Dear Editor,

On behalf of the Italian Society of Internal Medicine Ageno et al. [1] chose to use the database of a registry including 610 patients admitted with COVID-19 to five hospitals in Northern Italy in order to develop and validate a score that at the time of hospital admission would be able to predict a severe disease outcome, defined as the need of non-invasive ventilation, orotracheal intubation or death [1]. In the frame of their analysis meant to build the score and its accompanying validation the authors ultimately produced a score that, made of six clinical and laboratory variables, was able to identify patients at relatively low risk of a severe outcome who could be handled in the context of the low intensity of care offered by internal medicine wards [1]. During the selection and development of the best predictive score they chose to evaluate for inclusion data on demographic variables (age, sex, and time elapsed from disease onset), a few comorbidities (hypertension, diabetes, coronary artery disease, chronic obstructive pulmonary disease) but also an array of laboratory values obtained at admission, such as the white blood cells counts (WBC) (split by neutrophil and lymphocyte absolute counts), C-reactive protein (CRP), alanine amino transferase (ALT), aspartate amino transferase (AST), albumin, creatinine, D-dimer, and the neutrophil/lymphocyte ratio. Of these laboratory values CRP, AST, D-dimer, and the neutrophil/lymphocyte ratio were ultimately included in the actual score, with added clinical and demographic variables such as patient age and history of coronary artery disease [1].

In the context of our analysis of the database of another registry [2] implemented during the first wave of COVID-19 from February 2 to May 31 2020 in patients mainly admitted to two research hospitals acting as hubs in the metropolitan area of Milan (the capital city of Lombardy, the Italian region that was earlier and more severely hit by the SARS-CoV-2 pandemic) laboratory tests were carried out on admission in 1018 patients hospitalized with pneumonia and positive for SARS-CoV-2 RNA by RT PCR. After triage in the emergency room these patients were handled in the low- or intermediate care wards of pneumology, infectious disease or internal medicine. For our registry, we chose the laboratory tests with the main goal to ascertain the organs and body systems more compromised on admission in order to tailor on the basis of the corresponding results the type and intensity of medical support to organs and body systems other than the respiratory system. The latter, the principal one to be compromised by COVID-19, was supported by means of continuous positive airway pressure (CPAP), non-invasive ventilation (NIV) or high flow nasal cannula in patients admitted to pneumology wards, whereas those with relatively milder lung involvement were preferentially...
admitted to infectious disease and internal medicine wards, on oxygen supplementation provided mainly by means of nasal canula, ventimask and reservoir, more seldom by CPAP/NIV. The following tests were carried out: a complete blood count, including hemoglobin (Hb), WBC (split by neutrophil and lymphocyte absolute counts) and platelets. Moreover, to assess the degree of systemic inflammation, serum CRP, ferritin and interleukin 6 (IL 6) were measured. Non-specific markers of tissue damage such as lactate dehydrogenase (LDH), ALT, and AST were also evaluated. The coagulation system was broadly explored by means of the prothrombin time international normalized ratio (INR) and D-dimer, the kidney function through the measurement of serum creatinine and the plasma electrolytes sodium and potassium. Finally, blood glucose was measured as an index of metabolic competence.

Having obtained at admission this battery of laboratory values we chose to describe and discuss herewith the results obtained in the patients who survived (n = 798) compared with those who died in hospital (n = 220) (Table 1). This comparison was carried out first by means of logistic regression adjusted for sex and age but then also by multivariable logistic regression adjusted for all comorbidities in patients who died (P < 0.001) whereas the WBC and neutrophils were increased (P < 0.001), but the significance of the difference for Hb was lost in the fully adjusted multivariable analysis (Table 1). LDH was increased as non-organ specific marker of tissue damage. Regarding the tests exploring blood coagulation, the prothrombin time INR was higher (and thus more abnormal) in non-survivors (P = 0.013 after full adjustment). The mean values of D-dimer almost doubled in patients who ultimately died in hospital, a statistically significant difference from survivors maintained after full adjustment. Markers of systemic inflammation (CRP, ferritin and IL 6) were much higher in fatality cases, the statistical significance of the difference maintained after full adjustment (P < 0.02 for both). Pertaining to markers of renal function, serum creatinine was higher even after full adjustment in patients who died (P < 0.001) but there was no difference for potassium and a small difference for sodium (P = 0.033). Blood glucose was higher in non-survivors.

These laboratory results obtained at hospital admission in patients with COVID-19 can be discussed with the results obtained by Ageno et al. [1] in the context of both their derivation and validation cohorts, even though they did not use laboratory markers to specifically evaluate the condition of body organs and systems, but to build up the prediction score. Moreover, they did not compare survivors with non-survivors, but patients with severe with those with non-severe COVID-19. Notwithstanding these differences their baseline results reported in their Table 1 [1] are broadly similar to those obtained in the present registry.

| Laboratory test, median (1st–3rd quartiles) | Survivors (N=798) | Non-survivors (N=220) | p value (age and sex adjusted) | p value (fully adjusted) |
|--------------------------------------------|-------------------|-----------------------|-------------------------------|--------------------------|
| Haemoglobin (g/L)                          | n=739             | 13.0 (11.7–14.2)      | n=201                        | 11.9 (10.3–13.2)         | < 0.001                   | 0.234                     |
| Lymphocytes (mm³)                          | n=682             | 1055 (770–1450)       | n=186                        | 790 (550–1030)           | < 0.001                   | < 0.001                   |
| Neutrophils (mm³)                          | n=684             | 4475 (3020–6520)      | n=187                        | 5960 (4055–9810)         | < 0.001                   | < 0.001                   |
| WBC (mm³)                                  | n=737             | 6180 (4770–8330)      | n=202                        | 7475 (5530–10,840)       | < 0.001                   | < 0.001                   |
| Platelets (mm³)                            | n=739             | 230,000 (170,000–291,000) | n=201               | 196,000 (137,000–263,000) | 0.052                    | 0.096                     |
| PT (INR)                                   | n=563             | 1.1 (1.0–1.2)         | n=150                        | 1.2 (1.1–1.3)            | 0.004                    | 0.013                     |
| D-dimer (mg/L)                             | n=434             | 855 (471–1656)        | n=119                        | 1493 (953–3195)          | < 0.001                   | 0.006                     |
| LDH (IU/L)                                 | n=423             | 274 (215–356)        | n=119                        | 405 (293–512)            | < 0.001                   | < 0.001                   |
| AST (IU/L)                                 | n=377             | 46 (30–74)            | n=92                         | 55 (38–92)               | 0.194                    | 0.118                     |
| ALT (IU/L)                                 | n=610             | 37 (22–59)            | n=180                        | 30 (18–46)               | 0.663                    | 0.606                     |
| C reactive protein (mg/L)                  | n=654             | 6.1 (2.7–11.3)        | n=183                        | 11.9 (7.0–18.4)          | < 0.001                   | < 0.001                   |
| IL-6 (pg/mL)                               | n=154             | 25.2 (9.8–61.6)       | n=35                         | 75.9 (37.8–137.4)        | 0.004                    | 0.016                     |
| Ferritin (ng/ml)                           | n=318             | 754 (348–1397)        | n=96                         | 843 (450–1544)           | 0.119                    | 0.013                     |
| Creatinine (mg/dL)                         | n=718             | 0.9 (0.7–1.1)         | n=202                        | 1.2 (0.9–1.9)            | < 0.001                   | < 0.001                   |
| Sodium (mmol/L)                            | n=710             | 139 (137–141)         | n=193                        | 140 (136–144)            | 0.004                    | 0.033                     |
| Potassium (mmol/L)                         | n=699             | 4.1 (3.8–4.4)         | n=187                        | 4.2 (3.7–4.6)            | 0.116                    | 0.306                     |
| Glycemia (mg/dL)                           | n=612             | 105 (92–125)          | n=174                        | 113 (93–152)             | < 0.001                   | 0.009                     |

Fully adjusted: age, body temperature, heart rate, respiratory rate, systolic BP, diastolic BP, number of comorbidities, number of drugs, dementia, chronic heart failure, cancer, diabetes, COPD, asthma, previous myocardial infarction, previous stroke/TIA, atrial fibrillation, malnutrition
Our patients who died had higher WBC values, with a very high neutrophil to lymphocyte ratio owing to a decrease in lymphocytes. Inflammation markers such as CRP but also ferritin and IL6 were increased in non-survivors and this was also the case for creatinine, D-dimer and the PT INR, indicating that beside a marked inflammatory state COVID-19 is often associated with an impairment of renal function and activation of the coagulation system. The laboratory results obtained in the context of the present registry can also be compared with those obtained in China at the time of the first 2020 wave of COVID-19, because Zhou et al. [3] compared like us 137 hospital survivors with 54 non-survivors. Like us they found that the lymphocyte count was significantly lower in non-survivors and that serum levels of a non-specific marker of tissue damage, such as LDH, were elevated.

Limitations of this analysis are that the laboratory markers were evaluated only at the time of hospital admission with no follow-up except for information on survival in hospital. However, a strength is the large sample of survivors and non-survivors. From these data obtained at the time of the first wave of COVID-19 in the most heavily and earlier hit region of Italy, we learnt that COVID-19 is a multiorgan disease, and that kidney function, glucose metabolism and the coagulation system must be closely monitored in patients presenting on admission with laboratory markers of organ involvement.

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