Chronic interstitial lung diseases in children*

Doenças pulmonares intersticiais crônicas na criança

Maria Aparecida Soares de Souza Paiva, Sandra Mara Moreira Amaral

Abstract

Interstitial lung diseases (ILDs) in children constitute a heterogeneous group of rare diseases that have been described and classified according to experiences and research in adults. However, pediatric pulmonologists have observed that the clinical spectrum is broader in children than in adults, and that many of these disorders have different courses and treatment responses. In addition, probably due to the various stages of lung development and maturation, new clinical forms have been described, particularly in infants. This has broadened the classification of ILDs in this age bracket. The understanding that neither the usual definition nor the standard classification of these disorders entirely apply to children has prompted multicenter studies designed to increase knowledge of these disorders, as well as to standardize diagnostic and therapeutic strategies. We have reviewed the conceptualization of ILDs in children, taking into consideration the particularities of this group of patients when using the criteria for the classification of these diseases in adults. We have also made a historical review of several multicenter studies in order to further understanding of the problem. We have emphasized the differences in the clinical presentation, in an attempt to highlight knowledge of newly described entities in young children. We underscore the need to standardize management of laboratory and radiological routines, as well as of lung biopsy processing, taking such knowledge into account. It is important to bear in mind that, among the recently described disorders, genetic surfactant dysfunction, which is often classified as an idiopathic disease in adults, should be included in the differential diagnosis of ILDs.

Keywords: Lung diseases, interstitial; Lung diseases, interstitial/diagnosis; Lung diseases, interstitial/therapy; Child.

Resumo

As doenças pulmonares intersticiais (DPIs) da criança constituem um grupo heterogêneo de doenças raras que têm sido definidas e classificadas de acordo com as experiências e as pesquisas em adultos. Entretanto, os pneumologistas pediátricos vêm observando que o espectro clínico é mais amplo nas crianças, e que muitas destas doenças evoluem e respondem ao tratamento de forma diferente. Além disso, provavelmente devido a estágios diferentes de desenvolvimento e maturação pulmonares, novas formas clínicas têm sido descritas, principalmente em lactentes, ampliando a classificação nessa faixa etária. A compreensão de que nem a definição nem as classificações estabelecidas se aplicam inteiramente ao grupo pediátrico tem motivado a realização de estudos multicênicos com o objetivo de estudá-las melhor, unificando as estratégias diagnósticas e terapêuticas. Fizemos a revisão atualizando a conceituação das DPIs no grupo pediátrico, considerando as particularidades desse grupo na utilização do esquema de classificação dessas doenças para adultos e revendo o histórico dos esforços para uma melhor compreensão do problema com os estudos multicênicos. Foram ressaltadas as diferenças na apresentação clínica, procurando realçar os novos conhecimentos sobre as doenças recém descritas nas crianças pequenas. Alertamos também para a necessidade de ser seguida uma rotina padronizada de investigação laboratorial, radiológica e de processamento das biópsias à luz desses conhecimentos. É importante lembrar que, do grupo das novas doenças descritas, as alterações genéticas do surfactante devem constar também do diagnóstico diferencial das DPIs dos adultos, podendo se apresentar nesse grupo como uma das doenças classificadas como idiopáticas.

Descritores: Doenças pulmonares intersticiais; Doenças pulmonares intersticiais/diagnóstico; Doenças pulmonares intersticiais/terapia; Criança.

* Study carried out at the Rio de Janeiro State Worker’s Hospital, Rio de Janeiro, Brazil.

Correspondence to: Maria Aparecida Soares de Souza Paiva. Av. das Américas, 2300, Casa 37, Barra da Tijuca, CEP 22640-102, Rio de Janeiro, RJ, Brasil.
Tel 55 21 3431-1000. E-mail: mariaaparecida.paiva@gmail.com
Financial Support: None.
Chronic interstitial lung diseases in children


793

a period of 3 years (1995-1998) and identified 46 cases of ILD, symptom onset occurring before 1 year of age in 66% of the children. Of the 46 patients, 7 (16%) died, and 9 belonged to one of only four families.

A multicenter study involving 131 children (1995-1997), with questionnaires sent to 187 pulmonology centers in Europe, Australia and South Africa, obtained a response from 20.3% of these centers. In that study, noninvasive tests alone were able to establish the diagnosis in 5 patients (3.8%), and complementary invasive techniques—bronchoalveolar lavage (BAL) and biopsy—allowed the diagnosis of 117 patients (89%). Of these, 64% were males, and mean age ranged from 0.75 to 17.8 years.

Subsequently, a group of pediatric pulmonologists from the European Respiratory Society (ERS) formed a work group and sent questionnaires to all pulmonology centers in Europe, evaluating the records of 185 patients (1997-2002). Of these, 58 children were under 2 years of age. The prevalence was higher among males. Approximately 10% of the cases occurred in siblings. The clinical data and the invasive and noninvasive complementary tests allowed the diagnosis of 177 patients (95.6%). In 67 of the patients who underwent biopsy, the diagnosis reported in the records was not consistent with the classification of ILDs in adults. The authors raised various questions, principally regarding the need for a classification of ILDs in children. Based on the data collected, the patients were divided into four principal groups of diagnosis: 1) diffuse parenchymatous pulmonary disease of known causes (hypersensitivity pneumonitis, aspiration pneumonitis, etc.); 2) idiopathic interstitial pneumonia (desquamative interstitial pneumonia, lymphocytic interstitial pneumonia, nonspecific interstitial pneumonia [NSIP], etc.); 3) other forms of interstitial pneumonia (hemosiderosis, sarcoidosis); and 4) congenital disorders (surfactant dysfunction, lymphangiectasia, etc.). The authors did not clearly suggest a classification.

New forms of ILDs have been described in young children, diagnosed thanks to the advances in immunohistochemistry, genetic tests and electron microscopy (EM): neuroendocrine cell hyperplasia of infancy, pulmonary interstitial glycogenosis, genetic surfactant dysfunction, disorders of lung development.
In order to better define ILDs in children, the work group decided to incorporate clinical data to the definition of the "chILD syndrome" to aid in recognizing these disorders. For the diagnosis of ILDs, it was established that, in the absence of known causes of lung disease, at least 3 of the following criteria had to be present: 1) respiratory symptoms (cough, rapid or difficult breathing, or exercise intolerance); 2) signs (tachypnea at rest, crackles, retractions, digital clubbing, underdevelopment or respiratory failure); 3) hypoxemia; and 4) diffuse alteration on chest X-rays or CT scans. By applying the definition of the chILD syndrome, 3 to 4 criteria were met by 91% of the 218 cases studied in 11 centers in North America (1999-2004).

The group reviewed the biopsy results of 187 children (1999-2004) and developed a new clinical and histological classification, which principally organized the disorders in children under 2 years of age (Chart 1). This classification is probably not definitive. It should be pointed out that the duration should not be added to the definition because the disease progresses rapidly in some neonates. Some of the previous definitions, including that developed by the ERS work group, limited their scope to children presenting disease duration of 3 months.

With regard to older children, we believe that the consensus classification proposed by the American Thoracic Society (ATS) and the ERS (16) is appropriate. However, the experience gained from analyzing this age bracket should be considered. The usual interstitial pneumonia (UIP) and idiopathic pulmonary fibrosis (IPF) form, for instance, with the character-

<table>
<thead>
<tr>
<th>Chart 1</th>
<th>Most prevalent diffuse lung diseases in patients under two years of age.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diffuse diseases of development</td>
<td>Acinar dysplasia</td>
</tr>
<tr>
<td></td>
<td>Congenital alveolar dysplasia</td>
</tr>
<tr>
<td></td>
<td>Alveolar capillary dysplasia with misalignment of pulmonary veins</td>
</tr>
<tr>
<td>Abnormalities of development</td>
<td>Chronic lung disease of infancy</td>
</tr>
<tr>
<td></td>
<td>• related to chromosomal alterations</td>
</tr>
<tr>
<td></td>
<td>• related to congenital heart diseases</td>
</tr>
<tr>
<td></td>
<td>Pulmonary hypoplasia</td>
</tr>
<tr>
<td>Specific conditions of unknown etiology</td>
<td>Neuroendocrine cell hyperplasia of infancy</td>
</tr>
<tr>
<td></td>
<td>Pulmonary interstitial glycogenosis</td>
</tr>
<tr>
<td>Disorders caused by surfactant dysfunction</td>
<td>Mutations in the genes of SP-B, SP-C, ABCA3 and TTF-1</td>
</tr>
<tr>
<td></td>
<td>Other histological expressions (PAP, DIP, NSIP, CPI)</td>
</tr>
</tbody>
</table>

SP: surfactant protein; TTF-1: thyroid transcription factor-1; PAP: pulmonary alveolar proteinosis; DIP: desquamative interstitial pneumonia; NSIP: nonspecific interstitial pneumonia; CPI: chronic pneumonitis of infancy. Extracted from the flow chart created by the Children’s Interstitial Lung Disease (chILD) collaborative group and modified.
Chronic interstitial lung diseases in children

The clinical data are nonspecific and often less evident than they are in adults. The prognosis also seems to be different, the course of the disease being less favorable. Multiple organ involvement is more common and more severe in younger children. An excellent review of sarcoidosis in children was published in 2005.

Among the group of histologically well-defined diseases, we highlight, due to its severity in children, multisystem Langerhans cell histiocytosis, an immunologic disease that affects the bone marrow, spleen, liver, lymph nodes, thymus, skin, brain, bones, gastrointestinal tract and lungs. Lung disease might occur in isolation, principally in young individuals. It can cause spontaneous pneumothorax. Lung biopsy reveals Langerhans cell infiltrate, interstitial pneumonitis and honeycombing with cysts of different dimensions. Lytic lesions in the bones and cysts in the lungs are clues to the diagnosis.

**History**

The classifications in adult patients have also been evolving.

In 1944, Hamman and Rich reported cases of interstitial pneumonia. However, the study of ILDs began to advance in the 1960s, with the introduction of histologically defined diseases. A classification system was developed by the American Thoracic Society (ATS) in 1997, which included idiopathic pulmonary fibrosis (IPF) and other forms of idiopathic interstitial pneumonia (IIP).

**Figure 1** - Classification of diffuse lung diseases according to the American Thoracic Society consensus, modified by adding forms that are specific to children under two years of age, according to the Children’s Interstitial Lung Disease (chILD) collaborative group.
advances in thoracic surgery, making biopsies more common. Based on histopathological tests, the first classification was established by Liebow in 1975. However, subsequent questions led to revisions, because different terms occasionally designated the same disorder. The landmark review conducted by Katzenstein & Myers and published in 1998 divided the cases formerly classified as IPF into four different forms and established consistent histological criteria for each of them, with clinical, therapeutic and prognostic implications.

The need for internationally standardized diagnostic criteria and nomenclature led to a classification proposed by a multidisciplinary group of the ATS/ERS in a consensus concluded in 2001, in which the importance of the clinical, radiological and histopathological interaction in the study of ILDs was reinforced. This classification, the fruit of the accumulation of a great amount of experience, is still under debate, and it will certainly be modified. However, it organized the information available and allowed a standardized evaluation. Figure 1 shows a flow chart proposed by the ATS/ERS consensus, with the addition of the forms reported in children under 2 years of age aiming to place such disorders in the context of a classification used routinely, modified based on a study we have published recently. The different types of chronic idiopathic interstitial pneumonia are described in Chart 2, the diagnosis being generally suggested by a detailed history.

It becomes evident that it is not appropriate to classify all pediatric patients according to the classification in adults. The stage of lung development and maturation should be taken into account for a more appropriate approach.

Clinical forms specific to children

Persistent tachypnea of infancy with neuroendocrine cell hyperplasia

The neuroendocrine cells are present in the airway mucosa of mammals, in isolation or in combination with neuroepithelial bodies, and produce serotonin, bombesin and calcitonin.

A group of 15 children, of whom 12 were infants, presenting with persistent tachypnea, rales and hypoxemia, was investigated. Chest X-rays and HRCT scans showed hyperinflation and ground-glass opacities. Respiratory function tests confirmed obstruction with air trapping. Lung biopsy showed no inflammation or interstitial alterations. The most significant histological finding was neuroendocrine cell hyperplasia in the distal airways, revealed by bombesin-like immunoreactivity. There was a contrast between the unfavorable clinical picture and the minimal histological alterations observed. Many patients received the standard treatment, and the majority required supplemental oxygen for months or years. The etiology, pathogenesis and long-term prognosis remain unknown.

Of the 15 patients, 10 lived at an altitude of 1,600 m, 5 m at sea level, suggesting that hypoxia at elevated altitudes is a relevant factor in the pathogenesis of the disorder.

Neuroendocrine cell hyperplasia or dysfunction might occur in the following situations: pediatric asthma; cystic fibrosis; bronchopulmonary dysplasia; pulmonary hypertension; pneumonia; congenital malformations; sudden death syndrome; congenital central alveolar hypoventilation syndrome; and diffuse idiopathic neuroendocrine cell hyperplasia in adults. Therefore, the diagnosis of neuroendocrine cell hyperplasia of infancy requires a correlation among clinical, histological and radiological findings.

Pulmonary interstitial glycogenosis

One group of authors reported the cases of 7 children, 6 of whom were male, with nonin-

<table>
<thead>
<tr>
<th>Cause</th>
<th>Types</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aspiration syndromes</td>
<td>GER/swallowing disorders/ malformations</td>
</tr>
<tr>
<td>Chronic infections</td>
<td>Viral (EBV/CMV/adenovirus)</td>
</tr>
<tr>
<td></td>
<td>Bacterial (Chlamydia sp. and Mycoplasma sp.)</td>
</tr>
<tr>
<td>Physical and environmental agents</td>
<td>Hypersensitivity pneumonitis</td>
</tr>
<tr>
<td></td>
<td>Toxicity from oxygen and other gases</td>
</tr>
<tr>
<td></td>
<td>Mineral and organic dust</td>
</tr>
<tr>
<td></td>
<td>Drugs</td>
</tr>
<tr>
<td></td>
<td>Radiation</td>
</tr>
</tbody>
</table>

GER: gastroesophageal reflux; EBV: Epstein-Barr virus; and CMV: cytomegalovirus.
fectious respiratory symptoms in the neonatal period. The chest X-rays revealed tachypnea, retractions, hypoxemia, interstitial infiltrate and hyperinflation. Of the 7 children, 4 were premature neonates, the gestational age ranging from 25 to 33 weeks. The age at the onset of the symptoms ranged from 3 h to 4 weeks. Of the 7 children, 5 were submitted to mechanical ventilation, the duration of which ranged from 4 days to 6 months. All children underwent lung biopsy, which revealed uniform interstitial thickening, diffuse due to immature cells, which were similar to mesenchymal cells, showing abundant cytoplasmic glycogen detected by EM. This unusual finding of EM led the authors to conduct a review of 1,000 cases of lung biopsies in patients with different pathologies. None of the patients presented glycogen in mesenchymal cells, regardless of the gestational age. In contrast with other ILDs, there was no inflammation or alveolar cell hyperplasia. The course of the disease was favorable, and only one death occurred. Pulmonary interstitial glycogenosis is a rare neonatal disease and is presumably the same pathology that has been designated cellular interstitial pneumonitis. It seems that select interstitial cells are immature, and the etiology and pathogenesis are still unknown.

**Chronic pneumonitis of infancy**

One group of authors reported, in a samples consisting of 9 individuals (infants and toddlers), a rare interstitial disease that differed from the ILDs previously described. The histopathological features of this disease, designated chronic pneumonitis of infancy, included septal thickening intensified by primitive mesenchymal cells and marked pneumocyte hyperplasia, together with alveolar exudate containing numerous macrophages and eosinophilic cell remnants. Inflammatory cells were scarce. The analysis of these data suggested recurring pneumonia or slow-resolving pneumonia, affecting an immature or malformed lung. Other such cases have been reported. In general, it presents high mortality. The radiological findings are nonspecific. This histological pattern has been observed in certain patients with genetic surfactant dysfunction.

**Interstitial disease caused by genetic alterations in the surfactant proteins**

Surfactant is a phospholipid film that maintains alveolar stability, preventing alveolar collapse at the end of exhalation. It is composed of lipids (80-90%) and proteins (10-15%). Approximately 2-3% are surfactant proteins (SP-A, -B, -C and -D). Their metabolism involves other molecules, such as ABCA3 and thyroid transcription factor-1 (TTF-1).

For decades, surfactant deficiency has been known to cause hyaline membrane disease in premature neonates. However, only recently has this type of dysfunction been linked to other lung diseases. Mutations in the genes of the surfactant proteins, although rare, are increasingly reported as being the cause of ILDs in children and adults and should therefore be considered in any patient with the chILD syndrome. The form of presentation varies with age. In neonates and infants, the profile is usually severe, and mortality is high; in schoolchildren and adults, the disease is chronic (NSIP, UIP), requiring a high degree of suspicion.

**Mutations in the ABCA3 gene**

Mutations in the ABCA3 gene can manifest as fatal lung disease in neonates or as ILDs in older children and adults. The most common symptoms are cough, dyspnea, hypoxemia, digital clubbing and rales. In a study of 9 cases, 4 presented pectus excavatum. The HRCT scans show alterations that are commonly observed in ILDs and do not correlate with respiratory function tests, hypoxemia or the course of the disease. Lung biopsy might reveal various histological patterns: alveolar proteinosis, desquamative interstitial pneumonia, NSIP and chronic interstitial pneumonitis. The case of a 15-year-old teenager with ABCA3 deficiency

**Chart 3 - Severity score.**

1- asymptomatic
2- symptomatic without blood gas alterations
3- symptomatic with hypoxemia (oxygen saturation < 90%) only during sleep or exercise
4- symptomatic with hypoxemia (oxygen saturation < 90%) at rest
5- symptomatic with pulmonary hypertension

Taken from reference 7 (translated by permission).
Deficiency of surfactant protein B

A gene located on chromosome 2 encodes SP-B. The most common mutation, 121ins2, accounts for 70% of the cases of SP-B deficiency. Deficiency of SP-B follows an autosomal recessive inheritance pattern, and heterozygous are asymptomatic. The clinical and radiological profiles are similar to those of hyaline membrane disease. Deficiency of SP-B begins before 12 h of life and requires mechanical ventilation in term neonates. The response to exogenous surfactant is minimal or transitory. It does not respond to corticosteroids. Persistent pulmonary hypertension might occur, responsive or not to NO. Most patients with SP-B deficiency evolve to death. Survival rates are low in cases of partial SP-B deficiency.\(^{(30,31,37)}\)

The histological examination of the biopsy specimen might reveal alveolar proteinosis or desquamative interstitial pneumonia, or nonspecific findings such as interstitial fibrosis and alveolar cell hyperplasia. The use of EM aids in the differential diagnosis.\(^{(30)}\)

Deficiency of surfactant protein C

The gene that encodes SP-C (SFTPC) is located on chromosome 8. Approximately 55% of the mutations occur spontaneously, and the remainder follows an autosomal dominant inheritance pattern. The most common mutation is I73T.\(^{(30)}\)

Deficiency of SP-C is rare in the neonatal period. The onset of the symptoms occurs in childhood or in adulthood. The clinical profile is common to ILDs. In the neonatal period, the clinical profile might be similar to that of hyaline membrane disease, and it is indistinguishable from that of SP-B deficiency. Histological findings can present various patterns in children: NSIP, pulmonary proteinosis and chronic pneumonitis of infancy. In adults, the UIP/IPF pattern has also been reported. The severity of the disease varies. The disease might progress asymptptomatically, require transplantation (which is complicated in individuals of such an age) or progress to death.\(^{(18,30,38)}\)

The pathophysiology remains unclear. The accumulation of abnormal proteins in the endoplasmic reticulum activates the inflammatory cascade, favors apoptosis and interferes with normal protein synthesis. The release of cytokines and the recruitment of T lymphocytes and fibroblasts occur.\(^{(31)}\)

Deficiency of thyroid transcription factor-1

The TTF-1 plays a fundamental role in the formation and development of the lungs and controls the synthesis of SP-B, SP-C and ABCA3. Genetic mutations of the TTF-1 gene cause hypothyroidism, neurological symptoms (hypotonia and chorea) and neonatal respiratory distress syndrome or chronic lung disease. The BAL reveals a decrease in SP-B associated with an increase in pro-SP-C. The histological findings include enlargement of intercellular junctions, type II pneumocyte hyperplasia and intra-alveolar accumulation of material positive for periodic acid-Schiff staining.\(^{(31)}\)

Diagnosis

The clinical profile is variable and nonspecific. The onset can be acute or insidious. The most common signs and symptoms are dyspnea, cough, stunted growth (weight and height), crackles, digital clubbing, cyanosis, thoracic deformity and signs of pulmonary hypertension or cor pulmonale. Clinical history-taking should be thorough for pulmonary diseases and systemic diseases.\(^{(39)}\) Parents should be questioned about dyspnea and its progression and severity, as well as about weight loss or delayed somatic development of their children. Such complaints are usually not spontaneous and aid in the diagnosis of chronic hypoxemia. In infants, it should be observed whether there is respiratory effort during milk feedings or crying. In term or near-term neonates with persistent course, cough or difficulty in weaning from mechanical ventilation, the possibility of ILD should be considered.\(^{(6)}\) Cough might be the only symptom, and it is generally dry.\(^{(6)}\) Hemoptysis or bloodstained sputum can be a sign of pulmonary hemosiderosis, other vasculitis or Ehlers-Danlos
syndrome.\textsuperscript{23,40} In schoolchildren and adolescents, the profile is similar to that in adults.

History-taking should also include infections, environmental exposure to mineral or organic dust, ingestion of medications or mineral oil, symptoms suggestive of aspiration syndrome and symptoms related to systemic diseases such as joint, heart, skin, kidney and sinus diseases, as well as neurological diseases. Nonrespiratory symptoms sometimes predominate, and ILD is diagnosed during screening for systemic disease.

Family history is quite important, principally in infants. Mortality at an early age or ill-defined lung disease in adults can be a clue to genetic diseases.

The height-weight curve, dyspnea at rest and on exertion and diffuse crackles, principally in the lung bases, should be evaluated during clinical examination. In a later phase of these disorders, thoracic deformity with flattening of the anteroposterior diameter, pectus excavatum and signs of chronic hypoxemia, such as digital clubbing and signs of cor pulmonale, are observed. Persistent pulmonary hypertension might be the only symptom indicative of alveolar capillary dysplasia with misalignment of the pulmonary veins, a disease of development involving the vasculature and the lobular parenchyma.\textsuperscript{(6)} Cyanosis is a late sign of severity. In sarcoidosis, general symptoms such as weight loss, fever and abdominal discomfort have been reported more often in children than in adults, as have cutaneous sarcoid lesions, erythema nodosum, peripheral adenopathies and symptoms related to the central nervous system, articular symptoms being less common.\textsuperscript{(21)}

Based on the clinical history and on the physical examination, the following tests should be performed, sequentially, beginning by the noninvasive tests: blood workup; blood gas analysis at rest and, when possible, after exertion; radiological test; serology for viruses such as EBV, HIV and cytomegalovirus; serology for mycoplasma and \textit{Legionella pneumophila}; immunological profile; screening for aspiration syndromes; tests for collagen diseases; precipitins to organic antigens; the sweat test; tests for sarcoidosis; cardiac evaluation; and respiratory function tests.

The genetic tests for surfactant alterations are expensive and should be requested only when necessary. The identification of mutations using blood DNA or DNA from oral swab samples is the definitive diagnostic method. The PCR for specific fragments of ABCA3 proteins and restriction enzyme analysis should be initially performed. If the diagnosis is not established, gene sequencing of the most common mutations or complete gene sequencing is required. In order to screen for SP-B, ELISA or Western blot of the tracheal aspirate or of the BAL fluid can be performed as an initial step. An algorithm of diagnosis of these disorders is found in the reference list (30).

Spirometry can show the three types of respiratory pattern, although the restrictive pattern is the most common one. The DLCO technique confirms diffusion impairment that translates to hypoxemia. Functional alterations are seen in most children with active connective tissue disease, even in the absence of radiological abnormalities or of symptoms. These patients should be monitored from a functional standpoint.\textsuperscript{[22]} Infants can also be evaluated using various respiratory function tests. These, however, are available in few health care facilities. Some centers take advantage of the sedation used for HRCT to perform such tests.

Hypercapnia appears later and indicates greater severity. However, it appears earlier in dermatomyositis due to respiratory muscle weakness.

Subsequently, we will recommend radiological tests and invasive tests such as BAL and biopsy, as well as the tests that should be performed on the material obtained.

\textbf{Radiology}

The conventional radiological test at the time of diagnosis generally shows bilateral interstitial infiltrate, as classically reported in the initial forms of ILDs. However, in a time before HRCT, it was reported that 9.6% of a group of 458 adults with histologically diagnosed disease showed normal radiological test results, biopsy being performed due to the results of blood gas analysis and functional tests of diffusion, which revealed significant alterations in gas exchange. Therefore, normal radiological test results do not exclude the diagnosis at the onset of these disorders.\textsuperscript{[41]}

An HRCT scan can diagnose even initial lesions. The experience gained from such tests
in adults has been well-documented since their emergence; however, the rarity of such disorders and the technical difficulties in performing HRCT in children under 5 years of age make it difficult to conduct studies involving that age bracket. One group of authors analyzed the HRCT scans of 20 children (1-16 years of age) with ILDs confirmed by biopsy and classified the images as follows: airway diseases; septal pathologies; infiltrative lung disease; airspace disease; and diseases accompanied by cysts.\(^{(42)}\) That and other studies showed the limitations of CT scans, when used in isolation, in diagnosing ILDs in children.\(^{(43,44)}\) There is a consensus that certain disorders present typical CT characteristics that, in combination with clinical characteristics and less invasive tests, facilitate the diagnosis of such disorders, without the need for a biopsy. Such disorders include alveolar proteinosis, congenital lymphangiectasia and idiopathic hemosiderosis. In adults, the recent advances toward a deeper understanding of the clinical and radiological features of ILDs using a standardized technique have greatly improved the diagnostic accuracy by narrowing the possibilities of differential diagnoses and allowing, in certain situations, a specific diagnosis to be established without the need for a biopsy.\(^{(45,46)}\) The standardization of the HRCT techniques in the pediatric radiology routine is essential to make the most of imaging tests in children. A protocol addressing the best technique and emphasizing the dangers of high doses of radiation in children has been suggested to pediatric radiologists.\(^{(47)}\) The controlled ventilation technique promises to be useful in pediatric tests in certain situations, reducing the inability of young children to cooperate.\(^{(48)}\) The protocols highlight that, in the supine position, opacities in the dependent regions of the lung (small atelectasis) are commonly observed on CT scans, opacities that disappear in the prone position. The tests performed during inhalation and exhalation increase the sensitivity to aeration disturbances.

A CT scan can aid in defining the best site from which the biopsy specimen is to be taken, sites presenting a ground-glass pattern being generally chosen. The decision of monitoring pediatric patients using HRCT should be made carefully. In adults, serial evaluation of HRCT scans has proven to be a useful tool for predicting the prognosis of UIP/IPF patients, with scores for classifying honeycomb lesions.\(^{(49)}\)

A detailed description of how the principal clinical and pathological forms of ILD appear on CT scans is found in the ATS/ERS consensus statement,\(^{(16)}\) as well as in excellent reviews conducted recently.\(^{(45,46)}\)

**Bronchoalveolar lavage**

BAL is indicated for screening for etiologic agents and for evaluating cell profiles. Its diagnostic possibilities, however, are far greater. In adults, BAL has been widely used in ILDs, and the role of differential analysis of cells with diagnostic and prognostic objectives is still controversial in many situations.

An excellent routine recommendation for performing BAL in children and evaluating the material collected has been published.\(^{(50)}\) The laboratory must be prepared for processing the test. In the following cases, the first invasive test to be performed should be BAL, which can preclude the need for a biopsy: in suspected cases of aspiration pneumonia caused by mineral oil, hemosiderosis or alveolar proteinosis; when screening for certain infectious agents; and in immunocompromised patients.\(^{(51)}\) Depending on the cellularity, BAL results might suggest other diagnoses. In suspected cases of SP-B deficiency, ELISA or Western blot should be performed on the BAL fluid (and on the tracheal aspirate). The detection of SP-B rules out this diagnostic hypothesis, and genetic tests for other surfactant abnormalities are indicated.\(^{(30)}\) For the diagnosis of Langerhans cell histiocytosis, the presence of more than 5% of typical cells in the BAL fluid is diagnostic if it is associated with a suggestive clinical profile.

Since children need to be sedated so that BAL can be performed, a biopsy is the method of choice when children present a high degree of hypoxemia, because it provides more objective information (including prognostic information) and it allows screening for fibrosis, which is an irreversible factor of severity.

**Biopsy**

A biopsy should be performed early in suspected cases of ILD, i.e., as soon as all noninvasive tests have been performed, even in severe patients on mechanical ventilation, prior to...
the onset of pulmonary fibrosis. It should be performed before anti-inflammatory treatment is initiated. Biopsy is considered the gold standard because it reveals the presence of interstitial inflammation, thickening of the alveolar wall with different patterns of inflammatory cells, alveolar filling and fibrosis. There should be a close understanding among pulmonologists, pathologists and pediatric radiologists in order to make the most of the biopsy evaluation. The use of specific staining methods, immunohistochemistry and EM will depend on the clinical hypotheses. The examination of specimens taken from different sites improves the yield of the test in adults, and should be performed whenever the functional conditions allow it. The cooperative group formed to study chILDs has proposed a protocol for processing biopsy specimens from young children. In that proposal, the group recommends that, since the tests for DNA analysis take long, a fraction of any biopsy specimen performed in order to evaluate ILDs be saved for a future EM analysis. Since the histological features of the three types of genetic surfactant deficiency might overlap, a biopsy might distinguish among the three entities.

It has been reported that video-assisted thoracoscopy performed by a trained surgeon decreases morbidity from lung biopsy, allowing more rapid recovery and reducing the number of complications.

**Treatment**

The physician–patient relationship is central to the monitoring of chronic diseases that generally demand many appointments, frequent tests and prolonged treatment. It is important that the patient remain in the unit, giving care and attention, and help the family understand the disease.

As occurs in adults, due to the lack of randomized clinical trials involving large samples, there is no standardization of treatment regimens in children. The current therapeutic approach is based on the experience gained from small pediatric groups and on information from studies involving adults. More often than not, these treatment strategies fail, and the reported mortality is still high.

Treatment involves support measures, pharmacological treatment and specific strategies. Oxygen therapy, essential in cases of hypoxemia, might last months or years. Other measures involve appropriate nutritional support, prevention of infections through active or passive immunization against viruses and bacteria and cardiac monitoring. If the cause is identified, specific treatment or removal of triggering factors (environmental or medication-related) should be recommended. Psychological support might be necessary. Most patients require anti-inflammatory or immunosuppressive treatment for months or years. There has been little progress in the pharmacological treatment of ILDs in the last decades.

Oral corticosteroid therapy or pulse therapy are the cornerstone of treatment. An initial course of corticosteroids should be administered for 6 to 8 weeks, and the dose should be gradually removed, generally over months or years, according to patient response, which varies. If significant collateral effects are observed, or if the response is not satisfactory, other agents such as hydroxychloroquine and immunosuppressants/cytotoxic agents (azathioprine, cyclophosphamide, cyclosporine or methotrexate), can be used as corticosteroid-sparing adjuvants or as substitutes. Treatment, however, is not standardized, and doubts remain regarding the dose of oral prednisone (1-2 mg/kg/day), the frequency and dose of pulse therapy with i.v. methylprednisolone (10-30 mg/kg/3 days/month), the criteria used for treatment discontinuation and the potential value of inhaled corticosteroids in these disorders. Our experience, with a group of 25 children under prolonged follow-up, was quite satisfactory, and only two deaths occurred.

There is no consensus regarding treatment in cases that progress to fibrosis, although various drugs targeted at the activity of cytokines, growth factors and oxidants are being tested, principally in adults. The results of a recent meta-analysis of IFN-γ are encouraging. Lung transplants, which are indicated for children with advanced-stage ILD or early in fatal diseases such as certain mutations of the SPs, are increasing in number. Survival is similar to that of other pathologies. The course of the chILD syndrome varies according to the cause. It can lead to death still in the neonatal period, or the patient can reach adulthood with minimal or no symptoms. A score for evaluating disease severity has been
proposed. Higher scores indicate a lower likelihood of survival (Chart 3). Patients who respond well or fairly well to treatment show improved growth (weight and height) and improved psychomotor development, important parameters in children. There are few data regarding the long-term progression of lung function in children with ILD.

Final considerations

Children present peculiar immunological characteristics of defense and repair. In addition, lung development is still not complete. Therefore, children, principally younger ones, should be distinguished from adults. Because they involve large cohorts, the international collaborative studies conducted recently represent an important step toward the definition and individualization of the various characteristics of ILDs. Such studies focus their recommendations on the diagnosis of and approach to pediatric patients in daily practice and search for new information regarding this group of patients.

References

Chronic interstitial lung diseases in children


About the authors

Maria Aparecida Soares de Souza Paiva
Pediatric pulmonologist. Rio de Janeiro State Worker's Hospital, Rio de Janeiro, Brazil.

Sandra Mara Moreira Amaral
Pediatric pulmonologist. Rio de Janeiro State Worker's Hospital, Rio de Janeiro, Brazil.