Introduction

The field of critical care ultrasound founded by Dr. Lichtenstein in the early 1990s evolved over three decades into an indispensable bedside tool for management of the critically ill. Out of Lichtenstein’s whole-body ultrasound paradigm, lung ultrasound was born. It allowed, for the first time, rapid parenchymal assessment of this vital organ, at the point of care. Safe, non-invasive, and easily repeatable, lung ultrasound could provide immediate actionable diagnoses of life-threatening conditions, which contributed to the development of sonographic evaluation protocols for acute respiratory failure (1), shock (2-4), and cardiac arrest (5-7). Semiquantitative ultrasound assessment of lung aeration has found its application in optimization of mechanical ventilation. Ultrasound guidance is the standard of care for the majority of bedside procedures in the intensive care unit (ICU). Point-of-care ultrasound thus shifted the paradigm of thoracic imaging by eliminating the dissociation between the treating physician and physician.
interpreting the imaging (8). Although the typical role of interventional pulmonologist (IP) in the ICU is to perform advanced procedures, rather than day to day management of the critically ill, it is necessary to discuss the universal principles of lung and pleural ultrasound before focusing on sonographic aspects of pleural drainage procedures, transthoracic biopsies, and percutaneous dilatational tracheostomy (PDT).

**Basic principles of ultrasonography**

For a more detailed consideration of applied ultrasound physics, comprehensive review is available (9). The fundamental principle of ultrasound imaging is the reflection of ultrasound waves from tissues in the path of the beam. Echogenicity of the tissue refers to its ability to reflect and/or transmit ultrasound waves in the context of surrounding tissues. Translated to a visual greyscale, echogenic, highly reflective structures appear white, in contrast to black, non-reflective, anechoic structures; “hypoechoic”, “isoechoic” and “hyperechoic” are relative descriptors of brightness of the compared structures, in shades of gray.

Whenever there is an interface of structures with different echogenic properties as defined by their acoustic impedances, a difference in brightness will be apparent on the screen. The difference in acoustic impedances between two tissues is directly proportional to the extent of reflection of ultrasound waves at their interface. Minimal acoustic impedance of air determines large impedance mismatch with surrounding tissues and results in almost complete wave reflection at lung-chest wall interface, lending the pleural line its distinctive echogenic appearance. Because only a small proportion of waves penetrates air, and air is an unfavorable medium for sound transmission, aerated lung appears as homogenous amorphic grayness rather than a discrete structural entity. As the air content in the lung decreases, acoustic mismatch also diminishes and the ultrasound beam gets partially reflected, repeatedly, as it reaches the deeper zones. With even further decrease in air content, acoustic window completely opens and lung can be directly visualized as solid parenchyma.

Pleuropulmonary pathologic processes conducive to the ultrasonographic examination such as effusions and peripherally extending consolidations provide real images, while the rest of the lung ultrasound is based on interpretation of artifacts derived by air/tissue interface.

**Equipment**

**Ultrasound machine**

Lung sonography is based on artifacts described by original investigators using technology available in the 1990s; examination can therefore be performed using any commercially available two-dimensional (2D) scanner (10-13). The lung and pleural structures are visualized in 2D B-mode (brightness mode, which displays the acoustic impedance of a tissue cross-section), while pleural movement can be assessed via M-mode (motion mode, which emits pulses in quick succession, taking an image each time). Image quality is determined by the individual machine characteristics, presets, and post-image processing. One caveat is that modern ultrasound machines use numerous techniques to minimize the presence of ultrasound artifacts while lung ultrasonography often relies on analysis of such artifacts. Therefore, experienced operator is expected to adapt machine settings for optimal visualization of said artifacts, unless lung presets are configured by the manufacturer. Typically, lung presets will turn off smoothing and artifact minimization algorithms such as compound and harmonic imaging, and will also reduce dynamic range. This is particularly helpful during assessment of lung sliding and B-lines. Abdominal presets may be more appropriate when evaluating pleural effusions or consolidations. Doppler ultrasound is not a part of diagnostic lung sonography paradigm, but may be useful as a procedural adjunct in order to avoid vascular injury. Preferred ultrasound devices for bedside use are easy to transport and clean, with short boot-up times and solid data storage and transfer capabilities (10).

**Transducers**

Ultrasound probes in use today are multiple element transducers, commonly called arrays. Various combinations of array construction (linear, convex), and array functional features such as beam scanning and steering (accomplished by electronic sequencing or electronic phasing) and beam focusing (accomplished by electronic phasing) define the probe footprint and image formats. Aside from technology used for beam manipulation which defines array type and consequently probe footprint and image format, other important probe feature is the ultrasound frequency range. Simplified terminology that partially describes probes as linear, phased array, or convex is pervasive (14) and
used interchangeably with their most prominent imaging indication—vascular, cardiac, and abdominal, respectively. Higher frequency probes are typically used to acquire high resolution images of more superficial structures, while lower frequency transducers capture lower resolution images of deeper structures. If multiple probes are available, a useful rule is to choose the one that will provide the best resolution for the required depth. Certain probes have wide frequency ranges allowing them to be more versatile and therefore cost-effective. This concept is illustrated by the whole-body critical ultrasound paradigm which was built around a universal wide range microconvex probe (10,11). Intercostal acoustic windows are narrow, making it much easier to acquire images free of rib artifact using a smaller footprint probe, although diverging ultrasound beam also means that only a small field of the pleural surface is examined (“tunneled vision”) and lateral resolution of the image is poor. A larger footprint enables examination of a larger field of the pleural surface and improves resolution, but it may be more difficult to acquire an image.

Linear (vascular) probes (Figure 1A) are characterized by high frequency (7.5–10 MHz) and large footprint; they are typically used to visualize the chest wall, pleural line, and subpleural space in greater detail, and evaluate for pneumothorax. Because this evaluation is dependent on ultrasound artifacts generated by air/tissue interface, it is particularly important to use configured lung presets.

Phased-array (cardiac) probes (Figure 1B) are characterized by low frequency and small footprint allowing for relatively easy image acquisition of deeper chest structures, albeit with less image resolution and narrow field of view. Therefore, this transducer is a reasonable selection for lung ultrasound if gross and diffuse pathology is suspected.

The convex (abdominal) probes (Figure 1C) are characterized by intermediate frequency values and large footprint, allowing decent visualization of the superficial structures without losing the overview of the chest while attempting to search the entire pleural surface for localized pathology.

Microconvex probes (Figure 1D) have a small footprint in combination with good resolution for both superficial and deep structures, making them the probe of choice for lung ultrasonography (10,11).

**Figure 1** Various ultrasound probes. (A) Linear (vascular) probe; (B) phased array (cardiac) probe; (C) convex (abdominal) probe; (D) microconvex probe.

**Basic ultrasonography technique**

**Probe and image orientation**

Radiologic convention dictates that for imaging in longitudinal planes—sagittal and coronal, probe marker should be oriented towards the head (cephalad), while for
imaging in the transverse plane probe marker should be oriented towards the patient’s right. Image orientation on the screen is determined by the location of the screen marker which corresponds to the probe marker. Radiologists keep the screen marker on the left side of the screen and cardiologists traditionally keep it on the right. Critical ultrasound pioneering work was performed using head-to-the-left whole body convention including the heart, while practices among critical care practitioners are varied (15).

**Depth**

Prior to beginning an examination, depth should be considered. Examiner can start by visualizing the deeper structures by selecting the higher numerical depth setting and then zoom in on more superficial structures by switching to lower depth setting.

**Gain**

The gain function serves to optimize image quality by compensating for attenuation (sound amplitude reduction). As sound waves travel deeper into the body the amplitude of the returning signal diminishes but can be amplified by the receiver, therefore creating a brighter and better visible image on the screen. Gain can be adjusted for the near field, far field, or the entire field (overall gain). If the gain is set too low, the image appears dark, while excessive gain makes the image too bright by adding “noise”.

**Patient positioning**

Image acquisition performed with the patient sitting up allows full access to the chest, but is rarely feasible in a critically ill patient. More often patients are imaged while supine, semi-recumbent, in lateral decubitus, or slightly rotated. Because the distribution of lung abnormalities does not rapidly change with different body positions, with the notable exception of free pleural effusions, it is possible to perform an adequate ultrasonographic exam in almost any position (13). The only true limitation is difficulty assessing the dorsal chest of a supine patient who is not easily mobilized.

**Scanning strategy**

By examining various ultrasound planes systematically, expert ultrasonographer can develop a 3D model of the chest with its associated pathology. Although organized approach is recommended, there is no superior scanning protocol and flexibility is allowed, to account for clinical context (13,16). Universally, higher patient acuity dictates simpler and faster methodology. Comprehensive 28 rib space technique allows for semi-quantification of the interstitial syndrome (16,17), while eight-zone anterolateral lung examination is often employed during qualitative evaluation for interstitial syndrome (16,18). Bedside Lung Ultrasound in Emergency (BLUE) protocol assumes clinically significant interstitial syndrome and free pneumothorax should always be detected anteriorly (1), while free pleural fluid can predictably be found posteriorly above the diaphragm regardless of the volume (19). These assumptions allow rapid identification of the cause of acute respiratory failure, by examining only three identical predetermined points (the upper BLUE-point, lower BLUE-point and the posterolateral alveolar and/or pleural syndrome—or PLAPS—point) in both left and right hemithorax (20).

**Semiotics of lung and pleural sonography**

Basic consideration of lung and pleural sonography is that real images always reflect pathology (pleural fluid and consolidated lung parenchyma) while artifacts can reflect both physiological and pathological state.

**Signs**

The field of lung ultrasound is built on the following visual concepts: pleural line (bat-wing sign), lung sliding and corresponding seashore sign, absence of lung sliding and corresponding stratosphere sign, lung pulse, lung point, A-lines, B-lines, sinusoid sign, shred sign, tissue-like sign, and air bronchograms (16,21). Seashore, stratosphere, and sinusoid sign are recorded in M-mode. Lung pulse and lung point can be sought for in both M- and B-modes. Remaining signs pertain to B-mode.

**Pleural line (bat-wing sign)**

The basic image of lung sonography is B-mode longitudinal view called the bat-wing sign (Figure 2) generated by upper and lower ribs (echogenic wings of the bat) and the pleural line (echogenic back of the bat). Pleural line is visualized approximately 0.5 cm below rib lines (Figure 2) and it always represents the parietal pleura, while visceral pleura may or may not contribute. Utilizing oblique instead of
longitudinal approach one can evaluate a larger portion of the pleural line, not interrupted by the rib shadows.

**Lung sliding (pleural sliding) and seashore sign**

In B-mode (Video 1), lung sliding is represented by the movement of the pleural line synchronously with respiration (13). Despite various, even contradictory definitions, lung sliding is intuitively recognized as homogeneous shimmering starting at the pleural line and spreading below (10). Characteristic appearance of lung sliding in M-mode is named seashore sign (Figure 3). Area above the pleural line with a wave-like pattern represents the chest wall and area below the pleural line with a granular/sandy beach-like pattern represents the expanded and ventilated lung. In physiologic terms, evident lung sliding suggests that parietal and visceral pleura are closely approximated, and regional ventilation is preserved. Presence of anterior lung sliding rules out pneumothorax with 95.3% sensitivity, 91.1% specificity and 100% negative predictive value (22). Normally, lung sliding is more subtle in the upper parts and easier to recognize towards lung bases. Certain pathologic conditions or clinical contexts can significantly reduce lung sliding; for instance, it may be quite difficult to ascertain whether lung sliding is present in a paralyzed patient, protectively ventilated with low tidal volumes (12). For optimal visualization of subtle findings during lung sliding assessment, the harmonic imaging and compound imaging filters should be disabled and dynamic range (compression) reduced, although one could consider disabling all filters (11).

**Absence of lung sliding (stratosphere or barcode sign)**

If lung sliding is absent, pleural line does not move synchronously with ventilation. Consequently, no shimmering is detected in B-mode (Video 2). In M-mode, the granular or sandy beach pattern below the pleural line is replaced by horizontal lines, while chest wall appearance above the pleural line is unchanged, creating the barcode or stratosphere sign (Figure 4). Detection of this sonographic pattern in the anterior chest implies that pleural surfaces are separated by air, and/or that regional ventilation is impaired by airway obstruction, hyperinflation, or pleural adhesions. Further investigation is performed by assessing for the presence of B-lines, lung point, and lung pulse (see below), while considering the clinical context. In trauma patients, for example, isolated absence of lung sliding has 98.1% sensitivity and 99.2% specificity for detection of...
pneumothorax (23). Absence of lung sliding can be observed in patients ventilated with high positive end-expiratory pressure (PEEP) but without pneumothorax (24).

**Lung pulse**
In the absence of lung sliding, subtle movements of the pleural line caused by the force of cardiac pulsation and synchronous with the pulse can be seen in both B-mode (Video 3) and M-mode (Figure 5). Presence of lung pulse rules out pneumothorax and confirms absence of regional ventilation with 93.0% sensitivity, and 100% specificity (25).

**Lung point**
Lung point marks the location on the chest wall where the collapsed lung achieves transient contact with the parietal pleura during inspiration. In B-mode, this phenomenon is appreciated as inspiratory presence and expiratory absence of lung sliding in a particular scanning location. M-mode depicts alternating seashore and stratosphere appearances in relation to the phase of the respiratory cycle (Figure 6). Sensitivity of lung point for pneumothorax detection is only 66%, while specificity is 100% (26). Notably, complete lung collapse cannot produce the lung point. Location on the chest wall where lung point is detected can be used for pneumothorax semi-quantification (27).

**A-lines**
A-lines are hyperechoic, horizontal lines seen at regular intervals from the pleural line (Figure 7). These intervals are equal to the distance between the transducer on the skin and the pleural line. A-lines represent a reverberation artifact and indicate the presence of air, physiologic or free, below the parietal pleura (28). Significance of A-lines is further interpreted in the context of lung sliding and lung pulse, as well as other signs.

**B-lines**
B-lines (Figure 8) are discrete, laser-like, vertical, hyperechoic reverberation artifacts that arise from the pleural line, extend to the bottom of the screen without fading, erase A-lines, and move synchronously with lung sliding (16). They belong to the comet-tail artifacts family and were referred to as lung comets before the nomenclature was standardized (16). One theory of B-line genesis is that

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**Figure 5** Lung pulse (M-mode). In the absence of lung sliding, a distinct visual pattern below the pleural line (arrows) can be seen, and is created by pleural movement synchronously with cardiac pulsations. This pattern is generated by the lack of regional ventilation and rules out pneumothorax.

**Figure 6** Lung point (M-mode). The arrow marks the transition from the seashore sign to the stratosphere sign with changes in the respiratory cycle. Seashore sign indicates that lung sliding is present during inspiration and stratosphere sign indicates that lung sliding is absent during expiration.

**Figure 7** A-lines (B-mode). Horizontal, hyperechoic lines below the pleural line, marked by white arrows, and representing reverberation artifact.
ultrasound beam interacts with edematous interlobular septa originating from the visceral pleura (29). Although anatomic and physical basis of B-lines is not certain (16), it is widely accepted that B-lines can only be seen when the visceral pleura is opposing the parietal pleura and that their presence rules out pneumothorax, even if lung sliding is absent (30). B-lines represent the densitometer of lung parenchyma, and with their increasing number lung density increases, as measured by CT (28). The very presence of B-lines is not abnormal and a maximum of 2 B-lines per scan (longitudinal plane between two ribs) is observed at the lateral bases in 27% of healthy subjects (29). Interstitial syndrome is defined as diffuse anterolateral presence of 3 or more B-lines per scan (1). B-line distribution is another cardinal feature to consider, since focal B-pattern can be correlated with various pathologies (pneumonia, atelectasis, mass, lung contusion, lung infarct) (31), while diffuse pattern of interstitial syndrome usually reflects cardiogenic pulmonary edema or ARDS.

**Sinusoid sign**

Sinusoid sign pertains to the M-mode recording of the lung and visceral pleura respiratory movement within free pleural fluid collection (Figure 9). Sinusoid sign also implies low viscosity of the fluid, which is important during thoracentesis deliberations (19).

**Shred (fractal) sign**

Shred sign is seen with smaller subpleural consolidations visualized as hypoechoic areas separated from the rest of the lung by irregular, echogenic borders (Figure 10).

These consolidations can be of infectious origin (32), due to sub-pleural infarcts (33) or diffuse lung disease (34).

**Tissue-like sign**

Tissue-like sign (Figure 11) represents direct visualization of the lung tissue after complete loss of aeration has occurred, typically in lobar pneumonia pattern (35,36). Term sonographic hepatization of the lung effectively describes echogenicity of the consolidated lung, which is similar to that of the liver.

**Air-bronchogram**

Hyperechoic structure recognized within a tissue-like consolidation, which corresponds to air-filled airway (Figure 12). If air bronchogram is absent or static, potential lack of airway patency is suggested and/or partial to complete air resorption. Presence of dynamic air...
bronchograms (Video 4) supports regional airway patency and rules out atelectasis (37).

**Interpretation**

While sensitivity of lung ultrasound is generally robust, signs discussed above are often not specific for a diagnosis. Absent lung sliding, multiple B-lines, or lung consolidation can all be linked to various pathologic conditions and what influences interpretation the most is patient’s clinical condition. Directly imaged pathologies, such as effusions and peripherally extending consolidations are usually straightforward umbrella diagnoses, while their further etiologic classification is nuanced. Artifact interpretation is typically more challenging and is based on sequential evaluations of the distribution of different signs. The final result is a sonographic lung profile, which in an appropriate clinical context is either consistent with a diagnosis, or able to rule one out. This approach is codified in several focused assessment protocols, intended for application to a correct clinical scenario and integration with standard clinical approach.

**Sonographic diagnoses**

**Normal sonographic appearance of the lung**

Presence of lung sliding in combination with A-lines and maximum of 2 B-lines per view in anterior lung fields suggests normally aerated lung parenchyma. However, the same profile can be seen with pathologies sparing the parenchyma, such as thromboembolic or airway disease, while posterior interstitial changes can be seen in the absence of true pathology, as a result of gravity alone (1).

**Pleural effusion**

Ultrasound can easily diagnose the effusion, suggest etiology, estimate volume, and image-guide fluid removal, making evaluation of pleural effusions the most established application of ultrasound in the chest.

With regards to pleural fluid detection, ultrasound is more accurate than clinical examination (38), since it can detect as little as 3–5 mL. (39) Better sensitivity and reliability of ultrasound in comparison with portable chest X-rays is of special interest in critically ill patients (40-42). Chest radiography often discriminates poorly between pleural thickening, effusion, atelectasis, pneumonia, mass, or any combination of these findings (43,44), while ultrasonography is superior in differentiating pleural fluid from parenchymal processes and hemidiaphragm elevation (39,45). Lung ultrasound yields diagnostic accuracy of 93% when compared with chest CT for pleural fluid detection (46), while it outperforms CT for complex effusion diagnosis due to its superior ability to detect septations (47,48).

Obeying gravity, free-flowing fluid in the pleural space accumulates in the dependent part of the chest. Posterior axillary line above the diaphragm is thus the optimal site to evaluate for a non-loculated fluid collection (16). In contrast, loculated effusions do not change with body position and can occupy non-dependent areas, warranting examination of the entire hemithorax. With probe marker oriented towards the head during longitudinal scan,
The exam starts by identifying the diaphragm as a curvilinear, hyperechoic structure, which moves with breathing. Pleural fluid is then typically seen as an anechoic or hypoechoic collection above the diaphragm (Figure 13, Video 5). In case of a moderate to large effusion with adjacent parenchymal consolidation, tissue-like structure is noted flapping in the fluid collection (flapping lung or jellyfish sign—Video 6). Alternatively, aerated lung may be seen moving over the pleural effusion while covering liver and/or spleen during inspiration, which is called the curtain sign (Video 7). If cellular or proteinaceous debris suspended in the pleural fluid gets agitated by respiratory or cardiac motion, a swirling pattern called the plankton sign, is created. The presence of significant cellular debris is responsible for a layering effect caused by gravity, with hypocellular, hypoechoic top layer and cellular, hyperechoic bottom layer, which constitutes the hematocrit sign. Previously described sinusoid sign in M-mode is characteristic for a pleural effusion and can help delineate small effusions from pleural thickening, since it represents floating motion of the lung within low viscosity fluid (19).

Sonographic appearance of the pleural fluid has value in separating transudates from exudates. Four sonographic effusion patterns have been described: (I) simple, anechoic (Figure 13); (II) complex non-septated, characterized by non-homogeneous echogenic material and no septations (Figure 14); (III) complex septated, with floating fibrin strands or septae in a lattice-like pattern (Figure 15) and (IV) homogeneously echogenic (Figure 16). Transudates are almost always anechoic, while exudates can have any of the four patterns (49,50). Parapneumonic effusions, empyemas, and malignant effusions can all have complex septated appearance, whereas empyemas and hemorrhagic effusions
may appear homogeneously echogenic (51). Additional sonographic clues towards etiologic diagnosis of pleural effusion include echogenic swirling pattern and/or presence of pleural nodules which imply malignancy (52), and adjacent consolidated lung with dynamic air bronchograms which indicates infectious origin.

Despite the existence of various sonographic methods for fluid volume quantification (41,42,53,54) reliable estimation of the effusion volume remains challenging for multiple reasons (55). Inspiratory interpleural distance of at least 15 mm with effusion also identifiable in both adjacent intercostal spaces has been suggested as a feasibility marker of safe thoracentesis in mechanically ventilated patients (19).

**Pneumothorax**

In mechanically ventilated or trauma patients the importance of rapid pneumothorax recognition cannot be overstated. Supine portable chest radiographs are widely available and routinely performed for this indication, albeit with a high rate of misdiagnosis (56-58). Chest CT, the gold standard for air detection in pleural space, is highly impractical for most critical care settings, not to mention the radiation exposure. The tendency of free air to collect in non-dependent anterior chest makes the supine position ideal for ultrasound exploration of pneumothorax. Bedside ultrasound is proven to be an effective tool to rule out pneumothorax (22,25,30,59). Presence of anterior lung sliding virtually rules out pneumothorax in a severely dyspneic patient (22). In other words, in case of severe dyspnea/acute respiratory failure the presence of anterior lung sliding has a 100% negative predictive value for pneumothorax causing such symptoms. Absence of lung sliding, in contrast, is all but specific for pneumothorax and can also be attributed to a host of other pathologies (airway occlusion with mucous plug, tumor or foreign body, ARDS, pneumonia, effusion etc.). In the absence of lung sliding, one can still rule out pneumothorax by detecting either lung pulse (25) or B-lines (30). If not ruled out by these sequential evaluations in a critically ill patient with high pretest probability, it is safe to assume pneumothorax is present and proceed with urgent management without confirmatory testing (16,60). Clinical context also weighs in heavily in post-procedural settings, where if previously present lung sliding disappears on repeated evaluation, pneumothorax is highly likely. Detection of lung point confirms the presence of pneumothorax with 100% specificity, while inability to identify lung point does not rule out pneumothorax due to insufficient sensitivity (66%) (26). Notably, in cases of complete lung collapse lung point will not be seen, although lack of sensitivity can also be explained by suboptimal operator skills, incomplete examination, and unfavorable anatomy. Lung point location on the chest wall denotes pneumothorax extension. In one study, 90% of patients with lateral lung-points required drainage versus 8% of patients with anterior lung-point (59).

**Alveolar consolidation**

Alveolar consolidation extends to the pleura in overwhelming majority of cases, while 90% of consolidations may be imaged from the PLAPS point or intersection of a horizontal line at the level of the lower BLUE-point and posterior axillary line (61). Ultrasound has 90% sensitivity and 98% specificity for the diagnosis of alveolar consolidation, when compared to CT (61), and may be superior in detection of abscesses and necrotizing areas within the consolidated lung (62). Two main consolidation patterns are the smaller, non-translobar, which yields the shred sign, and the more extensive, translobar, which produces impressive image resembling anatomical lung (tissue-like sign) (61,63). Since various pathological processes (compressive or obstructive atelectasis, pneumonia, ARDS, contusion, infarction or mass) can present as consolidation, additional sonographic clues may help determine the exact etiology. Different pulmonary lesions exhibit characteristic patterns of vascularization on qualitative color Doppler sonography (64). Lung pulse is reported to have a 93% sensitivity and 100% specificity for the diagnosis of complete atelectasis following right main stem intubation in patients without previous respiratory disorders (25). Identification of dynamic air bronchograms within consolidated lung indicates bronchial patency and argues against resorptive atelectasis, thus yielding 97% positive predictive value for the diagnosis of pneumonia (37). Dynamic linear–arborescent air bronchogram seems to be a specific sign for the diagnosis of ventilator-associated pneumonia (32,65).

**Peripheral lung masses**

Malignant lesions are hypoechoic or moderately echogenic, round or oval, inhomogeneous structures with well-defined margins, often serrated or with finger-like projections into the surrounding lung tissue (Figure 17) (51). Ultrasound assessment of the chest wall invasion by the tumor has been reported to be superior to CT scan (66). Sonographic criteria
for chest wall infiltration include rib destruction, pleural line interruption, and limited respiratory excursion of the mass.

**Alveolar-interstitial syndrome (AIS)**

The AIS is a non-specific sonographic manifestation of different acute and chronic parenchymal conditions, defined as 3 or more B-lines in a single view, diffusely disseminated over the antero-lateral thorax. In direct comparison to chest radiography, diffuse sonographic B-pattern identifies diffuse radiographic AIS with 93% sensitivity and 93% specificity (29). B-line density in a single view correlates with the severity of AIS documented by chest CT. B-lines that are 7 mm apart at the pleural line imply subpleural interlobular septal thickening, while B-lines that are only 3 mm apart at the pleural line correlate with subpleural ground glass opacities (67). Differentiation among potential causes of AIS, such as cardiogenic and noncardiogenic pulmonary edema, ARDS, interstitial pneumonia and lung fibrosis depends on additional sonographic features, but even more so on clinical context. Acute cardiogenic pulmonary edema is characterized by symmetric spatial distribution and usually progresses from the dependent zones to the anterior upper chest (13). Considering that presence of B-lines in the anterior chest denotes a severe degree of cardiogenic pulmonary edema, focused anterior scanning as suggested by the BLUE protocol is highly accurate in the critically ill, but may be insufficient to diagnose less symptomatic patients (68), which would benefit from extended examination to include the lateral chest (18). Nonhomogeneous distribution of B-lines, irregular thickened pleura with subpleural and posterior consolidations favor ARDS over cardiogenic pulmonary edema in critically ill patients (69). Semi-quantification of B-lines has found many applications in the critical care arena: cardiogenic pulmonary edema monitoring (70), assessment of hemodialysis effectiveness for extravascular lung water removal (71), fluid management in patients with ARDS and septic shock (72), predicting ARDS mortality (73), defining PEEP responders (74), identifying pronation responders (75,76) and anticipating weaning failure (77).

**BLUE protocol**

BLUE protocol is an algorithm for acute respiratory failure evaluation using bedside ultrasound which, when applied to critically ill patients yields a correct diagnosis in 90% of cases (1). Protocol combines lung and venous ultrasound in rapid, sequential assessment for lung sliding, A and B-line artifacts, venous thrombosis, alveolar consolidation and/or pleural effusion, performed in 3 standardized locations per hemithorax. Combination of findings designates a profile consistent with one of the following: pulmonary edema, pulmonary embolism, pneumonia, asthma/COPD and pneumothorax.

**Ultrasound procedural guidance**

**Pleural drainage procedures**

Ultrasound reduces complications and increases the yield of thoracentesis and is integral to the management of complex pleural disease by allowing targeted placement of chest tubes. It also allows sequential sampling of fluid pockets, if pH variability is expected (78), while sonographic appearance informs the decision on whether to use tissue plasminogen activator with DNase therapy (79,80) or recommend surgical intervention.

Rate of postprocedural pneumothorax for thoracentesis without image guidance is unacceptably high (20–39%) and drops significantly when ultrasound is used to identify the optimal puncture site (81-83). When compared to
clinically marked procedural site, ultrasound reduced complication rate while increasing yield, regardless of clinician seniority (38). Superiority of ultrasound-guided thoracentesis has been confirmed by meta-analysis (84) and a large retrospective cohort study (85) and adopted into practice guidelines (86). British Thoracic Society Pleural Disease Guidelines also recommend the assistance of ultrasound when placing any chest drain for fluid, although in the absence of robust data, and mostly extrapolating from thoracentesis experience.

The optimal site, angle, and depth of needle insertion portend procedure success. Lung sliding should be documented prior to needle insertion, since its absence post procedure implies iatrogenic pneumothorax. Needle trajectory must replicate the angle and position of the transducer during image acquisition, while the depth of insertion is calculated from a frozen image of the fluid pocket by using calipers. Two basic techniques are real-time guidance and site-marking technique. With real-time guidance the position of the needle is monitored constantly, allowing true targeted placement in cases of complex effusions, but is technically more challenging and not routinely performed. As long as the site marking is done correctly, and procedure performed immediately after (to avoid fluid redistribution from patient repositioning), site-marking technique has very low complication rate, even in mechanically ventilated patients (87). Intercostal arteries are not amenable to external compression and inadvertent trauma during chest drain insertion can cause bleeding into a large negative pressure space (88). The risk of injury is greatest within 6 cm from the spinal column (89,90), justifying guideline recommendation for mid-axillary placement within the “safe triangle” and along the superior rib border. This approach does not account for anatomical variation (90), the risk still being relatively high in elderly patients with tortuous arteries (91) or when obesity and subcutaneous edema render the recognition of surface landmarks difficult (92). Ability of Color Doppler to identify the intercostal vessels prior to needle insertion was reported (93) and, after testing against the gold standard CT angiogram, recommended prior to thoracentesis, when there is an increased risk of bleeding (94). This practice is not yet widely adapted, since prospective confirmation is lacking.

Ultrasound-guided transthoracic needle biopsies (US-TTNB)

US-TTNB is typically used to obtain tissue from the chest wall, parietal pleura, lung, and sometimes diaphragm or mediastinum. The technique can also be applied to the nearby extra-thoracic structures such as supraclavicular lymph nodes. US-TTNB of pleural-based lesions performed by pulmonologists is a safe procedure (95-98), with an acceptable diagnostic yield (95-97,99-101), and represents a time-saving, cost-effective alternative to CT-guided biopsy (98,102). The pneumothorax rate is lower than for CT-guided biopsies (98,102), possibly because only peripheral lesions in direct contact with the pleura are amenable to US-TTNB. Aside from being more readily available to the critically ill patient than CT scan, a significant advantage of ultrasound is real-time guidance that allows dynamic evaluation of the target, so vascular and necrotic areas are avoided, while biopsy of smaller lesions can be conducted in favorable respiratory phase with breath holding. The use of biopsy guide simplifies the diagnostic procedure for the operator by moving the needle in the US plane, while free hand technique should be reserved for experts (103). US-TTNB has a potential role in microbiological diagnosis and/or therapeutic drainage of a pleural-based lung abscess (104), provided there is evidence of inflammatory pleurodesis at the planned access site, so that pleural contamination is avoided. The use of ultrasound guidance during closed pleural biopsy in a challenging population of patients, including cases of failed thoracoscopy, resulted in high diagnostic accuracy (105). Although not a routine bedside procedure in the ICU setting, US-TTNB in our opinion should always be considered in critically ill patients with suspected, but, undiagnosed malignancies where a tumor mass is abutting the pleura or an enlarged lymph node is identified with US. Under such circumstances US-TTNB can be performed easily at the bedside and guide therapeutic options or help with patient prognosis.

Percutaneous dilatational tracheostomy

PDT is the procedure of choice in critically ill patients (106-108), although surgical approach may be preferred in cases of unfavorable anatomical landmarks, unstable cervical spine, malignancy at the site of insertion, and emergency situations (108,109). Guidelines (108) recommend bronchoscopy use during PDT for real-time confirmation of needle entry in midline position, endotracheal tube visualization, and prevention of posterior tracheal wall injury. Reported drawbacks of bronchoscopy in this context are increase in intracranial pressure and decrease in ventilation with alveolar de-recruitment (109). Sonography can be
used during PDT as a neck screen, to identify patients unsuitable for the procedure and assist with landmark recognition if anatomy is difficult (110,111), estimate the distance from the surface of the skin to the trachea (112), and prevent puncture of aberrant vessels. In addition, real-time ultrasound-guidance may be employed for visualization of the needle and dilator entering the trachea. Bleeding is a significant concern with PDT (113). Ultrasound allows preoperative identification of aberrant neck vessels, which in one case series resulted in changing the intended tracheal puncture site in every fourth patient (114). In our practice we routinely use US preoperatively to estimate the distance from the surface of the skin to the trachea, and to identify any aberrant vessels in the field. In real-time ultrasound-guided PDT technique, a high-frequency linear transducer is used to image proximal trachea, cricoid cartilage, cricothyroid membrane, thyroid gland, and thyroid cartilage, allowing accurate puncture of the target tracheal interspace at midline (Figures 18,19).

Ultrasound-guided PDT differs from bronchoscopic-guided PDT primarily by visualization of the needle insertion, laryngoscopic endotracheal tube adjustment, and the tracheostomy tube confirmation (115). Studies comparing real-time ultrasound with bronchoscopy as PDT image guide found ultrasound guidance to be non-inferior to slightly favorable over bronchoscopy (116-118). Standalone sonographic guidance has been used in a neurosurgical population, avoiding the transient increase in intracranial pressure reported during bronchoscopy (119). However, it seemed to have an extended learning curve, with at least 50 procedures required to develop competence (120).

Education and competence in chest ultrasound

Chest sonography for both diagnostic purposes and image-guidance depends on the operator competence in image acquisition and interpretation at the point-of-care. However, there are barely any validated tools to evaluate demonstrated skills (121), no universally accepted training methodology exists, and accreditation programs are typically geared toward local clinical environments (122). Paucity of relevant literature justifies, to some extent, extrapolation from 2011 guidelines on training methodology for general critical care ultrasound (123), which were developed around American College of Chest Physicians/La Société de Réanimation de Langue Française competence statement (124). Certification programs in critical care sonography generally acknowledge the necessity of longitudinal experience needed to master ultrasound techniques (8). In 2017 Interventional Pulmonology Special Interest Group of the Thoracic Society of Australia and New Zealand developed an expert-consensus program for recognition of thoracic ultrasound competence, where competence was defined as “the ability to accurately and safely use bedside ultrasound to evaluate a pleural effusion, determine a safe site for thoracentesis or chest tube insertion, to assess pleural pathology or parenchymal lung disease and rule out a pneumothorax” (122). From IP’s perspective competency requirements should be expanded to include proficiency in performing transthoracic biopsies, vascular and anterior neck imaging, with the ultimate goal being an integrated training methodology for both the use of ultrasound and the performance of ultrasound-guided procedures.

Ultrasound limitations

Since air is a poor acoustic medium, subcutaneous
emphysema, pneumothorax and aerated lung all degrade ultrasound images and block visualization of nearby structures. Image quality may also be poor in morbidly obese or anasaric patients. Critically ill patients tend to be immobile and can be sub-optimally positioned during exam, while the presence of catheters and dressings additionally interferes with image acquisition.

**Conclusions**

Past several decades have witnessed bedside ultrasonography evolve into a cornerstone of critical care medicine, while lung ultrasound gained momentum mainly in the last decade, as an integral part of rapid evaluation algorithms for acute respiratory failure, shock and cardiac arrest. Wide application of semiquantitative sonographic techniques for assessment of lung aeration in the modern ICU keeps the momentum going. Lack of radiation exposure, cost-effectiveness, high diagnostic accuracy for actionable life-threatening conditions, and point-of-care procedural image-guidance are difficult to match by other imaging technologies. In the critically ill, less invasive advanced diagnostic and therapeutic procedures involving the chest and airways are preferred over the standard surgical approach. This trend dictates increased bedside presence of the IP who must be proficient in pleural, lung, and vascular ultrasound. Ultrasound competence for the IP's scope of practice would be best enhanced by a validated training curriculum.

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