The Impact of Different Lung Ultrasound Protocols in the Assessment of Lung Lesions in COVID-19 Patients: Is There an Ideal Lung Ultrasound Protocol?

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Abstract
Background In the past months, several lung ultrasonography (LUS) protocols have been proposed, mainly on previously validated schemes independent of coronavirus disease 2019 (COVID-19).
Objectives The main purpose of this study was to determine the impact and accuracy of different LUS protocols proposed in COVID-19.
Methods Patients were evaluated with a standard sequence of LUS scans in 72 intercostal spaces along 14 anatomic lines in the chest. A scoring system of LUS findings was reported and then analyzed separately according to each proposed LUS protocol zones. This score was then correlated to a validated Pulmonary Inflammation Index (PII) on chest Computed Tomography (CT).
Results Thirty-two patients were enrolled. The most frequent pattern was ground-glass opacities in the chest X-ray (53.1%), chest CT (59.1%) and subpleural or lobar consolidations (40.8%) in the posteroinferior areas \((p < 0.001)\) on LUS. The Interclass Correlation Coefficient (ICC) was significantly correlated with almost every protocol analyzed except the 8-zone \((p = 0.119)\) and the 10-zone protocol that only included one posterior point \((p = 0.052)\). The highest ICC was obtained with a 12-zone protocol \((ICC 0.500; p = 0.027)\) and decreased as more points were included.
Conclusions In conclusion, our study results suggest that performing an ultrasound protocol with 12-zone scanning, including the superior and inferior areas of the anterior, lateral and posterior regions of the chest was consistent with higher ICC and higher degree of concordance with CT. We emphasize the need of a more standardization technique to further implement and develop this imaging modality in COVID-19.

Keywords Point-of-care ultrasonography (POCUS) · Lung ultrasonography (LUS) · Coronavirus disease 2019 (COVID-19) · Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2)

Introduction

Ten months after the declaration of a global pandemic due to the novel coronavirus disease (COVID-19), more than 100 million confirmed cases and over 2.3 million deaths have been reported globally [1].

These facts enhance the critical need for an accessible, reproducible, low-cost and safe imaging modality for the diagnosis, risk stratification, monitoring and management of COVID-19 patients. Lung ultrasonography (LUS) qualifies for such purpose, providing an evidence-based method which will aid to diagnose and stratify patients, as well as guide therapy and early detect complications [2].

However, contrasting the relevant role of LUS, there is no validated scanning protocol in COVID-19 patients. At the moment, the main recommendation is to use previously validated schemes in other conditions other than COVID-19.

In the past months, several imaging protocols had been proposed, based on the number of areas or points to explore (see Table 1): 8-zone protocol [3], 10-zone protocol [4, 5], 12-zone protocol [6, 7], 14-zone protocol [8], 16-zone...
Table 1: The different lung ultrasonography (LUS) protocols to assess lung abnormalities in COVID-19 patients

| Anatomical line       | 8-zone protocol [3] | 10-zone protocol [5]* | 10-zone protocol [4] | 12-zone protocol [6] | 12-zone protocol [7]* |
|------------------------|---------------------|------------------------|-----------------------|----------------------|-----------------------|
|                        | Right   | Left  | Right   | Left  | Right   | Left  | Right   | Left  | Right   | Left  |
| Anterior               | Parasternal |        |         |       |         |       |         |       |         |       |
|                        | Mid-clavicular   | 2nd and 5th ICS | 2nd and 4th ICS | 3rd ICS | 3rd ICS |       |         |       |         |       |
| Lateral                | Anterior axillary|         |       | 2nd–5th ICS | 2nd–4th ICS | 5th ICS | 4th ICS | 2nd and 5th ICS | 2nd and 4th ICS | 2nd–5th ICS | 2nd–4th ICS |
|                        | Mid-axillary     | 2nd and 5th ICS | 2nd and 4th ICS | 2nd–5th ICS | 2nd–4th ICS | 4th ICS | 4th ICS | 2nd and 5th ICS | 2nd and 4th ICS | 2nd–5th ICS | 2nd–4th ICS |
|                        | Posterior axillary|         |       | 2nd and 5th ICS | 2nd and 4th ICS | 5th ICS | 5th ICS | 2nd and 5th ICS | 2nd and 4th ICS | 2nd–5th ICS | 2nd–4th ICS |
| Posterior              | Sub-scapular     | 7th–10th ICS | 7th–10th ICS | 2nd–5th ICS | 2nd–5th ICS | 4th ICS | 4th ICS | 2nd and 10th ICS | 2nd and 10th ICS | 2nd–10th ICS | 2nd–10th ICS |
|                        | Paravertebral    |         |       | 2nd and 5th ICS | 2nd and 5th ICS | 3rd ICS | 3rd ICS | 2nd and 5th ICS | 2nd and 5th ICS | 2nd–5th ICS | 2nd–5th ICS |
| Anatomical line        | 14-zone [8]      | 16-zone [9]   | 18-zone protocol [10] | 72-zone protocol [11] |
|                        | Right   | Left  | Right   | Left  | Right   | Left  | Right   | Left  | Right   | Leftler |

ICS inter-costal space

*Visualization of several inter-costal spaces in each zone
protocol [9] and 18-zone protocol [10] and 72-zone protocol [11].

Finding the optimal balance between the acquisition time and accuracy represents one of the main challenges. Whenever possible, a lower number of acquisition areas is preferred, reducing the cross-contamination and exposure of healthcare professionals, but this could lead to underestimations.

Additionally, the development of a more standardized approach will allow comparisons and reproducibility across different studies and exams, as well as facilitate research on pattern recognition with artificial intelligence algorithms and telematic applications [8].

To our best knowledge, the formulation of such an approach is in need for more research. We aimed to determine the impact on accuracy of currently proposed LUS scanning protocols, in determining the presence and extension of lung lesions, in patients with confirmed COVID-19.

Patients and Methods

This was a prospective study performed in the emergency department (ED) of an academic hospital, conducted in accordance with the Declaration of Helsinki, and approved by the Research Ethics Committee of our University Hospital. We obtained informed consent from each patient.

Patient Selection

Patients admitted to the ED with a confirmed COVID-19 requiring a chest computed tomography (CT) for evaluation were included. We defined a confirmed case as any patient with clinical symptoms and positive reverse transcription polymerase chain reaction (RT-PCR) or antigen test for the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2).

The main indication for CT was a negative chest X-ray, respiratory failure (oxygen saturation below 92%) or high suspicion of pulmonary embolism (PE).

We excluded patients < 18 years or those who declined to participate. A sample of patients who met these inclusion criteria were enrolled and prospectively studied.

Study Protocol

Patients underwent a chest CT performed by fellowship-trained radiologists together with a LUS, performed by an ultrasound fellowship-trained internal medicine physician performed all the ultrasound exams. Therefore, an opportunity sampling method was implemented for patient selection.

Chest CT Data Collection

Non-contrast chest CT scans were obtained by a multidetector CT (SOMATOM go.Up, Siemens Healthliners, Erlangen, Germany). The examination was performed with the patient in the supine position and at the end of inspiration. Scans were acquired and reconstructed as axial images using the following parameters: 1.5-mm section thickness, 0.7-mm interval, 130 kVp. A low-dose protocol was implemented with a mean CT Dose Index volume (CTDIvol) of 2 mGy.

Blinded to the clinical information, one radiologist trainee with 2 years of experience (KSAV) and four senior radiologists with more than 10 years of experience (MMG, ADT, MLGP, SOV) reviewed all the images.

Images were interpreted using the lung and mediastinum window settings. Each lung was divided into 10 zones based on the anatomy of the lung segments (Fig. 1). The presence and distribution of the following radiological abnormalities were evaluated in each lung segment [12]: ground-glass opacities (GGO), interlobular septal thickening, crazy paving pattern, consolidation, subpleural line, pleural thickening and effusion.

Ultrasound Data Collection

The ultrasound exam was performed with the patient in supine or near-supine position for the anterior scanning, and in the sitting or lateral decubitus position for the posterior scanning.

The probe was positioned obliquely, along the intercostal spaces [13]. The LUS examination was obtained moving the probe along anatomical reference lines (Fig. 2), 2nd–4th intercostal space (ICS) of parasternal, midclavicular, anterior axillary and midaxillary line (on the right side to the 5th ICS), whereas for the posterior chest, the paravertebral (2nd–10th ICS), sub-scapular (7th–10th ICS) and posterior axillary (2nd–10th ICS) lines [11]. A video clip was recorded along each anatomical line, recording at least 3 s at each ICS.

The examinations were performed using a GE VENUE ultrasound system fitted with a phased and curvilinear array transducer (1.5–4.5 MHz) (General Electrics Healthcare, Madrid, Spain). The sonographer was blinded to the patient’s past medical history, vital signs, symptoms, laboratory measurements, CT scan results and therapy.

In our limited experience with COVID-19 patients, it takes less than 10 min to perform LUS, excluding cleaning time [7].

Outcome Measures and Definitions

The main purpose of this study was to determine the impact and accuracy of different LUS protocols and the correlation
with the PII in the chest CT in patients with confirmed coronavirus disease (COVID-19).

According to the evaluation criterion established by Chongqing Radiologist Association of China, the pulmonary inflammation index (PII) was obtained from each patient [14].

$$PII = \frac{\text{distribution score} + \text{size score}}{40} \times 100\%$$

**Distribution score**: scored according to the lesion distribution, one score for each lung segment, and 20 scores for left and right lung.

**Size score**: according to whether the lesion occupied ≥ 50% (score of 1) or < 50% (score of 0).

We developed a numeric Lung Score based on the pathological findings in each lung sonographic area. We differentiated the lung scanning areas (Fig. 2) of each protocol and gave each pathological finding (Table 2) a different score:

Fig. 1 Bronchopulmonary segments (10 per hemithorax). Right lung. Upper lobe: AP (1)—apical; P (2)—posterior; AN (3)—anterior. Middle lobe: L (4)—lateral; M (5)—medial. Lower lobe: S (6)—superior; MB (7)—medial; AB (8)—anterior; LB (9)—lateral; PB (10)—posterior. Left lung. Upper lobe: APP (1–2)—apicoposterior; AN (3)—anterior; SL (4)—superior lingula; IL (5)—inferior lingula.

Fig. 2 Anatomical reference lines of the Lung Ultrasonography (LUS) protocol. Second to fourth intercostal space of parasternal (dashed line), mid-clavicular (solid line), anterior axillary (solid line) and midaxillary (dashed line) (on the right side to the fifth). Second to tenth intercostal space of the paravertebral (solid line) and posterior axillary (dash-dotted) lines. Seventh to tenth intercostal space of the sub-scapular (dashed) line
We summed every area’s points, dividing into the maximum score (i.e., for a 10-zone protocol, the maximum score would be 3 points x 10 zones = 30) and multiplying by 100 obtaining the patient’s standardized Lung Score according to the assessed protocol, ranging from 0 to 100.

Statistical Analysis

Baseline characteristics are presented as mean and standard deviation (SD) for continuous variables and count and proportions for categorical variables.

The interclass correlation coefficient (ICC) was used to assess the degree of agreement between the LUS score of each protocol and the PII on chest CT. A sample of 30 patients was obtained as previously recommended in reliability studies [15]. An ICC of less than 0.50 was considered poor, from 0.50 to 0.75 moderate, 0.75–0.90 was considered good and 0.90–1 excellent [15].

We assumed an α-value of 0.05 for two-sided hypothesis testing. Analyses were conducted with the statistical IBM SPSS software v25.0 (SPSS Inc., Chicago, IL, USA).

Results

A total of 32 patients were enrolled between September and October 2020 (summarized in Table 3). The mean age was 67.9 years (SD 19.5) and 59.4% were female. 18 patients (56.2%) had an underlying comorbidity. They had an average of 8.03 days (SD 4.14) after symptom onset. The patients were normotensive and had low oxygen saturation (92.3%—SD 4.1, with a respiratory rate of 25.6 rpm—SD 1.2). The mean lymphocyte count was $1.21 \times 10^9$ (SD 0.52, normal value—NV: 1.0–4.0), CRP was 83.7 mg/dL (SD 87.2, NV: 0–10), d-dimer 2444 ng/mL (SD 4156, NV: < 400) and LDH was 312.8 U/L (SD 87.2; NV: 140–280) at admission.

Regarding the imaging studies, all the included patients had a chest X-ray, CT and underwent a LUS study (see Table 4 and Fig. 3). The most frequent pattern was ground-glass opacities (GGOs) in the chest X-ray (53.1% [17 patients out of 32]) and chest CT (59.1% [378 lobes affected out of 640 explored]). About the LUS, the most common findings were subpleural or lobar consolidations (40.8% [940 ICS out of 2304 explored]) in the posteroinferior areas ($p < 0.001$) (see Fig. 1).

The average PII value was 43.8% (SD 19.9). Interclass Correlation analysis (see Table 5) showed that the PII was significantly correlated with almost every protocol analyzed except the 8-zone ($p = 0.119$) and the 10-zone protocol with only one posterior point scanned ($p = 0.052$). The highest ICC was obtained with a 12-zone protocol (ICC 0.500; $p = 0.027$) and decreased as more scanning points were included.

Discussion

Easy to access and reliable diagnostic methods which can accurately guide management in COVID-19 are vital in non-hospital settings and areas with limited resources. Some studies start to point out that LUS could be a first-line diagnostic tool alternative to conventional chest X-ray and CT scan, including the critically ill patients, where LUS score had been already been proven to be useful [16]. Moreover, since there is no exposure to ionizing radiation, can be considered in vulnerable population such as pregnant women and children [17, 18].

In this pandemic context, a useful proposal for LUS standardization in COVID-19 was reported, defining different parameters such as the acquisition time, the type of

**Table 2** Definition and interpretation of the main findings on Lung Ultrasonography in COVID-19

| LUS finding                  | Definition                                                                 |
|-----------------------------|---------------------------------------------------------------------------|
| A-lines                     | Horizontal reverberation artifacts parallel to the pleural line            |
| B-lines                     | Hyperechoic vertical artifacts that arise from the pleural line, extending to the bottom of the screen without fading that erases the A-line artifact |
| Confluent                   | Multiple converging or coalescent B-lines                                  |
| Isolated                    | Discrete, well demarcated B-lines                                         |
| Irregular Pleural Line      | Indented or broken pleural line                                           |
| Pleural effusion            | Anechoic space between the parietal and visceral pleura                   |
| Subpleural consolidation    | Hypoechoic areas smaller than 1 cm in diameter, surrounded by a hyperechoic artifact tail |

LUS lung ultrasound
transducer, the imaging settings and the scanning points [8], mainly based on expert’s recommendation, with scarce adoption.

Subsequently, several imaging protocols had been proposed, based on the number of areas or points to explore (see Table 1), complicating the reproducibility of the results and research.

In our study, we selected PII as the CT score index comparator. PII is a validated index that in previous studies significantly correlated with the degree of pulmonary inflammation and the main clinical symptoms and laboratory results [14]. This would more appropriately address the pathophysiology of the disease, beyond the anatomic abnormalities, at a cost of weaker correlations than reported previously in the literature [6].

Our findings are comparable to a recently published study on this matter [19]. In this multicentric study, they analyzed the outcome of 3 different LUS imaging protocols (4, 8 and 12-zone acquisitions) from 88 patients, comparing it to the

| Table 3 | Demographics and clinical characteristics of patients included (N=32) |
|---|---|
| Gender (female)—N (%) | 19 (59.4) |
| Age (years) mean (SD) | 67.9 (19.5) |
| Caucasian—N (%) | 18 (56.3) |

| Past medical history—N (%) |
| Pulmonary disease | 5 (15.7) |
| Diabetes Mellitus | 6 (18.8) |
| Hypertension | 13 (40.6) |
| Dyslipidemia | 13 (40.6) |
| Obesity | 3 (9.4) |
| Heart disease | 10 (31.3) |
| Previous thromboembolic disease | 3 (9.4) |
| Dementia | 4 (12.5) |

| Mean days from Symptom Onset | 8.03 (4.14) |

| Table 4 | Imaging modalities (chest computed tomography, X-ray and Point-of-care ultrasound) findings of patients included (N=32) |
|---|---|

| Chest computed tomography |
| Total zones (10 per hemithorax, 20 per patient, N=640) | 640 (100) |
| Normal zones (N=640) | 177 (27.7) |
| Abnormal findings |
| GGO (N=640) | 378 (59.1) |
| Consolidation (N=640) | 247 (38.6) |
| Pleural effusion (N=32) | 5 (15.6) |
| Subpleural line (N=640) | 99 (15.5) |
| Crazy paving pattern (N=640) | 39 (6.1) |
| Interlobular septal thickening (N=640) | 27 (4.2) |
| Atelectasis (N=640) | 7 (1.1) |
| Pleural thickening (N=640) | 3 (0.5) |
| PII % (SD) | 43.8 (19.9) |

| Chest X-ray—N (%) | 32 (100) |
| Normal | 4 (12.5) |
| Abnormal findings |
| Ground-glass opacity (GGO) | 17 (53.1) |
| Interstitial pattern | 11 (34.4) |
| Bilateral | 23 (71.9) |
| Unilateral | 5 (15.6) |

| Point-of-care ultrasonography (POCUS) results |
| Total ICS explored (32 patients, 72 ICS per patient, N=2304) | 2304 (100) |
| Normal zones (N=2304) | 535 (23.2) |
| Abnormal findings |
| Consolidation (N=2304) | 940 (40.8) |
| Pleural effusion (N=32) | 10 (31.3) |
| Confluent B-lines zones (N=2304) | 502 (21.8) |
| Irregular pleural line or isolated B-lines zones (N=2304) | 327 (14.2) |

| GGO ground-glass opacity, ICS intercostal space, PII: Pulmonary Inflammation Index, POCUS point-of-care ultrasonography, SD standard deviation |

ALT alanine transaminase, ICU intensive care unit, LDH lactate dehydrogenase, LMWH low molecular weight heparin, rpm rate per minute, SD standard deviation, WBC white blood cell

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reference LUS protocol (14-zone). The level of agreement of each protocol where 57% in the 8-zone, 70% in the 10-zone and 99% in the 12-zone. They concluded that a 12-zone scanning protocol had a similar performance comparable to the reference protocol (14-zone).

Similar to previously reported by Smargiassi [20], we found that limiting our exam to the anterior and lateral regions (8-zone) leads to an underestimation, whereas the posterior regions seem to be the most affected areas, where as proven, the examination of an isolated point per lung is not enough (10-zone protocol with one posterior point [4]). This study along with our results highlights the importance of scanning the posterior areas.

It is important to highlight that LUS reflects the lesions in the lung surface, and not necessary, the degree of the whole lung alteration. This might explain why incorporating more points in the exam might not translate into a higher accuracy of the lung involvement.

These findings are novel and useful, since they suggest that a 12-zone protocol could have a high degree of

| Protocol                  | Mean (SD) | ICC (95% CI)    | p value |
|---------------------------|-----------|-----------------|---------|
| 8-zone protocol [3]       | 40.8 (23.3)| 0.353 (−0.340–0.686)| 0.119  |
| 10-zone protocol [5]      | 55.3 (20.1)| 0.470 (0.018–0.733) | **0.022** |
| 10-zone protocol [4]      | 55.0 (20.4)| 0.405 (−0.121–0.697) | 0.052  |
| 12-zone protocol [6]      | 49.0 (21.2)| 0.500 (0.007–0.754) | **0.027** |
| 12-zone protocol [7]      | 59.8 (20.0)| 0.427 (0.095–0.711) | **0.022** |
| 14-zone [8]               | 53.9 (20.5)| 0.493 (0.018–0.745) | **0.019** |
| 16-zone [9]               | 55.5 (21.1)| 0.425 (0.08–0.707) | **0.041** |
| 18-zone protocol [10]     | 67.7 (23.2)| 0.373 (0.183–0.686) | **0.018** |
| 72-zone protocol [11]     | 58.9 (19.9)| 0.476 (0.042–0.742) | **0.011** |

Statistically significant expressed in bold

STD standard deviation

Table 5 Descriptive statistics and Interclass correlation coefficients (ICC) of each of the assessed Lung Ultrasonography (LUS) protocols

Fig. 3 Coronal (B) and sagittal (F) chest CT scan in lung window view of a 55 year-old woman, showing bilateral ground-glass opacifications correlated with lung ultrasound confluent B-lines (A, white arrow), reticular pattern with isolated B-lines (G, white arrow), consolidations with air bronchogram (D, white arrow) and normal lung with A-lines (C, white arrow).
concordance with CT, with similar performance to more comprehensive and time-consuming exams.

Moreover, we want to highlight that LUS is not an anatomical ultrasound, but a clinical-interpretation of artifacts, that should not replace chest X-ray or CT, instead complement them, and be regarded as its use as a clinical and diagnostic complementary and monitoring tool [21].

We acknowledge some study limitations. First of all, the poor specificity of LUS, thus the results from this study provide insights in COVID-19, and an opportunity to further investigate the role of LUS when the prevalence of disease decreases. Secondly, a small and convenience sample of confirmed COVID-19 patients, the expert sonographer performed all ultrasound scans on a consecutive sample selected based on his availability, which limits the generalizability of our results, and ought to be validated in future studies. Another limitation is that this study did not analyze the dynamic changes of the imaging modalities in different stages, and the patient outcomes, which would have a higher clinical impact. Therefore, for this purpose, we suggest the study can be considered hypothesis generating.

Although, these limitations are important, we believe that our study results support the current recommendation of a simple but comprehensive exam, including the superior and inferior areas of the anterior, lateral and posterior regions of each hemithorax. We consider appropriate a 12-zone ultrasound protocol with 12-zone scanning, including the superior and inferior areas of the anterior, lateral and posterior regions of each hemithorax. We consider appropriate a 12-zone exam protocols, as described in this study, and raise the question of the cost-effectiveness of more comprehensive exams.

Conclusion

In conclusion, our study results suggest that performing an ultrasound protocol with 12-zone scanning, including the superior and inferior areas of the anterior, lateral and posterior regions of the chest was consistent with higher ICC and higher degree of concordance with CT. We emphasize the need of a more standardization technique to further implement and develop this imaging modality in COVID-19.

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Data access and responsibility The principal investigator, Yale Tung Chen, had full access to all of the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

Declarations

Conflict of interest The authors have declared no conflicts of interest.

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References

1. Johns Hopkins Coronavirus Resource Center. https://coronavirus.jhu.edu/map.html. Accessed 9 Feb 2021.
2. Liu RB, Tayal VS, Panbianco NL, Tung-Chen Y, Nagdev A, Shah S et al (2020) Ultrasound on the frontlines of COVID-19: report from an international webinar. Acad Emerg Med Off J Soc Acad Emerg Med 27(6):523–526
3. Moore S, Gardiner E. Point of care and intensive care lung ultrasound: a reference guide for practitioners during COVID-19. Radiography. 2020;S1078817420300572.
4. Cleveret D-A. Lung ultrasound in patients with coronavirus COVID-19 disease. SIEMENS. https://www.siemens-healthineers.com/ultrasound-lung-ultrasound-covid-19. Accessed 29 Nov 2020.
5. Volpicelli G, Lamorte A, Villén T (2020) What’s new in lung ultrasound during the COVID-19 pandemic. Intensive Care Med 46(7):1445–1448
6. Tung-Chen Y, Martí de Gracia M, Diez-Tascón A, Alonso-González R, Agudo-Fernández S, Parra-Gordo ML., et al. Correlation between chest computed tomography and lung ultrasonography in patients with coronavirus disease 2019 (COVID-19). Ultrasound Med Biol. 2020;S030156292030301X.
7. Manivel V, Lesnewski A, Shamiin S, Carbonatto G, Govindan T (2020) CLUE: COVID-19 lung ultrasound in emergency department. Emerg Med Australas 32(4):694–696
8. Soldati G, Smargiassi A, Inchingolo R, Buonsenso D, Perrone T, Briganti DF et al (2020) Proposal for international standardization of the use of lung ultrasound for patients with COVID-19: a simple, quantitative, reproducible method. J Ultrasound Med 39(7):1413–1419
9. Gargani L, Soliman-Aboumarie H, Volpicelli G, Corradi F, Pastore MC, Cameli M (2020) Why, when, and how to use lung ultrasound during the COVID-19 pandemic: enthusiasm and caution. Eur Heart J Cardiovasc Imaging 21(9):941–948
10. Vassalou EE, Karantanas AH, Antoniou KM (2020) Proposed lung ultrasound protocol during the COVID-19 outbreak. J Ultrasound Med 6:jum.15402
11. Gargani L, Doveri M, D’Errico L, Frassi F, Bazzichini ML, Sedie AD et al (2009) Ultrasound lung comets in systemic sclerosis: a chest sonography hallmark of pulmonary interstitial fibrosis. Rheumatology 48(11):1382–1387
12. Hansell DM, Bankier AA, MacMahon H, McLeod TC, Müller NL, Remy J (2008) Fleischner society: glossary of terms for thoracic imaging. Radiology 246(3):697–722
13. Gargani L, Volpicelli G (2014) How I do it: lung ultrasound. Cardiovasc Ultrasound 12(1):25
14. Wu J, Wu X, Zeng W, Guo D, Fang Z, Chen L et al (2020) Chest CT findings in patients with coronavirus disease 2019 and its relationship with clinical features. Invest Radiol. 55(5):257–261
15. Koo TK, Li MY (2016) A guideline of selecting and reporting intraclass correlation coefficients for reliability research. J Chiropr Med 15(2):155–163
16. Allinovi M, Parise A, Giacalone M, Amerio A, Delsante M, Odone A et al (2020) Lung ultrasound may support diagnosis and monitoring of COVID-19 pneumonia. Ultrasound Med Biol 46(11):2908–2917
17. Vitale V, Rossi E, Di Serafino M, Minelli R, Acampora C, Iacobellis F, D’Errico C, Esposito A, Esposito F, Vallone G, Zeccolini M (2018) Pediatric encephalic ultrasonography: the essentials. J Ultrasound. https://doi.org/10.1007/s40477-018-0349-7
18. Minella R, Minelli R, Rossi E, Cremone G, Tozzi A (2020) Gastroesophageal and gastric ultrasound in children: the state of the art. J Ultrasound. https://doi.org/10.1007/s40477-020-00471-w
19. Mento F, Perrone T, Macioce VN, Tursi F, Buonsenso D, Torri E et al (2021) On the impact of different lung ultrasound imaging protocols in the evaluation of patients affected by coronavirus disease 2019: how many acquisitions are needed? J Ultrasound Med 40(10):2235–2238. https://doi.org/10.1002/jum.15580
20. Smargiassi A, Soldati G, Torri E, Mento F, Milardi D, Del Giacco P et al (2021) Lung ultrasound for COVID-19 patchy pneumonia: extended or limited evaluations? J Ultrasound Med 40(3):521–528. https://doi.org/10.1002/jum.15428
21. Di Serafino M, Notaro M, Rea G et al (2020) The lung ultrasound: facts or artifacts? In the era of COVID-19 outbreak. Radiol med 125:738–753. https://doi.org/10.1007/s11547-020-01236-5

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