Clinical frailty, and not features of acute infection, is associated with late mortality in COVID-19: a retrospective cohort study

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Abstract

Background Coronavirus disease 2019 (COVID-19) is associated with excess mortality after hospital discharge. Identification of patients at increased risk of death following hospital discharge is needed to guide clinical monitoring and early intervention. Herein, we aimed to identify predictors of early vs. late mortality in COVID-19 patients.

Methods A total of 471 patients with polymerase chain reaction-confirmed COVID-19 were followed up for 9 months [median (inter-quartile range) of follow-up time: 271 (14) days] after hospital admission. COVID-19-related signs and symptoms, laboratory features, co-morbidities, Coronavirus Clinical Characterisation Consortium (4C) mortality and Clinical Frailty Scale (CFS) scores were analysed by logistic regression for association with early (28 day) vs. late mortality. Receiver operating characteristic (ROC) analysis was used to determine the discriminative value of 4C and CFS scores for early vs. late mortality.

Results A total of 120 patients died within 28 days from hospital admission. Of the remaining 351 patients, 41 died within the next 8 months. Respiratory failure, systemic inflammation, and renal impairment were associated with early mortality, while active cancer and dementia were associated with late mortality, after adjustment for age and sex. 4C mortality score and CFS were associated with both early [odds ratio (OR) (95% confidence interval—CI): 4C: 1.34 (1.25–1.45); CFS: 1.49 (1.33–1.66)] and late [OR (95% CI): 4C: 1.23 (1.12–1.36); CFS: 2.04 (1.62–2.56)] mortality. After adjustment for CFS, the association between 4C and late mortality was lost. By ROC analysis, 4C mortality score was superior to CFS for 28 day mortality [area under the curve (AUC) (95% CI): 0.779 (0.732–0.825) vs. 0.723 (0.673–0.773), respectively; P = 0.039]. In contrast, CFS had higher predictive value for late mortality compared with 4C mortality score [AUC (95% CI): 0.830 (0.776–0.883) vs. 0.724 (0.650–0.798), respectively; P = 0.007].

Conclusions In our cohort, late mortality in COVID-19 patients is more strongly associated with premorbid clinical frailty than with severity of the acute infection phase.

Keywords COVID-19; Late mortality; Frailty; 4C mortality score; Prognosis

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Introduction

As of February 2022, the ongoing coronavirus disease 2019 (COVID-19) pandemic has affected more than 400 million people and has caused more than 5.5 million deaths worldwide. Clinical severity of the disease can vary from mild, common cold symptoms to severe pneumonia, systemic inflammatory response syndrome, respiratory failure, and death. Thus, early and accurate patient risk stratification to guide clinical decision making is of utmost importance. Towards this goal, many clinical algorithms have been developed and validated in real-life settings, most notably the Coronavirus Clinical Characterisation Consortium (4C) mortality score. However, such algorithms focus upon initial treatment escalation and short-term prognosis, while longer-term effects of COVID-19 are now increasingly recognized.

Recent studies have shown that patients who recover from acute infection by the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) virus exhibit continuous symptoms such as fatigue, muscle weakness, sleep and psychological disturbances, as well as increased incidence of new-onset organ dysfunction, rates of hospitalizations, and mortality over subsequent months. In a recent nationwide UK study, 30% of discharged COVID-19 patients were readmitted to hospital over a 5 month period after discharge, while 1 out of 10 patients died. Whether this late morbidity and mortality is directly associated with COVID-19 (‘post-COVID syndrome’) or stems from deterioration of underlying co-morbidities remains largely unknown. Therefore, identification of patients at risk of developing multi-organ dysfunction or death after hospital discharge is needed to guide clinical monitoring and early intervention. Herein, we examined the clinical and laboratory predictors of early (28 day) vs. late mortality (>28 day till max. follow-up period) among patients with COVID-19 treated in a large UK tertiary academic medical centre. The main goals of our manuscript were to compare predictors of early (28 day) vs. late mortality in patients with COVID-19; to examine whether 4C mortality score had a prognostic value beyond the first 28 days; and to compare the value of the well-established 4C mortality score and Clinical Frailty Scale (CFS) for early (28 day) mortality vs. late mortality. We hypothesized that frailty is more important for late mortality (>28 day) compared with parameters related to the acute infectious disease.

Methods

Patient cohort

This is a retrospective analysis of real-world clinical data captured during the first-wave period of the COVID-19 pandemic in the UK. In the initial few weeks of the pandemic (the so-called ‘containment phase’), all patients testing positive for COVID-19 were admitted to a high consequence infectious diseases (HCID) centre—which included our hospital—regardless of disease severity. A total of eight patients were admitted during the containment phase period, all of whom were young with mild disease and all of whom survived. Following this period, only patients with physiological perturbations requiring inpatient care were admitted to hospital. The decision to admit was made on an individual basis by the medical team rather than on a priori criteria, based on a severity of disease sufficient to warrant inpatient treatment (typically based upon the need for supplemental oxygen and/or intravenous therapy, and/or the requirement for intensive monitoring). Reflecting this high disease severity, 45% of the patients had an oxygen saturation <92% on admission, and 48% were tachypnoeic (respiratory rate >20). The observed high 28 day mortality rate (120/471, 25%) also underscores the high severity of infection in our cohort.

Electronic patient records were searched to identify all patients admitted with COVID-19 to the Newcastle upon Tyne Hospitals NHS Foundation Trust between 31 January and 31 May 2020 inclusive. Only patients with a positive SARS-CoV-2 reverse transcription polymerase chain reaction (PCR) test based on combined nose and throat swab or sputum sample were included. We excluded patients under the age of 18 years, those who were already inpatients at the time of infection, or those who were asymptomatic and admitted for an unrelated reason (e.g. positive screening swab prior to surgery). Electronic clinical records were retrospectively reviewed to collect demographic, clinical, and laboratory data, with the aid of a version-controlled data collection template (Excel, Microsoft Corporation). Mortality data for participants who were discharged alive were collected in January 2021, 271 (14) [median (inter-quartile range—IQR)] days after admission. All deaths, including deaths in the community after hospital discharge, were captured by robust daily electronic system updates via primary care. The clinical management and 28 day outcomes of the first 316 patients admitted in this cohort have been previously published.

Risk score calculation

The CFS is a widely used clinical algorithm with values ranging from 1 (very fit, among fittest persons for their age) to 9 (terminally ill, life expectancy <6 months) (Table S1). A cut-off of 5 differentiates people who do not need help with everyday activities (CFS 0–4) from individuals who are dependent on others for everyday help to terminally ill patients (CFS 5–9) (Table S1). Therefore, a cut-off of 5 was used in our analyses to define frailty. In accordance with the official guidance, we estimated CFS corresponding to the patients’ clinical condition 2 weeks prior to acute
illness (COVID-19). More specifically, CFS at 2 weeks prior to admission was estimated by the admitting medical team based on available information in the medical records, supplemented by additional information from relatives or care home staff where applicable/available. The notes review was performed by qualified medical doctors (medical degree holders, registered with the General Medical Council, and working within the National Health Service). Where CFS was not documented in the medical notes, we retrospectively estimated CFS based on the information available in the medical records. All available information in the notes was used to estimate the CFS—this includes contemporaneous documentation in the medical and nursing admission notes as to physical function, assistance required for feeding/washing/dressing (all of which are mandatory fields on the nursing admission documentation at our hospital), previous clinic letters, social worker entries, and any documented discussions with relatives and/or carers. While CFS has been widely validated for individuals older than 65 years, a recent study showed that CFS can also predict in-hospital death in younger individuals with COVID-19. Therefore, we calculated CFS for all study participants.

Finally, we calculated the 4C mortality score for each patient at admission. 4C mortality score is a validated risk score to estimate risk of mortality in patients with COVID-19. It consists of eight weighted variables (age, sex, co-morbidities as defined in the Charlson Comorbidity Index, with the addition of obesity, respiratory rate, oxygen saturation on room air, Glasgow Coma Scale score, urea, and C-reactive protein at the point of hospital admission) (Table S2). Patients who were receiving supplementary oxygen at time of admission were considered as hypoxic (SpO₂ < 92%). When data in one of the eight weighted variables were unavailable, 4C mortality score was not calculated. In total, 4C mortality score was calculated in 435/471 (92.4%) patients. Based on a cumulative score (0–21), patients are stratified in low (0–3), intermediate (4–8), high (9–14), and very high (≥15) risk for in-hospital mortality.

Inpatients were examined every day up to and including 28 days after admission, and an adapted World Health Organization ordinal scale for COVID-19 was calculated. More specifically, this score has values from 1 to 8, with 8 corresponding to patients who are not hospitalized and do not require oxygen supplementation and 1 corresponding to death. For our analysis, we used a cut-off of 4, which discriminates those patients who need hospitalization and oxygen supplementation.

**Statistical analysis**

Normality of variable distribution was tested by Kolmogorov–Smirnov test. Continuous variables are presented as median (IQR) and categorical variables as absolute count (percentage). Pairwise comparisons between continuous variables were performed by Mann–Whitney U test, while Fisher’s exact test was used to compare categorical variables between contrast groups. Logistic regression was used to examine the association of patients’ signs and symptoms, as well as laboratory parameters, with early (28 day) and late (after 28 days) mortality, after controlling for age and sex. We further constructed receiver operating characteristic (ROC) curves for the occurrence of 28 day and late mortality across a range of values for continuous classifiers. To infer about superiority in discrimination between predictors of common outcome, we compared the corresponding area under the curve (AUC) using the trapezoidal rule. For comparison of independent ROC curves (i.e. prediction of the occurrence of 28 day and late mortality by the same continuous predictor), we followed the approach described by Gönen.

To guard against overfitting, we used two distinct set of confounders, one composed of clinical variables and another integrating laboratory parameters. This approach reduced the overall number of covariates in the final multivariable model. Only variables with biological plausibility and/or statistical significance in univariable analysis were included in the final multivariable model. Thus, we ensured a ratio of 5–10 events per covariate used in the multivariable regression model. Moreover, we assessed whether addition of CFS significantly contributed to the core set of clinical or laboratory variables (core model) by calculating (i) explained variation (adjusted McFadden’s R² index) (Table S3) and (ii) the Akaike information criterion and likelihood ratio tests for nested models with respect to 28 day and late mortality (Table S4). We also employed a cross-validation procedure (iterated 50 times) by random splitting of the study population to training and validation data set and calculated the AUC of the final multivariable model in the two data sets (Table S5). Then, we compared AUCs between the two data sets and checked for concordance, which indicates absence of overfitting.

We used restricted cubic splines with three knots—fixed at the 10th, 50th, and 90th percentile of the underlying distribution of CFS or 4C mortality score—to model the association of these scores as continuous variables with overall mortality (dose–response curves). The Wald test was used to assess the significance of the non-linear term. Finally, to increase the robustness of our results (internal validation), we employed resampling techniques and calculated normal-based confidence intervals after bootstrapping with 1000 replicates for (i) difference in AUC for 4C and CFS with respect to late mortality and (ii) coefficients of 4C and CFS in multivariable [including age, sex, and estimated glomerular filtration rate (eGFR)] logistic regression models of early and late mortality. All tests were two-tailed; results were considered significant when P < 0.05. Statistical analysis was performed using SPSS v. 26 and Stata v. 13.
**Ethics**

The study was registered as a clinical service evaluation with the Newcastle upon Tyne Hospitals NHS Foundation Trust (Reference 10040), was exempt from ethical approval, and was exempt from a need for patient consent as a study of COVID-19 under Regulation 3(4) of the Health Service Control of Patient Information Regulations 2002 (March 2020). Analysis of anonymized healthcare data was approved by the Caldicott Guardian (References 7523 and 7595).

**Results**

**Prognostic value of COVID-19-related clinical signs and symptoms for 28 day and late mortality**

A total of 471 consecutive adults admitted to hospital with PCR-confirmed COVID-19 were included in our analysis. Clinical and laboratory features of individuals who survived COVID-19 (n = 351) vs. patients who died during the first 28 days from hospital admission (n = 120; early mortality group), as well as patients who survived up to 9 months after hospital admission with COVID-19 (n = 310) vs. those patients who died after the first 28 days (n = 41; late mortality group), are summarized in Tables S6 and S7. The study cohort was aged 74 (25) years [median (IQR)], had a small male predominance (52.7%), and consisted mainly of British, Irish, or other White ethnicity (85.8%).

Our analysis showed that patients who survived beyond 28 days from admission more frequently presented with classical common cold symptoms such as pharyngitis and myalgia (Table S6). Regarding patient demographics, we did not find an association between male sex and mortality, as previously described in the literature; however, a strong association of age with both early [OR (95% CI): 1.07 (1.05–1.09) per 1 year increase] and late OR (95% CI): 1.07 (1.04–1.10) per 1 year increase] mortality was observed (Table 1).

Respiratory distress, as evidenced by increased respiratory rate, shortness of breath, and decreased oxygen saturation at admission (SpO₂ < 92%), was associated with increased early mortality (Table 1), a relationship that remained significant after adjustment for age and sex (Table 3 and Figure 1). In contrast, acute respiratory symptoms were not associated with late mortality. On the other hand, while confusion showed a significant association with both early and late mortality in univariate analyses (Table 1), no association was observed after adjustment for age and sex (Table 3).

In line with current evidence highlighting the presence of co-morbidities as determining factor for COVID-19 prognosis, we observed an association of arterial hypertension, congestive heart failure, and chronic kidney disease with early mortality (Table 1). On the other hand, active cancer and ischaemic heart were associated with late mortality (Table 1). Dementia was the only co-morbid condition associated with both early and late mortality in univariate analyses (Table 1); however, after adjusting for age and sex, its association remained significant only with late mortality (Table 3 and Figure 1). Similarly, only heart failure and active cancer retained their association with early and late mortality, respectively, after controlling for the effect of age and sex (Table 3 and Figure 1). Further adjustment for renal function (eGFR), as a significant factor associated with mortality in patients with COVID-19, did not significantly affect the results (Table S8).

**COVID-19-related laboratory parameters in the prediction of 28 day and late mortality**

In addition to the clinical signs and symptoms, a number of laboratory features at point of admission were associated with increased mortality in our cohort. Presence of leucocytosis or leukopenia, elevated neutrophil count, elevated C-reactive protein (CRP), decreased eGFR, increased urea, increased alkaline phosphatase (ALP), decreased albumin, and increased sodium were associated with 28 day mortality (Table 2). Of note, most of these associations remained unaffected by adjustment for age and sex: presence of leucocytosis/leukopenia, increased neutrophil count, elevated CRP, decreased eGFR, increased urea, decreased albumin, and elevated sodium retained their association with 28 day mortality (Table 3 and Figure 1). In contrast, only ALP and the cachexia-related markers haemoglobin and albumin were associated with late mortality after adjustment for age and sex (Table 3 and Figure 1). Further adjustment for renal function (eGFR), as a significant factor associated with mortality in patients with COVID-19, did not significantly affect the results (Table S8).

**4C mortality and Clinical Frailty Scale scores in risk stratification of hospitalized patients with COVID-19 for early and late mortality**

To examine the potential value of currently used COVID-19 prognostic scores for late mortality, we calculated the 4C mortality score, an established and validated risk stratification tool for early (28 day) COVID-19 mortality, and CFS, which has shown additive prognostic value over age and co-morbidities in COVID-19.

Both 4C mortality score and CFS, which positively correlated with each other in our cohort (rho = 0.610, P < 0.001), were associated with total 9 month mortality [OR (95% CI) 4C: 1.34 (1.26–1.44), P < 0.001, and CFS: 1.72 (1.54–1.93), P < 0.001] (Table S9). By dose–response curves, we found a linear association for both CFS (P = 0.303 for the
non-linear term, OR for 80th to 20th percentile = 15.39, 95% CI 8.52–27.78, $P < 0.001$ and $4C (P = 0.215$ for the non-linear term, OR for 80th to 20th percentile = 9.17, 95% CI 5.17–16.26, $P < 0.001$) with overall mortality in patients with COVID-19 (Figure S1).

Of interest, we observed that a high 4C mortality score was associated with both early mortality [OR (95% CI): 1.34 (1.25–1.45)] and late mortality [OR (95% CI): 1.23 (1.12–1.36)] (Table 2 and Figure 2A). After adjusting for age and sex, 4C remained significantly associated with early mortality [adj. OR (95% CI): 1.26 (1.15–1.37), $P < 0.001$], but lost its association with late mortality [adj. OR (95% CI): 1.13 (0.99–1.28), $P = 0.066$] (Table 3). Similar results were observed after additional adjustment for eGFR (Table S8). After bootstrapping, similar results were observed, as 4C remained significantly associated with early mortality [OR (95% CI): 1.25 (1.14–1.37), $P < 0.001$], but lost its association with late mortality [OR (95% CI): 1.14 (0.966–1.31), $P = 0.058$].

Clinical Frailty Scale was also associated with both early and late mortality [OR (95% CI): 1.49 (1.33–1.66) and 2.04 (1.83–2.28), $P < 0.001$, respectively].

Table 1  Clinical signs, symptoms, and co-morbidities as predictors of early (28 day) vs. late mortality

| Demographics | Early mortality | Late mortality |
|--------------|----------------|---------------|
| Age (years)  | OR (95% CI) | P-value | OR (95% CI) | P-value |
| Age >65 years | 1.07 (1.05–1.09) | <0.001 | 1.07 (1.04–1.10) | <0.001 |
| Male sex     | 1.19 (0.78–1.80) | 0.419 | 0.79 (0.41–1.52) | 0.477 |
| Symptoms and signs at presentation | | | |
| T > 38°C or T < 36°C | 1.11 (0.69–1.80) | 0.664 | 1.15 (0.55–2.42) | 0.709 |
| Cough        | 0.79 (0.51–1.23) | 0.300 | 0.77 (0.39–1.52) | 0.456 |
| Sputum       | 0.65 (0.38–1.11) | 0.113 | 0.85 (0.39–1.86) | 0.679 |
| Shortness of breath | 1.56 (1.003–2.42) | 0.049 | 0.73 (0.38–1.40) | 0.343 |
| Haemoptysis  | 0.37 (0.08–1.66) | 0.196 | 0.74 (0.09–5.92) | 0.774 |
| Rhinorrhoea  | —               | —     | 1.15 (0.25–5.30) | 0.856 |
| Pharyngitis  | —               | —     | 0.41 (0.09–1.77) | 0.231 |
| Myalgia      | 0.37 (0.19–0.73) | 0.004 | 0.36 (0.13–1.05) | 0.062 |
| Arthralgia   | —               | —     | 0.93 (0.11–7.62) | 0.945 |
| Fatigue      | 1.01 (0.66–1.54) | 0.972 | 0.93 (0.47–1.82) | 0.826 |
| Headache     | 0.60 (0.27–1.32) | 0.202 | —               | —     |
| Confusion    | 2.68 (1.71–4.20) | <0.001 | 2.68 (1.33–5.40) | 0.006 |
| Anosmia      | 0.53 (0.18–1.59) | 0.259 | 0.77 (0.17–3.44) | 0.734 |
| Abdominal pain | 0.69 (0.35–1.39) | 0.301 | 0.72 (0.24–2.12) | 0.546 |
| Diarrhoea    | 0.62 (0.34–1.11) | 0.109 | 0.66 (0.27–1.63) | 0.368 |
| Heart rate (b.p.m.) | 1.00 (0.99–1.01) | 0.438 | 0.99 (0.98–1.01) | 0.347 |
| Heart rate >90 b.p.m. | 1.13 (0.74–1.72) | 0.573 | 0.52 (0.26–1.05) | 0.069 |
| Systolic BP (mmHg) | 1.00 (0.99–1.01) | 0.778 | 1.00 (0.99–1.02) | 0.832 |
| Diastolic BP (mmHg) | 0.99 (0.98–1.01) | 0.291 | 0.98 (0.95–1.00) | 0.073 |
| Respiratory rate (breaths per minute) | 1.07 (1.03–1.11) | <0.001 | 1.00 (0.94–1.06) | 0.966 |
| RR > 20 (breaths per minute) | 2.07 (1.34–3.19) | 0.001 | 1.08 (0.56–2.11) | 0.811 |
| Hypoxia (SpO2 < 92%) | 2.22 (1.45–3.42) | <0.001 | 1.19 (0.62–2.31) | 0.598 |
| Mean BP (mmHg) | 1.00 (0.98–1.01) | 0.674 | 0.99 (0.97–1.01) | 0.386 |
| Co-morbidities | | | |
| Active cancer | 1.60 (0.88–2.91) | 0.122 | 3.36 (1.49–7.58) | 0.004 |
| Asthma       | 0.61 (0.30–1.26) | 0.184 | 0.50 (0.15–1.69) | 0.265 |
| COPD         | 1.07 (0.61–1.90) | 0.806 | 2.04 (0.93–4.47) | 0.074 |
| Other ILD    | 1.29 (0.52–3.20) | 0.590 | 1.07 (0.24–4.90) | 0.927 |
| Ischaemic heart disease | 1.28 (0.78–2.11) | 0.322 | 2.84 (1.41–5.73) | 0.004 |
| Congestive heart failure | 2.93 (1.72–4.99) | <0.001 | 1.90 (0.78–4.66) | 0.160 |
| Chronic kidney disease | 2.00 (1.26–3.16) | 0.003 | 1.71 (0.82–3.55) | 0.149 |
| DM diet-controlled | 1.47 (0.76–2.82) | 0.253 | 0.79 (0.23–2.72) | 0.708 |
| DM tabs-controlled | 1.11 (0.58–2.14) | 0.747 | 1.53 (0.60–3.93) | 0.375 |
| DM insulin | 1.12 (0.48–2.61) | 0.788 | 0.78 (0.17–3.47) | 0.741 |
| Hypertension | 2.14 (1.41–3.27) | <0.001 | 1.19 (0.61–2.32) | 0.619 |
| Liver disease | 0.29 (0.07–1.28) | 0.103 | 1.44 (0.40–5.17) | 0.576 |
| Dementia    | 2.60 (1.65–4.24) | <0.001 | 6.52 (3.19–13.33) | <0.001 |
| Adapted WHO ordinal scale at admission | 94 (0.04–0.82) | <0.001 | 0.95 (0.64–1.41) | 0.800 |
| Adapted WHO ordinal scale at discharge | — | — | 0.89 (0.58–1.38) | 0.606 |

b.p.m., beats per minute; BP, blood pressure; CI, confidence interval; COPD, chronic obstructive pulmonary disease; DM, diabetes mellitus; ILD, interstitial lung disease; mmHg, millimetres of mercury; OR, odds ratio; RR, respiratory rate; T, temperature; WHO, World Health Organization. Values in bold denote statistically significant results.

*Odds ratios are derived from binary logistic regression with early or late mortality as the dependent variable, and per 1-unit increase for continuous variables, or vs. the reference category for categorical variables, as the independent variable.

*OR not calculated because of zero occurrences in one of the groups.

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For early and late mortality, respectively (Table 2 and Figure 2B). However, CFS retained its association with both early [adj. OR (95% CI): 1.26 (1.10–1.46), $P = 0.001$] and late mortality [adj. OR (95% CI): 1.97 (1.52–2.54), $P < 0.001$] after adjustment for age and sex (Table 3), or additional adjustment for eGFR (Table S8). After bootstrapping,
similar results were observed [OR (95% CI) early mortality: 1.25 (1.06–1.47), \( P = 0.009 \), and late mortality: 1.96 (1.52–2.51), \( P < 0.001 \)]. Finally, multivariable logistic regression analysis including all factors associated with early mortality (Table 4), late mortality (Table 5), or total 9 month mortality (Table S10) showed an independent association of CFS with all study outcomes. Importantly, after bootstrapping, the association of CFS with both 28 day and late mortality remained significant after controlling for clinical or laboratory variables [28 day mortality: OR (95% CI) for CFS after adjustment for clinical characteristics = 1.34 (1.12–1.59) and OR (95% CI) for CFS after adjustment for laboratory parameters = 1.33 (1.14–1.56)/late mortality: OR (95% CI) for CFS after adjustment for clinical characteristics = 1.99 (1.52–2.59) and OR (95% CI) for CFS after adjustment for laboratory parameters = 1.93 (1.49–2.49)]. Similar results were obtained when we used a cut-off of 5 for CFS, which differentiates people who do not need help with everyday activities (CFS 0–4) from individuals who are dependent on others for everyday help to terminally ill patients (CFS 5–9) (Tables S11 and S12). Across the main multivariable logistic regression models for both 28 day and late mortality, no evidence of overfitting was shown (Tables S3–S5).

Next, by ROC analysis, we found that 4C mortality score and CFS had similar prognostic value for total 9 month mortality [4C AUC (95% CI): 0.781 (0.738–0.824) vs. CFS AUC (95% CI): 0.783 (0.740–0.826), \( P = 0.946 \) (Figure S2)]. On the other hand, 4C mortality score was superior to the CFS for predicting early mortality [4C AUC (95% CI): 0.779 (0.732–0.825) vs. CFS AUC (95% CI): 0.723 (0.673–0.773), \( P = 0.039 \); Figure 2C]. A reversed pattern was observed for late mortality, with the CFS outperforming the 4C mortality score [CFS AUC (95% CI): 0.830 (0.776–0.883) vs. 4C AUC (95% CI): 0.724 (0.650–0.798), \( P = 0.007 \), Figure 2D]. Moreover, CFS outperformed 4C for predicting late mortality after bootstrapping [OR (95% CI): 0.839 (0.820–0.846) vs. 0.726 (0.701–0.752), for CFS and 4C respectively, \( P < 0.001 \)]. Accordingly, adjustment for CFS did not affect the relationship between age, respiratory symptoms, inflammation, and renal impairment at admission with 28 day mortality (Table S13 and Figure 3). On the other hand, only the presence of active cancer and low haemoglobin were independently associated with late mortality after adjustment for CFS (Table S13 and Figure 3), indicating the importance of frailty and active cancer as independent predictors of late mortality in this cohort.

### Table 3 Association of disease features with early (28 day) and late mortality adjusted for age and sex

|                    | Early mortality          | Late mortality          |
|--------------------|--------------------------|-------------------------|
|                    | OR (95% CI)\(^a\)       | \( P\)-value            | OR (95% CI)\(^a\)   | \( P\)-value |
| Symptoms and signs at presentation |                         |                         |                         |              |
| Shortness of breath | 2.30 (1.40–3.76)         | 0.001                   | 0.92 (0.46–1.85)       | 0.819        |
| Myalgia            | 0.74 (0.35–1.55)         | 0.425                   | 0.80 (0.25–2.52)       | 0.702        |
| Confusion          | 1.30 (0.78–2.14)         | 0.314                   | 1.30 (0.60–2.81)       | 0.506        |
| Respiratory rate (breaths per minute) | 1.09 (1.04–1.13)         | <0.001                  | 1.01 (0.95–1.08)       | 0.716        |
| RR > 20 (breaths per minute) | 2.53 (1.57–4.09)         | <0.001                  | 1.23 (0.61–2.46)       | 0.562        |
| Hypoxia (SpO<sub>2</sub> < 92%) | 2.67 (1.65–4.30)         | <0.001                  | 1.31 (0.66–2.62)       | 0.444        |
| Co-morbidities     |                         |                         |                         |              |
| Active cancer      | 1.48 (0.78–2.82)         | 0.231                   | 3.15 (1.33–7.47)       | 0.009        |
| Ischaemic heart disease | 0.82 (0.48–1.40)         | 0.460                   | 1.84 (0.89–3.83)       | 0.102        |
| Congestive heart failure | 2.07 (1.18–3.65)         | 0.011                   | 1.26 (0.50–3.20)       | 0.624        |
| Chronic kidney disease | 1.05 (0.64–1.74)         | 0.844                   | 1.01 (0.47–2.18)       | 0.976        |
| Hypertension       | 1.40 (0.89–2.21)         | 0.149                   | 0.73 (0.36–1.49)       | 0.393        |
| Dementia           | 1.12 (0.65–1.94)         | 0.671                   | 2.94 (1.30–6.65)       | 0.010        |
| Laboratory parameters |                         |                         |                         |              |
| Haemoglobin (g/L)  | 0.997 (0.987–1.008)      | 0.608                   | 0.975 (0.959–0.991)    | 0.003        |
| WBCs < 4 or WBCs > 12 (× 10<sup>9</sup>/L) | 1.97 (1.15–3.34)         | 0.013                   | 1.21 (0.51–2.90)       | 0.661        |
| Neutrophils (× 10<sup>9</sup>/L) | 1.12 (1.05–1.20)         | <0.001                  | 1.10 (0.99–1.22)       | 0.068        |
| Urea (mmol/L)      | 1.04 (1.00–1.07)         | 0.026                   | 1.00 (0.95–1.05)       | 0.972        |
| eGFR (mL/min/1.73 m<sup>2</sup>) | 0.989 (0.979–0.998)      | 0.020                   | 1.00 (0.98–1.01)       | 0.674        |
| ALP (IU/L)         | 1.002 (1.000–1.005)      | 0.075                   | 1.006 (1.002–1.011)    | 0.005        |
| CRP (mg/L)         | 1.005 (1.003–1.008)      | <0.001                  | 0.999 (0.996–1.004)    | 0.807        |
| Albumin (g/L)      | 0.94 (0.90–0.99)         | 0.011                   | 0.92 (0.86–0.98)       | 0.016        |
| Sodium (mmol/L)    | 1.04 (1.01–1.08)         | 0.025                   | 0.97 (0.91–1.03)       | 0.343        |
| Risk scores        |                         |                         |                         |              |
| Clinical Frailty Scale | 1.26 (1.10–1.46)         | 0.001                   | 1.97 (1.52–2.54)       | <0.001        |
| 4C mortality score | 1.26 (1.15–1.37)         | <0.001                  | 1.13 (0.99–1.28)       | 0.066        |

\( ^a \) Odds ratios are derived from multivariable logistic regression with early or late mortality as the dependent variable, and per 1-unit increase for continuous variables, or vs. the reference category for categorical variables, plus age and sex as independent variables.
Finally, a sensitivity analysis showed that CFS was significantly associated with both early and late mortality in participants aged >65 years, as well as in younger individuals (Table S14).

Discussion

The main findings of the present study are as follows: (i) we show that late mortality, that is, in patients who survived the acute COVID-19 infection, is not dependent on any of the characteristics of the disease course (inflammatory markers, clinical signs, and symptoms), but is only dependent on the premorbid status of the patient as quantified by CFS, and (ii) although 4C mortality score was indeed the best prognostic score for 28 day mortality, however, it should not probably be used for risk stratification of COVID-19 survivors, as its prognostic value for late mortality was limited and significantly inferior to CFS.

Increasing evidence supports increased long-term morbidity and mortality in COVID-19 beyond the acute infection phase, though the epidemiology of post-COVID syndrome remains poorly defined. A recent study examining 47,780 patients who had survived acute COVID-19 showed that discharged patients had 8 times higher risk of death (increasing to 14 times higher risk for individuals aged <70 years) compared with a general population control group matched for age, sex, ethnicity, and major co-morbidities. Major complications among these patients included new-onset organ dysfunction such as diabetes, kidney and liver disease, and especially major adverse cardiovascular events. While data
Multivariable analysis of predictors for early (28 day) mortality

| Clinical characteristics | OR (95% CI) | P-value |
|--------------------------|-------------|---------|
| Clinical Frailty Scale   | 1.32 (1.13–1.55) | 0.001   |
| Age >65 years            | 3.31 (1.48–7.42)  | 0.004   |
| Shortness of breath      | 1.46 (0.81–2.61)  | 0.206   |
| Myalgia                  | 1.06 (0.46–2.45)  | 0.883   |
| Confusion                | 1.58 (0.88–2.86)  | 0.127   |
| RR >20                   | 2.01 (1.19–3.34)  | 0.009   |

(terms per minute)
- Hypoxia (SpO2 < 92%)
  1.65 (0.95–2.85)  0.074
- Congestive heart failure
  2.13 (1.17–3.90)  0.014
- Chronic kidney disease
  1.21 (0.71–2.06)  0.481
- Hypertension
  1.76 (1.06–2.91)  0.028
- Dementia
  1.02 (0.54–1.93)  0.948

(a) Clinical characteristics
(b) Laboratory parameters
- Clinical Frailty Scale
  1.33 (1.16–1.53)  <0.001
- WBCs < 4 or WBCs > 12 (× 10⁹/L)
  - Neutrophils (× 10⁹/L)
    1.07 (0.99–1.16)  0.103
  - Urea (mmol/L)
    1.00 (0.96–1.05)  0.843
  - eGFR (mL/min/1.73 m²)
    0.986 (0.973–0.999)  0.031
  - ALP (IU/L)
    1.002 (0.999–1.005)  0.168
  - CRP (mg/L)
    1.100 (1.001–1.007)  0.007
  - Albumin (g/L)
    0.98 (0.93–1.04)  0.484
  - Sodium (mmol/L)
    1.041 (0.998–1.085)  0.060

ALP, alkaline phosphatase; CI, confidence interval; CRP, C-reactive protein; eGFR, estimated glomerular filtration rate; OR, odds ratio; RR, respiratory rate; WBCs, white blood cells. Values in bold denote statistically significant results.

Odds ratios are derived from multivariable logistic regression with early mortality as the dependent variable, and CFS as well as (a) all clinical features, or (b) all laboratory parameters associated at univariable analysis with early mortality, as independent variables.

Our study verifies the significance of COVID-19-related risk factors for in-hospital mortality and validates the strong prognostic value of age and the 4C mortality score in the acute stage of disease. Respiratory dysfunction, strong acute inflammatory response, and renal impairment, along with major co-morbidities, were the strongest age-independent and sex-independent predictors of early death in line with current evidence in literature. On the other hand, we show that late mortality is not associated with the course of acute COVID-19, which is further consolidated by the good clinical status of patients at the time of discharge. Baseline CFS score was the most significant prognostic factor for late mortality, supporting that late mortality is more strongly associated with premorbid performance status rather than specific COVID-19-related characteristics or infection course. Of interest, markers of cachexia such as low albumin and haemoglobin were also associated with late mortality independent of age and sex. Interestingly, frailty, malnutrition, and cachexia often co-exist in older individuals; in a recent study examining more than 110 000 older individuals across the UK, frailty was detected in >90% of patients characterized by malnutrition and in 100% of patients characterized by cachexia, while recent studies suggest an inverse relationship between CFS and albumin or BMI. The association of CFS with various tissue loss syndromes (sarcopenia, frailty, cachexia, and malnutrition) as a general marker of baseline patient status is worth exploring in future studies regarding COVID-19 patients and older individuals in general.

A meta-analysis including 23 944 COVID-19 patients showed that frailty is a major risk factor for COVID-19 in-hospital mortality. Frailty, measured with CFS, was associated with two-fold to three-fold increased risk of in-hospital death by COVID-19 independent of age, sex, and major co-morbidities. Apart from in-hospital mortality, frailty has been associated with increased care needs after COVID-19 discharge, while in a recent study, CFS had additive prognostic value over age and SOFA score for cumulative 6 month mortality in 1830 patients with COVID-19. Herein, we expand this observation by showing that CFS is superior to the widely used 4C mortality score for the prediction of late (after day 28) mortality. Of note, the association of CFS with late mortality was independent of age, underlying the
importance of assessing the age-related accumulation of clinical deficits (biological age) rather than chronological age, to develop personalized rehabilitation and monitoring plans.29 A nationwide British study showed that frailty was associated with increased risk of sepsis development following an infection consultation. Of note, severely frail patients aged 55 years had a similar probability of developing sepsis as non-frail 85-year-olds, further underlining the significance of biological vs. chronological age.30 Frail patients (CFS ≥ 5) admitted in ICU with non-COVID suspected infection had two-fold increased risk of in-hospital death, as well as two-fold increased risk of long-term care needs or rehospitalization within 30 days.31 Similarly, a 1.6-fold to 3.5-fold increase in late mortality (31 days to 2 years post-infection) has also been documented in non-COVID-19 sepsis compared with uncomplicated infections or control population, respectively, matched for demographics, quality of life, and co-morbidities.32

The lack of validation cohort is a limitation of our study, along with its retrospective character. However, this is the first worldwide report on predictors of late mortality, results are definite, sample size is adequate because of large number of events, and documentation is complete. Moreover, to increase the robustness of our results (internal validation), we employed resampling techniques and calculated normal-based confidence intervals after bootstrapping with 1000 replicates for (i) coefficients of 4C and CFS in multivariable logistic regression models of early and late mortality and (ii) difference in AUC for 4C and CFS with respect to late mortality, which validated our results. Thus, these results at the least consist a first proof of concept that premorbid status is cardinal for late mortality, strongly supporting the concept of recognition of frail patients at risk for mortality due to COVID-19, and encouraging the application of strategies of aggressive preventive measures in these patients such as booster vaccination and social distancing.

In conclusion, late mortality in our cohort of patients hospitalized with COVID-19 was associated with baseline frailty status, and not with measures of severity of the acute infection phase, in contrast with early mortality. We suggest that clinically frail patients are at especially high risk of poor long-term outcomes and represent a key vulnerable group requiring close medical attention after hospital discharge.

Figure 3 Forest plots showing odds ratios and 95% confidence intervals of independent predictors for early (28 day) and late mortality after adjustment for Clinical Frailty Scale. Odds ratios correspond to 10 year increase in age, 10 breaths per minute increase in respiratory rate (RR), 10 g/L increase in haemoglobin levels, 10 × 10⁹/L increase in number of neutrophils, 100 mg/L increase in reactive protein (CRP), 10 g/L increase in albumin, 10 mmol/L increase in urea levels, 10 mL/min/1.73 m² increase in estimated glomerular filtration rate (eGFR), 10 mmol/L increase in sodium levels, and 10-unit increase in 4C mortality score. OR for white blood cells (WBCs) corresponds to the presence of leukocytosis (>12 × 10⁹/L WBCs) or leukopenia (<4 × 10⁹/L WBCs). X-axis is log₂-transformed for visual clarity. CHF, congestive heart failure; SOB, shortness of breath.
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Conflict of interest

None declared.

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Online supplementary material

Additional supporting information may be found online in the Supporting Information section at the end of the article.
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