M = 10.70 \)) than both those who reported no change (\( M = 8.62 \)) and those who found it easier to self-manage their pain (\( M = 8.05 \)). The same trend was observed in relation to difference in depression scores. Those who struggled more with pain-management reported significantly higher levels of depression (\( M = 8.89 \)) than both those reporting no change (\( M = 7.55 \)) and those finding pain-management easier during the pandemic (\( M = 6.78 \)).

The results of the third MANOVA presented in Table 5 yielded a significant main effect of
access to healthcare during the pandemic (unrelated to COVID-19) on global sleep quality, anxiety and depression; Pillai’s Trace = 0.04, $F(6, 1190) = 4.12$, $p < 0.001$, $n^2 = 0.02$. There were significant between subject differences for sleep, anxiety and depression. Scheffe’s pairwise comparisons revealed that individuals who had healthcare appointments cancelled during the pandemic reported significantly poorer sleep ($M = 13.50$) than those who had usual access to care ($M = 12.05$). The same trend was observed with regards to anxiety levels. Individuals with cancelled healthcare appointments reported higher levels of anxiety ($M = 10.30$) than those with usual access to care ($M = 8.83$). Finally, participants with usual access to care during the pandemic had significantly lower levels of depressive symptoms ($M = 8.79$) than both those who experienced cancelled healthcare appointments ($M = 8.79$) and those who had face to face appointments replaced with telephone or virtual appointments ($M = 8.32$).

**Discussion**

Our prospective study describes the impact on wellbeing outcomes in a sample of 638 adults with chronic pain in the United Kingdom during the first wave of the COVID-19 pandemic. Despite increasing COVID-19 cases and related deaths between Time 1 and Time 2, statistically significant improvements in global sleep quality, subjective sleep quality, sleep duration, daytime dysfunction, depressive symptoms, pain interference, and pain severity were reported. At initial glance, it may seem these results contradict recent research findings which have reported an increase in sleep problems and mental health distress across general and clinical populations during the COVID-19 pandemic (Pierce et al., 2020; Wang et al., 2020b). However, our findings may reflect the difference in methodology used in other studies. Our study collected baseline data in the immediate period before government mandated lockdown measures were implemented and whilst the trend in cases was increasing exponentially. Follow-up measures were collected during lockdown, past the peak of case rises from COVID-19. One study from China conducted by Wang et al. (2020a) collected baseline and follow-up survey data during similar time points and found statistically significant reductions in PTSD symptoms between Time 1 and Time 2, as well as no changes in depression or anxiety scores. Although the Wang et al. (2020a) study was conducted on a general adult population, the methodology is comparable to the current study with regards to how and when data were collected during the outbreak in each respective country. The small improvements in sleep, depression, and pain outcomes between Time 1 and Time 2 may reflect feelings of uncertainty and health-related

| Variable | Access to healthcare during the pandemic (M, SD) |
|----------|---------------------------------------------|
|          | HC appointments cancelled $n = 375$ | HC appointments remote/online $n = 138$ | No Changes/no appointments $n = 86$ | $F (2, 596)$ | $p$ Value | $n^2$ |
| Sleep    | 13.50 (3.54)$^a$ | 12.94 (3.92)$^{ab}$ | 12.05 (4.15)$^b$ | 5.71 | 0.003 | 0.02 |
| Anxiety  | 10.30 (4.75)$^a$ | 9.78 (4.79)$^{ab}$ | 8.83 (4.60)$^b$ | 3.47 | 0.032 | 0.01 |
| Depression | 8.79 (4.47)$^a$ | 8.32 (4.08)$^a$ | 6.41 (4.10)$^b$ | 10.62 | <0.001 | 0.03 |

HC: healthcare; M: mean score; SD: standard deviation. Superscript letters have been used to indicate significant differences between-group Scheffe-corrected significance. Where letters are the same across variables there is no difference, and where letters differ (i.e. a and b), this denotes a significant difference ($p \leq 0.05$). Participants with missing values for independent or dependent variables were excluded from this analysis.
fear at the start of the outbreak, whilst there were upward trends in COVID-19 cases and deaths. The implementation of a nationwide lockdown and a downward trend in cases after mid-April 2020 may have contributed to some sense of security among our pain population thus resulting in the observed temporal changes in sleep, depression and pain scores. It is also possible that pain scores improved due to decrease in work related activities (e.g. furlough) and increased home-working practices.

Despite these improvements, scores for global sleep quality, anxiety and depression were above clinical cut-off points at Time 2 (Backhaus et al., 2002; Olsson et al., 2005). Research has shown that individuals living with chronic pain are at higher risk for concomitant sleep problems, anxiety, and depression, with higher scores compared to the general population (Hooten, 2016; Smith and Haythornthwaite, 2004). Our results demonstrate that individuals who felt less dependent on others had fewer sleep problems, anxiety, and depressive symptoms compared to those who reported feeling more dependent on others for practical and emotional support during this time. Furthermore, those who reported difficulty managing their pain condition had higher levels of sleep problems, anxiety and depression than those who managed their condition with ease. Interestingly, the disparity between these two groups crossed the clinical threshold for depression scores; individuals who found it easier to self-manage their pain were, on average, below threshold for depressive symptoms (scores <8) and those who reported more difficulty in their pain self-management were above the clinical threshold (scores >8) (Zigmond and Snaith, 1994). Finally, regarding access to healthcare, individuals whose healthcare appointments had been cancelled reported higher levels of sleep problems, anxiety, and depression than those who had experienced no disruption to usual care. Once again for depression scores, the difference between those individuals whose appointments had been cancelled and those who had usual care crossed the clinical threshold (Zigmond and Snaith, 1994).

The above findings support previous studies which illustrate how problems in sleep, increased anxiety and depression may develop as a long-term sequela of chronic pain. The results also demonstrate the disruption to independence, self-management and access to healthcare provision during the pandemic among this population. Chronic pain poses a threat to individuals’ perceived independence, and research has highlighted the importance of retaining independence in order to carry out activities of daily living and social interactions (Robinson et al., 2013; Tollefson et al., 2011). The ability to self-manage for individuals with chronic pain has been shown to reduce psychological distress and disability (Nicholas et al., 2012). Finally, our findings also outline the importance of maintaining access to health and care services as part of the wider care management plan for many with chronic pain.

Implications and recommendations

Based on the findings discussed, we highlight implications related to psychosocial wellbeing, workforce, and healthcare practice along with practical recommendations which should be considered in the current climate. Firstly, it is clear that the lockdown measures implemented during the first wave of COVID-19 have impacted communities and support for special populations. Our research highlights the importance of retained independence on mental health in chronic pain populations. It is foreseen that future lockdowns will cause a social disconnection and a threat of increased loneliness (Karos et al., 2020). Thus, we recommended that considerations are made for allowing social ‘bubbles’ and social cohesion to continue in support for chronic pain communities during future public health restrictions. Secondly, a significant proportion of the UK workforce is affected by chronic pain and disability across all sectors and skills bases who contribute greatly to our economy (Fayaz et al., 2016b). It is therefore vital to ensure this portion of the workforce can continue their contribution whilst managing the
increased risk of disease severity as a result of COVID-19. Workplace managers (supported by policies to protect workers’ rights) should carry out necessary risk assessment and ensure that individuals most at risk can contribute via adaptable working plans, whether through remote working or redeployment, before taking steps to prevent individuals from working in any capacity. Thirdly, there is evidence that closure of non-urgent health services has impacted waitlists as they begin to reopen (Karos et al., 2020). We therefore recommend that services which begin to triage backlogs of cases based on clinical need ensure referral pathways to psychological and mental health services are in place. There is evidence that mental health has suffered as a result of halted health and social care services and it is reasonable to assume that once financial assistance programmes come to an end (such as furlough schemes), many more will be in need of these services (Witteveen, 2020).

Limitations

This research relied on self-report measures which are known to misalign with more objective measures and reporting. Despite this, self-report measures offer a quick and accessible solution to data gathering via remote means which was imperative during the pandemic. Secondly, our sample consisted of individuals with chronic pain conditions, and therefore it is not possible to relate our findings to the general population. However, it is hoped they will be relevant to informing ways to improve wellbeing for individuals with chronic pain during future periods of lockdown. Thirdly, the majority of participants identified as ‘White – any background’ ethnicity and is therefore limited in generalizability, future studies should seek to explore methods to engage more diversity in responses (Fryer et al., 2016). Fourthly, it was not possible to include SES in the analysis, future studies should consider SES as a control variable. Finally, our data were collected across two time points, only one of which occurred during the lockdown which resulted in some of our analyses being conducted on cross-sectional data. Future research should seek to collect data longitudinally throughout the pandemic to assess causal relationships between lockdown measures on wellbeing.

Conclusion

To our knowledge, at time of writing, our study is the first to assess changes in wellbeing outcomes in a chronic pain population before and during the first lockdown in the UK. The results of this study suggest that there were small but significant improvements post-lockdown in relation to global sleep quality, depression and pain outcomes. Groups were identified as more likely to experience poorer wellbeing during lockdown based on increased dependence on others, lesser ability to self-manage pain, and restricted access to healthcare. As such, governing bodies and healthcare providers should enable essential services for chronic pain individuals as well as ensuring restrictions do not impact the ability for individuals to support themselves in managing their condition.

The research and learnings from the COVID-19 outbreak must be used to inform policy and emergency planning responses for future pandemics. Policymakers should consult with a wide range of professionals such as healthcare, social workers and Third Sector workers to plan local strategies which meet individual as well as collective needs within the population.

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Author contributions

ZZ, DD and AF: Concept and study design, ZZ: Survey creations, ZZ: Data cleansing, ZZ and AF: Data analyses, ZZ, DD, EJH: Manuscript and editing.
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