Radiology

Fleischner Society: Glossary of Terms for Thoracic Imaging

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Conflicts of interest are listed at the end of this article.

See also the editorials by Bhalla and Powell in this issue.

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Members of the Fleischner Society have compiled a glossary of terms for thoracic imaging that replaces previous glossaries published in 1984, 1996, and 2008, respectively. The impetus to update the previous version arose from multiple considerations. These include an awareness that new terms and concepts have emerged, others have become obsolete, and the usage of some terms has either changed or become inconsistent to a degree that warranted a new definition. This latest glossary is focused on terms of clinical importance and on those whose meaning may be perceived as vague or ambiguous. As with previous versions, the aim of the present glossary is to establish standardization of terminology for thoracic radiology and, thereby, to facilitate communications between radiologists and other clinicians. Moreover, the present glossary aims to contribute to a more stringent use of terminology, increasingly required for structured reporting and accurate searches in large databases. Compared with the previous version, the number of images (chest radiography and CT) in the current version has substantially increased. The authors hope that this will enhance its educational and practical value.

All definitions and images are hyperlinked throughout the text. Click on each figure callout to view corresponding image.

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Supplemental material is available for this article.

The present glossary of terms for thoracic imaging is the fourth prepared by members of the Fleischner Society and replaces the previous glossaries of terms for chest radiography (1) and CT (2,3), respectively. The impetus to update the previous version arose from an awareness that new terms have emerged, others have become obsolete, and the usage of some terms has either changed or become inconsistent to a degree that warranted a new definition. The methodology used for compiling this glossary is described in Appendix S1. Appendix S2 summarizes the terms included, excluded, and rearranged, compared with the previous version of the glossary.

The intention of this latest glossary is not to be exhaustive but to focus on terms of clinical importance and those whose meaning may be perceived as vague or ambiguous. This focus is important to establish standardization of terminology and, thereby, to facilitate communications between radiologists and other clinicians. Moreover, the current glossary aims for more stringent, consistent terminology required by structured reporting and essential for reasonably accurate searches in large databases (4,5).

Controversial or misused terms are also included, with recommendations for appropriate usage. Whenever this is not supported by objective evidence, the consensus preference of the authors is clearly identified. In this regard, the current glossary is not only an enumeration of terms and their definitions, but also a commentary regarding their use. As in previous versions, terms and techniques not specific to thoracic imaging are not included. Despite sporadic references to fluorine 18 (¹⁸F) fluorodeoxyglucose PET/ CT, most definitions continue to refer to chest radiography and CT, which remain the chief modalities used in thoracic imaging. The overall number of terms was kept almost identical. However, several features are new.

First, the glossary preserves the alphabetical order of terms, but several related terms are now grouped under thematic headings to allow a rapid overview for comparing individual terms. The number of imaging signs was reduced, as these are comprehensively described elsewhere (6–8). Likewise, the number of eponyms was reduced to a minimum, and preference was given to descriptive terms, whenever possible. Most terms continue to have a limited number of pertinent and updated references, but not all terms are referenced.

Second, for brevity, the glossary no longer includes subdivisions into pathology, radiographic, and CT definitions. Each term is now labeled to either reflect normal anatomy (anatomy), a physiologic concept (physiology), or an abnormal finding (pathology). It is important to emphasize that this is a glossary of radiologic terms and that, while often grounded in radiology-pathology correlations, terms labeled as *pathology* do not necessarily faithfully reflect definitions used by pathologists. Some terms are labeled as descriptors to differentiate them from terms that imply a radiologic diagnosis. Terms described elsewhere are cross-referenced by an asterisk (*).

Third, the glossary includes images for virtually all terms, and we hope that this will increase its practical value. While images were chosen to be typical for a given term, they are merely examples and do not represent the full range of possible imaging appearances, which may be found in the references of this glossary and in comprehensive textbooks.

We hope that this fourth edition of the Fleischner Society Glossary of Terms for Thoracic Imaging will be helpful for both radiologists and the respiratory medicine community at large. Because terminology is evolving, the provided definitions should be understood as a consensus proposal rather than strict rules. It is anticipated that systematic evaluations of the utilization of this glossary will be undertaken.

Acinus (Pl. Acini) (Anatomy)

The acinus is a structural unit of the lung distal to the terminal bronchiole* and is supplied by a first-order respiratory bronchiole, with a diameter of approximately 4-8 mm. The secondary pulmonary lobule* contains between three and 25 acini. Within the acinus, all parts are involved in gas exchange (ie, they contain alveoli). The acinus consists of respiratory bronchioles, alveolar ducts, and alveoli. Normal acini are not visible radiologically but the accumulation of pathologic material within acini may be seen as small ill-defined rounded opacities (opacity*), sometimes called "air space nodules" (9,10).

Air Bronchogram (Pathology) [Descriptor] (Fig 1)

Air bronchogram is the descriptive term for air-filled airways^{*} within lung parenchyma^{*} that is partially or completely airless. The term implies (*a*) patency of proximal airways, and (*b*) evacuation of alveolar air by means of absorption (eg, in atelectasis^{*}) or replacement (eg, fluid, tumor), or a combination of these processes (11).

Air Crescent (Pathology) [Descriptor] (Figs 2, 3)

Air crescent is the descriptive term for a crescent-shaped air collection between the wall of a cavity^{*} and a mass^{*} within the cavity. It is often associated with the retraction of infarcted lung parenchyma^{*} in angio-invasive aspergillosis or with the colonization of preexisting cavities by *Aspergillus* or other fungi. In the latter case, the resulting fungus ball^{*} may be mobile and change position with changes in the patient's position. Air crescents can also occur due to necrotic tissue or a blood clot within a cavity^{*} (12,13).

Air Trapping (Physiology/Pathology) (Fig 4)

Air trapping is the retention of air in lung parenchyma^{*} distal to more proximal obstruction of one or several airways^{*}. Air trapping (also "gas trapping") is seen at CT performed during or at the end of expiration. It is characterized by (*a*) a lessthan-normal increase in attenuation and (*b*) a less-than-normal decrease in volume on expiration. Air trapping should only be called on expiratory CT examinations (14). Of note, areas of air trapping are typically sharply marginated, however distinction between air trapping and decreased attenuation due to vascular occlusion, for example in chronic thromboembolic disease, may be difficult (15,16). Air trapping is also frequently associated with decreased focal perfusion (eg, caused by reactive vasoconstriction) (17–19).

Airspace (Anatomy)

An airspace is the air- or gas-containing part of the lung parenchyma*, including the respiratory bronchioles (bronchiole*), alveolar ducts, and alveoli, but excluding purely conductive airways* such as terminal bronchioles. The term is used in conjunction with opacity* and nodule* to designate the accumulation of material within airspaces (20).

Airways (Anatomy)

The airways are structural parts of the respiratory system that allow airflow. Conducting airways extend from the nares and buccal opening to the terminal bronchioles (bronchiole*). Large airways include the cervical and thoracic part of the trachea, as well as the main, lobar, and segmental bronchi. Small airways include the subsegmental bronchi to bronchioles with an internal diameter of less than 2 mm. Diseases of the central airways can often be visualized directly at CT, whereas diseases of the peripheral airways often manifest through indirect changes (eg, air trapping*) (21–23).

Anatomic Distribution (Anatomy) (Figs 5–25)

- Bronchocentric: adjacent to or involving the bronchi in the axial interstitium* (Fig 5).
- Centrilobular: centered on the core structures of the secondary pulmonary lobule* (Figs 6, 7). Often used to describe the distribution of nodules (nodule*) or micronodules (micronodule*).
- Diffuse: distributed throughout a lobe* or lobes without particular regional predominance, for example between central and peripheral parts of the lung (Fig 8).
- Endobronchial: located within the lumen of a bronchus (Figs 9, 10).
- Extrapleural: external to the pleura, without directly involving the pleura* or the pleural space (Fig 11).
- Fissural: refers to a lesion or disease that arises from or involves the fissure* (Fig 12). A lesion or disease that has direct contact with the fissure*, but does not arise from it, should be called "perifissural."
- Focal: confined to one distinct area (Fig 13).
- Geographic: distributed in a manner resembling the outline of regions on a map. The term implies well-defined margins between areas of differing attenuation (Fig 14).

- Multifocal: involving two or more distinct areas (Figs 15, 16).
- Peribronchovascular: distributed along the bronchovascular bundles (Figs 17, 18).
- Perilobular: Arcade-like or curvilinear opacities that can be related to the margins of secondary pulmonary lobules (Figs 19, 20).
- Perilymphatic: distributed along lymphatic vessels (Figs 21, 22).
- Pleural: refers to a lesion or disease that arises from or involves the pleura*. A lesion or disease that has contact with the pleura*, but does not arise from it, should be called *pleura-based*. The term *subpleural* should only be used for lesions or diseases in proximity to the pleura*, but without direct contact to it. Because this term has been attributed variable meanings in the literature, the term *peripheral* is preferred to describe such lesions or diseases. Of note, imaging cannot always determine whether a given lesion is located between two layers of pleura or within the pleura, or in the parenchyma and directly contiguous to the pleura (Figs 23–25).

The prefix *juxta*- has been used inconsistently to describe lesions that are variously in proximity or in contact with the pleura* or a fissure* (juxtafissural, juxtapleural) (24). We recommend instead using the more stringent terminology described above.

Aortopulmonary Window (Anatomy) (Fig 26)

The aortopulmonary window is a mediastinal compartment delimited anteriorly by the ascending aorta, posteriorly by the descending aorta, cranially by the aortic arch, caudally by the left pulmonary artery, medially by the trachea and the lateral wall of the left main bronchus, and laterally by the medial pleura* covering the left lung. The aortopulmonary window contains lymph nodes (lymph node*) and connective tissue and is traversed by the left phrenic nerve, the left vagus nerve, the left recurrent laryngeal nerve, and the left bronchial arteries. It is separated into a medial and lateral part by the ligamentum arteriosum (25).

Apical Cap (Pathology) (Fig 27)

Apical caps are cap-like rims at the lung apices, often bilateral, usually caused by pulmonary and/or pleural fibrosis* displacing extrapleural fat. Chronic ischemia causing a plaque of elastotic fibrosis of the visceral pleura*, with or without subpleural lung parenchyma*, has been suggested as the potential pathogenesis, which is variably associated with aging. In addition, apical caps can be caused by fluid collections associated with infection, neoplasms, or trauma, either extrapleural or within the pleural space. A left apical cap can be caused by hematoma, such as resulting from aortic rupture (26–28). Moderate or severe apical pleural caps also can be seen in upper lobe fibrosis, including pleuroparenchymal fibroelastosis (29).

Architectural Distortion (Pathology) (Fig 28)

Architectural distortion is characterized by the focal or diffuse derangement of the normal pulmonary anatomy, with abnormal displacement or disruption of airways^{*}, vessels, and interstitium^{*}. It is typically accompanied by volume loss^{*} and is commonly seen in fibrotic lung diseases (30,31).

Asbestosis (Pathology) (Fig 29)

Asbestosis is an occupational lung disease characterized by lower-lobe predominant pulmonary fibrosis caused by inhalation of asbestos fibers. Once established, fibrosis can progress. However, even without progression, the risk of lung cancer is increased. In addition, asbestos exposure increases the risk of mesothelioma. However, asbestosis is usually associated with high levels of asbestos exposure, in distinction to mesothelioma. Although pleural plaques, calcified or noncalcified, may also be identified, their presence is not essential for the diagnosis (pleural plaque*) (32,33).

Aspiration (Pathology)

Aspiration is the unintended inhalation of solid and/or fluid substances ("aspirate"). Depending on its location and extent within the airways*, the aspirate can cause atelectasis*, inflammation ("aspiration pneumonitis/bronchiolitis"), or infection ("aspiration pneumonia"). Changes caused by aspiration are most commonly seen in the posterior portions of the lungs and predispose to infection with anaerobic bacteria. Sequelae of aspiration, including scarring and bronchiectasis, can be seen in variable anatomic locations but tend to follow the direction of gravity (34,35).

Atelectasis (Pathology) (Figs 30-40)

Atelectasis is a partial or complete collapse* of the lung. Atelectasis can be caused by loss of negative pressure in the pleural space and/or by compression of lung parenchyma* (eg, by pleural effusions*), by resorption of air following obstruction (eg, by a foreign body or a neoplasm), or by an increase in alveolar surface tension (eg, by surfactant dysfunction). Atelectasis shows lower volume and higher attenuation than normal lung tissue and is often associated with abnormal displacement of fissures (fissure*), bronchi, vessels, hyperinflation of adjacent lobes (lobe*), or obscuration of cardiac and mediastinal interfaces (36,37).

- Lobar: atelectasis of an entire pulmonary lobe* (Figs 30-37).
- Linear (formerly "plate-like" or "band-like"): descriptor for local atelectasis with a linear, plate-like, or bandlike shape (Figs 38, 39).

Often seen in dependent lung regions, paralleling the hemidiaphragms on chest radiographs and the chest wall on CT examinations. Of note, the geometric definition of "line" refers to a one-dimensional structure. While the cross-sectional appearance of an atelectasis may appear linear, the atelectasis itself has greater dimensionality than a line. Rounded: rounded portion of a collapsed lung, associated with invaginated fibrotic pleura* and thickened and fibrotic interlobular septa (interlobular septum*) (Fig 40).

Rounded atelectasis is the consequence of chronic or recurrent pleural effusions, as seen in asbestos-related disease, chronic cardiac or renal failure, or thoracic malignancies. At imaging, rounded atelectasis presents as a rounded mass* abutting the pleura, associated with adjacent pleural thickening. Distorted vessels and bronchi converging toward the atelectasis can have a curvilinear appearance ("comet tail sign"). It is most commonly seen in the posterior lower lobes.

- Segmental: atelectasis of an anatomical segment* of the lung.
- Subsegmental: atelectasis of an anatomical subsegment of the lung, which often appears linear.

Atoll Sign (Pathology) [Descriptor] (Figs 41, 42)

The atoll sign is the descriptive term for a focal ground-glass opacity^{*} surrounded by a partial or complete ring of consolidation^{*}. It is a nonspecific sign originally described in organizing pneumonia^{*} and reported subsequently in various other lung diseases (38). A synonymous term is the "reversed halo" sign.

Azygoesophageal Recess (Anatomy) (Figs 43, 44)

The azygoesophageal recess is the right posterior mediastinal recess into which the medial part of the right lower lobe* extends. It is limited cranially by the azygos arch, posteriorly by the azygos vein and the pleura* anterior to the vertebral bodies, and medially by the esophagus. On frontal radiographs, the medial part of the recess can be seen as a vertically oriented interface between the right lower lobe and the esophagus, and the superior part of the recess is seen as a smooth arc with convexity to the left at the interface with the azygos vein. Distortion or obliteration of the interface can reflect disease (eg, adenopathy or neoplasm). At CT, the recess are often invisible on radiographs (39).

Bleb (Pathology)

A bleb is a circumscribed air-containing cystic* structure, exclusively pleural in location and usually subcentimeter in size. Blebs have been associated with small airways disease* and the occurrence of pneumothorax, notably in patients with active tobacco use (40). Small blebs are more readily appreciated with pathology than with imaging.

Blood Flow Redistribution (Pathology)

Blood flow redistribution is the nonspecific term for changes in blood flow, seen on both radiographs and CT examinations, reflecting regional redistribution resulting from pathologic conditions such as volume overload or chronic heart failure (41).

Bronchiole (Anatomy)

Bronchioles are non–cartilage-containing peripheral airways^{*}. Terminal bronchioles are the most distal of the purely conducting airways. They give rise to respiratory bronchioles, from which alveoli arise. Respiratory bronchioles allow gas exchange, and branch into multiple alveolar ducts. Normal bronchioles are not visible using current CT techniques; however, in diseases of the peripheral airways, bronchioles with thickened walls and/ or plugged bronchioles can become visible as nodular opacities (opacity^{*}) on radiographs and/or as tree-in-bud^{*} opacities on CT examinations. Infection and inflammation of the bronchioles are commonly referred to as "bronchiolitis" (9,42).

Bronchiectasis (Pathology) (Figs 28, 45)

Bronchiectasis indicates a clinical condition of irreversible bronchial dilatation. In the absence of established chronicity, we recommend using the term "bronchial dilatation." Bronchiectasis can be secondary to chronic inflammation, congenital, or caused by chronic infection and obstruction of more central airways*. It is often accompanied by bronchial wall thickening and/or mucoid impaction*. At CT, bronchiectasis manifests as dilatation of the bronchus relative to the accompanying pulmonary artery (signet ring sign*), lack of tapering, and presence of visible bronchi within 1 cm of the pleural surface ("tram-tracking"). Bronchial walls can be thickened and accompanied by scarring of the adjacent parenchyma (43,44).

- Traction bronchiectasis (pathology): Dilatation of the bronchial lumen associated with thickened, irregular* bronchial walls, caused by fibrosis* (45). This term should only be used in the context of fibrosis (Fig 28).
- Traction bronchiolectasis (pathology): Dilatation of bronchioles (bronchiole*) within the central portions of the secondary pulmonary lobule* typically associated with fibrosis* and honeycombing* (46).

Bronchocele (Pathology) (Fig 46)

A bronchocele is a focal bronchial dilatation, often with mucoid impaction^{*}. Bronchoceles can be either congenital (eg, in bronchial atresia) or caused by retained secretions or proximal obstruction (eg, by an endobronchial tumor). At CT, bronchoceles are typically branching V- or Y-shaped structures that may have a finger-like appearance. In bronchial atresia, the surrounding lung parenchyma^{*} can have low attenuation because of reduced ventilation and, thus, decreased perfusion (3,47,48).

Broncholith (Pathology) (Fig 47)

A broncholith is calcified material within the bronchial lumen, generally caused by erosion of a peribronchial lymph node* into the lumen, and most commonly a result of previous granulomatous infection. Broncholiths manifest when calcified lesions directly adjacent to or within the bronchial wall erode into the lumen; they can be accompanied by distal atelectasis*, bronchiectasis*, or mucoid impaction* (49–51).

Bronchomalacia (Pathology) (Fig 48)

Bronchomalacia is the mechanical weakening of the bronchial cartilage, either congenital or caused by chronic inflammation and damage, leading to excessive bronchial collapsibility, of approximately 70%–80% of the original luminal diameter, and subsequent ventilatory impairment. Excessive collapsibility is best confirmed with dynamic CT performed during forced expiration. Bronchomalacia is often associated with the same phenomenon in the trachea (tracheomalacia*) and can be accompanied by thickening of the airway walls. Of note, a substantial degree of collapse may also occur in healthy individuals (52,53).

Bulla (Pl. Bullae) (Pathology) (Figs 49, 50)

A bulla is a circumscribed air-containing cystic* structure in the lung parenchyma* lined by a thin layer of collapsed lung parenchyma. It is caused by dilatation, destruction, and confluence of airspaces (airspace*) distal to terminal bronchioles (bronchiole*). At imaging, a bulla typically appears as a rounded lucency, with a very fine and usually smooth wall (54).

Cavity (Pathology) (Figs 51-53)

A cavity is an abnormal gas- or fluid-filled structure with a typically thick and often irregular* wall, usually produced by the expulsion or drainage of a necrotic part of the lesion via the bronchial tree. It may be isolated or associated with adjacent lung pathology, and a gas-fluid level may be present in the cavity (54,55). Cavities are to be differentiated from pseudocavities (56). A pseudocavity appears as an area of low attenuation in nodules (nodule*), masses (mass*), or consolidations (consolidation*) that represent spared parenchyma, normal or ectatic bronchi (bronchus*), or emphysema (emphysema*), rather than cavitation.

Collapse (Pathology)

In the lung, synonymous with atelectasis*.

Consolidation (Pathology) [Descriptor] (Fig 54)

Consolidation refers to the increased attenuation of lung parenchyma* resulting from evacuation of air from the alveoli, and its replacement by fluid or other material. On radiographs, consolidation appears as a homogeneous increase in attenuation. At CT, it is defined as opacity* causing complete obscuration of the underlying bronchi and vessels, the latter only on unenhanced CT examinations. Patent bronchi within consolidation may form air bronchograms*. The term is a descriptor that does not indicate the pathologic nature of the condition causing the consolidation.

Cyst (Pathology) (Fig 55)

A cyst is any circumscribed and well-defined air-containing structure in the lung parenchyma^{*}. Cysts can originate from mechanisms such as airway obstruction with distal airspace^{*} dilatation (check-valve mechanism), necrosis of airway^{*} walls, or lung parenchymal destruction by proteases. A small number of lung cysts may be seen with aging (57). At imaging, a cyst appears as a circumscribed and well-defined radiolucency and/or low attenuation area with a thin and usually regular wall. Most cysts contain air but they can occasionally contain fluid or solid material. Of note, the radiologic and pathologic definitions of "cyst" differ (54,58,59).

Cystic (Pathology) [Descriptor] (Figs 49, 50, 55)

Cystic is the descriptive adjective designating lesions characterized by central air-equivalent attenuation, surrounded by a wall of variable thickness and regularity. There is unavoidable morphologic overlap between structures commonly described as "cystic" (bulla*, bleb*, cyst*), due to the lack of objective criteria to quantify the wall thickness of these structures, and because of a multitude of currently proposed definitions, rather than a decisive consensus in the literature. As the cause of cystic lesions is often difficult to determine at CT, the term "cystic" can be used as a descriptor without implying etiology.

Embolism (Embolus, Emboli) (Pathology) (Fig 56)

Embolism refers to the pathologic lodging of partially or completely obstructing material (embolus) inside of a blood vessel. The embolus may be a blood "clot" (ie, a thrombus), fat (fat embolism), air or another gas (gas embolism), amniotic fluid (amniotic fluid embolism), infectious material (septic embolism), neoplastic tissue (tumor embolism), or a foreign body. On ventilation-perfusion scans, areas affected by embolism appear as a "mismatch" between the ventilation and perfusion scans, with a ventilated perfusion defect (60,61). Partial or complete vascular occlusion caused by embolism may cause necrosis of the lung distal to the embolus (infarction*) (62,63).

Emphysema (Pathology) (Figs 57-64)

Emphysema is characterized by irreversible enlarged airspaces (airspace*) distal to and originating from the terminal bronchioles (bronchiole*), with destruction of alveolar walls (64). The absence of fibrosis was historically regarded as an additional criterion, but this is no longer the case because local fibrosis is now recognized to coexist with emphysema both histologically and at imaging. Emphysema is pathologically classified according to the anatomic area predominantly affected (65,66).

- Centrilobular: Focal emphysematous destruction that originally affects the center of the secondary pulmonary lobule* and, if progressive, can eventually involve the entire lobule*. Even if severe, centrilobular emphysema tends to respect the borders of the lobule, which are not destroyed. On CT examinations, early centrilobular emphysema is characterized by small areas of low attenuation surrounded by normal lung, without clearly defined walls, often with upper lobe predominance. When progressive, larger areas of low attenuation can span several secondary pulmonary lobules (Figs 57–59).
- Panlobular: Diffuse emphysematous destruction across the secondary pulmonary lobule*, the structures of

which are effaced, sometimes with lower lobe predominance. On CT examinations, panlobular emphysema is characterized by larger and/or diffuse areas of low attenuation, where components of the normal lung architecture, and notably those of the secondary pulmonary lobule*, are no longer recognizable (Figs 60, 61).

Paraseptal: Focal emphysematous destruction of the distal acinus*, typically located near the pleural surface close to the chest wall, the mediastinum*, and interlobar fissures (fissure*). On CT examinations, paraseptal emphysema is characterized by foci of low attenuation separated by intact interlobular septa (interlobular septum*) thickened by associated mild fibrosis*. Architectural distortion* is typically absent (Figs 62, 63).

In addition to the pathologic definition of emphysema, visually defined CT subtypes of emphysema have been described (65) (Fig 64). The terms "interstitial emphysema" and "subcutaneous emphysema" are sometimes used to designate abnormal air collections in the pulmonary interstitium* and the soft tissues, respectively, as may result from barotrauma. These abnormalities are distinct processes and are not related to the destruction of lung parenchyma.

Empyema (Pathology) (Figs 65, 66)

Empyema is a collection of pus in the pleural space, most often occurring as a complication of pneumonia* or esophageal perforation. Empyema is commonly unilateral. On CT examinations, the fluid collection may contain locules of gas, and the pleura* is thickened and enhances after intravenous contrast administration. At the margins of the empyema, the pleura can be seen dividing into its visceral and parietal layers. This is referred to as the "split pleura sign," which can occasionally help to distinguish an empyema from a parapneumonic effusion or a peripheral lung abscess (67–69).

Fibrosis (Adj. Fibrotic) (Pathology) (Fig 67)

Fibrosis is the general term for a repair mechanism of the lung in which parenchyma* is permanently replaced by connective tissue, causing remodeling, architectural distortion*, and volume loss. Other CT signs of fibrosis include traction bronchiectasis* or bronchiolectasis* and, potentially, honeycombing*. Fibrosis can be the result of mechanical, inflammatory, infectious, or iatrogenic damage. It can be focal or diffuse in distribution and is often accompanied by functional impairment.

Fissure (Anatomy) (Figs 68-72)

A fissure is a structure consisting of two layers of infolded visceral pleura* that separates lobes (lobe*) (Fig 68). The left major (oblique) fissure separates the left upper lobe* and the left lower lobe. The right minor (horizontal) fissure separates the right upper lobe and the middle lobe. The right major (oblique) fissure separates the right lower lobe from the right upper and middle lobes. On CT examinations, normal fissures are seen as thin and regular lines (70–73).

- Accessory: Supernumerary fissures usually separating segments (segment*) rather than lobes (Figs 69, 70).
- Azygos: Developmental variant consisting of a fold of pleura* pulled through the lung towards the hilum* by a non-migrating right posterior cardinal vein, one of the embryological precursors of the azygos vein. Because the azygos fissure has two visceral and two parietal layers, it is in fact a pseudofissure (Fig 71).
- Incomplete: Partial absence of a fissure between two lobes. Incomplete fissures allow collateral ventilation between two lobes. Therefore, their detection is relevant, for example before emphysema* treatment with endobronchial occlusive devices (Fig 72).

Ground-Glass (Physiology/Pathology) (Figs 73, 74)

Ground-glass refers to an area of increased attenuation that does not completely obscure the underlying bronchial and vascular structures. Although historically first described on radiographs (74), specifics of this term have been defined for CT and it should only be applied in the context of CT. Accurate identification of a ground-glass opacity* requires thin CT sections (equal to or thinner than 1.5 mm) acquired at full inspiration, preferably reconstructed with a high spatial frequency algorithm, and displayed with wide window widths. The finding is nonspecific, as ground-glass can be caused by any abnormal process below the resolution of the CT scanner that does not completely displace alveolar air, for example, partial filling of alveoli by fluid or cellular material, an increase in capillary blood volume, histologic fibrosis, a lepidic pattern of adenocarcinoma, or a combination of the above (75).

Halo (Pathology) (Fig 75)

A halo is a rim of ground-glass opacity* surrounding a nodule*, mass*, or consolidation*. The presence of a halo is nonspecific but often reflects local hemorrhage around an infectious or neoplastic lesion (eg, invasive aspergillosis) or lepidic tumor growth (eg, primary adenocarcinoma) (76,77).

Hilum (Pl. Hila, Adj. Hilar) (Anatomy)

The hilum is the site at the medial aspect of the lung, which is not covered by visceral pleura*, adjacent to the mediastinum*, where vessels and bronchi enter and leave the lung. The hilum contains bronchi, arteries, veins, lymphatic vessels, lymph nodes, nerves, and other structures (78,79).

Honeycombing (Pathology) (Figs 76–78)

Honeycombing represents the destruction of lung parenchyma* with loss of architecture and well-defined adjacent cystic* structures, typically clustered in the subpleural region. The traditional definition requires several layers of cysts (cyst*), but in current practice, this threshold has been reduced to a single layer of cysts, provided other signs of fibrosis are present (30,80). The diameter and wall thickness of the cysts are variable but their presence is inevitably accompanied by the complete effacement of normal lung architecture and usually occurs with other signs of pulmonary fibrosis^{*}, including traction bronchiolectasis^{*}. Of note, honeycombing can also exist at the microscopic level and does not necessarily reflect the CT appearance of honeycombing (81). Care should be exercised when using the term *honeycombing*, as it implies substantial fibrosis (30,82), and the term should only be used if other signs of fibrosis are present.

Infarction (Pathology) (Fig 56)

Infarction is the result of parenchymal damage from vascular obstruction or occlusion, caused by either emboli or pathologic changes of the vessel wall. At imaging, a pulmonary infarct is typically a triangular, dome-, or wedge-shaped opacity*, with the base abutting the pleura* and the apex directed toward the hilum*. The opacity reflects local edema, hemorrhage, necrosis, late organization and/or scarring, or a combination of these, depending on the timing of imaging during the natural history of the vascular occlusion (embolus*). However, many "infarcts" seen in acute pulmonary embolism* are due to edema and hemorrhage without necrosis and hence are not true infarcts.

Interlobular Septum (Anatomy)

See septum* (9,83).

Interstitial Lung Abnormalities (Pathology)

Interstitial lung abnormalities, or ILAs, is an umbrella term used for often subtle interstitial CT abnormalities in nondependent parts of the lungs, detected incidentally. They often manifest as variable combinations of peripheral ground-glass*, and reticular opacities, which should affect more than 5% of the upper, middle, or lower lung zone, as demarcated by the levels of the inferior part of the aortic arch and the right inferior pulmonary vein, respectively. The term is not necessarily associated with the presence of respiratory symptoms (84,85).

Interstitial Lung Disease (Pathology)

Interstitial lung disease is the umbrella term used for a large group of lung diseases that involve the interstitium and are typically associated with restrictive pulmonary impairment. These conditions are in fact very often multicompartmental and may then also involve the pulmonary airspaces (airspace*), airways*, and vasculature to varying degrees, and some may result in pulmonary fibrosis* (86–88).

Interstitium (Anatomy)

The interstitium consists of a continuum of connective tissue, extending from the most central to the most peripheral components of the lung parenchyma^{*}, which mechanically supports airways^{*}, vascular structures, and alveoli. Anatomically, it forms a single interconnected space encompassing the entire lung. For didactic reasons, it is divided into three compartments (10).

• Axial: Portion of the interstitium that envelops airways*, vascular structures, and lymphatic vessels, the combination of which are sometimes referred to as "bronchovascular bundle."

- Interlobular (peripheral): Portion of the interstitium that extends from the visceral pleura* and forms the envelope of the secondary pulmonary lobule* and is visible as interlobular septa (septum*) on CT examinations.
- Intralobular (septal): Portion of the interstitium within the secondary pulmonary lobule* that serves as the support structure for the alveolar walls.

Irregular (Anatomy/Pathology) [Descriptor] (Fig 79)

Irregular is a general term describing the margination of a structure (eg, a pulmonary nodule* or its components) as not smooth and/or uneven. More precise descriptors falling under this general term include lobulation* and spiculation*.

Juxtaphrenic Peak (Anatomy) (Fig 80)

A juxtaphrenic peak is a small triangular opacity* based on either the right or the left hemidiaphragm, with its tip pointing toward the lung apex, caused by upward retraction of an inferior accessory fissure* or an intrapulmonary septum* associated with the pulmonary ligament (pleura*). The retraction reflects volume loss in the ipsilateral upper lobe, for example after radiation or surgery. The peak is best visualized on frontal radiographs and coronal CT reconstructions (89–91).

Lobe (Anatomy)

The lobes are the principal divisions of the lungs. The right lung has three lobes and the left lung has two lobes. Each lobe is enveloped by visceral pleura*, except at the hilum* or when an interlobar fissure* is incomplete (92).

Lobulation (Anatomy/Pathology) [Descriptor] (Fig 81)

Lobulation is the descriptive term referring to a lobule-like and often asymmetric protrusion at the margins of a structure (eg, a pulmonary nodule* or its components). Pathologically, lobulations often reflect different rates of tissue growth or the presence of different tissues within the same structure, and typically have smooth margins. Although lobulation is often seen in malignant nodules (nodule*) or masses (mass*), it can also occur in benign lesions.

Lobule (Anatomy)

See secondary pulmonary lobule*.

Lymph Node (Anatomy) (Figs 82–84)

Lymph nodes are organs of the lymphatic system surrounded by a fibrous capsule and divided into an outer cortex and inner medulla. The hilum* is an indentation on the concave surface of the lymph node where lymphatic vessels exit, and blood vessels both enter and exit. Normal lymph nodes are best seen on CT examinations where they appear as rounded, oval, or bean-shaped structures, sometimes with a visible fatty hilum. For purposes of cancer staging, mediastinal, hilar, and intrapulmonary nodes are classified according to anatomic location ("stations"). Lymph nodes present in the lung parenchyma* are called pulmonary or intrapulmonary lymph nodes; they are typically lobular or triangular in outline and occur along pleural surfaces, interlobular septa (interlobular septum*), and at vascular branch points. Aggregates of lymphatic tissue without a capsule, as commonly seen along pleural fissures (fissure*), are, strictly speaking, not lymph nodes (93).

Lymphadenopathy (Pathology) (Fig 85)

Lymphadenopathy refers to any pathologic transformation of lymph nodes (lymph node^{*}) with regard to size, shape, and morphology. In more common usage, the term refers to the abnormal enlargement of lymph nodes. Therefore, the synonym "lymph node enlargement" is more precise and thus preferable. On CT examinations, and depending on the clinical scenario, the somewhat arbitrary threshold of either 10 mm or 15 mm short-axis diameter is often regarded as the upper limit of normal. Because lymph nodes commonly show a wide range of sizes according to location, differentiation between normal and diseased lymph nodes based on size alone is unreliable and should be complemented by consideration of location and morphologic features, or with the assessment of metabolic activity by ¹⁸F fluorodeoxyglucose PET/CT (94,95).

Mass (Pathology) (Fig 86)

A mass is any circumscribed lesion greater than 30 mm in diameter. The term usually refers to a predominantly solid soft tissue structure but cavitary, cystic*, or calcified components may be present to various degrees. Shape and margin descriptors are similar to those used for pulmonary nodules (nodule*). In addition, distortion or invasion of adjacent structures may be present. The adjectival term "mass-like," for example, used in "masslike consolidation*," may be used to describe the appearance of a given lesion, without implying a neoplastic etiology (also see consolidation*).

Mediastinum (Anatomy)

The mediastinum is the anatomic space located between the lungs and extending from the thoracic inlet to the diaphragm. The current definition of mediastinal compartments is based on CT criteria and includes the prevascular (anterior), visceral (middle), and paravertebral (posterior) compartments.

Boundaries of the prevascular (anterior) compartment are superiorly, the thoracic inlet; inferiorly, the diaphragm; anteriorly, the sternum; laterally, the parietal mediastinal pleura*; and posteriorly, the anterior aspect of the pericardium as it wraps around in a curvilinear fashion. Thus, any vessels contained within the pericardium are located in the visceral (middle) mediastinum. The major contents of the prevascular compartment are the thymus, fat, lymph nodes (lymph node*), and the left brachiocephalic vein.

Boundaries of the visceral (middle) mediastinum are superiorly, the thoracic inlet; inferiorly, the diaphragm; anteriorly, the anterior aspect of the pericardium (which envelops the distal aspect of the superior vena cava, the proximal aspect of the ascending aorta and lateral rim of the aortic arch, and the intrapericardial pulmonary arteries); posteriorly, a vertical line connecting a point on the thoracic vertebral bodies 1 cm posterior to the anterior margin of the spine. The major contents of this compartment fall into two main categories: first, a vascular category (ie, heart, superior vena cava, ascending thoracic aorta, aortic arch, descending thoracic aorta, intrapericardial pulmonary arteries, thoracic duct), and second, the trachea, carina, and esophagus, which share an embryological origin (the endoderm), as well as lymph nodes.

Boundaries of the paravertebral (posterior) compartment are superiorly, the thoracic inlet; inferiorly, the diaphragm; anteriorly, the visceral compartment; and posterolaterally, a vertical line along the posterior margin of the chest wall at the lateral aspect of the transverse processes. The major contents of this compartment are the thoracic spine and paravertebral soft tissues (96,97).

Micronodule (Pathology) (Figs 21, 22, 87)

A micronodule is a circumscribed and typically round opacity*, previously defined as less than 3 mm, but currently defined as less than 6 mm in average diameter. This size threshold serves to distinguish micronodules, if found incidentally, from larger potentially actionable nodules, consistent with current management guidelines (98). In patients with a high risk for cancer or with an underlying history of cancer, micronodules can become actionable, notably if newly appeared. Micronodules can show the same shape and margin, as well as structure and location characteristics as nodules (nodule*). However, in the case of micronodules, these characteristics can be difficult or impossible to assess accurately because of their small size (98,99). Of note, the concept of individual micronodules is different from the concept of a micronodular pattern*, described elsewhere in the text.

Mosaic Attenuation (Pathology) (Fig 88)

For the description, see pattern, mosaic*. The mosaic attenuation CT pattern* can be caused by obliterative small airways disease*, occlusive vascular disease, multifocal interstitial lung disease*, or by a combination of the above. In obliterative small airways disease and in occlusive vascular disease, the lower attenuation areas are abnormally hypodense and the higher attenuation abnormally hyperdense, reflecting differences in vascular perfusion, whereas in multifocal interstitial lung disease* the lower attenuation areas are normal and the higher attenuation areas abnormal. Mosaic attenuation should only be described on CT examinations performed in full inspiration. "Mosaic perfusion" refers to the mosaic pattern caused by local variations in lung perfusion (14,16,100,101).

Mucoid Impaction (Pathology) (Fig 89)

Also referred to as "mucus plugging," mucoid impaction represents airway filling by mucoid secretions and can be obstructive or nonobstructive. It is often combined with signs of chronic airways* disease (102,103).

Mycetoma (Syn. Fungus Ball) (Pathology) (Figs 90, 91)

A mycetoma is a mass* of intertwined hyphae, usually of an *Aspergillus* species, matted together by mucous, fibrin, and cellular debris, colonizing a preexisting cavity*. At imaging, the mycetoma appears as a rounded structure within the cavity, often showing an air crescent* between the mass and the inner wall of the cavity. On CT examinations, mycetoma can have a sponge-like structure and calcifications may be present. Mycetomas are often mobile and changes in position within the cavity can be confirmed by imaging the patient in different body positions (eg, supine, prone) (104,105).

Nodule (Pathology) (Figs 92–114)

A nodule is a circumscribed, typically round opacity*, less than or equal to 30 mm in average diameter. A rounded lesion larger than 30 mm is referred to as a mass*. A rounded lesion less than 6 mm in diameter should be called micronodule*, a term which serves to distinguish it from larger potentially actionable nodules, consistent with current management guidelines (98,99). The terminology described below is intended to standardize the description and reporting of nodules, while acknowledging that nodules represent a morphologic spectrum, encompassing both benign and malignant features (106,107).

- Shape and margin: Nodules can be described as round (Fig 92), ovoid (Fig 93), or polygonal in shape, and as spherical, flat, or elongated in configuration, based on their appearance in various imaging planes. The margin of a nodule can be well-defined (Fig 94) or ill-defined (Fig 95), depending on the appearance of the interface with the surrounding lung. A smooth nodule border is even and free of substantial protrusions or indentations, as opposed to an irregular* nodule border. Irregular* borders can be lobulated (Fig 81), with lobule-like protrusions (lobulation*), and/or spiculated (Figs 86-89), with multiple radiating small sharp projections (spiculation*). Linear components extending from the nodule to a pleural surface are called "pleural tags" (pleural tag*) (Fig 99). Combinations of the above shapes and margins may occur (Figs 100, 101).
- Attenuation and structure: Based on *attenuation*, nodules with pure ground-glass* attenuation are referred to as "ground-glass" nodules (Fig 95). Nodules with pure soft tissue attenuation are referred to as "solid" nodules (Figs 92, 94). Nodules with both ground-glass and soft tissue attenuation components are referred to as "part-solid" nodules (107). In the context of lung adenocarcinoma, the ground-glass compo-

nents generally correspond to a lepidic growth pattern and solid components generally correspond to invasive growth patterns. The terms ground-glass, solid, and part-solid nodule do not necessarily encompass the entire complex morphology of pulmonary nodules and are also limited by low inter- and intra-observer reproducibility among radiologists (108,109). Nonetheless, these terms are established in the literature and it is likely that they will continue to be used despite their limitations, notably for purposes of management and big data categorization. Note that thin-section CT is essential for the accurate characterization of small nodules. Based on structure, nodules are either "simple" or "complex." Simple nodules show a single structural component (soft tissue, ground-glass*, calcium, fat) (Figs 93, 95, 102). Complex nodules have two or more structural components (soft tissue, ground-glass, calcium, fat, air) (Figs 103-110). Of note, simple nodules can develop into complex nodules (Figs 111-113), and vice versa (Fig 114).

Location: Nodules can be described as peribronchovascular or centrilobular, depending on their location in relation to other structures (see anatomical distribution*). Nodules arising from a fissure* should be called "fissural." Nodules with direct contact to a fissure*, but not arising from it, should be called "perifissural." Nodules arising from the pleura* should be called "pleural." Nodules with direct contact to the pleura*, but not arising from it, should be called "pleura-based." The term "subpleural" should only be used for nodules in proximity to the pleura, but without direct contact to it. Because this term has been attributed variable meanings in the literature, the term "peripheral" is preferred to describe such nodules.

The prefix *juxta*- has been used inconsistently to describe nodules that are variously in proximity or in contact with the pleura* or a fissure* (juxtafissural, juxtapleural) (24). We recommend instead using the more stringent terminology described above.

Oligemia (Anatomy/Physiology/Pathology) (Figs 115, 116)

Oligemia is a focal or diffuse reduction in blood flow and/or blood volume. At imaging, pulmonary oligemia appears as a decrease in attenuation, and/or as a decrease in dimension and number of pulmonary vessels. Imaging signs of oligemia can be subtle and may require comparison with normally perfused lung for accurate detection (110).

Opacity (Anatomy/Pathology) [Descriptor] (Fig 74)

Opacity refers to any focal or diffuse nonspecific area of increased attenuation. The term is a general descriptor that does not indicate the nature of the condition causing the opacity. The term has sometimes been used synonymously with the now obsolete and nonrecommended term "infiltrate" (111).

Organizing Pneumonia (Pathology) (Figs 117, 118)

Organizing pneumonia is a histologic, clinical, and radiographic entity, reflecting a repair mechanism of the lung parenchyma* after exogenous or endogenous injury. Histologically, organizing pneumonia is characterized by loose plugs of organizing connective tissue in distal airways* and alveoli. The imaging presentation is variable, with sharply defined peripheral or bronchocentric consolidation* with or without ground-glass*, air bronchograms*, curvilinear perilobular opacities (opacity*), nodular opacities, and the atoll sign* being common manifestations. However, numerous additional manifestations and patterns (pattern*) have been described, including peripheral sparing (112).

Paratracheal Stripe (Anatomy) (Fig 119)

- Right paratracheal stripe: The interface between the right tracheal wall, the adjacent pleura*, and the mediastinal fat between them, as seen on frontal chest radiographs. It begins superiorly at the level of the clavicles and extends inferiorly to the right tracheobronchial angle at the level of the azygos arch.
- Left paratracheal stripe: The interface between the left upper lobe* and either the mediastinal fat adjacent to the left tracheal wall or the left tracheal wall itself. It extends superiorly from the aortic arch to join with the reflection from the left subclavian artery.

Thickening of the stripe on either side may reflect enlarged lymph nodes (lymph node*), hematoma, or another mediastinal pathology (113,114).

Parenchyma (Anatomy)

Parenchyma refers to the part of the lung tissue involved in gas transfer. The lung parenchyma includes the alveoli, alveolar ducts, and respiratory bronchioles (bronchiole*).

Parenchymal Band (Pathology) (Fig 120)

A parenchymal band is a thin linear opacity paralleling the pleura*, occasionally described as extending to the pleural surface. It can be associated with architectural distortion* and is commonly seen in the context of asbestos exposure (115). More recently, it has been described as a sequela of COVID-19–related pneumonia* (116) and organizing pneumonia* (117).

Pattern (Anatomy/Pathology) (Figs 121–136)

- Crazy paving: Focal areas of ground-glass opacities (opacity*) with well-defined margins within which septal thickening and intralobular reticular lines can be identified, in a pattern similar to crazy paving stonework (118) (Figs 121, 122).
- Interstitial: Descriptor referring to the underlying connective tissue framework of the lung (119) (Fig 123).
- Miliary: Subcategory of nodular pattern, designating profuse, diffuse, and randomly distributed well-de-

fined micronodules (micronodule*) of uniform size, with diameters of 1 to 2mm resembling millet seeds (119) (Figs 124, 125).

- Mosaic: Patchwork of regions with normal and abnormally increased attenuation (mosaic attenuation*) (Fig 126).
- Nodular: Pattern including nodules (nodule*) (119) (Figs 127, 128).
- Reticular: Collection of intersecting linear opacities that produce an appearance resembling a net, sometimes referred to as "reticulation" (119) (Fig 129).
- Reticulonodular: Combination of a reticular and a nodular pattern.
- Three-attenuation (three-density): Combination of sharply demarcated zones with normal, increased, and decreased attenuation on CT images acquired in inspiration (14) (Fig 130).
- Tree-in-bud: Micronodules connected to linear opacities with one or more branching sites, thus resembling a budding tree (120) (Figs 131–136).

Pleura (Anatomy) (Figs 137–139)

The pleura is a two-layered serous membrane folding back on itself at the hilum* that covers both lungs. The two layers of the membrane are separated by a small amount of viscous lubricant called pleural fluid, which causes surface tension and, together with negative pressure, pulls the parietal and visceral pleura adjacent to each other. Each pleural layer consists of a monolayer of mesothelial cells and a layer of underlying mesenchymal tissue. Both layers form the virtual "pleural space." At imaging, the normal pleura is best seen where it forms the interlobar fissures (fissure*) or in the presence of extrapleural fat (see anatomic distribution*).

The parietal pleura is the pleural layer that lines the inner surfaces of the thorax on each side of the mediastinum^{*}, and can be subdivided into mediastinal (covering the lateral surfaces of the fibrous pericardium, esophagus, and thoracic aorta), diaphragmatic (covering the upper surface of the diaphragm), and costal (covering the inside of the rib cage). The visceral pleura is the pleural layer that covers the surface of each lung and inserts between the lobes (lobe^{*}) of the lung, forming the interlobar fissures (121,122).

The adjectival form "pleural" is often associated with the following terms:

• Effusion (pathology): Excessive pleural fluid caused by mechanical factors, edema, infection, inflammation, or malignancy (Fig 137).

On radiographs, small pleural effusions manifest as blunting of the costophrenic angles, often meniscus-shaped, and larger effusions as homogeneous opacities (opacity*) in dependent regions. On CT examinations, effusions are of variable attenuation and widen the pleural space. CT can depict loculations, as well as pleural thickening and hyperemia when performed with contrast enhancement.

• Plaque (pathology): Fibrohyaline focal lesions arising in the parietal pleura.

Pleural plaques are usually associated with previous exposure to natural fibrous silicates, usually asbestos. At imaging, pleural plaques are well-defined elevated flat or plateau-like areas of pleural thickening that often contain calcifications (Figs 138, 139).

- Tag (pathology): One or more coarse linear or band-like strands extending from a structure (eg, a nodule or its components) and having direct contact with the pleura or a fissure* (Figs 79, 89, 100, 101). Pathologically, pleural tags are often associated with thickened interlobular septa (interlobular septum*) connecting a structure to the pleura. While usually non-specific, pleural tags have also been described as a potential indicator for visceral pleural invasion (123).
- Thickening (pathology): Focal or diffuse increase in thickness of the pleura, commonly caused by fibrosis following chronic* or recurrent mechanical irritation (eg, by rib fractures) and by pleural effusion or pneumothoraces (pneumothorax*) (Figs 23, 40, 65, 66).

Pleural thickening can also be caused by asbestos exposure or infection, as well as by neoplasms such as metastases, lymphoma, or mesothelioma. Pleural thickening is visible at CT as a soft tissue curvilinear stripe, passing internally to the ribs and innermost intercostal muscles. Thickened pleura often enhances and is best seen after intravenous contrast administration.

Pneumatocele (Pathology) (Fig 140)

A pneumatocele is a thin-walled, gas-filled cystic* structure in the lung parenchyma*. Pneumatoceles can be caused by pneumonia*, trauma (including barotrauma), or aspiration* of hydrocarbon fluid (eg, in "fire eating"). Pneumatoceles can be transient and are likely caused by a combination of parenchymal necrosis and a check-valve airway obstruction. At imaging, pneumatoceles appear as rounded, thin-walled cystic lesions (cyst*) in the lung parenchyma, sometimes with an air-fluid level (124).

Pneumomediastinum (Pathology) (Figs 141, 142)

Pneumomediastinum is the abnormal collection of air or gas in the mediastinum^{*}, outside of the esophagus and the tracheobronchial tree. It can be caused by spontaneous rupture of alveoli, by trauma, or by esophageal rupture with a subsequent incursion of air along the axial interstitium^{*} into the mediastinum^{*}. It is most often associated with surgical trauma, barotrauma, or conditions related to prolonged cough, such as asthma or interstitial lung disease^{*}. At imaging, pneumomediastinum appears as one or more low-attenuation areas of linear or band-like air collections outlining mediastinal structures (125,126).

Pneumonia (Pathology)

Commonly used diagnostic term that typically designates infection of lung parenchyma*. Pneumonia can be caused by bacteria, viruses, fungi, parasites, or other microorganisms, and can be focal* or multifocal*. Pneumonia can manifest as opacity*, potentially accompanied by air bronchograms (air bronchogram*), tree-in-bud* opacities, or pleural effusions (pleural effusion*) (127,128).

Pneumonitis (Pathology)

Pneumonitis is a diagnostic term that designates inflammation of lung parenchyma*, both infectious and noninfectious. Infectious pneumonitis is generally referred to as pneumonia*. Noninfectious pneumonitis can be caused by a wide range of conditions, including aspiration*, hypersensitivity, radiation, autoimmune diseases, and drugs.

The terms "pneumonia" and "pneumonitis" are sometimes used interchangeably but the above definitions are preferred (129,130).

Pneumopericardium (Pathology) (Figs 143, 144)

Pneumopericardium is the accumulation of air in the pericardial cavity. Pneumopericardium has been described in preterm neonates or in the context of mechanical ventilation in infants. In adults, it is usually due to trauma, or cardiac surgery. It can be associated with other pathologic air collections, such as pneumomediastinum* or pneumothorax*. At imaging, pneumopericardium manifests as air or gas attenuation in the structureless space separating the two layers of the pericardium (131).

Pneumothorax (Pathology) (Figs 145, 146)

Pneumothorax refers to the presence of air or gas in the pleural* space that separates the parietal and the visceral pleura*. Therefore, parts of the lung can partially or completely collapse. Pneumothorax can be caused by spontaneous rupture of blebs*, or by mechanical trauma, including iatrogenic damage. The diagnosis of tension pneumothorax is determined by clinical and physiologic criteria that typically correlate radiographically with contralateral mediastinal shift and depression of the ipsilateral hemidiaphragm, most often based on a check-valve mechanism. Ultimately, a definitive diagnosis of tension pneumothorax is determined by clinical and/or physiologic criteria. Pneumothorax is typically identified on upright chest radiographs by the presence of a curvilinear visceral pleural line indicating the demarcation between pleural air and aerated lung. However, on radiographs acquired in the supine position, when the separated pleural surfaces are not tangential to the x-ray beam, pneumothorax may be difficult to detect. In these cases, ancillary findings, such as unusually sharp interfaces between lung and mediastinal or cardiac structures can be helpful. The "deep sulcus sign" refers to an abnormally deepened costophrenic angle, which has low attenuation and is abnormally sharply defined (132,133).

Saber-sheath Trachea (Pathology) (Fig 147)

Saber-sheath trachea refers to the abnormal shape of the trachea, with an increased sagittal (anteroposterior) and a decreased coro-

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nal (left-to-right) diameter, which resembles the sheath of a saber. To meet the definition, the internal coronal diameter of the trachea must be two-thirds or less than the sagittal diameter at the same anatomic level, without any extrinsic cause of compression. Saber-sheath trachea is most commonly seen in patients with chronic obstructive pulmonary disease and is assumed to be the result of chronic inflammation and subsequent remodeling of the tracheal wall (65,134,135).

Secondary Pulmonary Lobule (Anatomy) (Fig 148)

A secondary pulmonary lobule is the smallest structural unit of the lung parenchyma* surrounded by a connective tissue sheath. It is irregularly polyhedral or pyramidal in shape, with borders ranging from 5 mm to 25 mm in length. It contains three to 25 acini (acinus*). The structures in the center of the secondary pulmonary lobule ("core structures," "centrilobular structures") include bronchioles (bronchiole*) and their accompanying pulmonary arterioles and lymphatic vessels. The connective tissue envelope of the secondary pulmonary lobule contains veins and lymphatic vessels. The secondary pulmonary lobule also has an internal support structure, the intralobular interstitium*. CT can help visualize the three basic components of the normal secondary pulmonary lobule, namely the central structures, the region where acini are located, and the connective tissue sheath that forms the interlobular septa (interlobular septum*) (9,136).

Segment (Anatomy)

A segment is a structural unit of a lobe* ventilated by a segmental bronchus, perfused by a segmental pulmonary artery, and drained by intersegmental pulmonary veins. The right lung has 10 segments, with three in the upper lobes (lobe*), two in the middle lobe, and five in the lower lobes. The left lung has either eight or 10 segments, depending on whether the apicoposterior upper lobe segment and the anteromedial segment in the lower lobe are considered to represent one or two segments. At imaging, individual segments are identified inferentially based on the position of the supplying segmental bronchus and artery. Occasionally, segments can be separated by accessory fissures (accessory fissure*) (137).

Septum (Pl. Septa) (Anatomy) (Figs 149-155)

A septum is a sheath-like connective tissue structure surrounding secondary pulmonary lobules (secondary pulmonary lobule*). Because they separate lobules (lobule*), they are also referred to as "interlobular septa" (interlobular septum*). They contain veins and lymphatic vessels. Being oriented perpendicularly to the pleura*, isolated septal lines can be occasionally seen on CT examinations in otherwise healthy patients in the lung apices and the bases as thin linear structures separating secondary pulmonary lobules. They are more conspicuous when thickened (eg, in interstitial pulmonary edema) (9,10).

The adjectival form "septal" is commonly used with the following terms:

• Line (anatomy): Linear structure representing an interlobular septum (Fig 151). • Thickening (pathology): Seen on chest radiographs and CT examinations as thin linear structures at right angles to and in contact with the lateral pleural surfaces (Figs 152–155).

Formerly referred to as "Kerley lines" on radiographs, an obsolete and nonrecommended term, they reflect the accumulation of fluid or tumor in the interstitium* (eg, in pulmonary edema, lymphangitic carcinomatosis, and/or mechanical lymphatic obstruction). On CT examinations, septal thickening can be smooth or nodular, which may help to differentiate edema from lymphatic infiltration or obstruction.

Signet Ring Sign (Pathology) (Fig 156)

The signet ring sign refers to the ring-shaped structure representing a dilated bronchus imaged perpendicularly, accompanied by a smaller adjacent rounded and dense structure representing the accompanying pulmonary artery. The combination of both resembles a signet or pearl ring. It is considered typical of bronchiectasis* but can also be seen in diseases with abnormally decreased pulmonary arterial flow (eg, chronic thromboembolic disease or proximal interruption of the pulmonary artery) (138–141).

Silhouette Sign (Pathology) (Fig 157)

The silhouette sign refers to the obscuration of a normal silhouette or interface (often incorrectly described as "silhouetting") and results from the juxtaposition of structures with similar radiographic attenuation that obliterate normal anatomic borders or interfaces. However, it is not always indicative of pulmonary disease; for example, obscuration of the heart border can sometimes be caused by anatomic variants such as pectus excavatum (6,142,143).

Small Airways Disease (Pathology)

This umbrella term summarizes diseases of the peripheral airways^{*}. The definitions of the term vary. Some radiologists consider small airways as those with an internal diameter of less than or equal to 2 mm and a wall thickness of less than 0.5 mm. Others place the demarcation between central and peripheral airways at the level of the bronchiole^{*}. Although the two definitions are not consistent, the first one appears more practical and is more commonly used in the literature. On CT examinations, these diseases can manifest as peribronchiolar and/or bronchiolar abnormalities, including tree-in-bud opacities^{*} (opacity^{*}), centrilobular micronodules (micronodule^{*}), air trapping^{*}, bronchiolectasis^{*}, oligemia^{*}, or a combination of these findings (101,144,145).

Spiculation (Pathology) [Descriptor] (Fig 158)

Spiculation is a descriptive term referring to multiple fine linear strands extending from a structure (eg, a nodule* or its components) into the surrounding lung parenchyma* in a stellate manner. Pathologically, spiculations are often associated with a desmoplastic process (eg, an invasive malignancy) within the nodule, resulting in fibrotic strands radiating into the surrounding lung. Spiculations can also be associated with direct infiltration into adjacent bronchial or vascular structures or focal lymphangitic extension. Spiculation in a nodule increases the likelihood of cancer, although it is not specific (146).

Tracheomalacia, Tracheobronchomalacia (Pathology) (Figs 159, 160)

The terms tracheomalacia and tracheobronchomalacia are used when there is excessive inward movement of the trachea and/ or main bronchi during expiration, due to pathologic weakness of the cartilaginous airway wall (147). The closely related term "excessive dynamic airway collapse" (Fig 161) is used when tracheobronchial collapse is due to excessive invagination of the posterior muscular tracheal or main bronchial walls during expiration. Both entities can be diagnosed with dynamic expiratory CT. However, it is important to be aware that in healthy individuals the trachea may collapse by 70% or more during a forced expiration (53,148).

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Figure 1: Air bronchogram: (A) Coronal CT image reconstruction of the right lung and (B) frontal chest radiograph of the left lung show air bronchograms (arrows). *click to return to page 2*



Figure 2: Air crescent: Transverse CT image of the right upper lobe shows air crescent (arrow) caused by angioinvasive aspergillosis. *click to return to page 2*



Figure 4: Air trapping: Transverse CT image at the level of the carina in (A) inspiration and (B) expiration. While the lung parenchyma in inspiration (A) is homogeneous, expiration (B) reveals failure of lung density to increase in areas of air trapping (arrows). click to return to page 2



Figure 5: Bronchocentric: Transverse CT image of the left upper lobe shows bronchocentric opacity (arrow).

click to return to page 2



Figure 6: Centrilobular: Transverse CT image of the lungs shows diffuse centrilobular micronodules. *click to return to page 2*



Figure 7: Centrilobular: Transverse CT images of (A) right and (B) left lungs in two different patients show diffuse centrilobular micronodules. *click to return to page 2*



Figure 8: Diffuse: Transverse CT image of the right lung shows diffuse increase in lung attenuation caused by confluent ground-glass nodules. *click to return to page 2*

Figure 9: Endobronchial: Transverse CT images in (A) lung and (B) mediastinal window through the right lung show partially obstructing endobronchial lesion in the bronchus intermedius (arrow). *click to return to page 2*





Figure 10: Endobronchial: **(A)** Transverse and **(B)** coronal contrast-enhanced CT images through left lung of the same patient show endobronchial lesion in left upper lobe bronchus (arrow). *click to return to page 2*



 Figure 11:
 Extrapleural: Transverse CT image of the left lower lobe shows extrapleural fat (arrows).

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Figure 12: Fissural: Transverse CT images of the right lung in two different patients show (A) solitary and (B) multiple perifissural micronodules (arrows).



Figure 13: Focal ground-glass opacities: Transverse CT image of the upper lobes shows subtle, focal ground-glass opacities in the right and left lungs (arrows).



Figure 14: Geographic: **(A, B)** Transverse contrastenhanced CT images of the right lung in two different patients show ground-glass opacities with geographic distribution.



Figure 15: Multifocal: Transverse CT image shows multifocal opacities (arrows). *click to return to page 3*



Figure 16: Multifocal: Transverse CT image shows multifocal opacities (arrows). *click to return to page 3*



Figure 17: Peribronchovascular: Transverse contrastenhanced CT image of the right lung shows radiation reaction with peribronchovascular thickening (arrow).

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Figure 18: Peribronchovascular: Transverse CT image of the right lung shows nodules with peribronchovascular distribution (arrow).



Figure 19: Perilobular: Transverse CT images of the **(A)** right and **(B)** left lung in two different patients show opacities with perilobular distribution (arrows).



Figure 20: Perilobular: Transverse CT image of the left lower lobe shows opacities with perilobular distribution. *click to return to page 3*



Figure 21: Perilymphatic micronodules: **(A, B)** Transverse CT images of the left lung in the same patient at different levels show micronodules with perilymphatic distribution (arrows).

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Figure 23: Pleural: Transverse CT images of the left lung in (A) lung and (B) soft tissue window show diffuse nodular pleural thickening (arrows). *click to return to page 3, page 11*

Figure 22: Perilymphatic micronodules: Transverse CT image of the left lung shows micronodules with perilymphatic distribution along the fissure (open arrow) and the bronchus (solid arrow). *click to return to page 3, page 8*



Figure 24: Pleural: Transverse CT images of the right lower lobe in (A) lung and (B) soft tissue window show focal pleural lesion (arrow).

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Figure 25: Pleural: **(A, B)** Transverse CT images of the lung bases show focal pleural nodules (solid arrow), some of which are partially calcified (open arrow).

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Figure 26: Aortopulmonary window: (A) Frontal chest radiograph and (B) coronal CT image reconstruction in the same patient show aortopulmonary window (arrow).

Figure 27: Apical cap: **(A)** Frontal chest radiograph and **(B)** coronal CT image reconstruction in the same patient show bilateral apical caps (arrows).

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Figure 28: Architectural distortion: Transverse CT images of the (A, B) right and (C) left lung of three different patients show traction bronchiectasis (solid arrows). Note the presence of coexisting fibrosis and architectural distortion (open arrow in B and C). *click to return to page 3, page 4*

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Α



Figure 29: Asbestosis: Transverse CT images of the lower lobes show fibrosis following asbestos exposure. Note that the presence of pleural plaques is not required for the diagnosis.



Figure 30: Atelectasis: Frontal chest radiographs of two different patients show (A) partial and (B) complete right upper lobe atelectasis (arrows). *click to return to page 3*



Figure 31: Atelectasis: Frontal chest radiographs of the right lung in the same patient show (A) complete and (B) partial atelectasis (arrow) of the right upper lobe. *click to return to page 3*



Figure 32: Atelectasis: (A) Frontal chest radiograph and (B) coronal CT image reconstruction show right upper lobe atelectasis (solid arrow). Note presence of juxtaphrenic peak in B (open arrow).



Figure 33: Atelectasis: (A) Frontal chest radiograph and (B, C) coronal CT image reconstructions in mediastinal and lung windows show right upper lobe atelectasis (solid arrow) caused by central tumor (open arrow). Note presence of juxtaphrenic peak in A (open arrow).



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Figure 34: Atelectasis: **(A)** Frontal and **(B)** lateral chest radiograph of the same patient show middle lobe atelectasis (arrow). *click to return to page 3*

upper lobe atelectasis (solid arrow) caused by central tumor (open arrow) Note presence of juxtaphrenic peak ir **A** (open arrow). click to return to page 3



Figure 35: Atelectasis: Frontal chest radiograph shows right lower lobe atelectasis (arrows).

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Figure 36: Atelectasis: (A) Frontal and (B) lateral radiograph shows complete left upper lobe and lingula atelectasis (solid arrows). Note presence of juxtaphrenic peak (open arrow). *click to return to page 3*



Figure 37: Atelectasis: Frontal chest radiographs of two different patients with complete left lower lobe atelectasis (arrow) show that the size of the atelectatic lobe can vary from (A) large to (B) small. *click to return to page 3*



Figure 38: Atelectasis: (A, B) Frontal chest radiographs of two different patients show bilateral linear atelectasis (arrows). click to return to page 3



Figure 39: Atelectasis: Transverse image of the left upper lobe shows linear atelectasis along the major fissure (arrow).



Figure 40: Atelectasis: Transverse CT image of the right lower lobe shows rounded atelectasis (solid arrow) adjacent to thickened pleura and a small pleural effusion, and "comet tail" (open arrow).

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Figure 41: Atoll sign: Transverse CT images of the (A) right and (B) left lower lobes in two different patients show focal groundglass opacities surrounded by a thin ring of consolidation (arrows).

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Figure 42: Atoll sign: Transverse CT image of the left lower lobe shows focal ground-glass opacities surrounded by a thin ring of consolidation (arrows). *click to return to page 4*



Figure 44: Azygoesophageal recess: Transverse CT image of the right lung shows the azygoesophageal recess (arrow), bordered by the azygos vein and the neighboring esophagus. *click to return to page 4*



Figure 43: Azygo-esophageal recess: Frontal chest radiograph of the mediastinum shows the azygo-esophageal recess (arrows). click to return to page 4



Figure 45: Bronchiectasis: Transverse CT images of the lungs in two different patients show severe bronchiectasis (solid arrow) in **(A)** right upper lobe and milder bronchiectasis (solid arrow) in **(B)** left lower lobe. Note the smaller diameter of the accompanying pulmonary artery (signet ring sign) (open arrow).

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Figure 46: Bronchocele: Transverse CT images of the right lung in two different patients show localized bronchial dilatation (A) with and (B) without mucoid impaction within a hyperlucent lobe, due to congenital bronchial atresia (arrow). *click to return to page 4*



Figure 47: Broncholith: Transverse CT image through the right lung (A, lung window; B, bone window) show broncholith (arrow).

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Figure 48: Bronchomalacia: Transverse CT image at the level of the main bronchi in **(A)** inspiration and **(B)** expiration. Expiration causes substantial collapse of the main bronchi (arrows). *click to return to page 5*



Figure 49: Bulla: **(A)** Frontal chest radiograph and **(B)** transverse CT image of the right upper lobe show an air-containing cystic structure lined by a thin layer of collapsed lung parenchyma (arrow).



Figure 50: Bulla: Transverse CT image of the right upper lobe shows an air-containing cystic structure lined by a thin layer of collapsed lung parenchyma (arrow). *click to return to page 5*



Figure 51: Cavity: Transverse CT image of the (A) right and (B) left upper lobe in two different patients show gas-filled structures with thick and partially irregular walls (arrow).



Figure 52: Cavity: Transverse CT image of the left lower lobe shows two round gas-filled structures (arrows). *click to return to page 5*



Figure 54: Consolidation with peripheral groundglass: Transverse CT image of the left lower lobe shows consolidation (solid arrow) and ground-glass opacities (open arrow). *click to return to page 5*



Figure 53: Cavity: Transverse contrast-enhanced CT image of the left lower lobe shows (A) soft tissue and (B) lung window of gas-filled structure with thickened and irregular walls (arrow). *click to return to page 5*



Figure 55: Cyst: Transverse CT images of the **(A)** right and **(B)** left lung of two different patients. Cysts are solitary (arrow) in **A** and diffuse in **B**. *click to return to page 5*



Figure 56: Embolism with infarction: Transverse contrast-enhanced CT images in (A) soft tissue and (B) lung window show emboli (solid arrows) and corresponding infarction (open arrows). (C) Coronal CT image reconstruction and (D) frontal chest radiograph in the same patient show central embolus (solid arrow) and peripheral infarction (open arrow). Emphysema. *click to return to page 5, page 7*



Figure 57: Centrilobular: (A, B) Transverse CT images of the right upper lobes of two different patients show centrilobular emphysema. *click to return to page 5*



Figure 58: Centrilobular: (A, B) Transverse CT images of the right upper lobes of two different patients show centrilobular emphysema. *click to return to page 5*



Figure 59: Centrilobular: (A, B) Transverse CT images of the right upper lobes of two different patients show centrilobular emphysema. *click to return to page 5*



Figure 60: Panlobular: Transverse CT image shows panlobular emphysema. *click to return to page 6*



Figure 61: Panlobular: **(A, B)** Transverse CT images of the right upper lobes of two different patients show panlobular emphysema (arrow). *click to return to page 6*



Figure 62: Paraseptal: Transverse CT image of the left upper lobe shows paraseptal emphysema (arrows). Mild centrilobular emphysema is also present.



Figure 63: Paraseptal: Transverse CT image of the upper lobes shows paraseptal emphysema (arrows). *click to return to page 6*



Figure 64: Visual subtypes of centrilobular emphysema: Transverse CT images of the right lung in different patients show visual subtypes of centrilobular emphysema according to reference 63: (A) trace emphysema, (B) mild emphysema, (C) moderate emphysema, (D) confluent emphysema, and (E) advanced destructive emphysema.



Figure 65: Empyema: Transverse contrast-enhanced CT image of the right lower lobe shows empyema (asterisk) surrounded by thickened and enhancing pleura (solid arrows), with adjacent atelectasis (open arrow). click to return to page 6, page 11



Figure 67: Fibrosis: Transverse CT image of the right lower lobe shows extensive fibrosis with honeycombing and reticular opacities.

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Figure 66: Empyema: Transverse contrastenhanced CT image of the left lower lobe shows empyema (asterisk) with air-fluid level, surrounded by thickened and enhancing pleura (solid arrow), with adjacent atelectasis (open arrow).

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Figure 68: Fissure: Transverse CT image of the right lung shows major fissure (arrows).

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Figure 69: Accessory: Transverse CT image of the right lung shows two accessory fissures in the right lower lobe (arrows).



Figure 70: Accessory: (A) Transverse and (B) sagittal CT images of the left lung of the same patient show accessory fissure in the left lower lobe (arrow). *click to return to page 6*



Figure 71: Azygos: Transverse CT image of the right upper lobe shows azygos fissure (arrows). This structure is in fact a pseudofissure, as it consists of four rather than the normal two pleural layers. *click to return to page 6*



Figure 72: Incomplete: Transverse CT image of the left lung shows absence of the medial portion (solid arrow) of the left major fissure (open arrow). *click to return to page 6*



Figure 73: Groundglass opacity: (A–C) Transverse CT images of the left lung of three different patients show ground-glass opacities. Opacity is subtle and focal in A (arrow), subtle and diffuse with centrilobular nodules in B, and severe and diffuse with mild reactive airway dilatation in C.



Figure 74: Ground-glass opacities and consolidation: Transverse CT image of the right lung shows ground-glass opacities and focal consolidation (arrow). *click to return to page 6, page 9*



Figure 76: Honeycombing: Transverse CT image of the left lung shows honeycombing (arrow).



Figure 75: Halo: Transverse CT image of the right lung shows a ground-glass halo (arrow) surrounding central nodular consolidation.

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Figure 77: Honeycombing: Transverse CT image of the right lung shows mild honeycombing (arrow).

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Figure 78: Honeycombing: **(A, B)** Transverse CT image of the right lung in two different patients shows honeycombing (arrow). *click to return to page 6*



Figure 79: Irregular margination of pulmonary nodule: Transverse CT image of the right lower lobe shows an irregular nodule with spiculations and pleural tags (arrow).

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Figure 80: Juxtaphrenic peak: Frontal chest radiograph of the left lung shows juxtaphrenic peak (arrow).

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Figure 82: Lymph node: Transverse CT image of the mediastinum shows mediastinal lymph node (arrow) with fatty hilum.

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Figure 83: Lymph node: Transverse CT image of the mediastinum shows mediastinal lymph node (arrow) with fatty hilum.



Figure 84: Lymph node: Transverse CT images in (A) soft tissue and (B) lung window show calcified mediastinal lymph node (arrow).

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Figure 85: Lymphadenopathy: (A–C) Transverse CT images of the mediastinum show mediastinal lymph node enlargement (arrows).

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Figure 86: Mass: Transverse CT image of the right upper lobe shows mass with spiculations (open arrow) and pleural tags (solid arrow).



Figure 87: Micronodule: Transverse CT image of the right upper lobe shows multiple micronodules (arrows). click to return to page 8

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Figure 88: Mosaic attenuation: Transverse CT image of the left lung shows geographic regions of normal and abnormally increased attenuation.

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Figure 89: Mucoid impaction: Transverse CT image of the left upper lobe shows bronchus impacted by mucus (arrow).

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Figure 90: Mycetoma: Transverse CT images of the left upper lobe in two different patients show mycetomas (arrow) in preexisting cavities with air crescent surrounded by (A) abnormal and (B) normal lung parenchyma. The mycetoma in B shows sponge-like appearance.

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Figure 91: Mycetoma: Transverse CT images of the right upper lobe in (A) supine and (B) prone position show mycetoma (arrow) within a preexisting cavity surrounded by abnormal lung parenchyma, changing position with gravity.



Figure 92: Nodule: Transverse CT image of the right upper lobe shows a round, well-defined, solid soft tissue nodule.



Figure 93: Nodule: Transverse CT images in (A) lung and (B) soft tissue windows of the right lung show an ovoid soft tissue nodule (arrow).

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Figure 94: Nodule: Transverse CT images in (A) soft issue and (B) lung windows of the right lung show a well-defined, solid soft tissue nodule (arrow). Note the small pleural effusion. *click to return to page 9*



Figure 95: Nodule: Transverse CT image of the (A) left upper and (B) left lower lobes of two different patients show ground-glass nodules with irregular margins.

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Figure 96: Nodule: Transverse CT image of the right lung shows a mildly spiculated nodule (arrow).

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Figure 97: Nodule: Transverse CT image of the right upper lobe shows an irregular nodule with mild spiculation.

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Figure 98: Nodule: Transverse CT image shows a markedly spiculated nodule in the right lower lobe.

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Figure 99: Nodule, pleural tag: Transverse CT images of the left lung show complex nodule with single pleural tag (arrow), (A) thin at first appearance, and (B) thickening over time. *click to return to page 9*



Figure 100: Nodule: Transverse CT images of the (A) left upper and (B) right lower lobes show spiculated nodules with pleural tags (solid arrow) and fissural involvement (open arrow).

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Figure 101: Nodule, spiculation, pleural tag: Transverse CT image of the left upper lobe shows a spiculated nodule with pleural tags (arrows). *click to return to page 9, page 11*



Figure 102: Nodule: Transverse CT images in (A) lung and (B) soft tissue window of the left upper lobe show calcified nodule (arrow). *click to return to page 9*



Figure 103: Nodule, complex morphology: Transverse CT image of the left lower lobe shows a complex nodule with cystic (open arrow) and ground-glass (solid arrow) components. *click to return to page 9*



Figure 104: Nodule, complex morphology: Transverse CT image of the right lower lobe shows a ground-glass nodule with a cystic component (arrow). *click to return to page 9*



Figure 105: Nodule: Transverse CT image of the left lower lobe shows a complex nodule with fat (open arrow) and soft tissue (solid arrow) components.



Figure 106: Nodule: Transverse CT images in (A) soft tissue and (B) lung windows of the right lower lobe show a complex nodule with a calcified (solid arrow) and soft tissue (open arrow) component.

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Figure 107: Nodule: (A-C) Transverse CT images of the right lung of three different patients show a complex nodule with solid and ground-glass components (part-solid nodules). While the ground-glass component is predominantly present in **A**, the solid component is larger in **B** and becomes the dominant component in **C**. *click to return to page 9*



Figure 108: Nodule: Transverse CT image of the right upper lobe shows a complex nodule with fat (solid arrow), calcium (open arrow) and soft tissue (dashed arrow) components.

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Figure 109: Nodule: Transverse CT image of the left lower lobe shows a complex nodule with fat (solid arrow), calcium (open arrow) and soft tissue (dashed arrow) components.

Figure 110: Nodule: Transverse CT image of the left upper lobe shows a complex nodule with cystic (open arrow), ground-glass (dashed arrow) and soft tissue (solid arrow) components.



Figure 111: Nodule: Transverse CT images of the right upper lobe show the evolution of a nodule from (A) soft tissue (arrow) to (B) a nodule with cystic (open arrow) and soft tissue (solid arrow) components. *click to return to page 9*



Figure 112: Nodule: Transverse CT images of the right lung show the evolution of a nodule from **(A)** a ground-glass nodule (arrow) to **(B)** a nodule with ground-glass (solid arrow) and soft tissue and cystic (open arrow) components. *click to return to page 9*



Figure 113: Nodule: Transverse CT images of the right upper lobe show the evolution of a nodule with **(A)** cystic (open arrow) and ground-glass (dashed arrow) components to **(B)** a nodule with cystic (open arrow) and soft tissue (solid arrow) components. *click to return to page 9*



Figure 114: Nodule: Transverse CT images of the left lung show pulmonary nodule that evolves from (A) a complex to (B) a simple morphology. The complex morphology in A consists of a ground-glass (open arrow) and a solid (solid arrow) component. *click to return to page 9*



Figure 115: Oligemia: Transverse CT image shows areas of oligemia caused by small airways disease. Note the reduced number of vessels and thickened bronchial walls. *click to return to page 9*



Figure 116: Oligemia: Transverse contrast-enhanced CT image shows oligemia caused by vascular arterial disease. Note the substantially dilated central pulmonary arteries (asterisk). *click to return to page 9*



Figure 117: Organizing pneumonia: Transverse CT image of the left lower lobe shows consolidation with perilobular and peribronchovascular distribution. *click to return to page 10*



Figure 119: Paratracheal stripe: Frontal chest radiograph shows right paratracheal stripe (arrows). *click to return to page 10*



Figure 118: Organizing pneumonia: Transverse CT image of the lower lobes shows groundglass opacities and peribronchial consolidation. *click to return to page 10*



Figure 120: Parenchymal band: Transverse CT images of the **(A)** left and **(B)** right lung of two different patients show linear parenchymal bands paralleling the pleura (arrows). Pattern. *click to return to page 10*



Figure 121: Crazy paving: (A, B) Transverse CT images of the left lung of two different patients show crazy-paving pattern.

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Figure 122: Crazy paving: Transverse CT image of the right lung shows areas of crazy paving with reticular lines. *click to return to page 10*



Figure 123: Interstitial: Transverse CT image of the right upper lobe shows interstitial pattern caused by thickened interlobular septa.

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Figure 124: Miliary: Transverse CT image of the right lung shows miliary micronodules.

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Figure 126: Mosaic: Transverse CT image of the right lung shows mosaic pattern. click to return to page 10



Figure 125: Miliary: Transverse CT image of the lungs show diffuse distribution of micronodules. *click to return to page 10*



Figure 127: Nodular: Transverse CT images of the right lung of two different patients show (A) moderate and (B) extensive nodular pattern. *click to return to page 10*





Figure 128: Nodular: Transverse contrast-enhanced CT image of the lungs shows diffuse nodular pattern.



Figure 129: Reticular: Transverse CT images in three different patients show (A) subtle, (B) moderate, and (C) severe reticular opacities, with ground-glass opacities. *click to return to page 10*



Figure 130: Three Attenuation: Transverse CT images in two different patients through the (A) right and (B) left lung show lobules with normal, increased, and decreased attenuation.



Figure 131: Tree-in-bud: Transverse CT images of the middle lobe. (A) While the standard reconstruction shows several micronodules only, (B) the maximum intensity projection reveals the tree-in-bud appearance of these micronodules (arrow). *click to return to page 10*



Figure 132: Tree-in-bud: Sagittal CT image reconstruction of the left lower lobes shows focal tree-in-bud opacities (arrows). click to return to page 10



Figure 133: Tree-in-bud: Transverse CT image of the right lower lobe shows multifocal tree-in-bud with branching opacities. *click to return to page 10*



Figure 134: Tree-in-bud: Transverse CT image of the right lung in (A) thin-section and (B) maximum intensity projection reconstructions show focal tree-in-bud opacities (arrow).

Figure 135: Tree-in-bud: (A, B) Transverse CT images of the right lung in two different patients show focal tree-in-bud opacities. *click to return to page* 10







Figure 136: Treein-bud: Transverse CT image of right lung shows focal tree-in-bud opacities (arrows).





Figure 138: Plaque: Transverse CT image of the left upper lobe shows calcified pleural plaque (arrow).

Figure 137: Effusion: Frontal chest radiograph shows left pleural effusion (arrow). *click to return to page 10*



Figure 139: Plaque: (A, B) Transverse CT images of the lower thorax show partially calcified pleural plaques (arrows). *click to return to page 11*



Figure 140: Pneumatocele: Transverse CT image through the left lung shows well-defined pneumatocele (arrow). click to return to page 11

Figure 141: Pneumomediastinum: **(A)** Frontal and **(B)** lateral chest radiographs show mediastinal air collection (arrows).

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Figure 142: Pneumomediastinum: (A, B) Transverse CT images in two patients show mediastinal (solid arrow) and extrapleural (open arrow) air collections.



Figure 143: Pneumopericardium: (A, B) Frontal chest radiographs in two patients show air in the pericardial cavity (arrows).

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Figure 144: Pneumopericardium: Transverse contrast-enhanced CT image shows ventral pericardial air collection (arrow).



Figure 145: Pneumothorax: (A) Frontal chest radiograph and (B) transverse CT image reconstruction of the left lung show air in the pleural space (arrows). *click to return to page 11*



Figure 146: Pneumothorax: (A) Frontal chest radiograph, and (B) coronal and (C) transverse CT image reconstructions of the lung show air in the left pleural space loculated due to pleural adhesions. *click to return to page 11*



Figure 147: Sabersheath trachea: Transverse CT image of the mid trachea shows saber-sheath configuration.

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Figure 148: Secondary pulmonary lobule: Transverse CT image of the right lower lobe shows secondary pulmonary lobule surrounded by thickened interlobular septa (arrows).

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Figure 149: Septum: (A, B) Transverse CT images of the right upper lobe in two different patients show thickened interlobular septa (arrow).







Figure 150: Septum: Transverse CT image of the right upper lobe shows thickened interlobular septa (arrows). *click to return to page 12*



Figure 151: Septal lines: (A) Frontal chest radiograph and (B) coronal CT image reconstruction show septal lines (arrows), formerly called Kerley B lines.

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Figure 152: Septal thickening: Frontal chest radiograph shows septal lines (arrow). *click to return to page 12*



Figure 153: Septal thickening: Transverse CT image of the right lower lobe shows septal thickening and intralobular lines (arrows) in an area of crazy paving.

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Figure 154: Septal thickening: Transverse CT image of the right upper lobe shows irregular ground-glass opacities (arrows) with interlobular septal thickening and intralobular lines in an area of crazy paving.

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Figure 155: Septal thickening: Coronal CT image of the right lung shows septal thickening and intralobular lines (arrow) in an area of crazy paving.

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Figure 156: Signet ring sign: Transverse CT images of the (A) left and (B) right lower lobe in two different patients show dilated bronchi accompanied by smaller pulmonary arteries (arrow).

Figure 157: Silhouette sign: (A, B) Frontal chest radiographs of two different patients show obscuration of right-sided heart border by adjacent consolidation in the middle lobe. *click to return to page 12*

Figure 158: Spiculation: Transverse CT image of the right upper lobe shows a spiculated mass.

Figure 159: Tracheobronchomalacia: Transverse CT images at the level of the (A, C) trachea and (B, D) the main bronchi in (A, B) inspiration and (C, D) expiration. Expiration causes nearcomplete collapse of both trachea and main bronchi (arrows). *click to return to page 13*

Figure 160: Tracheomalacia: Transverse CT images of the upper thorax. (A) In inspiration, the shape of the trachea (arrow) is abnormal, with an increased transverse diameter. (B) Expiration shows near-complete tracheal collapse (arrow). *click to return to* page 13

Figure 161: Excessive dynamic airway collapse: Transverse CT images at the level of the aortic arch in (A) inspiration and (B) expiration. Expiration causes an excessive invagination of the posterior tracheal wall (arrow). *click to return to page 13*