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Imbalance of von Willebrand factor and ADAMTS13 axis is rather a biomarker of strong inflammation and endothelial damage than a cause of thrombotic process in critically ill COVID-19 patients

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Abstract
Background: Critically ill patients with coronavirus disease 2019 (COVID-19) are prone to developing macrothrombosis and microthrombosis. COVID-19 has been reported to be rarely associated with thrombotic microangiopathies. A disintegrin and metalloprotease with thrombospondin type I repeats, member 13 (ADAMTS13) severe deficiency, the hallmark of thrombotic thrombocytopenic purpura (TTP), induces the formation of platelet, unusually large von Willebrand factor (VWF) multimer microthrombi. In immune-mediated TTP, ADAMTS13 adopts specifically an open conformation. The VWF/ADAMTS13 couple may contribute to the microthrombi formation in pulmonary alveolar capillaries in COVID-19.

Objective: To investigate clinical features, hemostatic laboratory parameters, VWF/ADAMTS13 axis, and ADAMTS13 conformation in critically ill COVID-19 patients at admission.

Methods: Fifty three critically ill COVID-19 patients were enrolled between March 18 and May 9 2020 in a monocentric hospital.

Results: The median age was 59 years and the male-to-female ratio was 2.8/1. We reported seven pulmonary embolisms and 15 deaths. Biological investigations showed increased fibrinogen and factor V levels, and strongly increased D-dimers correlated with mortality. No patient presented severe thrombocytopenia nor microangiopathic hemolytic anemia. An imbalance between high VWF antigen levels and normal or slightly decreased ADAMTS13 activity levels (strongly elevated VWF/ADAMTS13 ratio) was correlated with mortality. Three patients had a partial quantitative deficiency in ADAMTS13. We also reported a closed conformation of ADAMTS13 in all patients, reinforcing the specificity of an open conformation of ADAMTS13 as a hallmark of TTP.
1 | INTRODUCTION

ADAMTS13 (a disintegrin and metallopeptase with thrombospondin type I repeats, member 13) is the specific cleaving protease for von Willebrand factor (VWF), a multimeric glycoprotein released in blood from endothelial cells and that mediates platelet adhesion and aggregation. Under physiological conditions, ADAMTS13 regulates the size of VWF multimers to prevent platelet-rich thrombi formation in the blood microvessels and it circulates in a closed conformation through a CUB-spacer (complement component C1r/C1s, Uegf, and bone morphogenetic protein 1) domains interaction, which is temporarily disrupted upon binding to VWF. The pathologic development of autoantibodies to ADAMTS13 induces a severe ADAMTS13 deficiency (activity $< 10$ IU/dl), leading to the accumulation of ultra-large VWF multimers and subsequent systemic microvascular thrombosis that causes a specific thrombotic microangiopathy (TMA) named immune-mediated thrombotic thrombocytopenic purpura (iTT). Moreover, specifically in acute iTT, ADAMTS13 adopts a sustained open conformation, likely mediated by ADAMTS13 autoantibodies. In addition, bacterial and viral infections are triggers of TTP but their role on ADAMTS13 conformation is not known yet.

Interestingly, a high frequency of both macrovascular and microvascular thrombosis affecting lungs, kidneys, heart, and potentially complicated by multorgan failure has been reported in prospective cohort studies led in critically ill patients with coronavirus disease 2019 (COVID-19). This novel coronavirus, termed severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), binds to angiotensin-converting enzyme 2, highly expressed in lung alveolar cells but also in the vascular endothelium. The aggression of the endothelial host cells by SARS-CoV-2 explains both the endotheliopathy and the thromboinflammatory processes supporting thrombotic microvascular events.

Since 2020, to further understand the pathophysiology for SARS-CoV-2-related microthrombosis, several studies have focused on the VWF/ADAMTS13 axis in COVID-19 patients admitted on general wards and/or intensive care units (ICUs). In summary, most results converge to an imbalance between VWF and ADAMTS13 showing markedly increased VWF antigen levels and VWF collagen-binding capacity but normal or slightly decreased ADAMTS13 activity, resulting in the formation of large VWF multimers with the latter biomarkers sometimes correlating with mortality. This VWF/ADAMTS13 imbalance may support the concept of a secondary TMA-like syndrome potentially present in some critically ill COVID-19 patients and contributing to the microthrombi formation in pulmonary alveolar capillaries. Surprisingly, however, $-20$ TTP cases likely triggered by a SARS-CoV-2 infection were recently reported. Also, the link between a TMA process and the COVID-19-associated microthrombosis may be supported by the clinical improvement and the reduction of circulating thromboinflammatory markers (i.e., the VWF/ADAMTS13 ratio) induced by plasma exchange undertaken in some critically ill COVID-19 patients. One ex vivo study suggested that purified recombinant ADAMTS13 should be considered as a potential therapy for COVID-19 patients. To date however, no study has investigated ADAMTS13 conformation in COVID-19 patients, although infectious triggers are candidates to explain the initial disruption of the interaction between self-domains of ADAMTS13 leading to its switch from a folded to an open conformation.

In the current study, our major aim was thus to investigate hemostatic laboratory parameters and plasmatic ADAMTS13 features including ADAMTS13 conformation in critically ill COVID-19 patients at ICU admission.

2 | PATIENTS AND METHODS

2.1 | COVID-19 patients

The protocol was approved by the local institutional ethical committee. All consecutive patients admitted for acute respiratory distress syndrome (ARDS) by SARS-CoV-2 in the ICU of Saint-Louis hospital
(AP-HP, Paris, France) between March 18 and May 9, 2020, were enrolled. Real-time quantitative PCR assay for SARS-CoV-2 RNA on nasopharyngeal swab specimens was positive in all patients. Clinical data and outcome were analyzed from medical records.

2.2 Biological investigations

The following biological parameters were measured at admission: hemoglobin, platelet count, lactate dehydrogenase (LDH), fibrinogen, D-dimers, and factor V in the context of care. Measurement of VWF antigen (VWF:Ag) and ADAMTS13 investigation were also performed on citrated plasma samples collected at admission and stored at −80°C until use: VWF:Ag was measured with an ELISA (Asserachrom VWF:Ag), ADAMTS13 activity with FRETS-VWF73 assay (Institute Inc), anti-ADAMTS13 IgG titration with ELISA (TECHNOZYM ADAMTS-13 INH, Technoclone), ADAMTS13 antigen with a homemade 3H9-ELISA, and ADAMTS13 conformation with a homemade 1C4-ELISA, before and after incubation with the anti-CUB1 antibody 17G2 inducing an open conformation in ADAMTS13 (defined by a conformation index >0.5).2,3

2.3 Statistical analysis

Quantitative variables were expressed as medians [25th, 75th percentiles] (minimum-maximum) and categorical variables as numbers and percentages. Comparisons were performed using Mann-Whitney and Fisher’s exact tests, as appropriate, using GraphPad Prism v8.4.2 software (GraphPad Software). Any p values <0.05 were considered significant.

3 RESULTS AND DISCUSSION

Characteristics and outcome of the 53 patients enrolled are shown in Table 1. The median age at admission was 59 years [53, 66] and the male-to-female ratio was 39/14 (73.6% of men). Forty-three patients (81.1%) had at least one risk factor for cardiovascular disease such as overweight (body mass index ≥ 25–29; n = 18, 34.0%) and obesity (body mass index ≥ 30, n = 16, 30.2%), dyslipidemia (n = 11, 20.7%), diabetes (n = 13, 24.5%), and hypertension (n = 24, 45.3%). The overall in-hospital mortality rate was 28.3% (n = 15) and was higher in males (n = 13/39, 33%). Pulmonary embolism (PE) was diagnosed by computed tomography pulmonary angiography in seven patients (13.2%) with no history of thromboembolic event, a male-to-female ratio of 4/3, and associated with death in four cases (57% of patients with PE).

At ICU admission, three patients had a moderate thrombocytopenia over 80 × 10^9/L and 10 patients had an anemia with a hemoglobin level <10 g/dL without hemolysis. No patient had clinical nor biological features of TMA syndrome (platelet counts: 207 × 10^9/L [168, 275]; hemoglobin levels: 11.9 g/dL [10.2, 13.3]). All patients presented elevated LDH levels (844 U/L [699, 939]) and a marked increase of fibrinogen (6.68 g/L [5.87, 7.60]; >8 g/L in 9 patients [17%]), factor V (140% [118, 154]; >120% in 31 patients [59%]) and D-dimers (1120 ng/ml [770, 2840]; >3000 ng/ml in 13 patients [25%]) (Table 1).28 Interestingly, nonsurvivor patients had significantly higher LDH, IL-6, and D-dimer concentrations when compared with survivor patients (LDH: 909 U/L [853, 1000] vs. 806 U/L [660, 870], p = 0.026; IL-6: 113 pg/ml [79.8, 299.5] vs. 81 pg/ml [38.5, 109.0], p = 0.031; D-dimers: 2310 ng/ml [1125, 3655] vs. 985 ng/ml [675, 2185], p = 0.018) (Table 1). These results are the most consistent biological features with intense inflammatory response and hypercoagulability in COVID-19. However, no difference was reported when comparing PE with non-PE COVID-19 patients (data not shown). Our results, especially highly elevated D-dimer levels predictive of mortality, are in line with those previously reported in critically ill COVID-19 patients at admission.29

VWF:Ag levels were markedly increased in both survivor and nonsurvivor patients (326 [284, 378], vs. 416 IU/dl [355, 554], respectively) compatible with severe endothelial damage, but with borderline significant difference (p = 0.05) (Table 1). In our 53 critically ill COVID-19 patients, ADAMTS13 activity ranged from 39 to 150 IU/dl (108 IU/dl [79, 132]), without detectable anti-ADAMTS13 IgG (<15 U/ml in all patients), ADAMTS13 antigen ranged from 0.453 to 2.030 μg/ml (1.125 μg/ml [0.887, 1.269]), and only three patients (6%) had a partial quantitative deficiency in ADAMTS13 (both ADAMTS13 activity <50 IU/dl [42, 39, and 39 IU/dl] and ADAMTS13 antigen <0.50 μg/ml), ruling out TTP diagnosis (Figure 1). No significant difference of either VWF or ADAMTS13 was identified between patients with PE and those with no PE (data not shown). In contrast, interestingly, median ADAMTS13 activity was significantly lower in nonsurvivor patients when compared with survivor patients (85 vs. 115 IU/dl, p = 0.026) (Figure 1). As a consequence, median VWF:Ag/ADAMTS13 activity ratio was strongly elevated in nonsurvivors (4.94 [3.13, 7.21] vs. 3.01 [2.47, 3.69], p = 0.025) and was overall increased in the 53 patients (3.18 [2.59, 4.88], normal range 0.5–2). In contrast, ADAMTS13 antigen levels were not significantly different between survivors and nonsurvivors (1.127 vs. 1.020 μg/ml, p = 0.025) (Table 1). In our 53 critically ill COVID-19 patients at admission.30

Interestingly, in all 53 COVID-19 patients, ADAMTS13 was not captured by the monoclonal antisapser ADAMTS13 antibody 1C4, defining a closed conformation of ADAMTS13 (conformation index <0.5). As expected, in a control experiment, addition of the monoclonal anti-CUB1 ADAMTS13 antibody 17G2 was able to induce an open conformation of ADAMTS13 in vitro in all 53 patients (Figure 2).

In this monocentric retrospective study, we confirm that SARS-CoV-2 infection leads to severe inflammatory response and hypercoagulability with markedly increased fibrinogen, factor V, D-dimers, and IL-6 levels in critically ill COVID-19 patients at ICU admission, with both latter biologic parameters correlating with mortality.9–12 Endothelitis lesions were highly suggested by increased VWF levels. Previous studies supported a potential link...
| Parameters                              | Methodology       | Reference Values          | COVID-19 Patients (n = 53) | Survivors (n = 38, 71.7%) | Nonsurvivors (n = 15, 28.3%) | p    |
|----------------------------------------|-------------------|---------------------------|---------------------------|---------------------------|-----------------------------|------|
| Age (y)                                | Medical records   | 59 [53, 66] (29–76)       | 60 [50, 64] (29–76)       | 58 [56, 67] (49–70)        | 0.77                        |      |
| Gender                                 | Medical records   | 14F / 39 M                | 12F / 26 M               | 2F / 13 M                 | 0.18                        |      |
| BMI (kg/m²)                            | Medical records   | <25                       | 28 [24, 31] (20–47)      | 28 [25, 32] (20–47)       | 28 [25, 29] (20–37)         | 0.66 |
| ≥2 cardiovascular risk factors         | Medical records   | n = 22 (42%)              | n = 15 (39%)             | n = 7 (47%)               | 0.63                        |      |
| Pulmonary embolism                     | CTPA              | n = 7 (13%)               |                          |                          | 0.07                        |      |
| Time of hospitalization (in days)      | Medical records   | 196 [189, 202] (158–210) | 197 [190, 202] (158–208) | 193 [183, 203] (174–210) | 0.56                        |      |
| Hemoglobin (g/dl)                      | CBC (XN10, Sysmex)| ≥13–17 (5.9–16.3)         | 11.9 [10.2, 13.3] (5.9–16.3) | 12.4 [10.8, 13.3] (5.9–16.3) | 11.6 [10.0, 12.4] (6.1–14.6) | 0.28 |
| Platelet count (×10⁹/L)                | CBC (XN10, Sysmex)| 150–450 (85–635)          | 207 [168, 275] (85–635)  | 214 [179, 275] (85–641)   | 183 [153, 286] (94–635)     | 0.26 |
| LDH (U/L)                              | Enzymatic (Roche Cobas)| 135–225 (380–1597)   | 844 [699, 939] (380–1597) | 806 [660, 870] (519–1597) | 909 [853, 1000] (380–1420)  | 0.03 |
| IL-6 (pg/ml)                           | ECLIA (Roche Cobas)| <7                      | 83.5 [40.5, 150.0] (3.3–2470.0) | 81.0 [38.5, 150.0] (3.3–319) | 113.0 [79.8, 299.5] (14.5–2470.0) | 0.03 |
| Fibrinogen (g/L)                       | Chronometric (STA-FIB Liquid, Stago)| 2–4                               | 6.68 [5.87, 7.60] (3.92–10.47) | 6.68 [5.96, 7.42] (3.92–9.47) | 6.70 [5.74, 7.98] (4.91–10.47) | 0.72 |
| D-dimers (ng/ml)                       | Immunoturbidimetry (STA-Liatest D-Di Plus, Stago)| <500                               | 1120 [770, 2840] (270–20000) | 985 [675, 2185] (270–20000) | 2310 [1215, 3655] (730–20000) | 0.02 |
| Factor V (%)                           | Chronometric (STA-Deficient V, Stago)| 70–120                            | 140 [118, 154] (56–200) | 141 [123, 158] (56–200) | 127 [115 145] (92–195) | 0.21 |
| VWF antigen (IU/dl)                    | ELISA (Asserachrom VWF:Ag, Stago)| 50–150                             | 354 [285, 429] (144–704) | 326 [284, 378] (144–704) | 416 [355, 554] (145–596) | 0.05 |

Note: Quantitative variables are expressed as medians [25th, 75th percentiles], (min-max) and categorical variables as numbers and percentages. Comparisons were performed using Mann-Whitney and Fisher’s exact tests, as appropriate. p-values <0.05 were considered significant. Abbreviations: BMI, body mass index; CBC, complete blood count; CTPA, computed tomography pulmonary angiography; ICU, intensive care unit; VWF, von Willebrand factor.
between severe COVID-19-related microthrombosis and secondary TMA-like syndrome enhanced by partial consumption and/or inhibition of ADAMTS13 by inflammatory cytokines as IL-6, in line with a process similar to that observed in sepsis. In COVID-19, a phenomenon of thrombotic pulmonary capillaritis supporting ARDS could be partially explained by the unbalance between VWF and ADAMTS13 and play a crucial role in short-term prognosis. In our group of 53 critically ill patients with COVID-19, both ADAMTS13 activity and the VWF:Ag/ADAMTS13:activity ratio were correlated to mortality, in agreement with other studies. However, this result needs to be nuanced because ADAMTS13 activity remained in the normal or subnormal range, so it cannot support a strong thrombotic tendency in the pathogenesis of ARDS. Also, none of our patients exhibited diagnostic hallmarks of classic TMA because there is no severe thrombocytopenia or microangiopathic hemolytic anemia. Of course, our study may be limited by its monocentric and its 8-week prospective enrollment. However, when considering our data combined with those of the recent literature, it seems more reasonable to think that both VWF and ADAMTS13 are biomarkers reflecting the severity of the endothelial disease caused by SARS-CoV-2 infection (and collateral prognosis biomarkers) rather than strong pathophysiologic actors of the microthrombotic process of COVID-19-associated ARDS. This interpretation is further supported by our main result showing that SARS-CoV-2 is not an infectious agent able to induce an open conformation of ADAMTS13, a biological feature that could be one of the triggers for the preliminary step to immune-mediated ADAMTS13 deficiency. Altogether, our results also reinforce the specificity of an open conformation of ADAMTS13 as a hallmark of iTTP.

In conclusion, COVID-19-associated TMA is likely mostly restricted to the pulmonary microcirculation. This study suggests that normal or slightly low ADAMTS13 activity and highly elevated VWF are rather biomarkers reflecting strong inflammation and endothelial damage rather than drivers of the thrombotic process reported in COVID-19. Further studies are definitely needed to consider the VWF/ADAMTS13 axis as a rational therapeutic option in severe COVID-19.

ACKNOWLEDGMENTS
This work was partly funded by KU Leuven grant COVID19-Thrombosis in COVID19. We thank Sandrine Benghezal, Sophie Capdenat, Adeline Delton, and Sylvaine Savigny for expert technical assistance.

CONFLICT OF INTEREST
None of the authors has any conflict of interest to declare.
AUTHOR CONTRIBUTIONS
Bérangère S. Joly performed and supervised the experiments, analyzed the data, and wrote the manuscript. Bérangère S. Joly and Michael Darmon performed the statistical analysis. Charlotte Dekimpe was highly involved in the ADAMTS13 conformation experiments, analyzed data, and critically reviewed the manuscript. Elie Azoulay, Michael Darmon, Thibault Dupont, Guillaume Dumas, and Elise Yvin enrolled the patients, provided the clinical data, and critically reviewed the manuscript. Nicolas Beranger performed some experiments and critically reviewed the manuscript. Karen Vanhoorelbeke, Elie Azoulay, and Agnès Veyradier supervised the study and critically reviewed the manuscript. All authors accepted the final version of the manuscript.

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How to cite this article: Joly BS, Darmon M, Dekimpe C, et al. Imbalanced of von Willebrand factor and ADAMTS13 axis is rather a biomarker of strong inflammation and endothelial damage than a cause of thrombotic process in critically ill COVID-19 patients. J Thromb Haemost. 2021;19:2193–2198. https://doi.org/10.1111/jth.15445