Examining temporal interactions between loneliness and depressive symptoms and the mediating role of emotion regulation difficulties among UK residents during the COVID-19 lockdown: Longitudinal results from the COVID-19 psychological wellbeing study

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Introduction

The general public, academic community, and World Health Organisation have raised concerns about the mental health impact of the COVID-19 crisis (Cowan, 2020; Holmes et al., 2020; WHO, 2020). Physical distancing restrictions and subsequent loss of social interaction, in addition to other challenges associated with living through a global pandemic, represent a significant stressor. The rapid onset of such a stressor may trigger loneliness (Lim et al., 2020) and depressive symptoms (Nolen-Hoeksema et al., 1992). This is concerning as loneliness and depression impact physical health and increase mortality (Holt-Lunstad et al., 2015; Holwerda et al., 2016). Relative to loneliness, objective isolation is a more robust predictor of mortality (Steptoe et al., 2013), whereas loneliness (i.e., subjective isolation) is more robustly associated with depression (Matthews et al., 2016). Successive waves of transmission have led to reintroductions of physical distancing measures to restrict the spread of the virus, including local and regional lockdowns. In the absence of an effective vaccine, mitigation policies are likely to be in place indefinitely. A priority for research is to understand the psychosocial impact of the pandemic and potential mechanisms accounting for any change in mental health symptoms (Holmes et al., 2020). This is in regard to the current COVID-19 global pandemic, and to future-proof the mental health response to future pandemics.

Loneliness is defined as a negative emotional experience resulting from the perception that the quantity or quality of one’s social relationships are insufficient (Perlman & Peplau, 1981). In contrast to the objective nature of social isolation (i.e., having few connections or infrequent contact), loneliness is the unpleasant subjective experience of feeling isolated which has been posited to involve social pain, sadness, and emptiness (Weiss, 1973). Loneliness and depression are distinct but closely related constructs (Mahon et al., 2006; Weeks et al., 1980). Establishing the nature of this relationship has been challenging. The interpersonal theory of depression (Coyne, 1976) and the stress-generation theory (Hammen, 1991) posit that depressive symptoms lead to interpersonal conflicts that damage relationships, thus leading to loneliness. This position has some empirical support from studies of older adulthood (Houtjes et al., 2014) and adolescence (Lasgaard et al., 2011). However, the balance of evidence suggests that either loneliness precedes depression or that the relationship is reciprocal. For older adults, loneliness predicted depressive symptoms 2-12 years later (Cacioppo et al., 2010; Holvast et al., 2015; Jaremka et al., 2014; S. L. Lee et al., 2020). Similar results were found with younger adults (Wei et al., 2005) and adolescents (Qualter et al., 2010). A rapid review of 63 studies concluded that loneliness has a negative impact on the mental health of children, adolescents, and young adults for up to 9 years, with the strongest impact found on
Numerous longitudinal studies have found a bidirectional relationship between loneliness and depression (Cacioppo et al., 2006; Domènech-Abella et al., 2019; Hsueh et al., 2019; Santini et al., 2020; Vanhalst, Luyckx, Raes, et al., 2012; Vanhalst, Luyckx, Teppers, et al., 2012). One study found the relationship was stronger with depression as the initial symptom (Hsueh et al., 2019), whereas others found it was stronger and more robust with loneliness as the initial symptom (Domènech-Abella et al., 2019; Vanhalst, Luyckx, Raes, et al., 2012; Vanhalst, Luyckx, Teppers, et al., 2012).

A substantial body of research implicates emotion dysregulation in the development of depression (Aldao et al., 2010; Ehring et al., 2010; Joormann, 2010; Joormann & Gotlib, 2010; Joormann & Stanton, 2016; Martin & Dahlen, 2005). Although loneliness is a negative emotional experience, less research has focused on its link with emotion regulation. Research has found an association between adaptive regulation strategies (i.e., reappraisal) and lower loneliness (Kearns & Creaven, 2017). Difficulty regulating emotions has been associated with greater loneliness (Wols et al., 2015) and was identified as a risk factor for loneliness during the COVID-19 pandemic (Groarke et al., 2020). There is some evidence that the relationship between loneliness and depressed mood is mediated by the use of maladaptive regulation strategies (e.g., rumination) (Vanhalst, Luyckx, Teppers, et al., 2012; Zawadzki et al., 2013). However, further research is required to understand the role of emotion dysregulation in the development of loneliness, and as a potential mediator of the relationship between loneliness and depression.

Mental health in the COVID-19 pandemic
Studies carried out during the pandemic have reported slight increases in depression (Canet-Juric et al., 2020; Li et al., 2020), whereas, others found depression and loneliness remained stable (Bu et al., 2020; Fancourt et al., 2020; Wang et al., 2020) or gradually decreased (Mata et al., 2020). Research has also examined the impact of the pandemic by comparing mental health before and after the introduction of mitigation policies. In the UK, distress rose by 8.4% one month into the lockdown compared to one year earlier (Pierce et al., 2020). Loneliness increased among Dutch older adults, but other mental health outcomes remained stable (van Tilburg et al., 2020). In Central European countries there was a five-fold increase in depressive symptoms, and those with high loneliness had a greater severity of depressive symptoms (Novotny et al., 2020). Moreover, loneliness was the strongest predictor of depression during the pandemic in an Israeli sample (Palgi et al., 2020). There was no significant change in loneliness measured before the outbreak, during the initial stages of the pandemic, or during the lockdown among American adults (Luchetti et al., 2020). Yet other studies
in the US reported higher loneliness and depression following the onset of the pandemic, and found that change in loneliness was associated with change in depression among older adults (Krendl & Perry, 2020) and younger adults (C. M. Lee et al., 2020). However, no studies have examined emotion regulation difficulties as a mechanism of this effect between isolation, loneliness, and depression in the context of COVID-19. This is warranted as research suggests the relationship between loneliness and depressed mood is mediated by the use of maladaptive regulation strategies (Zawadzki et al., 2013). Indeed, an ability to effectively regulate one’s emotions plays a key role in alleviating the potential negative impact of difficult circumstances many individuals are likely to experience during the pandemic (Restubog et al., 2020).

The current study
Few longitudinal studies have examined the temporal association between mental health outcomes during the COVID-19 outbreak. Drawing upon pre-pandemic research, it is predicted that the relationship between loneliness and depressive symptoms will be reciprocal, with the possibility of stronger or more robust relationships from loneliness to depressive symptoms. The bidirectional nature of the relationship has been overlooked in existing research during COVID-19. Furthermore, existing studies have focused exclusively on older adults (75 years old on average) or younger adults (18-23 years) in the US, with loneliness and depression assessed before and after but not also during shelter-in-place restrictions (Krendl & Perry, 2020; C. M. Lee et al., 2020). The current study expands upon this small body of work using structural equation modelling, within a UK context, and with a broader age range of adults. This study will determine the longitudinal relationship between loneliness and depressive symptoms over three timepoints during a period of acute stress (i.e., the four months following the announcement of the UK national lockdown on March 23rd 2020), and will examine, for the first time, the mediating role of emotion dysregulation in this relationship. Studies carried out prior to the pandemic have typically examined these relationships over prolonged periods (1-14 years), therefore, it is important to determine how associations between loneliness and depressive symptoms may manifest over a briefer timeframe and in a pandemic context. This study addresses key research priorities for the pandemic identified by the academic community, public health experts, and the general public (Cowan, 2020; Holmes et al., 2020; WHO, 2020).

Method
Design and Procedure
We conducted a longitudinal structural equation modelling analysis with data from the COVID-19 Psychological Wellbeing Study. Full methodological details of the study are reported elsewhere
(Armour et al., 2020). The study was approved by the Research Ethics Committee at Queen’s University Belfast (Reference: EPS 20_96). Participants were recruited via social media platforms and using a panel of UK residents hosted by Prolific. Prospective participants read a participant information sheet and if they provided informed consent proceeded to the online survey, administered through Qualtrics. Baseline data was collected between March 23rd and April 24th 2020 (Time 1). Participants were sent a follow-up survey one month post-baseline (Time 2), and again one month later (Time 3: two months post-baseline). One email reminder was sent to non-responders.

Participants
There were 2511 responses to the survey. Participants were excluded if they did not provide informed consent, were not resident in the UK, or were under 18 years of age. Responses were also screened for data quality (i.e., not completing any measures, or having a completion time less than half the median completion time), with 522 respondents removed from the dataset (see Figure 1). Of these participants, 98.4% were eligible for follow-up, that is, they provided valid contact information. 84.8% of those eligible responded to the follow-up assessment one month after baseline, and 79.1% responded to the final follow-up. There was an overall attrition rate of 20.9%. There were some significant yet slight differences across sociodemographic factors and baseline scores for those with and without complete data and by recruitment strategy (See supplementary appendix 1). Participants with complete data had slightly lower depressive symptoms, but there were no differences in loneliness or emotion regulation difficulties. Participants recruited through Prolific had greater emotion regulation difficulties, but there were no differences in loneliness or depressive symptoms.
The response categories often statements like ‘(Kaufman et al., 2016) Emotion Regulation Difficulties: from 0 (not at all) to 3 (nearly every day)..bothered by of major depressive disorder. Depressive symptoms: current Items were summed to produce a total loneliness score companionship?’. The items were rated “hardly Loneliness Measures Notes; Figure 1. Flowchart of the study sample by recruitment strategy at each timepoint. SMC = social media campaign. Measures Loneliness: The UCLA Three-Item Loneliness Scale (Hughes et al., 2004) assessed feelings of loneliness. Participants responded to items such as ‘how often do you feel that you lack companionship?’ The items were rated “hardly ever” (0), “some of the time” (1) or “often” (2). Items were summed to produce a total loneliness score. Internal consistency of the measure in the current sample was very high (Cronbach α T1 = .82, T2 = .85, T3 = .88). Depressive symptoms: The Patient Health Questionnaire (Kroenke et al., 2001) assessed symptoms of major depressive disorder. The 9-item questionnaire measures how often respondents have been bothered by symptoms such as ‘Feeling down, depressed, or hopeless’. Response options range from 0 (not at all) to 3 (nearly every day). In the current sample reliability coefficients were high (Cronbach’s α T1 = .91, T2 = .92, T3 = .88). Emotion Regulation Difficulties: The Difficulties in Emotion Regulation Scale - Short Form (Kaufman et al., 2016) contains 18 items rated on a 5-point Likert scale. Respondents are asked how often statements like ‘When I’m upset, I have difficulty controlling my behaviours’ apply to them. The response categories ranged from, ‘almost never’ (1) to ‘almost always’ (5). Cronbach’s alpha coefficients were high across each measurement time point (Cronbach α T1 = .91, T2 = .92, T3 = .92).
At baseline participants were asked to report their experiences of loneliness and emotion regulation difficulties over the past week and their depressive symptoms over the past two weeks. At Time 2 and Time 3, measures were rephrased to ask participants to reflect on their experiences over the previous month.

Statistical Analysis

Across study variables, less than 1% of data was missing at baseline, with 15.8% missing at Month 1, and 20.1% missing at Month 2. Little’s MCAR test ($\chi^2(708) = 750.836, p = .128$) showed the data were missing completely at random. Missing data were addressed using the Expectation Maximization algorithm. The 1958 participants who were eligible for follow-up are the focus of all subsequent analyses.

We tested the associations between loneliness, depression symptoms, and emotion regulation difficulty using Pearson’s product moment correlations, and descriptive statistics were calculated for all study variables. Cross-lagged structural equation modelling was used to examine the temporal relationships between variables using Amos version 23 (Arbuckle, 2014). Cross-lagged panel models allow for testing bidirectional predictive effects and assessing the relative strength of cross-lagged effects (i.e., the standardised regression coefficients), while also controlling for cross-sectional associations and autoregressive effects. Given this level of statistical control, effect sizes were expected to be small to moderate.

Maximum likelihood estimation was used. Following best practice, a range of indicators were used to assess model fit. Specifically, the chi-square test, the normed chi-square index (Q), the comparative fit index (CFI), the Tucker-Lewis index (TLI), the standardised root mean square residual (SRMR) and the root mean square error of approximation (RMSEA) (Byrne, 2010; Ullman, 2001), with non-significant $\chi^2$, Q values ranging from 2 to 5, CFI and TLI greater than .90, and SRMR and RMSEA less than .08 indicating good model fit (Hu & Bentler, 1999; Kenny & McCoach, 2003).

Structural equation modelling was carried out in two stages. The validity of the measurement model was tested first, followed by the validity of the structural model. The measurement model specified included 36 observed variables, that is, the 3 items of the UCLA loneliness measure and the 9 items of the PHQ-9 loading on their 6 respective latent variables (loneliness and depression) at each timepoint (Time 1, Time 2, Time 3) (Model A). Covariances between errors of the repeated measures
were allowed across timepoints on the basis of high modification indices. In the next step, the autoregressive effects were investigated (Model B). Error terms were co-varied for autoregressive effects, which can be thought of as indicators of the constructs’ temporal stability. Then, the structural models examining cross-lagged effects for loneliness and depression separately, with exogenous variables covaried (Models C and D) and the full reciprocal model, including all covariances, autoregressive paths, and cross-lagged paths (Model E) were assessed. Subsequently, for parsimony, the best fitting model was trimmed by excluding non-significant paths (Model F). Finally, we estimated an expanded model including age as a control variable (Model G) and a modified model including emotion dysregulation as a mediator (Model H) and non-significant paths were trimmed from that model (Model I). Due to the complexity of the model and the length of the DERS-SF (18 items), total scores on the DERS-SF were added to the model as 3 observed (i.e., measured) rather than latent variables.

Results

Baseline characteristics of the study sample are presented in supplementary appendix 1 and in the methodological paper describing the COVID-19 Psychological Wellbeing Study (Armour et al., 2020). In summary, participants (N = 1958) were predominantly Caucasian (92.7%) females (69.8%) with no religion (57.6%) residing in England (38%). Age ranged from 18-87 years (M = 37.01, SD = 12.81). Regarding COVID-19, 58.6% were self-isolating and 37.4% were in keyworker positions. Further, 33.8% of the participants met clinical criteria for probable depression (scores above 10) (Kroenke et al., 2001, 2007), and 26.8% of participants were classified as lonely (scores above cut-off of 6) (S. L. Lee et al., 2020; Steptoe et al., 2013).

Table 1 presents descriptive statistics and bivariate correlations among variables included in the longitudinal analysis. There were significant associations between loneliness and depression cross-sectionally and longitudinally with Pearson’s r coefficients ranging from .50 to .59 (all p < .001). Emotion dysregulation was significantly correlated with loneliness and depression (r = .45 – .71, all p < .001), however, correlations with depression were stronger overall (r = .61 – .71, all p < .001). Across each timepoint age was inversely associated (all p < .001) with loneliness (r = -.18 – -.21), depressive symptoms (r = -.23 – -.25), and difficulties regulating emotions (r = -.27 – -.28). Gender was not associated with loneliness or difficulties regulating emotions (p > .05). There were weak correlations between gender and depressive symptoms, indicating that symptoms were more severe for females (r = .09 – .11).
**Table 1.** Means, standard deviations, and Pearson’s correlation coefficients between variables included in the longitudinal analyses (N = 1958).

|                  | Loneliness |                  | Depressive Symptoms |                  | Emotion Regulation Difficulties |                  |
|------------------|------------|------------------|---------------------|------------------|---------------------------------|------------------|
|                  | T1         | T2               | T3                   | T1               | T2                | T3               | T1               | T2                | T3               |
| M                | 5.36       | 5.28             | 5.28                 | 7.85             | 7.76              | 7.50             | 42.52            | 41.64             | 41.37             |
| SD               | 1.92       | 1.89             | 1.89                 | 6.46             | 6.15              | 5.93             | 13.13            | 12.71             | 12.85             |
| Loneliness      |            |                  |                      |                  |                   |                  |                  |                   |                   |
| T1               | 1          |                  |                      |                  |                   |                  |                  |                   |                   |
| T2               | .699**     | 1                |                      |                  |                   |                  |                  |                   |                   |
| T3               | .699**     | .899**           | 1                    |                  |                   |                  |                  |                   |                   |
| Depressive       |            |                  |                      |                  |                   |                  |                  |                   |                   |
| Symptoms         |            |                  |                      |                  |                   |                  |                  |                   |                   |
| T1               | .552**     | .491**           | .491**               | 1                |                   |                  |                  |                   |                   |
| T2               | .523**     | .564**           | .564**               | .796**           | 1                 |                   |                  |                   |                   |
| T3               | .495**     | .594**           | .594**               | .742**           | .850**            | 1                 |                  |                   |                   |
| Emotion          |            |                  |                      |                  |                   |                  |                  |                   |                   |
| Regulation       |            |                  |                      |                  |                   |                  |                  |                   |                   |
| Difficulties     |            |                  |                      |                  |                   |                  |                  |                   |                   |
| T1               | .512**     | .465**           | .465**               | .670**           | .642**            | .614**           | 1                |                   |                   |
| T2               | .467**     | .487**           | .487**               | .626**           | .710**            | .670**           | .817**           | 1                 |                   |
| T3               | .447**     | .509**           | .509**               | .616**           | .665**            | .710**           | .793**           | .860**            | 1                 |

*Notes:*** p < .001; Time 1: Baseline (March 23rd-April 24th), T2: one month later, T3: two months later.
Fit of the measurement model (Model A) was acceptable. All standardised regression coefficients (i.e., factor loadings) were above .53, and statistically significant (all $p < .001$). Parameter estimates are presented in Supplementary Appendix 2. There was an improvement in fit indices with the addition of autoregressive effects (Model B). There was further improvement with the addition of cross-lagged effects (Models C – E). The measurement model and the structural models (Models A – F) were all deemed to have acceptable fit on the basis of CFI, TLI, SRMR and RMSEA criteria (See Table 2). The significant Chi-Square test should not lead to rejection of the model as this is frequently violated when large samples are used (Hu & Bentler, 1999). Based on fit indices Model F is the best fitting and most parsimonious model, that is, the full reciprocal model with non-significant paths trimmed (see Figure 2). The non-significant paths trimmed from Model F were cross-lagged effects between exogenous variables and criterion variables 2 months later. Specifically, loneliness at baseline did not significantly predict depression at Time 3 and depression at baseline did not significantly predict loneliness at Time 3.

**Table 2. Fit indices for alternate models**

| Goodness of fit index: | $\chi^2$ (df) | Q | CFI | SRMR | RMSEA (95% CI) | TLI |
|------------------------|----------------|---|-----|------|----------------|-----|
| Suggested value:       | $p > .05$      | 2-5 | > .90 | < .08 | < .08 | > .90 |
| Model A: measurement model with error terms covaried across timepoints | 4058.49(548) | 7.406 | 0.947 | 0.236 | 0.057 (.056,.059) | 0.932 |
| Model B: measurement model with autoregressive effects | 3315.58(549) | 6.039 | 0.953 | 0.073 | 0.051 (.049,.052) | 0.946 |
| Model C: model B with cross-lagged effects from loneliness to depression | 3220.96(546) | 5.899 | 0.955 | 0.054 | 0.050 (.048,.052) | 0.948 |
| Model D: model B with cross-lagged effects from depression to loneliness | 3253.905(546) | 5.960 | 0.954 | 0.060 | 0.050 (.049,.05) | 0.947 |
| Model E: model B with all cross-lagged effects (full reciprocal model) | 3176.150(543) | 5.849 | 0.956 | 0.048 | 0.050 (.048,.051) | 0.948 |
| Model F: trimmed model (model E = best fitting model) | 3176.24(545) | 5.828 | 0.956 | 0.048 | 0.050 (.048,.051) | 0.949 |
| Model G: Model F with age as control variable | 3888.829(576) | 6.751 | 0.944 | 0.140 | 0.054 (.053,.056) | 0.936 |

*Notes; CI = confidence intervals*
Further inspection of Model F (Figure 2) revealed high temporal stability in loneliness ($b = .79-.81$) and depression ($b = .73-.85$) over the first four months of the lockdown. The cross-lagged path between depression and subsequent loneliness was positive and significant from Time 1 to Time 2 ($b = .09$) and from Time 2 to Time 3 ($b = .11$). The cross-lagged path between loneliness and subsequent depression was also positive and significant at Time 2 ($b = .17$) and Time 3 ($b = .06$). The model supports the hypothesis that the relationship between loneliness and depression is reciprocal. From baseline to Time 2 the effect of loneliness on depression is stronger ($b = .17$) than the effect of depression on loneliness ($b = .10$), but from Time 2 to Time 3 it is weakened ($b = .06$) while the effect of depression on loneliness remains the same ($b = .11$).

A multigroup analysis indicated that Model F was different for males and females ($\text{CMIN}_{(74)} = 226.711$, $p < .001$). When individual paths of the structural model were constrained to be equal for males and females there were no significant differences for any of the autoregressive or cross-lagged effects ($\text{all } p > .05$). This suggests the source of invariance is in the measurement model. A full psychometric validation of the measurement instruments is beyond the scope of this study. Model F was expanded to include age as a control variable (Model G). Older age predicted lower loneliness ($b = -.23$) and depression ($b = -.23$) at baseline, and lower depression at Time 2 ($b = -.04$) only. Including age reduced model fit (see Table 2) and did not alter the pattern of temporal associations between loneliness and depressive symptoms (see Table 3).

![Figure 2. Cross-lagged structural model showing significant paths only.](image)

*Notes; all $p < .001$; L = UCLA three-item loneliness scale; D = PHQ-9; Time 1: Baseline (March 23rd-April 24th), T2: one month later, T3: two months later.*
Are the relationships between loneliness and depression mediated by emotion regulation difficulties?

The fit of the mediation model (Model H) was inferior to the best fitting model (Model F), but still represented a good fit of the data ($\chi^2_{(647)} = 4384.016$, $p < .001$; $Q = 6.776$, $CFI = .944$, $TLI = .936$, $SRMR = .049$, $RMSEA = .054$ [CI: .053, .056]). There was evidence of high temporal stability in emotion dysregulation across the three months of the study ($b = .68-.76$). The reciprocal association between loneliness and depression was replicated in the modified model, but we did not find support for the prediction that emotion dysregulation mediated the relationship. Loneliness predicted greater difficulties regulating emotion one month later ($b = .06$), which in turn predicted greater depressive symptoms ($b = .11$), and there was a direct effect of depressive symptoms at baseline on subsequent emotion regulation difficulties ($b = .16$), but no direct effect of emotion regulation difficulties on subsequent loneliness. There was, however, support for bidirectional temporal associations between emotion dysregulation and depression independent of loneliness. Difficulties regulating emotions predicted greater depressive symptoms one month later ($b = .11$), and depressive symptoms predicted higher emotion regulation difficulties one month later ($b = .13-.16$). Trimming the non-significant paths resulted in a more parsimonious model but did not improve fit (Model I: $\chi^2_{(649)} = 4385.208$, $p < .001$; $Q = 6.757$, $CFI = 0.944$, $TLI = .936$, $SRMR = 0.049$; $RMSEA = .054$ [CI: .053,.056]).

Figure 3. Cross-lagged structural mediation model showing significant paths only.

Notes; all $p < .001$; L = UCLA three-item loneliness scale; D = PHQ-9. DERS = Difficulties in Emotion Regulation Scale; Time 1: Baseline (March 23rd-April 24th), T2: one month later, T3: two months later.
### Table 3. Parameter estimates for structural and mediation models

| Predictor | Outcome | Model B | Model C | Model D | Model E | Model F | Model G | Model H | Model I |
|-----------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| L1        | L2      | .852*** | .866*** | .761*** | .795*** | .795*** | .790*** | .790*** | .795*** |
| D1        | D2      | .846*** | .715*** | .848*** | .734*** | .734*** | .788*** | .678*** | .679*** |
| L2        | L3      | .875*** | .879*** | .804*** | .807*** | .806*** | .815*** | .805*** | .806*** |
| D2        | D3      | .884*** | .841*** | .885*** | .847*** | .846*** | .845*** | .784*** | .784*** |
| L1        | D2      | .188*** | .167*** | .167*** | .182*** | .145*** | .146*** |         |         |
| L1        | D3      | -.006   | -.006   | -.006   | -.006   | -.006   | -.006   | -.006   | -.006   |
| L2        | D3      | .075*   | .067*   | .063*** | .058*** | .050**  | .049**  | .050**  | .049**  |
| D1        | L2      | .114*** | .087*** | .086*** | .187*** | .079**  | .090*** | .079**  | .090*** |
| D1        | L3      | -.007   | -.008   | -.007   | -.008   | -.007   | -.008   | -.007   | -.008   |
| D2        | L3      | .122*** | .115*** | .109*** | .103*** | .101*** | .109*** | .101*** | .109*** |
| Age       | L1      |         |         |         | -.234***|         |         |         |         |
| Age       | D1      |         |         |         | -.234***|         |         |         |         |
| Age       | L2      |         |         |         | -.013   |         |         |         |         |
| Age       | L3      |         |         |         | -.045** |         |         |         |         |
| Age       | D3      |         |         |         | -.009   |         |         |         |         |
| DERS 1    | DERS 2  |         |         |         |         | .676*** | .675*** |         |         |
| DERS 2    | DERS 3  |         |         |         |         | .761*** | .761*** |         |         |
| DERS 1    | L2      |         |         |         |         | .020    |         |         |         |
| DERS 1    | D2      |         |         |         |         | .108*** | .107*** |         |         |
| L1        | DERS 2  |         |         |         |         | .056**  | .057**  |         |         |
| D1        | DERS 2  |         |         |         |         | .158*** | .158*** |         |         |
| DERS 2    | D3      |         |         |         |         | .108*** | .108*** |         |         |
| DERS 2    | L3      |         |         |         |         | .012    |         |         |         |
| L2        | DERS 3  |         |         |         |         | .033*   | .032*   |         |         |
| D2        | DERS 3  |         |         |         |         | .127*** | .128*** |         |         |

**Notes:** ***p < .001  **p < .01 *p < .05; β = standardised beta coefficient; L = UCLA three-item loneliness scale; D = PHQ-9. DERS = Difficulties in Emotion Regulation Scale; Time 1: Baseline (March 23rd-April 24th), T2: one month later, T3: two months later.
Discussion

There was high stability in loneliness and depressive symptoms over the first 4 months of the UK lockdown. As predicted, there was a bidirectional positive association between loneliness and symptoms indicative of major depressive disorder. Loneliness predicted higher depressive symptoms one month later, and depressive symptoms predicted higher loneliness one month later. Age was inversely associated with loneliness and depressive symptoms cross-sectionally, but had limited impact over time. Age and gender did not impact the temporal interaction between loneliness and depressive symptoms. The relationship between loneliness and symptoms of depression was not mediated by emotion regulation difficulties. Difficulties regulating emotions and depressive symptoms were related reciprocally. To our knowledge this is the first study to apply longitudinal structural modelling to understand the temporal relationship between key mental health outcomes during the COVID-19 pandemic.

The finding of a reciprocal relationship between loneliness and depressive symptoms is in line with many pre-pandemic studies (Cacioppo et al., 2006; Domènech-Abella et al., 2019; Hsueh et al., 2019; Santini et al., 2020; Vanhalst, Luyckx, Raes, et al., 2012; Vanhalst, Luyckx, Teppers, et al., 2012). The findings are also consistent with US research which demonstrated that loneliness predicted depression during the pandemic (Krendl & Perry, 2020; C. M. Lee et al., 2020). However, these studies did not simultaneously test the hypothesis that depressive symptoms also predicted loneliness. A recent synthesis of qualitative studies with young people with depression offers a framework for understanding this bidirectional relationship (Achterbergh et al., 2020). Specifically, depressive symptoms, like fatigue and low self-esteem, prompt social withdrawal. Fear of disclosing depression contributes to feelings of alienation. Subsequently, social withdrawal and non-disclosure lead to loneliness, and the negative emotional experience of loneliness contributes to depression.

The current study suggests that although the relationship is bidirectional, loneliness influenced depression less consistently. Parameter estimates reveal that the effect of loneliness on depression is stronger initially but may diminish over time. This aligns with a pre-pandemic study of older adults (Hsueh et al., 2019), but diverges from studies that found the relationship is more robust with loneliness as the initial symptom (Domènech-Abella et al., 2019; Vanhalst, Luyckx, Raes, et al., 2012; Vanhalst et al., 2012). It is also surprising that the cross-lagged paths between loneliness and depression were significant at the subsequent timepoint, but not over longer time periods (two months later). Bidirectional associations have been observed between these variables over much
longer intervals (i.e., 1-5 years). This suggests that longitudinal patterns may operate differently in the short term or in the context of a novel stressor. The pattern of relations between mental health outcomes may also alter with continued exposure and adaptation to a stressor, or due to changes in the stressor, such as, the relaxing and reintroducing of physical distancing restrictions. Future studies exploring relations between loneliness and depression across shorter timeframes (e.g., days/weeks), or during periods of acute and unpredictable stress are thus warranted.

Contrary to expectations, the longitudinal relationship between loneliness and depressive symptoms was not mediated by difficulties in emotion regulation. It may be that the use of particular regulation strategies (e.g., ability to seek out social support, rumination, or reappraisal), rather than general difficulties regulating one’s emotions, mediates the relationship (Zawadzki et al., 2013). Few studies have examined the link between emotion regulation and loneliness, and the current study did not support the idea that they are significantly related in individuals experiencing lockdown due to the COVID-19 pandemic. Although regulation strategies have been associated with loneliness (Wols et al., 2015), and difficulties regulating emotions was a risk factor for loneliness in the initial phase of the lockdown (Groarke et al., 2020), in the current study emotion regulation difficulties did not predict loneliness over time and loneliness had limited impact on emotion dysregulation. However, there was an independent bidirectional association between emotion regulation difficulties and depression, supporting a significant body of pre-pandemic research (Aldao et al., 2010; Ehring et al., 2010; Joormann, 2010; Joormann & Gotlib, 2010; Joormann & Stanton, 2016; Martin & Dahlen, 2005).

There was no significant change in loneliness and depression over the three study timepoints, consistent with some international research (Bu et al., 2020; Fancourt et al., 2020; Luchetti et al., 2020; Wang et al., 2020). Loneliness and depressive symptoms remained stable despite significant shifts in disease-containment policies over the same timeframe. While stricter lockdown measures were in place at Time 1, from May 8th to 18th (Time 2) the UK announced easing of restrictions to allow, for example, outdoor exercise more than once per day and some outdoor gatherings. Notably, average scores on the PHQ-9 were below the threshold indicative of probable depression (Kroenke et al., 2001), and mean scores on the UCLA loneliness measure were also below standard cut-offs for loneliness (Groarke et al., 2020; Steptoe et al., 2013). Despite exposure to a major stressor, most respondents did not suffer significant impairment in mental health. That being said, the four-month timeframe may not capture a delayed impact of the pandemic, which may be mediated by short-term coping strategies. Further, research has identified more susceptible populations such as the
LGBTQ community, people in lower socioeconomic positions, minority ethnic groups, and people with disabilities or chronic physical and mental health conditions (lob et al., 2020; Salerno et al., 2020; Wright et al., 2020). Future studies should examine whether the temporal relationships observed in the current study differ among at-risk groups.

This study has important implications for the amelioration of subsyndromal mental health symptoms in the pandemic. The reciprocal relationship observed implies that interventions to reduce loneliness may alleviate depressive symptoms and vice versa. There are four main approaches to reducing loneliness: improving social skills, increasing social support, providing opportunities for social interaction, and addressing maladaptive social cognitions. Meta-analytic and systematic review evidence suggests that interventions that modify social cognitions are most effective (Jarvis et al., 2020; Ma et al., 2020; Masi et al., 2011). For mild-moderate symptoms of depression, clinical guidelines recommend low-intensity psychosocial interventions (NICE, 2009), such as low-intensity Cognitive Behavioural Therapy (CBT) a form of guided self-help that can be facilitated remotely. Online and telephone-based delivery of interventions can reduce depression (Gilbody et al., 2017; Lawn et al., 2019) and loneliness (Bouwman et al., 2017; Weinert et al., 2008), which is encouraging given the pandemic context. Clinical guidelines for treatment of moderate depressive symptoms also recommend physical activity (NICE, 2009). Reviews have concluded that walking and exercise reduce depressive symptoms (Robertson et al., 2012; Stathopoulou et al., 2006), as well as, loneliness (Hwang et al., 2019; Pels & Kleinert, 2016). Therefore, public health approaches that emphasise increasing physical activity may be particularly helpful for mitigating the mental health impact of the pandemic. Low intensity CBT that addresses social cognitions may also be of benefit. Treatments for depressive symptoms that address emotion regulation difficulties such as Dialectical Behaviour Therapy may be helpful, and adding emotion regulation skills training did improve the efficacy of CBT for major depression (Berking et al., 2013). However, a concurrent focus on loneliness may be necessary to fully address the psychosocial impact of the pandemic.

Strengths and Limitations

Strengths of the study include the large sample size, the use of validated measures, and the longitudinal design which permitted the examination of temporal relationships and mediation. This study addresses a gap in the research regarding associations between key mental health outcomes in the early stages of the lockdown in a UK context. However, limitations are acknowledged. The sample was not representative, males and older age groups were underrepresented (Armour et al., 2020). Although the psychometric properties of the measures used are well-established (Hughes et
al., 2004; Kaufman et al., 2016; Kroenke et al., 2001), clinical interviews may have been superior (Levis et al., 2019), but would be unfeasible given the large sample size. Consistent with other studies, loneliness, depression, and emotion dysregulation were stable during the lockdown. There was no pre-pandemic assessment, thereby limiting the conclusions that can be drawn regarding the mental health impact of the COVID-19 crisis. However, that was not the primary objective of the current study. The longitudinal design allows us to determine the temporal relationship between variables, not the causal relationship. Alternative explanations for the association cannot be ignored, for example loneliness and depression may share a common cause not measured in this study. The model was a good fit of the data. As with all structural equation models there may be alternative models that also fit the data equally well, that include other variables and covariates not accounted for in this structural model. Previous research, however, has not found the relationship between loneliness and depression to be significantly impacted by demographics, physical health, or related variables like perceived stress, social isolation, or social support (Cacioppo et al., 2010; Hsueh et al., 2019; S. L. Lee et al., 2020). Confidence in the structural model is enhanced, in that it was initially grounded in empirical research and theory.

Conclusion

Despite concerns about the negative impact of the pandemic on mental health, findings of the current study suggest that, on average, loneliness and depressive symptoms were mild-moderate and remained stable over the first four months of the UK lockdown. Results demonstrate that feelings of loneliness were positively associated with subsequent depressive symptoms, and that depressive symptoms were also positively associated with subsequent feelings of loneliness. Difficulties in emotion regulation predicted higher depression symptoms and vice versa, but difficulty regulating emotions did not mediate the relationship between loneliness and depression. The findings suggest that loneliness should be considered a potentially important part of case conceptualisation and formulation for individuals presenting with depressive symptoms during this time. Additionally, a measure to screen for loneliness could support assessment and identify those at-risk. Clinical efforts to improve mental health during the pandemic could focus on interventions that target either loneliness, or depression, or both. Such approaches include increasing physical activity or low-intensity cognitive therapies delivered remotely.
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