Increased number of pulmonary megakaryocytes in COVID-19 patients with diffuse alveolar damage. An autopsy study with clinical correlation and review of the literature.

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Method Article

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Abstract

Megakaryocytes are normally present in the lung where they play a role in platelet homeostasis. The latter are well known to participate in the pathogenesis of lung damage, particularly in acute lung injury. Although megakaryocytes are usually not mentioned as a characteristic histopathologic finding associated to acute pulmonary injury, a few studies point out that their number is increased in the lungs of patients with diffuse alveolar damage. In this autopsy study we have observed a relevant number of pulmonary megakaryocytes in COVID-19 patients dying with acute respiratory distress syndrome. We have studied pulmonary tissue samples of 18 patients most of which died after prolonged disease and use of mechanical ventilation. Most samples showed fibroliprotective or fibrotic diffuse alveolar damage and an increased number of megakaryocytes. In six, thrombi of the pulmonary microcirculation were seen. We compare our findings with previous published autopsy reports, mainly focusing on the description of megakaryocytes. Our patients showed abnormal coagulation parameters with high levels of fibrinogen, D-dimers and variable thrombocytopenia. Since the lung is considered an active site of megakariopoiesis, a prothrombotic status leading to platelet activation, aggregation and consumption may trigger a compensatory pulmonary response. An increased number of pulmonary megakaryocytes suggests and supports a relation with the thrombotic events so often seen in COVID-19. Regardless of its etiology, future studies of patients dying with acute pulmonary injury should include pulmonary megakaryocytes as a histologic variable of interest.

Introduction

Megakaryocytes (MKs) are normally present in the human lung and play a role in platelet homeostasis [1,2]. Originally described by Aschoff in 1893, the interest in pulmonary MKs and their platelet production has raised considerably during the last decade [1-3]. Several studies have shown that platelets participate in lung damage, particularly in acute lung injury [2,3]. MKs are usually not mentioned as a characteristic histopathologic finding associated to acute pulmonary injury [4-7]. However, a few studies and textbooks point out that their number is increased in the lungs of patients with diffuse alveolar damage (DAD), burns, shock or sepsis [8-11]. Inflammatory injury to alveolar epithelium and endothelial cells, resulting in intra-alveolar edema, deposition of fibrin and formation of microthrombi are important pathogenic mechanisms of DAD. There is experimental and clinical evidence that platelets contribute to alveolar damage and repair in acute respiratory distress syndrome (ARDS) and other forms of acute lung injury [2,3]. Similarly, abnormalities in platelet number and function influence the natural history of ARDS [2,3].

Most deaths related to severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2) infection are due to pulmonary damage. Autopsy studies performed on these patients usually reveal DAD in different evolutionary phases [7,12-36]. Although no specific pulmonary pathologic findings have been observed, thrombosis of major vessels and microcirculation are often highlighted. It is well-known that COVID-19 patients often show prothrombotic coagulation abnormalities [37-39], being thrombocytopenia associated with a poor clinical outcome [40]. In this autopsy study we report that pulmonary MKs are a common finding in COVID-19 patients with pulmonary damage. In addition to numerous pulmonary MKs our patients suffered from coagulation abnormalities, including thrombocytopenia. This is not an isolated observation since other autopsy reports on COVID-19 patients mention in their microscopic descriptions an elevated number of pulmonary MKs [29-36]. A prothrombotic status leading to platelet activation, aggregation and consumption may trigger a compensatory response of pulmonary MKs. In our work we compare our autopsy findings with previous reports, mainly focusing of pulmonary MKs. Our series includes analytical parameters related to patient's coagulation status including blood platelet counts. Since the lung is considered an active site of megakariopoiesis, an increased number of pulmonary MKs suggests and supports a relation with the thrombotic events and thrombocytopenia so often seen in severe COVID-19.
Methods

Patient selection

We analyzed pulmonary autopsy specimens from 17 patients who died from respiratory failure caused by SARS-CoV-2 infection. A further infected patient who died because of end-stage malignant lymphoma (but without ARDS) and showed pulmonary COVID-19 involvement was also evaluated (case 8). The study was approved by the Ethics Committee of the University Hospital Gregorio Marañón, Madrid (code: EcoBCOV). All the autopsies were performed after informed consent from the closest relatives. All patients fulfilled the World Health Organization criteria for COVID-19 and presented with fever and acute respiratory symptoms, dyspnea, and hypoxia. All tested positively for SARS-CoV-2 RNA by polymerase chain reaction (PCR) assay at the time of hospital admission. In all patients’ molecular tests for common respiratory viruses and bacteria disclosed negative results. Relevant clinical information (table 1) was retrieved from the electronic files of the different hospitals information systems.

Autopsies and histologic examination

In 11 patients, complete or limited autopsy procedures were performed by pathologists at the Departments of Pathology of University Hospitals Puerta de Hierro and 12 de Octubre. Both medical centers are equipped with autopsy rooms that fulfill the recommendations of the Spanish Society of Pathology for this type of autopsies. In these cases, either complete lungs or extensive samples from the different lobes were obtained. The remaining 7 patients underwent ultrasound-guided minimally invasive autopsies. These were done by anesthesiologists at University Hospital la Princesa. Several needle core biopsies of both lungs and other organs were obtained. The procedure was done using ultrasound guidance. All tissue samples were processed routinely after 10% buffered formalin fixation for at least 48 hours. For a better detection of MKs immunohistochemical evaluation of CD61 was performed in all cases. Initial histologic evaluation was independently done by pathologists from each hospital. Afterwards, they were reviewed together by two expert pulmonary pathologists (CS, JAJH). After the initial histologic analysis further examination was made putting emphasis on the detection of thrombi and MKs. Pulmonary MKs were quantified according to previously published methodology [8,35]. A high value was defined as the presence of more than four MKs per 25 high-power fields (hpf)(x40). We further stratified MK counting: absent or rare (≤4/25 hpf), slightly increased (5-7/25 hpf), moderate (8-11/25 hpf) and abundant (≥12 hpf). As mentioned by Carsana et al [35] each sample was initially inspected at low magnification to identify areas in which MKs were most easily recognizable and then were counted in these areas. MK morphology is so characteristic that mature cells are easily recognizable with hematoxylin and eosin staining. Nevertheless, CD61 immunoeexpression was used for confirmation. The latter can also be expressed by platelets and platelet-rich thrombi so correlation with morphology is essential to avoid errors. In most cases MKs were a remarkable finding and even in core needle biopsies quantification was easy. As seen in figure 1 in many cases there were areas with three or more MKs in a single high-power microscopic field. Apart from MKs no other myeloid or erythroid bone marrow cells were observed. Several other relevant histologic parameters were also evaluated (table 2).

Results

Clinical findings

Table 1 presents the main clinical data of the eighteen patients evaluated. The patients were 10 men and 8 women, with a mean age of 61 years (range: 41-75). Regarding comorbidities, eight had hypertension, eight had diabetes, seven had dyslipidemia, six had current or past malignancies, six were smokers and two had hyperuricemia or gout.
The most relevant initial symptoms were fever, cough, dyspnea or tachypnea and diarrhea. At the time of hospitalization, image studies showed bilateral ground glass infiltrates in seventeen patients.

They were firstly admitted on a regular medical ward and afterwards were derived to an intermediate medical ward or intensive care unit, according to the severity of disease. Except for one patient (case 8) who died of end-stage angioimmunoblastic T-cell lymphoma, all patients were treated with mechanical ventilation. The mean hospital stay was 42 days (range: 22 to 73) and the time on mechanical ventilation 31 days (range: 13 to 56). Laboratory findings during the last week before death are summarized in table 1. Fourteen patients had lymphopenia (lymphocytes count <1,0 x 10^9/L). Abnormal coagulation values were seen in the majority of patients: nine (50%) had thrombocytopenia (< 150 platelets x 10^9/L), eighteen (100%) had high D-dimers (>0,50 mg/ml) and fifteen (83%) had high fibrinogen (>400 mg/mL). Rarely, prolonged levels of international normalized ratio (INR) were observed (4 patients, 22%). All patients were treated with heparin.

**Pulmonary histologic findings**

Main histologic findings are summarized in table 2. Except for patient 8 all cases showed histologic evidence of DAD. The predominant lung pattern was DAD in fibroproliferative or fibrotic stages. The most common pattern of fibrosis was interstitial with occasional doughnut-like areas. One case showed extensive areas of organizing pneumonia with numerous intraalveolar fibroblastic plugs. Intermixed with the fibrotic areas most cases had alveolar septal congestion and varying degrees of intraalveolar hemorrhage, fibrin, edema and desquamation of pneumocytes. Hyperplastic pneumocytes many of which showed dense cytoplasm and large nuclei with prominent nucleoli were also a constant finding. Almost all samples had foci of squamous metaplasia near bronchial or bronchiolar structures. In one case Kuhn’s hyaline was present in the cytoplasm of reactive pneumocytes. We observed no viral inclusions. Multinucleated giant cells were present in 12 of the 18 cases. Three cases had associated areas of bronchopneumonia with numerous neutrophils and focal necrosis. Subpleural fibrotic cysts, most probably due to invasive mechanical ventilation were seen in five cases. Other findings were alveolar corpora amylacea and focal lesions of osseous metaplasia. In patient 8, no established lesions of DAD were present. The lungs showed intense edema, some degree of pneumocyte hyperplasia and focal presence of alveolar fibrin. Thirteen of the 18 patients (72.2%) analyzed showed an increased number of MKs, as previously defined (>4/25 high-power fields) (Figs. 1,2). In patients 2, 7 and 9, MKs were present but their number was below the established threshold. In patient 14 despite a long clinical course and use of mechanical ventilation the core needle tissue sample only revealed areas of exudative DAD with numerous hyaline membranes and no fibrosis or relevant number of MKs. We believe it may be due to a sampling problem and that the more consolidated pulmonary areas were not biopsied. In three cases the increase in MK number was slight (5-7/25 hpf), in five moderate (8-11/25 hpf) and in five abundant (≥12/25 hpf). Thrombi were an evident finding in six cases (Fig. 2). Morphologic detection of small thrombi in the microcirculation can be very difficult and we only considered those cases in which concordance among pathologists was present. It is important to note that all patients were receiving anticoagulant therapy with heparin. Five of the cases in which thrombi were detected corresponded to large pulmonary tissue samples from “open” autopsy procedures. In contrast, thrombi were present in only one of the samples obtained using ultrasound-guided minimally invasive procedures.

**Discussion**

Pulmonary histopathologic changes related to ARDS are usually similar regardless of its etiology [4-7]. This observation can be extended to cases caused by SARS-CoV-2 infection [7]. In 2007 Mandal et al reported abnormalities in platelet homeostasis including an increase in the number of pulmonary MKs in patients with DAD [8]. In their series of 21 patients, those with thrombocytopenia had a worse prognosis. In this autopsy study, we have
shown that MKs are a common finding in the lungs of COVID-19 patients dying with DAD. Our patients showed abnormal coagulation parameters with high levels of fibrinogen, D-dimers and variable thrombocytopenia.

Numerous autopsy studies describing patients dying from COVID-19 have been published [7,12-36]. In seven of them, pulmonary MKs are mentioned in the autopsy reports. Three of these studies relate their presence to the hypercoagulability status so characteristic of these patients [29,30,36]. They describe general autopsy findings and do not quantify MKs or include a correlation with coagulation parameters. Four other reports refer to pulmonary MKs as a relevant finding [31-35]. Carsana et al quantified them revealing an increased number in 33 of their 38 patients [35]. Similarly, these reports describe general findings and the presence of MKs is not further commented. In the remaining autopsy studies, including a review article there are no references to MKs [12-28,41]. Various reasons could explain this absence. As mentioned before, MKs are not a histological variable usually associated to DAD so pathologists may not be tempted to perform a specific search or to report them. Lung MKs are rare and even if their number is increased the counts in absolute terms can still be low. If a specific search is not performed, MKs can easily be overlooked. Because MKs are trapped in the pulmonary microcirculation their morphology differs from that seen in bone marrow. Pulmonary MKs show less cytoplasm and fewer nuclear lobulations. The nucleus tends to be elongated as if adapted to the vessel diameter. Although hematoxylin and eosin stain permits a confident recognition of MKs, their detection can be facilitated by immunohistochemical analysis. CD61 is expressed by MKs, but it can also be expressed by platelets and platelet-rich thrombi [42]. Therefore, to avoid overcounting a close correlation between histology and immunohistochemistry is advisable. Another possible reason for the lack of references to MKs is that some reports describe patients with early or incidental pulmonary lesions but no ARDS [19,20]. In this sense, one of our patients without MKs had pulmonary edema but no definitive histologic findings of DAD. This patient died because of end-stage malignant lymphoma without ARDS. Other reports describing deaths in non-hospitalized patients focus on macroscopic findings, mainly thromboembolic events and although pulmonary histologic findings are mentioned they are not described in-depth. Size tissue sample is another potential limiting factor for the detection of MKs. However, it should be noted that we have easily found them in trucut biopsies. Similarly, the previously mentioned study by Duarte-Neto et al [29] is based on trucut biopsies. Finally, not all patients showed an increase number of MKs. Our series and that of Carsana et al [35] revealed no significant number of MKs in 27.8% and 13.2% of the patients, respectively.

In addition to SARS-CoV-2, two other members of the coronavirus family: SARS-CoV and Middle East respiratory syndrome (MERS-CoV) cause pulmonary injury. The autopsy studies performed describe pulmonary damage consistent with different phases of DAD but do not mention or illustrate pulmonary MKs [43-48]. Their absence could be attributed to a greater tendency to thrombotic events in COVID-19 patients but similar procoagulation abnormalities have been described in SARS and MERS [37].

Regarding pathogenesis, the observed increment in pulmonary MKs may obey to a compensatory response. It is well-known that such responses occur in the bone marrow because of thrombocytopenia. Knowing that the lung is a normal site of megakaryopoiesis it is tempting to believe that the increased number of MKs observed in our patients is, in part, secondary to thrombotic events, platelet activation, aggregation and consumption. In this sense, a recent study shows that COVID-19 significantly alters platelet gene expression, triggering a robust platelet hyperreactivity [38]. In addition, COVID-19 patients have elevated plasma levels of thrombopoietin, a well-known megakaryocyte growth factor [38]. In addition to DAD-related thrombosis of the pulmonary microcirculation, COVID-19 patients have a systemic procoagulatory status.

Finally, another interesting aspect of MK biology concerns its fibrotic capacity. MKs produce transforming growth factor-beta and participate in bone marrow fibrosis [49]. A similar pro-fibrotic capacity has been demonstrated for
pulmonary MKs in an experimental model of lung fibrosis [50]. Precisely, diffuse lung fibrosis is one of the greatest complications of ARDS. Therefore, pulmonary MKs must be considered as potential contributors to fibrosis in this precise context.

**Conclusion**

This study shows that pulmonary MKs are increased in COVID-19 patients with evolved DAD. There are still few pathologic studies relating MKs and acute pulmonary injury but it seems that such increase is a characteristic of DAD regardless of its cause. The relative abundance of pulmonary MKs in COVID-19 patients may reflect a greater prothrombotic tendency in these patients. Future studies of patients dying with acute pulmonary injury should include pulmonary MKs as a histologic variable of interest. Similarly, the review of previous autopsy studies on COVID-19 patients looking for pulmonary MKs may help us to further define their role in the pathogenesis of pulmonary damage.

**Declarations**

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**Conflicts of interest/Competing interests:** The authors have no conflict of interest to declare

**Ethics approval:** The study protocol was approved by the Ethics Committee of University Hospital Gregorio Marañón, Madrid, Spain (code: EcoBCOV)

**Consent to participate/for publication:** For all patients, informed consents were obtained from closest relatives

**Availability of data and material:** All data and material available

**Code availability** (software application or custom code): not applicable

**Authors' contributions:** Mariel F. Valdivia Mazeyra: study design, data analysis, literature review, writing. Clara Salas: study design, data analysis, literature review, editing of text. Jesús M. Nieves-Alonso: ultrasonographic autopsy performance, data analysis. Luz Martín-Fragueiro: data analysis, review of histopathology. Carmen Bárcecena: autopsy performance, review of histopathology, data analysis, editing of the text. Patricia Muñoz-Hernandez: review of histopathology, literature review. Karen Villar: ultrasonographic autopsy performance, review of histopathology. Javier Martín-López: data analysis, review of histopathology. Fernando Ramasco-Rueda: ultrasonographic autopsy performance, data analysis. Javier Fraga: study design, data analysis, review of histopathology, editing of the text. José A. Jiménez-Heffernan: study design, writing, data analysis, review of histopathology.

Mariel F. Valdivia Mazeyra and Clara Salas contributed equally to this work. Javier Fraga and José A. Jiménez-Heffernan share senior authorship.

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Table 1. Main clinical and analytical data

| Parameter                                   | Number of cases (when applicable) | % or range |
|---------------------------------------------|-----------------------------------|------------|
| Sex: male-female ratio                      | 10 - 8                            | -          |
| Age (years)                                 | 61                                | 41 - 75    |
| Length of hospital stay (days)              | 42                                | 22 - 73    |
| Days on mechanical ventilation             | 31                                | 13 - 56    |
| **Comorbidities**                           |                                   |            |
| Hypertension                                | 8                                 | 44%        |
| Diabetes mellitus                           | 8                                 | 44%        |
| Dyslipidemia                                | 7                                 | 39%        |
| Smoker or previous smoker                   | 6                                 | 33%        |
| Malignancy                                   | 6                                 | 33%        |
| Hyperuricemia or gout                       | 2                                 | 11%        |
| **Initial clinical presentation**           |                                   |            |
| Fever                                       | 16                                | 44%        |
| Cough                                       | 8                                 | 89%        |
| Dyspnea / tachypnea                         | 8                                 | 44%        |
| Diarrhea                                     | 6                                 | 33%        |
| **Radiological findings**                   |                                   |            |
| Ground glass infiltrates                    | 17                                | 94%        |
| **Mean laboratory findings during last week before death** | | |
| Lymphopenia (<1.0 x 10^9/L)                 | 14                                | 78%        |
| Low platelets (<150 x 10^9/L)               | 9                                 | 50%        |
| High D-dimers (>0.50 µg/mL)                 | 18                                | 100%       |
| High Fibrinogen (>400 mg/ml)                | 15                                | 83%        |
| Prolonged INR* (>1.30)                      | 4                                 | 22%        |
| Treatment with heparin                      | 18                                | 100%       |

* INR: international normalized ratio

Table 2. Main pulmonary pathologic findings
| Case | Increased megakaryocytes | Predominant DAD phase | Alveolar Fibrin | Fibrosis | Thrombi | MGCs | Other |
|------|--------------------------|-----------------------|----------------|----------|---------|------|-------|
| 1    | Yes (8/25 hpf)           | Fibroproliferative     | Residual HM    | Interstitial | +       | +    | Subpleural cysts, osseous metaplasia, BP foci |
| 2    | No                       | Fibroproliferative     | No             | Interstitial | -       | +    | Pulmonary infarct, Osseous metaplasia |
| 3    | Yes (13/25 hpf)          | Proliferative          | Residual HM    | Interstitial | -       | +    | Osseous metaplasia |
| 4    | Yes (8/25 hpf)           | Fibroproliferative     | No             | Doughnut-like and interstitial | - | + | Osseous metaplasia |
| 5    | Yes (15/25 hpf)          | Fibrotic               | Residual HM    | Interstitial | +       | +    | Osseous metaplasia |
| 6    | Yes (6/25 hpf)           | Fibrotic               | No             | Interstitial, Doughnut-like | + | + | Subpleural cysts, osseous metaplasia |
| 7    | No                       | Fibrotic               | Fibrin "balls" | Interstitial | -       | -    | Subpleural cysts |
| 8a   | No                       | Exudative              | Fibrin "balls" | Interstitial, Doughnut-like | - | + | Prominent edema |
| 9    | No                       | Fibrotic               | Fibrin "balls" | Interstitial | -       | +    | |
| 10   | Yes (10/25 hpf)          | Fibroproliferative     | Fibrin "balls" | Interstitial | +       | -    | Osseous metaplasia, Corpora amylacea |
| 11   | Yes (18/25 hpf)          | Fibrotic               | Fibrin "balls" | Interstitial | +       | +    | Subpleural cysts |
| 12   | Yes (5/25 hpf)           | Fibroproliferative     | Residual HM    | Interstitial | +       | +    | Corpora amylacea, BP foci |
| 13   | Yes (6/25 hpf)           | Fibrotic               | No             | Interstitial | -       | +    | Osseous Metaplasia |
| 14   | No                       | Exudative              | Numerous HM    | No         | -       | -    | |
| 15   | Yes (8/25 hpf)           | Proliferative          | Fibrin "balls" | Minimal Interstitial | - | - | Corpora amylacea, BP foci |
| 16   | Yes (8/25 hpf)           | Fibrotic               | No             | Organizing Pneumonia | - | - | |
| 17   | Yes (15/25 hpf)          | Fibroproliferative     | No             | Interstitial | -       | +    | |
| 18   | Yes (13/25 hpf)          | Fibroproliferative     | Fibrin "balls", HM | Interstitial, Doughnut-like | - | + | |

DAD: diffuse alveolar damage; MGCs: multinucleated giant cells; hpf: high-power fields; HM: hyaline membranes; BP: bronchopneumonia foci; *Only patient who received no mechanical ventilation

**Figures**
Figure 1

(A-D) Pulmonary histologic sections from different COVID-19 patients with diffuse alveolar damage showing a relevant increase in the number of megakaryocytes. In contrast to those of the bone marrow, pulmonary megakaryocytes often show an elongated nuclear morphology, scarce cytoplasm and few nuclear lobulations (hematoxylin-eosin (HE), x600). (E) A megakaryocyte is visible within the lumen of a larger vessel (HE, x600). (F) As expected, megakaryocytes showed immunoexpression of CD61 (immunoperoxidase, x600).
Figure 2

Pulmonary histologic samples from four COVID-19 patients showing thrombosis and megakaryocytes. (A) An intravascular thrombus is visible at the left of the image. In the same high-power field, a megakaryocyte is clearly visible (arrow) (HE, x600). (B) The image reveals an intravascular thrombus at a pulmonary artery (lower right) (HE, x200). The inset highlights the presence of a megakaryocyte in the vicinity. (C,D) In each high-power image intravascular thrombi and megakaryocytes (arrows) are present (HE, x600).