Illustrated Brazilian consensus of terms and fundamental patterns in chest CT scans*

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Abstract
The objective of this new Brazilian consensus is to update and to continue the standardization of the principal terms and fundamental patterns in chest CT scans in Portuguese. There is a succinct definition of the principal terms used to describe chest CT findings, as well as illustrations of classic examples. The group of authors comprised radiologists specializing in chest radiology and holding membership in the Brazilian College of Radiology and Diagnostic Imaging, as well as pulmonologists having a special interest in diagnostic imaging and holding membership in the Brazilian Thoracic Association.

Keywords: Lung; Consensus; Tomography.

Introduction
In the evaluation of patients with thoracic diseases, CT scans are a diagnostic tool of great importance. In recent years, new CT patterns have been recognized, new terms have been coined in order to describe CT findings in English, and some terms have become obsolete.\(^{1-4}\) Despite the great dissemination of the Brazilian Consensuses published in 2002 and in 2005, respectively, in the journal Radiologia Brasileira (Brazilian Radiology), the official organ of the Brazilian College of Radiology and Diagnostic Imaging, and in the Brazilian Journal of Pulmonology, the official organ of the Brazilian Thoracic Association, some terms have yet to be stand-

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Note to readers: The original document was composed in Portuguese, and this (translated) glossary of terms in English is therefore alphabetized by the names of the corresponding terms in Portuguese.
Glossary

Air trapping (aprisionamento aéreo)

Air trapping is the retention of excess gas (air) in part of or all of the lung (especially upon exhalation), resulting from partial or total airway obstruction or secondary to focal abnormalities of lung compliance. It is recognized during the expiratory phase as reduced attenuation of the lung parenchyma, especially apparent as lower-than-usual density (Figure 1) and absence of volume reduction. With regard to distribution, the following types of atelectasis can be found: subsegmental, segmental, lobar or whole lung. Atelectasis can also be qualified, according to its form, as plate-like (discoid) or rounded. In studies using iodinated contrast material, the homogeneous enhancement of the lung parenchyma can aid in the differentiation between atelectasis and consolidation (Figure 3). The term “collapse” can be used in cases of total atelectasis of one lobe or whole lung atelectasis.

Plate-like atelectasis (atelectasia laminar)

Plate-like atelectasis is a focal area of subsegmental atelectasis with linear or discoid configuration, almost always extending to the pleura (Figure 4). It is generally horizontal or oblique, but it can also have a vertical orientation. The thickness of laminar atelectasis can range from a few millimeters to over 1 cm. Laminar atelectasis is synonymous with discoid atelectasis.

Figure 1 - Axial HRCT scan of the chest during inhalation (a), revealing no significant changes. HRCT scan of the chest during exhalation (b), revealing air trapping (arrows).
Rounded atelectasis (atelectasia redonda)

Rounded atelectasis is a type of oval-shape atelectasis caused by the adherence of the adjacent lung parenchyma to the area of pleural thickening, for example, in cases of asbestos-related pleural disease or empyema resolution.\(^{15}\) It manifests as a rounded focal opacity to which bronchovascular structures converge (comet tail artifact) with a pleural base near the area of pleural thickening (Figure 5).\(^{16,17}\) It presents homogeneous enhancement when iodinated contrast material is used.

Parenchymal bands (banda parenquimatosa)

Parenchymal bands are linear opacities, generally peripheral and often in contact with the pleural surface, which might be thickened and retracted at the site of contact (Figure 6). They are usually 1-3 mm thick and at least 5 cm long.\(^{13}\) Parenchymal bands often have horizontal distribution (perpendicular to the pleural surface), but they can also have an oblique distribution. They generally indicate pleuroparenchymal fibrosis. Lung architecture distortions are often identified. Parenchymal bands are frequently found in patients exposed to asbestos.\(^{18}\)

Fungus ball (bola fúngica)

A fungus ball results from the fungal colonization of preexisting pulmonary cavitation, generally secondary to tuberculosis or sarcoidosis, but it can also occur within cysts (e.g., bronchogenic cyst), bullae or dilated bronchi. Colonization by *Aspergillus* spp. is the most common type of colonization, and in this case, the term “aspergilloma” is used. A fungus ball is characterized by a mass-like collection of intertwined hyphae associated with mucus, fibrin, and cellular debris. On CT scans, a fungus ball is observed as a rounded or oval mass, which tends to hang down when images are acquired with the patient in different horizontal positions (Figure 7).\(^{19}\) Other common findings in this condition include the presence of the “air crescent sign”, amorphous calcification within the lesion, sponge-like appearance of the lesion and adjacent pleural thickening. The term “fungus ball” should not be used as a synonym.
Bulla (bolha)

A bulla is a hypodense focal area that presents well-defined and smooth walls that do not exceed 1 mm in thickness (Figure 8). Bullae generally have gaseous content, but they can also have an air-fluid level. In general, they are associated with other signs of pulmonary emphysema and have a paraseptal location (findings that aid in the differentiation from cysts on CT scans). They typically measure ≥ 1 cm in diameter. Bullae that are < 1 cm and located in the visceral pleura or in the subpleural region of the lung are known for “mycetoma”, since they represent different processes.

See also Air crescent sign and Mycetoma.
Broncholith (broncolito)

A broncholith is a calcified peribronchial lymph node that erodes into the adjacent bronchus. It is often secondary to infections by Histoplasma sp. or Mycobacterium tuberculosis. On CT scans, a broncholith is seen as a small

Air bronchogram (broncograma aéreo)

An air bronchogram is the radiological translation of air-filled bronchi, surrounded by sick lung parenchyma, in which the air of the airways has been substituted by any pathological material, radiologically denser than air (e.g., transudate, exudate, blood, accumulation product or neoplastic cells). In general, this is the expression used when a lucent (gas-filled) tubular region is seen within an area of opacified lung (Figures 3 and 10). This tubular image should be of a size and orientation typical of a bronchus or multiple bronchi, presumably representing a segment of the bronchial tree.
focus of calcification within or adjacent to the airway, being more common in the middle lobe bronchus (Figure 11). Mucoid impaction, bronchiolectasis or atelectasis might be observed distally.\(^{24,25}\)

**Bronchiectasis (bronquiektasia)**

Bronchiectasis is irreversible bronchial dilatation, which can be focal or diffuse. It generally results from chronic infection, proximal airway obstruction or congenital bronchial abnormalities. The morphological findings seen on HRCT scans (Figure 12) are as follows: the internal diameter of the bronchus is larger than the diameter of the adjacent pulmonary artery (signet-ring sign); loss of normal bronchial tapering, defined as the maintenance of the bronchial diameter for more than 2 cm, distally to the bifurcation ("tram tracks"); and identification of airway less than 1 cm from the pleural surface. Bronchiectasis is frequently associated with the thickening of the bronchial walls, mucoid impaction and small airway changes.\(^{26,27}\) Pathology defines three types of bronchiectasis, according to the appearance of the affected bronchus, namely cylindrical, varicose and vesicular (or cystic).

See also Signet-ring sign.

**Bronchiolectasis (bronquiolectasia)**

Bronchiolectasis refers to the dilatation of a bronchiole. It is analogous to bronchiectasis, but it affects an airway of much smaller diameter, identified in the periphery of the lung (Figure 14). Bronchiolectasis manifests as rounded or tubular structures, generally located in the periphery of the lung, with thick walls or filled with secretion (see also Tree-in-bud pattern).\(^{3,7}\) It can also be combined with other pulmonary opacities and to the distortion of the parenchyma in cases of fibrosis.

See also Traction bronchiectasis and traction bronchiolectasis.

**Traction bronchiectasis and traction bronchiolectasis (bronquiektasia e bronquiolectasia de tração)**

Traction bronchiectasis and traction bronchiolectasis refer, respectively, to bronchial and bronchiolar dilatation secondary to the retraction of the parenchyma resulting from fibrosis (Figure 13).\(^{13}\) They manifest as bronchial and bronchiolar dilatations, often irregular, associated with the distortion of the parenchyma due to fibrosis and other lung alterations (principally reticular opacities, ground-glass opacities and consolidations). They have a tubular, cystic or microcystic aspect (peripheral bronchioles), depending on the relationship between the bronchial axis or the bronchiolar axis and the CT scan slice. This last aspect can be confused with honeycombing, another alteration frequently associated with pulmonary fibrosis.\(^{28}\)

**Cavitation (cavidade or escavação)**

Cavities are gas-filled spaces, with or without air-fluid level, within a nodule, mass or pulmonary consolidation. They are typically produced by the expulsion or drainage of a necrotic part through the airway (Figure 15) or to the pleural space. The walls of a cavity are usually irregularly outlined and measure over 1 mm in thickness. The word “cavity” is not synonymous with “abscess”. Although the term “cavitation” can be translated to Portuguese as cavitación or escavação, the term cavitão has a different meaning in Portuguese and therefore should not be used as a synonym for escavação in this context.\(^{7}\)

See also Signet-ring sign.
**Cyst (cisto)**

A cyst is any rounded, well-circumscribed space surrounded by an epithelial or fibrous wall of variable thickness. On CT scans, a cyst is seen as a rounded area with low attenuation coefficient on the lung parenchyma, having a well-defined interface with the adjacent normal lung (Figure 16). The cyst wall is usually thin (< 2 mm), but it can vary in thickness. Cysts are usually filled with air but can also contain liquid (e.g., bronchogenic cyst) or even a solid material. Diseases accompanied by multiple pulmonary cysts include lymphangioleiomyomatosis, Langerhans cell histiocytosis, lymphocytic interstitial pneumonia and Birt-Hogg-Dubé syndrome.

**Collapse (colapso)**

Collapse is generally used as a synonym for total atelectasis of one lobe or whole lung atelectasis (Figure 2). See also Atelectasis.

**Consolidation (consolidação)**

Consolidation is when the air in the alveolar spaces is supplanted by any type of pathological product, such as inflammatory exudate (pneumonia), transudate (edema), blood (alveolar hemorrhage), lipoprotein (alveolar proteinosis),

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**Figure 12** - Bronchiectasis characterized by the “signet-ring sign” (long arrow) and “tram tracks” appearance (short arrow).

**Figure 13** - Axial HRCT slices of the chest showing the right upper lobe (a) and the base of the right lung (b), revealing traction bronchiectasis (long arrows) and traction bronchiolectasis (short arrows) in a patient with interstitial fibrosis (nonspecific interstitial pneumonia), associated with distortion of the lung parenchyma and reticulate parenchyma, as well as with ground-glass opacities.
Architectural distortion (distorção da arquitetura)

Architectural distortion is characterized by displacements in the path and/or distortions in the morphology of anatomical structures such as bronchi, vessels, fissures, or interlobular septa; it is usually related to caused by diffuse parenchymal diseases, particularly interstitial fibrosis. On HRCT scans, the loss of anatomical definition of the secondary lobule and the local volume reduction are considered signs of architectural distortion of the lobule and can be considered indirect signs of fibrosis (Figure 13).

Pulmonary emphysema (enfisema pulmonar)

Pulmonary emphysema is characterized by permanently enlarged airspaces distal to the...
terminal bronchiole with destruction of the alveolar walls.\textsuperscript{(35)} The additional histological criterion of absence of “obvious fibrosis” has been questioned because some degree of interstitial fibrosis might be present due to smoking.\textsuperscript{(36)} Emphysema is classified according to the acinar region affected: proximal (centriacinar or centrilobular emphysema), distal (paraseptal emphysema), or whole acinus (panacinar or panlobular emphysema).\textsuperscript{(37)} The tomographic findings are areas of low attenuation, typically without visible walls.\textsuperscript{(37,38)}

**Bullous emphysema (enfisema bolhoso)**

Bullous emphysema is the bullous destruction of the parenchyma in combination with centriacinar emphysema, distal (paraseptal) emphysema or panacinar emphysema (Figure 17). It is designated giant bullous emphysema when the bullae, which might vary from 1 to over 20 cm in diameter, occupy at least one third of the hemithorax.\textsuperscript{(7,39)}

See also Bulla.

**Centriacinar emphysema (enfisema centroacinar)**

Centriacinar emphysema is the destruction of the walls of the centriacinar alveoli combined with an increase in respiratory bronchioles and associated alveoli. Centriacinar emphysema is the most common form of emphysema in cigarette smokers. Tomographic findings are centrilobular areas of decreased attenuation, usually without visible walls, having nonuniform distribution and predominantly located in the upper pulmonary regions (Figure 18).\textsuperscript{(37,38)} The centrilobular arteries can frequently be identified within the hypodense areas. The term centrilobular emphysema is commonly used as a synonym in CT.

**Interstitial emphysema (enfisema intersticial)**

Interstitial emphysema is characterized by the air dissection of the pulmonary interstitium, typically located on the bronchovascular sheaths, interlobular septa and visceral pleura; it is more frequently identified in neonates receiving mechanical ventilation (Figure 19). Interstitial emphysema is seldom recognized radiographically in adults and is rarely seen on CT scans. The process appears as areas of air density with perivascular distribution, peribronchovascular distribution or distributed along the interlobular septa (Figure 20), or as rounded areas of low attenuation simulating small bullae or cysts (Figures 19 and 20).\textsuperscript{(40,41)}

**Panacinar emphysema (enfisema panacinar)**

Panacinar (panlobular) emphysema involves all portions of the acinus and, reasonably uniformly, the secondary pulmonary lobule.
It predominates in the lower lobes and is the form of emphysema associated with alpha-1 antitrypsin deficiency. Panacinar emphysema is seen on CT scans as a generalized decrease in the attenuation of the lung with a reduction in the diameter of the blood vessels in the affected areas, with or without distortion of these vessels (Figure 21). Severe panacinar emphysema can coexist and merge with severe centriacinar emphysema. On CT scans, it can be indistinguishable from the findings of severe constrictive bronchiolitis.

Paraseptal, or distal acinar, emphysema (enfisema parasseptal/acinar distal)

Paraseptal emphysema affects predominantly the distal alveoli and their ducts and sacs. It is characteristically delimited by any pleural surface or interlobular septa. On CT scans, it is characterized by areas of low attenuation in the subpleural and peribronchovascular regions, separated by intact interlobular septum (Figure 22). It is sometimes associated with bullae.

Air spaces (espaços aéreos)

Air spaces correspond to the spaces bounded by the alveolar walls, including the alveolar sacs, alveolar ducts and the alveoli on the walls of the respiratory bronchioles. This term can be used in association with consolidation, opacity or nodules to characterize the origin of such lesions, which fill these spaces with liquid or cells. The air bronchogram sign (Figures 3 and 10) is the trademark of air space filling.

See also Air bronchogram.

Interlobular septal thickening (espessamento de septos interlobulares)

Interlobular septal thickening is the thickening of the connective tissue septa separating the secondary pulmonary lobules. It is radiographically characterized by thin linear opacities, also designated Kerley B lines. It is usually in close contact with the lateral pleural surface, near the costophrenic sulci, at a right angle to the pleural wall. Currently, the terms “septal lines” or “septal thickening” have gained favor over Kerley lines. On HRCT scans (Figure 23), the thickening of the interlobular septa is characterized by the presence of linear opacities that...
delimit the secondary pulmonary lobules, more easily characterized in the subpleural region, in which it has the aspect of lines at right angles to the pleural surface. In the central regions of the lungs, the thickening of the septa of adjacent lobules results in the aspect of polygonal arches. Septal thickening can be secondary to changes in any of its components (veins, lymphatic vessels or connective tissue) and is a finding common to various lung alterations; however, its presence is particularly highlighted in cases of pulmonary edema and lymphangitic carcinomatosis. Septal thickening can be smooth (Figure 23), nodular (Figure 24) or irregular. This differentiation can aid in the differential diagnosis.

**Centrilobular region (estruturas centrallobulares)**

The centrilobular region comprises the central portion of the secondary pulmonary lobule, consisting of the pulmonary artery, bronchiole and surrounding lung interstitium. On HRCT scans of normal patients, it corresponds to a small nodular or linear image, located 3-10 mm from the pleural surface or the interlobular septum, representing the intralobular pulmonary artery and measuring approximately 1 mm in diameter (Figure 25). The corresponding bronchiole, when normal, has walls with approximate thickness of 0.15 mm, below the resolution limits of the HRCT. Therefore, the bronchiolar disease

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**Figure 22** - Paraseptal, or distal acinar, emphysema in the right upper lobe (arrows).

**Figure 23** - Interlobular septal thickening (arrows) of the smooth type in a patient with pulmonary edema. Pleural effusion (asterisk).

**Figure 24** - Interlobular septal thickening of the nodular type (arrows), with a beaded appearance (beaded septum sign).
Honeycombing, or honeycomb lung (faveolamento, or favo de mel)

Honeycombing is characterized by pulmonary cysts created as a result of the destruction of the distal air spaces due to parenchymal fibrosis, with loss of acinar and bronchiolar architecture. Honeycombing cysts are covered with metaplastic bronchiolar epithelium and represent the end stage of a number of pulmonary diseases.\textsuperscript{1,48} On HRCT scans, honeycombing is characterized by multiple cysts, generally subpleural and of comparable diameters (typically 0.3-1 cm), grouped on layers and sharing well-defined walls of 1-3 mm in thickness (Figure 27).\textsuperscript{49} Honeycombing is a tomographic marker of pulmonary fibrosis. The principal causes of honeycombing include idiopathic pulmonary fibrosis, collagenoses, chronic hypersensitivity pneumonitis, drug-induced pulmonary reactions and asbestosis.\textsuperscript{28} It should be differentiated from paraseptal emphysema and traction bronchiolectasis.

Progressive massive fibrosis (fibrose maciça progressiva)

Progressive massive fibrosis is the conglomeration of small pulmonary nodules, generally accompanied by fibrosis (Figure 28). It is frequently bilateral and predominates in the upper lobes. It can be accompanied by an irregular increase in the size of the air space periphery. Progressive massive fibrosis is generally found in patients with a history of heavy exposure to inorganic dust (e.g., coal worker's pneumoconiosis and silicosis).\textsuperscript{50} The tomographic findings of progressive massive fibrosis can be similar to those of sarcoidosis and talcosis.\textsuperscript{51}

Fissure (fissura or cissura)

Fissure is the invagination of the visceral pleura that covers the outer surface of the lung to the parenchyma.\textsuperscript{77} Each interlobar fissure is formed by the juxtaposition of two layers of visceral pleura. In general, we can identify major (oblique) fissures, which separate the lower lobes from the others, and the lesser (horizontal) fissure, distinguishing the middle lobe from the right upper lobe (Figure 29). Supernumerary fissures are frequently found.
An interface is the separation between two structures or spaces. When two thoracic structures with different radiological densities are juxtaposed, their boundaries are clear. For example, vessels presenting soft tissue density coming into contact with the air density of the surrounding ventilated lung. The “sign of the interface” defines the irregularity of the margins of different intrathoracic structures, such as vessels, bronchi and pleural surfaces, generally resulting from interstitial disease that causes fibrosis (Figure 30).\(^1\)

**Interstitium (interstício)**

The interstitium is a net of connective tissue that spreads over the lungs and is subdivided into: (a) axial (bronchovascular) interstitium—surrounding the bronchi, arteries and veins from the hila to the level of the respiratory bronchioles; (b) peripheral interstitium—composed...
of the connective tissue contiguous to the pleural (subpleural) surfaces and interlobular septa; and (c) intralobular interstitium (also designated acinar or parenchymal interstitium)—composed of the alveolar walls (alveolar septa), and supporting the structure of the secondary pulmonary lobule.\(^{[2,3,7]}\)

**Lymph node enlargement (linfonodomegalia)**

Lymph node enlargement is characterized by the increase in volume in one or more lymph nodes, due to any cause, surpassing the dimensional limits considered normal for the lymph node chain in question (Figure 31).\(^{[52,53]}\) The term “adenomegaly” is not an acceptable synonym, since lymph nodes are not true glandular structures. The term “linfonodopatia” (lymphatic disease) is reserved for situations in which a disease, such as necrosis, has been identified within a lymph node.

**Subpleural curvilinear line (linha curvilínea subpleural)**

The subpleural curvilinear line is a curvilinear opacity of 1-3 mm in thickness, located in the subpleural region and having a parallel distribution over the pleural surface (Figure 32). It is a nonspecific sign for atelectasis, edema, fibrosis or inflammation.\(^{[1,7]}\)

**Intralobular lines (linhas intralobulares)**

Intralobular lines are seen on HRCT scans as thin linear images within the secondary pulmonary lobule. When numerous, they can have a fine reticular aspect (Figure 33). This finding can be seen in different conditions, especially in cases of fibroses (e.g., usual interstitial pneumonia) and diseases associated with the crazy-paving pattern (e.g., alveolar proteinosis).\(^{[3,54]}\)

See also Reticular pattern and Crazy-paving pattern.

**Secondary pulmonary lobule (lóbulo pulmonar secundário)**

The secondary pulmonary lobule is the smallest anatomic unit of lung delimited by a septum of connective tissue. It is polyhedral in shape, measures 1.0-2.5 cm in diameter and contains a variable number of acini.\(^{[45]}\) The
center of the lobule is formed by the bronchiole and its accompanying pulmonary artery, lymphatic vessels and adjacent interstitium. In its periphery, the connective tissue forms thin septa designated interlobular septa, which contain small pulmonary veins and lymphatic vessels (Figure 25). In normal patients, the interlobular septa are better identified in the anterior, lateral and paramediastinal peripheral regions of the upper and middle lobes, as well as in the peripheral region of the anterior diaphragmatic region of the lower lobes, tending to be incomplete or absent in the remaining lung regions. The interlobular septa are easily recognized on HRCT scans when the septal interstitium is affected (Figures 23 and 24).\[7\]

**Mass (massa)**

A mass is any expansive pulmonary, pleural, mediastinal or chest wall lesion presenting

density of soft, fatty or bony tissue; greater than 3 cm in diameter; with at least partially defined contours; outside of the topography of the fissures and independent of the characteristics of its contours or the heterogeneity of its content (Figure 34).\[7\]

**Mycetoma (micetoma)**

A mycetoma characteristically represents a group of subcutaneous chronic infections caused by the traumatic inoculation into the skin of material contaminated with actinomycetes, principally *Nocardia brasiliensis*, or eumycetes, resulting in actinomycetoma and eumycetoma, respectively.\[20\] It tends to invade adjacent tissues, forming nodules or masses with cavities and fistulous pathways, with the elimination of suppurative secretion containing grains constituted by masses of hyphae and filaments. In most cases, it is located in the lower limbs and can cause deformities and fractures. Pulmonary and pleural involvement is rare.\[55\] When the lung is affected, the aspect is of consolidation with necrosis, and pleural effusion can be seen.\[56\] Mycetoma generally affects agriculturists and is endemic in Latin America, India and Africa.\[20,57\] Mycetoma is not a colonization of a preexisting pulmonary cavity; therefore, the use of this term as a synonym for “fungus ball” should be avoided.

See also Fungus ball.

**Nodule (nódulo)**

A nodule is a focal opacity that is rounded, or at least partially delineated, smaller than 3.0

Figure 34 - Pulmonary mass in the right upper lobe measuring 5.5 cm in diameter.

Figure 35 - Axial HRCT slices of the chest in three patients revealing pulmonary nodules (arrows) with different attenuation: a) solid (soft tissue density); b) non-solid (ground-glass attenuation); and c) part-solid or semisolid.
cm in diameter and generally presenting soft tissue or calcified tissue density (Figure 35). When the opacity is smaller than 10 mm, it is recommended that the term “small nodule” be used. When the opacity is smaller than 3 mm, it is recommended that the term “micronodule” be used.\(^{(3,7)}\) Nodules should be described according to the characteristics of their borders (well- or ill-defined), to their location or to their distribution (random, perilymphatic, centrilobular or pleural). With regard to attenuation, nodules can be classified as solid (Figure 35a), when they completely obscure the parenchyma; as non-solid (ground-glass attenuation), when they do not obscure the vascular margins and bronchial walls (Figure 35b); or as part-solid or semisolid (ground-glass opacity with solid areas), when they partially obscure the vascular margins and the bronchial walls (Figure 35c).\(^{(7)}\)

See also Mass.

**Oligemia (oligoemia)**

Oligemia is a focal, regional or generalized reduction in pulmonary blood volume. It appears as a decrease in the diameter and number of pulmonary vessels (regional or widespread), indicating less than normal blood flow (Figure 36).\(^{(3,7)}\)

**Opacity (opacidade)**

Opacity is an image which, due to its greater density, is at least partially distinguishable from the surrounding or superimposed structures. On chest X-rays, this term does not specify the pathological nature, size or specific location of the image. An opacity can be caused by an abnormality in the lung, pleura or chest wall, as well as by an external object. On CT scans, the pulmonary opacities can represent ground-glass opacity or consolidation.

See also Consolidation.

**Ground-glass opacity/attenuation (opacidade/atenuação em vidro fosco)**

On CT scans, ground-glass opacity corresponds to increased density of the lung parenchyma in which it is still possible to identify the contours of the vessels and bronchi within the area affected by a pathological process (Figure 37). This image pattern is related to interstitial thickening, partial filling of air spaces, partial collapse of alveoli, increased capillary blood volume or a combination of all of these mechanisms.\(^{(58,59)}\) Ground-glass attenuation should be distinguished from “consolidation” (Figures 10 and 38), in which the vessels are not identifiable within the affected area of the lung.

See also Consolidation.
**Linear opacity (opacidade linear)**

Linear opacity is characterized by a thin, elongated linear image with soft tissue density. Rarely, calcification or foreign material, which can increase the attenuation, is seen. It is a nonspecific term used for a number of etiologies. It is recommended that more specific terms be used whenever possible, such as “laminar atelectasis”, “parenchymal band” and “interlobular septal thickening”.

**Hanging opacity (opacidade pendente)**

A hanging opacity is defined as a subpleural opacity in one of the “hanging pulmonary regions”, which are areas of atelectasis that appear when the patient is lying down. Hanging opacities are seen in posterior regions when the patient is in the supine position (Figure 39a) and in anterior regions when the patient is in the prone position. They disappear when the patient is no longer in a horizontal position (Figure 39b).

**Parenchymal opacification (opacidade/ opacificação parenquimatosa)**

Parenchymal opacification is the increased attenuation in the lung parenchyma that can potentially obscure the contours of the vessels and bronchi. The term “consolidation” indicates that the margins of these structures are not visible (except for air bronchograms), and “ground-glass attenuation” indicates that, despite the altered lung density, the vessels and airways remain identifiable. It is recommended that the more specific terms “consolidation” and “ground-glass opacity” be used (Figure 38).

**Tree-in-bud pattern (padrão de árvore em brotamento)**

The tree-in-bud pattern represents centrilobular branching opacities with small nodules at the extremities that resemble the budding of certain trees (Figure 26). In most cases, this pattern represents dilated bronchioles filled with pathological material, although it might also be associated with infiltration of the peribronchial connective tissue in the centrilobular vasculature or, occasionally, with dilatation or filling (e.g., intravascular metastases) of the centrilobular arteries. This finding is often indicative of disease affecting the airways and is particularly common in infectious processes (e.g., tuberculosis, bronchopneumonia and infectious bronchiolitis) but it can also be found in a number of other diseases (e.g., bronchiectasis, cystic fibrosis and panbronchiolitis).

**Mosaic attenuation/perfusion pattern (padrão de atenuação/perfusão em mosaico)**

The mosaic attenuation pattern appears as patchwork of regions of differing attenuation that can represent infiltrative lung disease, obliterate small-airways disease or occlusive vascular disease (Figure 40). Air trapping secondary to bronchial or bronchiolar obstruction can produce parenchymal foci of lower attenuation, which become more evident in expiratory CT scans. In obliterative small-airways disease and vascular occlusive disease, the areas of low attenuation are abnormal; in general, the number and size of pulmonary vessels in these areas are reduced when compared with those in the adjacent normal lung, which might present normal or increased attenuation (due to redirec-
tion of blood flow). Mosaic attenuation pattern can also be caused by parenchymal pulmonary disease, characterized by ground-glass opacity; in this case, the areas with increased attenuation are the affected regions and the remaining regions characterize foci of preserved lung.\(^{[3,7]}\)

See also Air trapping.

**Crazy-paving pattern**

*padrão de pavimentação em mosaico*

The crazy-paving pattern presents superimposition of ground-glass opacities, interlobular lines and thickened interlobular septa (Figure 41). The interface between the normal and the affected lungs tends to be well-delimited in this pattern of pulmonary lesion. This pattern was initially identified in patients with pulmonary alveolar proteinosis, but it can also be seen in other diffuse pulmonary diseases in which the interstitial and alveolar compartments are affected (e.g., pulmonary hemorrhage).\(^{[1,2,64]}\)

**Centrilobular nodules**

*padrão nodular centrolobular*

Centrilobular nodules are small nodules located in the center of the secondary pulmonary lobule, generally related to bronchiolar diseases, pulmonary artery diseases or peribronchovascular bundle diseases.\(^{[7]}\) The principal tomographic finding is that these nodules are located at a distance of a few millimeters from the pleural surface and the fissures (Figure 42a).\(^{[45-47]}\) The most common cause of this condition is inhalation-related diseases (e.g., hypersensitivity pneumonitis, silicosis and respiratory bronchiolitis). If it is accompanied by the tree-in-bud pattern (Figure 26), infectious causes should be considered (e.g., tuberculosis and bronchopneumonia).
Perilymphatic nodules
(padrão nodular perilinfático)

The perilymphatic nodular pattern is characterized by a pattern of distribution of small nodules along the pulmonary lymphatic system (interlobular septa, peribronchovascular bundle and pleural surface) (Figure 42b). The principal diseases accompanied by perilymphatic nodules are sarcoidosis and lymphangitic carcinomatosis.\(^{(45,65)}\)

Miliary nodules—See Random, or miliary, nodules

Random, or miliary, nodules
(padrão nodular randômico, or miliar)

The random distribution of small nodules throughout the lungs (Figure 42c) is most often caused by miliary tuberculosis, miliary histoplasmosis or hematogenous metastases.\(^{(3,45,65)}\)

Perilobular pattern (padrão perilobular)

The perilobular pattern is characterized by the distribution of abnormality along the structures that border the pulmonary lobules, that is, the interlobular septa, the visceral pleura and major pulmonary vessels.\(^{(66)}\) The term is more frequently used in the context of diseases (e.g., perilobular organizing pneumonia) that are distributed principally around the periphery of the secondary lobule. On CT scans, it is characterized by the presence of thick and irregular polygonal opacities in the periphery of the secondary pulmonary lobule (Figure 43).\(^{(67)}\) It should be distinguished from the interlobular septal thickening (septal pattern), because it is thicker and more irregular; and from the reversed halo sign.

Figure 43 - Perilobular pattern (arrows).

Figure 44 - Reticular pattern in a patient with pulmonary fibrosis.

Figure 45 - Pleural plaques with calcification (arrows) in a patient exposed to asbestos.
of its characteristic mechanism of check-valve airway obstruction. (71) Pneumatocele is usually associated with infectious diseases, especially those caused by Staphylococcus sp. in children and Pneumocystis sp. in adults. On CT scans, it manifests as a rounded air space bounded by thin walls, within the lung (Figure 46a). (1,3,7) It can resolve spontaneously, even if tardily, while the infection is being treated (Figure 46b).

Pseudocavity (pseudocavidade)

A pseudocavity is a round or oval-shaped area of low attenuation in nodules, pulmo-
is more commonly found in sarcoidosis, silicosis and coal workers’ pneumoconiosis.

**Signet-ring sign (sinal do anel de sinete)**

The signet-ring sign is a sign composed of a ring-like opacity representing a dilated bronchus, together with a smaller, rounded opacity contiguous to the bronchial wall, representing its artery (pulmonary artery or, rarely, the bronchial artery), resembling a “signet ring” or a “pearl ring”. It is the basic tomographic sign of bronchiectasis (Figure 12). Occasionally, the signet-ring sign can also be found in diseases characterized by an abnormal reduction in pulmonary arterial flow, such as chronic pulmonary thromboembolism and proximal interruption of the pulmonary artery.

See also Bronchiectasis.

**Air crescent sign (sinal do crescente aéreo)**

The air crescent sign corresponds to a collection of air of variable size and in the form of a crescent or half moon, located in the periphery of a nodule or mass and presenting soft tissue density; in the proper clinical context, this finding is suggestive of angioinvasive aspergil-
Halo sign (sinal do halo)

The halo sign is characterized by a ground-glass opacity surrounding a nodule, mass or round area of consolidation (Figure 50). It was initially described as a sign of hemorrhage surrounding a focus of angioinvasive aspergillosis. The halo sign appears early and is highly specific to neutropenic patients with fever. It can be caused by hemorrhage associated with other types of disease (e.g., candidiasis and Kaposi's sarcoma) or by local pulmonary infiltration by neoplasm (e.g., adenocarcinoma).

Reversed halo sign (sinal do halo invertido)

The reversed halo sign is a ground-glass opacity surrounded by a complete or partially complete ring of consolidation (Figure 51). Although it was initially described as a sign of organizing pneumonia, it has also been associated with other diseases, such as paracoccidioidomycosis.

Beaded septum sign (sinal do septo nodular em contas/rosário)

The beaded septum sign is an irregular and nodular thickening of the interlobular septa resembling a row of beads (Figure 24). Although it was initially described as a sign of lymphangitic carcinomatosis, it is more frequently found in sarcoidosis.

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References


Figure 50 - Nodule showing the halo sign (arrows).

Figure 51 - Reversed halo sign (arrows).
Illustrated Brazilian consensus of terms and fundamental patterns in chest CT scans

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