Lung Ultrasound in Community-Acquired Pneumonia and in Interstitial Lung Diseases

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Introduction

Community-acquired pneumonia (CAP) remains the leading cause of death from infections in most developed countries [1]. In Germany, the overall incidence of CAP is estimated to reach 400,000–600,000 cases per year, with a mortality rate of up to 13–14% for hospitalized patients [2]. In adults, the diagnosis of CAP is confirmed if a new infiltrate on chest X-ray is established. Nevertheless, due to the limitations of chest radiographs, computed tomography (CT) of the thorax is the gold standard for diagnosing CAP.

CAP also plays an important role in children because it is responsible for many deaths in children under 5 years of age in developing countries and for a substantial portion of diseases in developed countries [3]. In contrast to adults, the diagnosis of CAP in children is only based on clinical criteria. Chest radiographs are only recommended
in severe cases treated in hospital or if complications occur [4, 5]. Therefore, it has been shown that >80% of children with clinical suspicion of CAP cannot be confirmed radiology [6, 7], leading to unnecessary antibiotic therapy. Therefore, lung ultrasound (LUS) has become important for the diagnosis and follow-up of pneumonia in adults as well as in children in the last years [8, 9].

Although CT would be the best reference standard for all sonographically diagnosed pneumonia, it is only applied in adults in case of discrepant X-ray results in two planes [9]; in children, normally chest X-ray in only one plane serves as the reference [8].

In this article, interstitial lung diseases cover diffuse parenchymal lung diseases (DPLD; different kinds of fibrosis) as well as interstitial pneumonia, acute respiratory distress syndrome (ARDS) and acute heart failure because LUS becomes more and more important in rapid decision making regarding these entities.

**Technique of LUS**

LUS may be performed with a usual ultrasound machine. At present, there are no data for establishing the best scanner settings in order to better visualize abnormal lesions and artifacts. For the examination of adult patients, convex, microconvex and linear probes are generally employed, while linear probes are preferred in children. The linear probe is the best choice for studying the dynamics of breath-dependent motion as well as pleural line abnormalities.

The patient may be evaluated in a sitting or supine position. This depends on the clinical condition of the patient as well as on the clinical problem. In most cases, dorsal regions are evaluated in the sitting patient, while anterior regions are assessed in the supine position. Pleural effusion is best detected in the sitting position, while the supine position is favored for a pneumothorax.

In general, image acquisition may be performed for the evaluation of the whole lung or as focal investigation when a clinicopathological finding is detected. Panoramic evaluation is achieved by performing longitudinal scans, from top to bottom, along the anatomical lines of the thorax (parasternal, midclavicular axillary and para-vertebral lines) as well as horizontal scans along the intercostal spaces. A focal examination is performed with different scans (longitudinal, transverse and oblique views) and sometimes also with different probes. Often, the examination begins in symptomatic areas, for example, where the patient complains of pleuritic pain or where a pathological finding is present on auscultation or percussion. In patients with a forced supine position, posterior areas may be explored in the lateral decubitus position.

Some areas of the lung (retroscapular, periclavicular and superior portions of the axilla) are hidden by anatomic structures. These regions could be explored through appropriate tilting of the probe but about 20% of the lung surface is not visualized on transthoracic ultrasound examination.

Certain anatomical characteristics in children, such as a thinner chest wall and smaller thoracic width and lung mass, facilitate ultrasound imaging and ensure good-quality images of the lung. In noncooperative young patients, the examination is more time consuming but is usually feasible. Performance of a chest ultrasound scan requires less skill than other sonographic scans (e.g. of the abdomen or heart) and the learning curve is faster. Bedetti et al. [10] have demonstrated that beginners are able to detect the presence of the pulmonary interstitial syndrome (IS) after fewer than 10 examinations and a total training time of 30 min. Ultrasound diagnosis of lung consolidation may be considered as a basic sonographic technique with a steep learning curve [11]. For the evaluation of B-lines, it is recommended to set the focus at the level of the pleural line.

**LUS Compared with Other Methods in the Diagnosis and Follow-Up of CAP in Adults**

**Sonomorphology**

Sonomorphology of CAP may be divided into parenchymal, pleural and vascular criteria (table 1).

The most important parenchymal criterion is the positive air bronchogram within an echopoor area, which may be found in about 70–97% of cases [9, 12–14] (fig. 1), while among pleural criteria basal effusion was the most frequent detection (in about 34–61% cases) [9, 12–15]. Determination of vascularization is very helpful, especially for differential diagnosis, but it is not essential for the diagnosis of uncomplicated CAP. Table 2 gives an overview of studies involving sonography in the diagnosis of adult CAP.

**Differential Diagnosis**

Lung cancer, pulmonary embolism and atelectasis are the most important differential diagnoses of pneumonia. Lung cancer may be accompanied by pneumonia following obstruction of the airways. In this case, no complete recovery will occur during the follow-up of pneumonia. Lung carcinoma normally has no positive air
bronchogram. Malignancies are polycyclic, not well de-
marcated and sometimes infiltration into the thorax wall
may be discovered that avoids breath-dependent motion
of the lesion. In spectral curve analysis, various vessels are
detectable, particularly new blood vessels in tumor tissue
(neoangiogenesis).

Pulmonary embolism may also cause pneumonia due
to embolic occlusion, which might be suspected if pneu-
monia is triangular. Characteristic sonographic signs of
peripheral pulmonary embolism are multiple (mean 2–3), mostly triangular, hypoechoic, subpleural lesions
with breath-dependent motion, which prefered the dorsal
and basal regions of the lung [19, 20]. Peripheral pulmo-
{}nary embolism typically reveals no flow signals in color
doppler sonography. Pleurisy and pleural lesions of un-
known causes may be further differentiated by contrast-
enhanced sonography [21]. For further reading on this
topic, we like to refer to the review by Kreuter et al. [22]
in this Thematic Review Series.

Atelectasis, compressive atelectasis as well as resorptive
atelectasis have to be differentiated from pneu-
monia. Compressive atelectasis is normally induced by a
large pleural effusion and moves within the effusion like
a ‘waving hand’. It is moderately echoic, sharp demar-
cated and concave. Occasionally, it is possible to notice
breath-dependent ventilation within the atelectasis. Af-
after thoracocentesis or diuretic therapy, compressive at-
electasis becomes smaller or even disappears. Resorptive

Table 1. Sonomorphology of adult CAP

<table>
<thead>
<tr>
<th>Parenchymal criteria</th>
<th>Pleural criteria</th>
<th>Vascular criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Area of echopoor echogenicity and an inhomogeneous echotexture with blurred margins</td>
<td>Pleural line attenuation corresponding to the affected area</td>
<td>Color Doppler sonography: enhanced, tree-like vascularity</td>
</tr>
<tr>
<td>Positive air bronchogram: reflects residua of air within consolidated areas</td>
<td>Localized pleural effusion</td>
<td>Spectral curve analysis: pulmonary and bronchial artery</td>
</tr>
<tr>
<td>Positive fluid bronchogram: reflects fluid-filled airways</td>
<td>Basal pleural effusion</td>
<td>Contrast-enhanced sonography: short time to enhancement and marked extent of enhancement</td>
</tr>
</tbody>
</table>

Necrotic areas within pneumonic lesions possible

Superficial fluid alveologram: subpleural echopoor region without bronchogram and fluid bronchogram

Fig. 1. A 28-year-old woman with clinical signs of pneumonia. 

a LUS of the left side reveals an echopoor lesion with typical positive air bronchogram. b Corresponding chest X-ray performed on the day of her hospital admission shows infiltration on both sides. c LUS of the left side on day 16 is normal, where- as chest X-ray on day 16 still shows residual disease on both sides (d).
atelectasis is a consequence of airway obstruction, mostly caused by cancer, exudate or aspiration. The accompanying effusion is small in comparison with the atelectasis, and thoracocentesis does not influence the size of resorptive atelectasis. Its echogenicity is comparable to that of the liver and the shape is very variable. Sometimes a fluid bronchogram reflecting exudate-filled airways may be detected. In contrast to pneumonia and compressive atelectasis, resorptive atelectasis does not reveal a dynamic air bronchogram [23]. Using contrast-enhanced sonography, resorptive atelectasis may be demarcated from the inducing cancer or metastasis [24], which is very helpful to determine the optimal site for tumor biopsy.

Table 2. Overview of studies on the sonographic diagnosis of CAP in adults [16, 17]

<table>
<thead>
<tr>
<th>Feature</th>
<th>Gehmacher et al. [12], 1995 (n = 143)</th>
<th>Reissig and Kroegel [13], 2007 (n = 30)</th>
<th>Parlamento et al. [15], 2009 (n = 49)</th>
<th>Sperandeo et al. [14], 2011 (n = 342)</th>
<th>Reissig et al. [9], 2012 (n = 362)</th>
<th>Cortellaro et al. [18], 2012 (n = 120)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Study design</td>
<td>Patient with X-ray-established pneumonia</td>
<td>Patient with suspected CAP; LUS follow-up was required</td>
<td>Patient with suspected CAP admitted to ED</td>
<td>Inclusion of X-ray-established CAP; LUS follow-up</td>
<td>Patient with suspected CAP; LUS follow-up</td>
<td>Patient with suspected CAP in ED; CT if clinically indicated (n = 30); most X-rays in supine position; final diagnosis as surrogate</td>
</tr>
<tr>
<td>Established diagnosis of pneumonia</td>
<td>143 (100%)</td>
<td>LUS and X-ray follow-up</td>
<td>32/49 (65.3%)</td>
<td>342 (100%)</td>
<td>229/362 (63.3%)</td>
<td>81/120 (67.5%)</td>
</tr>
<tr>
<td>LUS positive</td>
<td>127/143 (88.8%)</td>
<td>Not reported</td>
<td>31/32 (96.9%)</td>
<td>314/342 (92%)</td>
<td>211/229 (92.1%)</td>
<td>80/81 (98%)</td>
</tr>
<tr>
<td>X-ray positive</td>
<td>Reference test</td>
<td>Reference test</td>
<td>24/32 (75%)</td>
<td>Reference test</td>
<td>199/229 (86.9%)</td>
<td>54/81 (67%)</td>
</tr>
<tr>
<td>Interstitial pattern</td>
<td>Not reported</td>
<td>Not reported</td>
<td>22/32 (68.8%)</td>
<td>Not reported</td>
<td>Not reported</td>
<td>7/80 (8.8%)</td>
</tr>
<tr>
<td>Positive air bronchogram</td>
<td>112/127 (88.1%)</td>
<td>32/32 (97%)</td>
<td>16/32 (50%)</td>
<td>220/314 (70%)</td>
<td>183/211 (87.7%)</td>
<td>71/80 (88.8%)</td>
</tr>
<tr>
<td>Fluid bronchogram</td>
<td>Not reported</td>
<td>0/33 (0%)</td>
<td>Not reported</td>
<td>100/314 (31%)*</td>
<td>17/211 (8.1%)</td>
<td>28/80 (35%)</td>
</tr>
<tr>
<td>Basal pleural effusion in LUS</td>
<td>78/143 (54.5%)</td>
<td>20/33 (61%)</td>
<td>11/32 (34.4%)</td>
<td>120/34 (35.1%)</td>
<td>105/193 (54.4%)</td>
<td>31/80 (38.8%)</td>
</tr>
<tr>
<td>Size (LUS), cm</td>
<td>Not reported</td>
<td>3.37×0.94 and 9.12×4.53</td>
<td>Not reported</td>
<td>6.63×3.11</td>
<td>3.2×3.7</td>
<td>Not reported</td>
</tr>
<tr>
<td>Sensitivity for LUS</td>
<td>Not applicable</td>
<td>Not reported</td>
<td>Not reported</td>
<td>Not applicable</td>
<td>93.4%; CI: 89.2–96.3%</td>
<td>98%; CI: 93.3–99.9%</td>
</tr>
<tr>
<td>Specificity for LUS</td>
<td>Not applicable</td>
<td>Not reported</td>
<td>Not reported</td>
<td>Not applicable</td>
<td>97.7%; CI: 93.4–99.6%</td>
<td>95%; CI: 82.7–99.4%</td>
</tr>
<tr>
<td>Likelihood ratio for positive LUS</td>
<td>Not applicable</td>
<td>Not reported</td>
<td>Not reported</td>
<td>Not applicable</td>
<td>40.5; CI: 13.2–123.9%</td>
<td>19.3; CI: 4.99–74.2%</td>
</tr>
<tr>
<td>Likelihood ratio for negative LUS</td>
<td>Not applicable</td>
<td>Not reported</td>
<td>Not reported</td>
<td>Not applicable</td>
<td>0.07; CI: 0.04–0.11%</td>
<td>0.01; CI: 0.002–0.09%</td>
</tr>
</tbody>
</table>

ED = Emergency department; *= patients with millimetric hypoechoic tubular formations.
LUS Compared With X-Ray, CT and Auscultation

The sensitivity of LUS in detecting CAP varies between 93.4 and 98%, and the specificity between 97.7 and 95%; the positive likelihood ratio ranges from 19.3 to 40.5, and the negative likelihood ratio from 0.01 to 0.07 [9, 18]. In several studies, LUS was at least as good as X-ray evaluation in detecting CAP [8, 9, 15]. Nevertheless, the gold standard CT was never used in all patients due to the hazards of radiation exposure [8, 9, 15]. Sonography detects peripheral pneumonic lesions, even small ones, whereas X-ray covers also central localized processes, but small ones may escape radiological detection. During the follow-up, LUS may already appear normal because the lesion no longer reaches the peripheral pleura. However, both techniques have advantages and disadvantages. They are comparable as well as complementary. It has been demonstrated that a combination of LUS and auscultation findings resulted in a higher positive (42.9, 95% confidence interval, CI, 10.8–17.0) as well as in a lower negative likelihood ratio (0.04, CI 0.02–0.09) [9]. LUS enables the early detection of complications such as empyema and lung abscesses.

LUS Compared with Other Methods in the Diagnosis and Follow-Up of CAP in Children

At present, only a few papers have been published on this topic [8, 25–27], but all studies have demonstrated that LUS is able to diagnose pneumonia in children and young adults with high accuracy (table 3). It will not completely replace chest radiography, but with increasing use and experience by ultrasonographers combined with improvements in LUS techniques, it has the potential to establish as a very important diagnostic imaging option for pneumonia, especially in children [28].

Sonomorphology

The ultrasonographic appearance of pneumonia does not differ between children and adults [11]. Pneumonia appears as a hypoechogenic area with poorly defined borders and with the presence of B-lines at the far-field margin. The pleural line is less echogenic in the area affected by lung consolidation and lung sliding is reduced or absent. In the case of consolidations, branching echogenic structures – representing air bronchograms – are seen in the infected area (fig. 2 and fig. 3). Air bronchograms may show intrinsic dynamic centrifugal movements due to breathing. This finding is called dynamic air bronchogram: it attests bronchial patency and rules out obstructive atelectasis. Multiple lenticular echoes, representing air trapped in the smaller airways, are also frequently observed. Fluid bronchograms, described in postobstructive pneumonia, are identified as anechoic tubular structures with hyperechoic walls but without color Doppler signals. Fluid bronchograms are frequently observed in pneumonia in children (fig. 4, 5). Pleural effusion is easily detected on LUS and appears as an anechoic area in the pleural space. A honeycomb organization of fibrin is observed in pleural empyema (fig. 6).

Differential Diagnosis

Like in adults, pulmonary atelectasis may simulate pneumonia. The dynamic ultrasound signs are often very useful for the differential diagnosis and may be monitored.
at bedside. Ultrasonographically, atelectasis is characterized by a liver-like appearance of the lung with ‘lung pulse’, absence of lung sliding and a parallel course of air bronchograms, in agreement with results in adult patients [29]. The evidence of dynamic air bronchograms rules out obstructive atelectasis [23]. Fluid bronchograms are often present.

**LUS Compared with X-Ray and CT**

Children and infants with pneumonia may present with a number of clinical symptoms and signs, such as fever, cough and tachypnea. A minority of children present with fever of unknown origin and may have no respiratory symptoms or signs. In these cases, a ra-

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**Fig. 2.** Transverse thoracic scan of a 9-year-old child with clinically suspected pneumonia. LUS shows the presence of consolidation with evidence of air bronchograms.

**Fig. 3.** Lobar pneumonia in an 11-year-old child. Air bronchograms are seen in the infected area as branching echogenic structures.

**Fig. 4.** Fluid bronchograms, which are frequently observed in post-obstructive pneumonia, appear as anechoic tubular structures with hyperechoic walls without any color Doppler signal.

**Fig. 5.** Lobar pneumonia in a 10-year-old child admitted to the emergency department for fever and cough. LUS reveals air and fluid bronchograms within the affected area.
diological examination may be necessary. In the guidelines, chest X-ray is still considered to be the first imaging step for diagnosing pneumonia in children [4, 5, 30]. CT has a high level of accuracy but cannot be used as a first-line radiological examination due to its high exposure to ionizing radiation, reduced availability and costs. In adults, LUS has been shown to be a very promising technique due to its high sensitivity in detecting pleural effusions, lung embolism, pneumonia, pneumothorax, IS and atelectasis [31], but research on these topics in children is scarce. Ultrasound examination can be done at the patient’s bedside and repeated without any radiation risk. Due to these characteristics, LUS is an ideal tool for diagnosis as well as the detection of complications and for the follow-up of pneumonia.

LUS is a clinically useful diagnostic tool in pediatric patients with suspected pneumonia. The ultrasound signs of lung and pleural diseases described in adults are also found in pediatric patients. Although LUS is a clinically useful tool for the diagnosis of pneumonia, it does not rule out consolidations that do not reach the pleura [11]. These data suggest that in case of a clinical suspicion of pneumonia, a positive LUS excludes the need to perform chest radiography. Nevertheless, a negative chest radiograph does not rule out pneumonia.

**LUS Compared with Other Methods in the Diagnosis of Interstitial Lung Diseases**

**Diffuse Parenchymal Lung Diseases**

According to the American Thorax Society and European Respiratory Society, DPLD are classified into DPLD of known causes (e.g. drug induced), idiopathic interstitial pneumonia (e.g. idiopathic pulmonary fibrosis, IPF), granulomatous DPLD (e.g. sarcoidosis) and DPLD of other causes [32]. The main symptom in IPF patients is increasing dyspnea on exertion. Diagnosis is based on clinical history and examination, serological tests, as well as lung function, CT and, if necessary, bronchoscopy with bronchoalveolar lavage or transbronchial biopsy. In some rare cases, thoracic surgery is included in the differential diagnosis. Current studies show the significance of LUS in diagnosing interstitial lung diseases [33–37].

The most important ultrasound sign for interstitial diseases is B-lines. The generation of B-lines is still debated. It is supposed that the creation of B-lines depends on increased density of the lung due to a decrease in the amount of air, an increase in interstitial tissue or both. These phenomena generate an acoustic permeability of the pleural line that in normal conditions acts a specular reflector as a consequence of the high difference in acoustic impedance between the chest wall tissue and aerated lung [38, 39]. Nevertheless, B-lines are artifacts and may also appear in the healthy lung [33].

In general, IS may be diffuse bilateral (e.g. pulmonary edema of various causes, interstitial pneumonia or pneumonitis, or DPLD) or focal (e.g. accompanying pneumonia or pneumonitis, atelectasis, pulmonary contusion or infarction, pleural disease or neoplasia). Furthermore, B-lines may have a homogeneous or nonhomogeneous distribution and the localization as well as the gradient (ventral-dorsal/apical-caudal) and the alterations in the pleural line (fragmentation, irregularity and swelling) should be assessed for further differential diagnostic considerations. For DPLD (e.g. IPF), table 4 summarizes characteristic sonomorphologic features of DPLD (fig. 7).

In interstitial pneumonia, an interstitial LUS pattern combined with spared areas is strongly suggestive for viral pneumonia and correlates with the findings on CT scans [40, 41]. Nevertheless, some studies lack the ‘gold standard’ for diagnosis or have a time lag between LUS and CT [37]. Using LUS, it is possible to detect and follow up interstitial abnormalities. Nevertheless, to determine the cause of the history of B-lines, clinical examination and sometimes follow-up and CT are necessary.
To date, the following issues remain open to discussion:

- counting of B-lines (8-zone technique vs. 28 rib interspaces) [11, 34];
- the effect of the ultrasound equipment, the scanner (convex or linear) and penetration depth used on the number of B-lines and pleural line abnormalities;
- differences in scoring systems between various studies (>6 B-lines per scan; ≥3 B-lines in a longitudinal plane between 2 ribs; ≥3 B-lines in ≥2 adjacent scanning sites or >5 B-lines) [11, 33, 37].

In general, it has to be taken into account that counting of B-lines is best performed in real-time mode as they may disappear on a frozen image.

Increase in Extravascular Lung Water: Acute Heart Failure and ARDS

In acute cardiopulmonary clinical conditions, bilateral and diffuse B-lines are more frequently an ultrasonographic sign of increased extravascular lung water. The sensitivity of B-lines in detecting the increase in extravascular lung water is very high, ranging from 96.9 to 100% in different studies, and their absence in a dyspneic patient virtually rules out pulmonary edema [42, 43].

In critically ill patients, the differential diagnosis between acute cardiogenic pulmonary edema (APE) and ARDS may be very difficult [44]. Differential diagnosis using CT has been well described, although the accuracy of chest X-ray is unclear. Chest X-ray is a poor at detecting the cause or location of excess extravascular lung water [45].

The pathophysiology of ARDS is completely different from that of APE and this generates different pleuropulmonary ultrasound patterns with a different distribution of IS [11, 42]. In ARDS, the damage of the alveolar capillary membrane causes an early, diffuse, heterogeneous alveolar flooding ranging from a ‘ground glass’ appearance to lung consolidation. The heterogeneous involvement of the lungs in ARDS explains the presence of ‘spared areas’. Pleural line abnormalities are very common in patients with ARDS and may be well visualized with a linear probe. It is possible to observe areas with a reduction or absence of ‘lung sliding’ and often areas in which B-lines are coalescent (‘white lung’) are observed. In addition, the ‘lung pulse’ sign could be observed. The pleural line appears irregular, thickened and coarse because of the presence of multiple small subpleural consolidations. Involvement of the pleural line is not homogeneous and corresponds to the distribution and degree of IS (fig. 8; table 5).

In contrast, APE is a hydrostatic edema and IS shows a homogeneous distribution involving both anterior and posterior lung fields. Superior lung fields might be less affected but ‘spared areas’ cannot be observed, which reflects the pathophysiology. B-lines initially prefer the lung...
### Table 5. Sonomorphological differentiation of DPLD, APE and ARDS

<table>
<thead>
<tr>
<th></th>
<th>DPLD</th>
<th>APE</th>
<th>ARDS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical setting</td>
<td>Chronic</td>
<td>Acute</td>
<td>Acute</td>
</tr>
<tr>
<td>IS</td>
<td>Always present</td>
<td>Always present</td>
<td>Always present</td>
</tr>
<tr>
<td>Distribution of IS</td>
<td>Nonhomogeneous distribution, mostly in the dorsal and lower areas</td>
<td>Bilateral and symmetric distribution</td>
<td>Nonhomogeneous distribution, presence of spared areas</td>
</tr>
<tr>
<td>Effect of diuresis on B-lines</td>
<td>No effect</td>
<td>Reduction</td>
<td>No effect</td>
</tr>
<tr>
<td>Pleural line abnormalities</td>
<td>Present, typical</td>
<td>Absent</td>
<td>Present, typical</td>
</tr>
<tr>
<td>Reduction or absence of lung sliding</td>
<td>Absent</td>
<td>Absent</td>
<td>Present</td>
</tr>
<tr>
<td>Lung pulse</td>
<td>Present</td>
<td>Absent</td>
<td>Present</td>
</tr>
<tr>
<td>Consolidations</td>
<td>May occur</td>
<td>Absent</td>
<td>Frequent in the posterior areas</td>
</tr>
<tr>
<td>Pleural effusion</td>
<td>Often localized effusions</td>
<td>Very frequent and large</td>
<td>Common but small</td>
</tr>
</tbody>
</table>
bases but with increasing capillary venous pressure extend to the medium and superior fields. The pleural line is rarely involved and appears as a hyperechoic band without sliding impairment (fig. 9; table 5).

Conclusion

The published data show that LUS performs well in a lot of clinical scenarios. Nevertheless, at present, CT still represents the gold standard for most pulmonary diseases. However, in comparison with chest X-ray, especially in the critical care setting, LUS at bedside performs better than chest radiography in numerous cases.

LUS is an accurate tool in the diagnosis of pneumonia in adults as well as in children. Regarding pneumonia, LUS is at least as accurate as chest radiography.

The presence of multiple diffuse bilateral B-lines on lung examination indicates IS. For further differential diagnosis, an integrated consideration of history, clinical examination, LUS (B-lines, their distribution as well as variations in the pleural line) and echocardiography should be performed. LUS is an excellent tool for IS screening. Especially in pulmonary edema, repeated LUS control examinations may reflect the dynamics of the disease under therapy and so LUS may serve as a therapy guide.

At present, LUS is not considered in any guideline for the diagnosis of CAP or IS. Nevertheless, we think that future guidelines should include LUS.

Ultrasound avoids the use of ionizing radiation. Therefore, the use of ultrasound – especially in pediatric patients – needs to be encouraged not just as a valid diagnostic alternative but as a necessary ethical choice. Further research regarding the role of color Doppler sonography, spectral curve analysis and contrast-enhanced ultrasound is necessary, especially with respect to differential diagnosis of lung consolidations and early detection of complications.

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