**Recovery from Covid-19 critical illness: a secondary analysis of the ISARIC4C CCP-UK cohort study and the RECOVER trial**

**Ellen Pauley, Thomas M Drake, David M Griffith, Louise Sigfrid, Nazir I Lone, Ewen M Harrison, J Kenneth Baillie, Janet T Scott, Timothy S Walsh, Malcolm G Semple, Annemarie B Docherty on behalf of the ISARIC4C and RECOVER Investigators**

**Author Affiliations**

**Ellen Pauley,** BMedSci, Medical Student,1 <https://orcid.org/0000-0001-6994-9722>

**Thomas M Drake,** MBChB BMedSci**,** Clinical Research Fellow,2 <https://orcid.org/0000-0002-9334-6494>

**David M Griffith,** MBChB, MD Reader and honorary consultant in critical care.3 <https://orcid.org/0000-0001-9500-241X>

**Louise Sigfrid**, MD PhD, Clinical Research Fellow, Public Health Specialist,. <https://orcid.org/0000-0003-2764-1177>

**Nazir Lone**, MBChB MSc PhD, Senior Clinical Lecturer and honorary consultant in Critical Care,3,5 <https://orcid.org/0000-0003-2707-2779>

**Ewen M Harrison,** MBChB MSc PhD, Professor of surgery and data science,2 UK. <https://orcid.org/0000-0002-5018-3066>

**J Kenneth Baillie,** MBChB PhD, Senior Clinical Lecturer and honorary consultant in critical care3.6 <https://orcid.org/0000-0001-5258-793>

**Janet T Scott,** MBChB PhD, Clinical Lecturer in Infectious Disease,7 <https://orcid.org/0000-0001-8030-5223>

**Timothy S Walsh,** MBChB, MD Professor ofAnaesthesia, Critical Care & Pain Medicine,3 <https://orcid.org/0000-0002-3590-8540>

**Malcolm G Semple** MBChB PhD, Professor of outbreak medicine and child health and consultant physician in paediatric respiratory medicine**,8** [**https://orcid.org/0000-0001-9700-0418**](https://orcid.org/0000-0001-9700-0418)

**Annemarie B Docherty,**MBChB MPH PhD, senior clinical lecturer and honorary consultant in critical care 2,3<https://orcid.org/0000-0001-8277-420X>

1. University of Edinburgh Medical School, Edinburgh, UK.
2. Centre for Medical Informatics, The Usher Institute, University of Edinburgh, Edinburgh, UK.
3. Anaesthesia, Critical Care & Pain Medicine, University of Edinburgh, Edinburgh, UK.
4. Centre for Tropical Medicine and Global Health, University of Oxford, Oxford, UK
5. Centre for Population Health Sciences, The Usher Institute, University of Edinburgh, Edinburgh, UK.
6. Roslin Institute, University of Edinburgh, Edinburgh, UK
7. MRC-University of Glasgow Centre for Virus Research, Glasgow, UK.
8. NIHR Health Protection Unit in Emerging Infectious Diseases, Institute of Infection, Veterinary and Ecological Sciences, Faculty of Health and Life Sciences, University of Liverpool, Liverpool, UK

Corresponding author: Annemarie B Docherty

Anaesthesia, Critical Care & Pain Medicine

Deanery of Molecular, Genetic, and Population Health Sciences

Usher Institute, University of Edinburgh

Room S8208, 2nd Floor

The Royal Infirmary of Edinburgh

NHS Lothian, 51 Little France Crescent

Edinburgh BioQuarter

Edinburgh EH16 4SA

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**Abstract**

**Background** We aimed to compare the prevalence and severity of fatigue in survivors of Covid-19 versus non-Covid-19 critical illness, and to explore potential associations between baseline characteristics and worse recovery.

**Methods** We conducted a secondary analysis of two prospectively collected datasets. The population included was 92 patients who received invasive mechanical ventilation (IMV) with Covid-19, and 240 patients who received IMV with non-Covid-19 illness before the pandemic. Follow-up data was collected post-hospital discharge using self-reported questionnaires. The main outcome measures were self-reported fatigue severity and the prevalence of severe fatigue (severity >7/10) 3 to 12-months post-hospital discharge.

**Results** Covid-19 IMV-patients were significantly younger with less prior comorbidity, and more males, than pre-pandemic IMV-patients. At 3-months, the prevalence (38.9% [7/18] vs. 27.1% [51/188]) and severity (median 5.5/10 vs. 5.0/10) of fatigue was similar between the Covid-19 and pre-pandemic populations respectively. At 6-months, the prevalence (10.3% [3/29] vs. 32.5% [54/166]) and severity (median 2.0/10 vs. 5.7/10) of fatigue was less in the Covid-19 cohort. In the total sample of IMV-patients included (i.e. all Covid-19 and pre-pandemic patients), having Covid-19 was significantly associated with less severe fatigue (severity <7/10) after adjusting for age, sex, and prior comorbidity (adjusted OR 0.35 (95%CI 0.15-0.76, p=0.01).

**Conclusion**. Fatigue may be less severe after Covid-19 than after other critical illness.

**Introduction**

Post Intensive Care Syndrome (PICS) describes the constellation of physical, psychological and cognitive symptoms affecting 25% of critical illness survivors, which include fatigue, muscle weakness, and posttraumatic stress1. PICS may persist for longer than 5 years and is associated with high hospital resource use and readmission rates2. Implementation of post-ICU rehabilitation guidelines is inconsistent, and the best ways to support intensive care unit (ICU) survivors are uncertain3–5; Identifying those at greatest risk of PICS and providing evidence-based strategies to enhance their recovery remain priorities for critical care research5. The Covid-19 pandemic heralded a dramatic increase in ICU admissions: in one study including over 20,000 UK hospitalised patients, 17% required admission to ICU or high dependency units , over half of whom survived to discharge, thus a vast new cohort of patients at risk of PICS has emerged6,7. This new cohort is relatively novel in terms of younger age, less previous comorbidity, and longer ICU stays than ICU admissions before the Covid-19 pandemic8,9.

 Many patients who survive acute-Covid-19 experience persistent symptoms beyond 4 weeks, known as ‘long-Covid’10. This includes a wide range of symptoms, such as breathlessness, fatigue, muscle pain, many of which overlap with PICS and other post-viral syndromes1,11–18. The most common symptom of long-Covid described following both community and hospital-managed acute-Covid-19 was fatigue: fatigue was reported in 97.7% of community-managed cases who reported symptoms lasting over 28 days, and in 83% and 98% of UK and China patients respectively greater than 3-months after hospital discharge following admission with acute-Covid-1913,18,19. Recent studies have shown requiring invasive mechanical ventilation (IMV) is associated with experiencing worse fatigue and other recovery, but have not attempted to compare ‘long-Covid’ to recovery from all critical illness and were without control groups11,13,14.

The prevalence of fatigue in ICU-survivors ranges from 13.8% to 80.9%20. This has a profound negative impact on survivors’ quality of life (QOL), and is one of the most prevalent and debilitating challenges of ICU-survivorship20,21. Fatigue may lead to delayed return to employment and pre-critical illness physical fitness, difficulty in completing activities of daily living, social isolation, depression, and many other negative sequalae 20–22. It is unclear if recovery is different in survivors of Covid-19 critical illness compared to other critical illness, requiring distinct risk assessment tools or interventions to support their recovery.

We aimed to characterise the prevalence and severity of fatigue in Covid-19 ICU-survivors, compared to survivors of non-Covid-19 critical illness, and to explore potential associations between baseline characteristics and fatigue severity in these populations to identify potential groups at greater risk.

**Methods**

*Study design and population*

This was a secondary analysis of two prospectively collected datasets.

*Covid-19 cohort: International Severe Acute Respiratory Infection Consortium Coronavirus Clinical Characterisation Protocol - United Kingdom (ISARIC-4C CCP-UK):*

Patients aged 18 years and over, admitted to hospital between 17th January and 5th October 2020 with confirmed or highly suspected SARS-CoV-2 infection at 31 UK centres, who consented to be contacted for follow-up studies, who completed a follow-up questionnaire, and who were discharged at least 90 days ago at the time of data collection were eligible for inclusion. Patients who did not survive to follow-up were excluded. We restricted this secondary analysis to patients who received IMV.

*Pre-pandemic cohort: Evaluation of a Rehabilitation Complex Intervention for Patients Following Intensive Care Discharge (RECOVER) trial:*

Patients aged 18 and over, discharged from ICU between 1st December 2010 and 31st January 2013 at 2 hospitals in Edinburgh, Scotland, who received a minimum of 48-hours of IMV23. Patients randomised to the intervention group received enhanced hospital and community-based physical rehabilitation, the control group received routine care23. Both groups were included in this analysis, as there were no significant differences at baseline or follow-up23.

Full details of the populations included in CCP-UK and RECOVER, and study information, have previously been published13,23.

*Clinical variables*

Both datasets captured demographic information, including age, sex, pre-existing comorbidities, admission diagnosis, length of hospital stay, and severity of acute illness. Comorbidities were defined in the ISARIC4C CCP-UK protocol for the Covid-19 cohort24, and by the Functional Comorbidity Index (FCI) for the pre-pandemic cohort23. For the pre-pandemic cohort, we combined ‘Angina’, ‘Congestive heart failure or heart disease’, and ‘Heart Attack’ from the FCI under the heading ‘Chronic Cardiac Disease’ to compare to the Covid-19 cohort (for which this was a category in the ISARIC4C protocol); similarly 'Chronic Obstructive Pulmonary Disease’ on the FCI was compared to ‘Chronic Pulmonary Disease’. Diagnosis category was captured from the RECOVER data in order to stratify the RECOVER cohort by diagnosis, to allow those with Respiratory diagnoses only to be compared to the Covid-19 cohort. Variables indicating acute illness severity differed: CCP-UK included ISARIC-4C Mortality and World Health Organisation Severity Scores25,26; RECOVER included the Acute Physiology and Chronic Health Evaluation-II (APACHE-II) score27, duration of IMV, duration of ICU-admission, total and Respiratory Sequential Organ Failure Assessment (SOFA) scores28. Thus, and given the different underlying disease processes, acute illness severity, including duration of IMV and ICU admission, was not directly compared. RECOVER captured survival at 3, 6, and 12-months post-hospital discharge: patients alive at each timepoint were contacted for follow-up.

*Outcomes*

Both studies used patient-completed questionnaires to collect outcome data13,23. CCP-UK completed follow-up assessment via postal questionnaires (or telephone where this was not possible), which patients completed once and returned13. RECOVER completed follow-up assessment face-to-face or by telephone at 3-months, and by post at 6 and 12-months post-hospital discharge23.

In this secondary analysis, the primary outcome was patient-reported fatigue severity. Secondary outcomes were: ‘severe fatigue’, breathlessness, and measures of health-related QOL.

In both studies, fatigue severity was measured using a 10-point Visual Analogue Scale (VAS), where zero is no fatigue and ten is worst possible fatigue. ‘Severe fatigue’ was defined as fatigue of at least 7/10 on the VAS.

*Statistical methods*

Covid-19 IMV-patients were compared to the pre-pandemic IMV-patients at baseline, follow-up, and potential associations with outcomes within each group were compared.

Covid-19 and pre-pandemic IMV-patients, and the pre-pandemic subset with Respiratory diagnosis category, were compared at baseline in terms of age, sex, prior comorbidity, deprivation. Fatigue severity and the prevalence of severe fatigue was compared. Fatigue in Covid-19 IMV-patients was stratified by time since hospital discharge that patients responded to the CCP-UK questionnaire: approximately 3 (90-120 days) and 6-months (150-210 days), in order to compare fatigue at equivalent timepoints to that measured in the RECOVER study. Thus a smaller subset of the Covid-19 sample was compared to the pre-pandemic group for fatigue at each timepoint. Fatigue severity (VAS scores) and ‘severe fatigue’ were compared as continuous and binary outcomes respectively.

Where direct comparison was not possible, i.e. where the same variables measured with the same tools were not available in the two datasets, outcome variables from both datasets were presented separately.

Categorical data were summarised as frequencies and percentages, continuous data as median and interquartile range (IQR). To test for differences across comparison groups in categorical data, we used Fisher’s exact test and for continuous data, using the Wilcoxon rank-sum test for two-sample testing and Kruskall-Wallis where there were more than 2 groups.

Using ordinal logistic regression and logistic regression, we explored if significant association existed between Covid-19 and severe fatigue in all patients included (Covid-19 and pre-pandemic group), after adjustment for patient demographics (age, sex, presence of comorbidity). With ordinal regression we explored fatigue in categories: 0-2/10, 2-4/10, 4-6/10, 6-8/10, 8-10/10, and with logistic regression we explored more severe reported fatigue, taking scores less than 7/10 or not as a binary outcome. Pre-pandemic outcomes at 6 months were included. We repeated these regression analyses including the Covid-19 group and the pre-pandemic subset with Respiratory diagnosis category. Effect estimates are presented as mean differences alongside 95% confidence intervals (95%-CI). Statistical analyses were performed using R version 3.6.3 (R Foundation for Statistical Computing, Vienna, AUT) with the tidyverse, finalfit, eq5d and Hmisc packages. Statistical significance was taken at the level P≤0.05.

**Results**

*Patients included*

335 Covid-19 patients were included in the CCP-UK study: 27.5% (92/335) received IMV (Figure-1). 240 patients were included in RECOVER (pre-pandemic patients). A total of 332 patients (92 Covid-19, 240 pre-pandemic patients) who received IMV were included in this secondary analysis (Figure-1). 29.2% [70/240] and 35.0% [84/240] of the pre-pandemic cohort were admitted with Cardiovascular and Respiratory diagnosis categories respectively (Table-1).

Covid-19 IMV-patients were significantly younger (median 59.7 years, IQR:51.1 to 64.5, p=0.017) than pre-pandemic IMV-patients (median 62.0 years, IQR:52.0 to 70.0, Table-1). A higher proportion of Covid-19 IMV-patients were males (70.7% [65/92]) than pre-pandemic (57.1% [137/240]). Significantly fewer Covid-19 IMV-patients had prior comorbidity (62.0% [57/92] vs 87.9% [211/240], p<0.001), and the median number of comorbidities was significantly lower (1, IQR:0 to 2 vs 2, IQR:1 to 4, p<0.001) Significantly fewer Covid-19 IMV-patients had chronic pulmonary disease (5.8% [5/86] vs 15.4% [37/240], p=0.036). Both groups were similar in terms of socioeconomic status, and the prevalence of obesity, diabetes, chronic cardiac disease, and asthma. These associations were consistent when only Respiratory category pre-pandemic patients were included (Supplementary Table-1). Covid-19 IMV-patients responded a median 185 days, IQR:137 to 241, after hospital discharge (Table-1).

Pre-pandemic IMV-patients received a median 8 days of IMV, IQR:5 to 15; the median APACHE-II score was 20, IQR:16 to 25 (Table-1).

*Primary outcome*

The median fatigue severity reported by Covid-19 IMV-patients at the timepoint they responded was 5.0, (n= 85), IQR:2.0 to 7.0, Table-2). At 3-months, fatigue severity was similar between Covid-19 (n=18) and pre-pandemic IMV-patients (n=188). At 6-months, pre-pandemic IMV-patients (n=166) reported significantly greater fatigue than Covid-19 IMV-patients (n=29), median 5.7/10, IQR:3.5 to 7.3/10 vs median 2/10, IQR:1.0 to 5.0/10, p<0.001). These associations were again consistent when only Respiratory category pre-pandemic patients were included (Supplementary Table-2): at 3-months, fatigue severity was similar between Covid-19 (n=18) and Respiratory pre-pandemic IMV-patients (n=62); at 6-months, Respiratory pre-pandemic IMV-patients (n=54) reported significantly greater fatigue than Covid-19 IMV-patients (n=32), median 6.6/10, IQR:4.6 to 7.8/10, p<0.001. Fatigue at 12-months was not compared because Covid-19 IMV-patients had not accrued follow-up to this time point.

Severe fatigue at the timepoint they responded was reported in 28.2% [24/85] of Covid-19 IMV-patients who responded (Supplementary Table-3). At 3-months post-hospital discharge, the prevalence of severe fatigue was similar in Covid-19 (38.9%, [7/18]) and pre-pandemic IMV-patients (27.1%, [51/188]), and at 6-months significantly less Covid-19 IMV-patients experienced severe fatigue (10.3% [3/29] vs 32.5% [54/166], p=0.015). This pattern was again consistent when only Respiratory category pre-pandemic patients were included (Supplementary Table-4): at 3-months and 6-months respectively, 35.5% [22/62] (p=0.789) and 44.4% [24/54] (p=0.0014) of Respiratory category pre-pandemic IMV-patients reported severe fatigue.

*Predictors of fatigue*

At a univariable level, in the Covid-19 group, females under 50 reported more severe fatigue (mean 5.20, 95%CI: 2.93 to 7.47); In pre-pandemic IMV-patients, sex did not affect outcomes (Supplementary Figure-1, Supplementary Table-5).

In the total sample included (Covid-19 and pre-pandemic patients), having Covid-19 was significantly associated with less severe fatigue (cut-off 7/10, adjusted OR 0.35 (95%CI:0.15 to 0.76, p=0.01) (Figure-2, Supplementary Table-5). When the Covid-19 IMV-patients and the Respiratory category pre-pandemic IMV-patients were included, having Covid-19 (adjusted OR 0.15, 95%CI:0.05 to 0.41) and female sex (adjusted OR 0.34, 95%CI:0.12 to 0.88) were significantly associated with less severe fatigue (Supplementary Figure-2, Supplementary Table-6). Ordinal logistic regression analysis showed that ventilated patients with Covid-19 (adjusted OR 0.50, 95%CI:0.29to 0.84) had less severe fatigue than patients ventilated due to other aetiologies (total pre-pandemic cohort). One or more comorbidities (adjusted OR 3.17, 95%CI:1.75 to 5.81) was associated with greater fatigue, and age 50-69 (adjusted OR 0.56, 95%CI:0.29 to 1.05), and age over 70 (adjusted OR 0.39, 95%CI:0.18 to 0.84) were associated with less fatigue (Supplementary Table-7). When the Covid-19 IMV-patients and the Respiratory category pre-pandemic IMV-patients were included, ordinal logistic regression also showed that having Covid-19 (adjusted OR 0.29, 95%CI:0.13 to 0.58) was significantly associated with less severe fatigue (Supplementary Table-8); having prior comorbidity was associated with greater fatigue (adjusted OR 2.18, 95%CI:1.06 to 4.54).

**Discussion**

We found a high prevalence and severity of patient-reported persistent fatigue following hospital discharge in ventilated patients with both Covid-19 and non-Covid-19 critical illness. At 3-months post-hospital discharge, the prevalence and severity of fatigue reported was similar following Covid-19 and non-Covid-19 critical illness, however by 6-months, patients ventilated for Covid-19 had significantly less severe fatigue than patients ventilated for other critical illness.

Both Covid-19 and non-Covid-19 critical illness survivors experienced high levels of fatigue, adding to existing evidence that finding the best rehabilitation services to support ICU-survivors is a research priority4,5. In our study, the difference in fatigue prevalence and severity at 6-months suggests that the recovery trajectory from Covid-19 critical illness may be different to general PICS. Covid-19 critical illness survivors are also more likely to suffer additional respiratory sequelae29. In our analysis patients who survived Covid-19 were younger with less prior comorbidity than those surviving other critical illness - this may explain why less fatigue at 6-months was reported after Covid-19 critical illness, as coping with the challenges of comorbidity has been identified as a barrier to ICU-recovery30,31. The lower fatigue in Covid-19 patients at 6-months is surprising as Covid-19 patients may have been expected to suffer worse fatigue than pre-pandemic patients if they were more likely to be discharged at a lower functional level due to greater high-demand for ICU beds throughout the pandemic and a focus on survival as opposed to rehabilitation. A multitude of different factors interact to drive fatigue in critical illness survivors, including deconditioning secondary to extended immobilisation, anaemia, poor sleep, depression, post-traumatic stress, respiratory illness (which may be due to prolonged ventilation), chronic disease, and drugs1,20,30,32–35. It is possible that once acute illness severity exceeds the threshold for requiring IMV and ICU-admission, further illness severity and underlying disease have little impact on persistent fatigue. This is consistent with recent research which found that post-ICU rehabilitation requirements were unaffected by Covid-19 infection status36.

Our data provides important insights into these complex syndromes, which need far greater research to assess the true impact they have on patients. To our knowledge, no previous study has included an appropriate control group with similar illness severity to a Covid-19 cohort. There are several important limitations to our study that must be considered. Firstly, our findings may be subject to responder and survivor bias. The sample sizes included by both CCP-UK and RECOVER were relatively small, with the RECOVER sample being substantially larger, and individuals with milder symptoms may have felt less compelled to respond, and those with the most severe symptoms or who died would have been unable to respond. Self-reported fatigue may have been affected by patients’ perceptions of their function or fatigue level prior to ICU admission, which may be affected by their age or previous comorbidities. As the Covid-19 group was significantly younger with less comorbidity, and therefore may have had a self-perceived lower baseline fatigue, an ideal study would have compared more objective measurements of fatigue level. Secondly, the overall Covid-19 cohort was included in our logistic regression models, which may have introduced bias from differing questionnaire response times post-hospital discharge. However, we controlled for CCP-UK and RECOVER measuring outcomes at different timepoints by stratifying fatigue by time since hospital discharge when comparing fatigue prevalence and severity, which allowed equivalent timepoints to be compared but further restricted Covid-19 sample sizes, therefore limiting the external validity of our findings to the general ICU population. Thirdly, our results may not be fully representative of all survivors of Covid-19 or non-Covid critical illness, as only patients admitted to a few UK hospitals were included (31 hospitals were included in CCP-UK and only 2 in RECOVER, therefore CCP-UK is likely to have captured a much wider variation in population and recovery services available to the patients included). Forthly, as CCP-UK and RECOVER did not use the same outcome measures, we were unable to explore differences in indicators of acute illness severity, including duration of IMV and length of ICU stay, between Covid-19 and pre-pandemic patients, which may have been an important risk factor for fatigue and therefore confounded our results, however a recent systematic review found no significant difference in mortality, length of hospital stay, or IMV-free days between Covid-19 and the general Acute Respiratory Distress Syndrome population37. Also, pre-critical illness factors, such as prior comorbidity, may be stronger predictors of hospital readmission and resource use than acute illness factors in ICU survivor patients31. An ideal study design would have used retrospective measurements of pre-critical illness functional levels, utilised repeated measures (much like the RECOVER study) of those with Covid-19 ,and would also feature a non-Covid-19 contemporaneous control group, to control for other factors which may impact post-ICU recovery, such as the ‘lockdown’ restrictions in place during CCP-UK’s study period; restrictions such as office closures may have meant survivors’ lives were less physically demanding, allowing for greater rest, and the pandemic has heralded the formalisation of ICU-recovery clinics in some areas38. As the RECOVER study was completed 8 years prior to CCP-UK, the level of follow-up care available is likely to have changed over this time prior to the Covid-19 pandemic, particularly as rehabilitation after critical illness was made a NICE quality standard in 2017, recommending a review at 2-3 months post ICU-discharge4.

We found high levels of fatigue following both Covid-19 and non-Covid-19 critical illness. Survivors of Covid-19 critical illness experienced less severe fatigue at 6-months post-hospital discharge than survivors of non-Covid-19 critical illness, potentially due in part to the comparatively younger and less comorbid Covid-19 ICU cohort versus the non-Covid-19 cohort. Research targeting interventions to best support critical illness survivors is required in order to optimise recovery.

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Study registration ISRCTN66726260.

The ISARIC WHO CCP-UK study was registered at <https://www.isrctn.com/ISRCTN66726260> and designated an Urgent Public Health Research Study by NIHR.

# Ethical considerations

Ethical approval was given by the South Central - Oxford C Research Ethics Committee in England (Ref 13/SC/0149), the Scotland A Research Ethics Committee (Ref 20/SS/0028), and the WHO Ethics Review Committee (RPC571 and RPC572, 25 April 2013.

# Data availability

This work uses data provided by patients and collected by the NHS as part of their care and support #DataSavesLives. The CO-CIN data was collated by ISARIC4C Investigators. ISARIC4C welcomes applications for data and material access through our Independent Data and Material Access Committee (<https://isaric4c.net>).

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**Tables and Figures**

**Table 1: Patient characteristics of Covid-19 (n=92) and pre-pandemic IMV-patients (n=240).**

IQR – Interquartile range, presented as 25th to 75th centiles. Numbers are presented as N (%), unless otherwise denoted as a continuous variable. ICU – Intensive Care Unit. IMV – Invasive Mechanical Ventilation. SOFA – Sequential Organ Failure Assessment. APACHE-II – Acute Physiology and Chronic Health-II.

|  |  | Covid-19 cohort | Pre-pandemic cohort | p-value |
| --- | --- | --- | --- | --- |
| Total N (%) |  | 92 (27.7) | 240 (72.3) |  |
| Age | Median (IQR) | 59.7 (51.1 to 64.5) | 62.0 (52.0 to 70.0) | 0.017 |
| Sex | Male | 65 (70.7) | 137 (57.1) | 0.032 |
|  | Female | 27 (29.3) | 103 (42.9) |  |
| Number of comorbidities | Median (IQR) | 1.0 (0.0 to 2.0) | 2.0 (1.0 to 4.0) | <0.001 |
| Comorbidities or not | No comorbidities | 35 (38.0) | 29 (12.1) | <0.001 |
|  | One or more comorbidities | 57 (62.0) | 211 (87.9) |  |
| Deprivation | Most deprived 1 | 10 (10.9) | 33 (13.8) | 0.277 |
|  | 2 | 19 (20.7) | 60 (25.0) |  |
|  | 3 | 26 (28.3) | 48 (20.0) |  |
|  | 4 | 22 (23.9) | 45 (18.8) |  |
|  | Least deprived 5 | 15 (16.3) | 54 (22.5) |  |
| Obesity | No | 62 (72.9) | 187 (77.9) | 0.434 |
|  | Yes | 23 (27.1) | 53 (22.1) |  |
| Diabetes | No | 66 (75.9) | 203 (84.6) | 0.097 |
|  | Yes | 21 (24.1) | 37 (15.4) |  |
| Asthma | No | 71 (82.6) | 197 (82.1) | 1.000 |
|  | Yes | 15 (17.4) | 43 (17.9) |  |
| Chronic Pulmonary Disease (Not Asthma) | NoYes  | 81 (94.2)5 (5.8) | 203 (84.6)37 (15.4) | 0.036 |
| Chronic Cardiac Disease | NoYes | 78 (90.7)8 (9.3) | 194 (80.8)46 (19.2) | 0.052 |
| ICU admission diagnosis category | Cardiovascular | - | 70 (29.2) | - |
|  | Respiratory | 92 (100.0) | 84 (35.0) | - |
|  | Gastrointestinal tract | - | 59 (24.6) | - |
|  | Renal | - | 3 (1.3) | - |
|  | Trauma | - | 8 (3.3) | - |
|  | Neurologic |  | 12 (5.0) |  |
|  | Miscellaneous diagnoses | - | 4 (1.7) | - |
| ISARIC-4C Mortality Score | Median (IQR) | 8.0 (5.0 to 10.0) | - | - |
| Length of hospital stay | Median (IQR) | 26.0 (15.5 to 40.0) | 32.0 (20.0 to 50.0) | 0.02 |
| ICU length of stay | Median (IQR) | - | 11.0 (6.8 to 18.0) | - |
| Duration of IMV | Median (IQR) | - | 8.0 (5.0 to 15.0) | - |
| Total SOFA Score | Median (IQR) | - | 3.0 (2.0 to 4.0) | - |
| Respiratory SOFA Score | 0 | - | 23 (9.6) | - |
|  | 1 |  | 73 (30.4) |  |
|  | 2 |  | 144 (60.) |  |
| APACHE-II Score | Median (IQR) | - | 20.0 (16.0 to 25.0) | - |
| Time from discharge to completing CCP-UK questionnaire (days) | Median (IQR) | 185.0 (136.8 to 241.2) | - | - |
| 3-month survival | Yes  | - | 216 (94.7) | - |
|  | No |  | 12 (5.3) |  |
| 6-month survival | Yes  | - | 169 (89.4) |  |
|  | No |  | 20 (10.6) |  |
| 12-month survival | Yes | - | 163 (88.1) |  |
|  | No |  | 22 (11.9) |  |

**Table 2: Fatigue severity in Covid-19 and pre-pandemic IMV-patients.**

|  |  | Covid-19 cohort | Pre-pandemic cohort | p-value |
| --- | --- | --- | --- | --- |
| Total N |  | 18  | 188  |  |
| Fatigue severity at ~3 months | Median (IQR) | 5.5 (2.0 to 6.2) | 5.0 (2.8 to 7.0) | 0.809 |
| Total N  |  | 32  | 166 |  |
| Fatigue severity at ~6 months | Median (IQR) | 2.0 (1.0 to 5.0) | 5.7 (3.5 to 7.3) | <0.001 |
| Total N  |  |  | 159 |  |
| Fatigue severity at ~12 months |  |  | 5.0 (3.0 to 7.0) |  |
| Total N |  | 85 |  |  |
| Fatigue severity across follow-up period | Median (IQR) | 5.0 (2.0 to 7.0) |  |  |

Fatigue severity according to Visual Analogue Scale in self-reported questionnaires. Covid-19 cohort: ~3 months – 90-120 days, ~6 months – 150-210 days, Fatigue at 12 months was not compared as the Covid-19 cohort had not accrued follow-up to this timepoint.



**Figure 1: Patient inclusion flowchart. Red: RECOVER trial participants, n=240; Blue: ISARIC4C participants, n=92**



**Figure 2: Logistic regression model for severe (>=7/10) fatigue at 6-months including all patients (Covid-19 and pre-pandemic group). OR, 95% CI, p-value.** Pre-pandemic patients included at 6-months follow-up. Number in model = 251,  AIC = 262.4, C-statistic = 0.676, H&L = Chi-sq(8) 2.26 (p=0.972) Group = Covid-19 status. 0= pre-pandemic group, non-Covid. 1= Covid-19 cohort. No\_comorbid\_factor = Number of comorbidities. Male, Group 0, No Comorbidities, Age Under 50 are marked with ‘-‘ as these were used as the reference level for each respective variable.