INFECTIONS AND NONINFECTIONOUS PULMONARY COMPLICATIONS IN PATIENTS WITH PRIMARY IMMUNODEFICIENCY DISORDERS

Yazdani R1,2,3, Abolhassani H2,4, Asgardoon M2, Shaghagi M2,5, Modaresi M6, Azizi G7, Aghamohammadi A2

1Department of Immunology, School of Medicine, Isfahan University of Medical Sciences, Isfahan, Iran
2Research Center for Immunodeficiencies, Children’s Medical Center, Tehran University of Medical Sciences, Tehran, Iran
3Molecular Immunology Interest Group (MIIG), Universal Scientific Education and Research Network (USERN), Isfahan, Iran
4Division of Clinical Immunology, Department of Laboratory Medicine, Karolinska Institutet at Karolinska University Hospital Huddinge, Stockholm, Sweden
5Network of Immunology in Infections, Malignancy and Autoimmunity (NIIMA), Universal Scientific Education and Research Network (USERN), Tehran, Iran
6Department of Pediatric Pulmonary and Sleep Medicine, Children Medical Center, Tehran University of Medical Sciences, Tehran, Iran
7Imam Hassan Mojtaba Hospital, Alborz University of Medical Sciences, Karaj, Iran

Abstract

Primary immunodeficiency disorders (PIDs) are caused by 1 or more defects of the immune system. Patients are more likely to experience recurrent and/or severe infections and tend to develop a wide range of complications. Respiratory diseases are the main and initial manifestation in most cases and the most common complication. Pulmonary complications cause significant morbidity and mortality in patients with PIDs. Early diagnosis and appropriate treatment can prevent or at least slow the development of respiratory complications. Since the spectrum of pulmonary complications in PIDs is broad, we divided pulmonary complications into upper respiratory complications (eg, sinusitis, otitis media, and laryngeal angioedema) and lower respiratory complications (eg, pneumonia, bronchitis, bronchiectasis, interstitial lung diseases, organizing pneumonia, pulmonary adenopathies and malignancies, hyperreactive airway diseases, pulmonary dysgenesis, and adverse reactions to treatment). This review covers the main respiratory manifestations in patients with PIDs.

Key words: Primary immunodeficiency disorders. Pulmonary complications. Upper respiratory tract. Lower respiratory tract.

Resumen

Las inmunodeficiencias primarias (PIDs) son enfermedades causadas por uno o más defectos del sistema inmunológico. Estos pacientes presentan con frecuencia infecciones recidivantes y/o severas así como otro tipo de complicaciones. Las patologías respiratorias son la principal y más frecuente manifestación y complicación de las PIDs. Las complicaciones de estas patologías pulmonares constituyen una de las principales causas de morbimortalidad entre los pacientes que sufren PIDs. El diagnóstico temprano y el tratamiento adecuado pueden prevenir, o al menos retrasar, la aparición de las complicaciones respiratorias en estos pacientes. Dado que el espectro de las enfermedades pulmonares es muy amplio, hemos dividido estas complicaciones entre aquellas que afectan a las vías aéreas superiores (sinusitis, otitis media y angioedema laringeo, etc.) y las que afectan a las vías aéreas bajas (neumonia, bronquitis, bronquiectasias, enfermedades pulmonares intersticiales, neumonía organizada, adenopatías pulmonares y neoplasias, hiperreactividad bronquial, disgenesia pulmonar y las debidas a los efectos secundarios del tratamiento instaurado). Este artículo revisa las manifestaciones respiratorias que se observan más frecuentemente en los pacientes con PIDs.

Introduction

Primary immunodeficiency disorders (PIDs) are a heterogeneous group of genetic disorders caused by 1 or more abnormalities of the immune system. PIDs range from mild to severe forms of immunodeficiency such as severe combined immunodeficiency (SCID). Although the main clinical presentations are recurrent and chronic infections, patients also have an increased incidence of autoimmunity, lymphoproliferative disorders, and cancers [1-4]. The incidence and prevalence of PIDs are unknown, although they are estimated to affect between 1:10 000 and 1:600 individuals in over 40 countries [5]. Recent advances in molecular genetics and immunology have led to the detection of several causative genes and a better knowledge of the pathogenesis of PIDs. More than 300 PIDs have been identified to date, thus expanding the range of deficiencies to immune dysregulation, autoinflammatory disorders, and defects in innate and intrinsic immunity [6]. Immunoglobulin replacement therapy is a lifesaving intervention for many patients with PID, although antibiotics, immunomodulators, and hematopoietic stem cell transplantation (HSCT), can be used under specific conditions [7].

Pulmonary complications are common in patients with PID and contribute significantly to morbidity and mortality. Recurrent respiratory infections are often the first warning sign and are major causes of death. The spectrum of respiratory manifestations is extremely wide, including acute and chronic infections (eg, recurrent, severe, persistent infections affecting various locations, and opportunistic or unusual pathogens), immune dysregulation (eg, autoimmunity, allergy, and lymphoproliferative disorders), structural abnormalities, and malignancies. As survival from infection improves (mortality from PID is still 30%-45% worldwide), noninfectious pulmonary complications are more frequently observed among patients with PIDs [8,9]. Standard management of the respiratory complications of PID is complemented by more specific therapeutic modalities, including surgery (otitis, sinusitis, adenoiditis, and lung infection), lung transplantation, and symptomatic therapies (anti-inflammatory drugs, mucolytics, bronchodilators, and inhalers). Since respiratory disorders are a significant cause of morbidity and the leading cause of death (30%-65%) in both children and adults with PIDs, timely diagnosis and appropriate therapy can improve or at least decelerate the progression of these complications [10-12]. The aim of this study was to review upper and lower respiratory tract complications in patients with PID.

Upper Respiratory Tract

The upper respiratory tract complications of PID include acute and chronic infections (rhino/sinusitis, otitis media, pharyngitis, laryngitis, tonsillitis, and epiglottitis), upper airway anatomical defects (obstruction due to angioedema, obstruction and destruction of the nasal septum mostly secondary to fungal infections in combined immunodeficiency), atopic reactions (allergic rhinitis, mostly in antibody deficiencies), and lymphoproliferative disorders (adenitis or lymphadenitis, mostly in antibody deficiency and immune dysregulation). The most frequent complications are sinusitis, otitis media, and laryngeal angioedema, which are described below.

Sinusitis

Sinusitis (also known as rhinosinusitis) is defined as a common inflammatory condition of the paranasal sinuses and nasal cavity that can be acute or chronic (>12 weeks). Sinusitis is diagnosed based on clinical signs and symptoms, including chronic nasal obstruction, mucopurulent rhinorrhea, postnasal discharge, and cough [13]. Serious sinus infections (more than 2 in childhood and 4 in adulthood) within 1 year are 1 of the 10 warning signs of PID proposed by the Jeffrey Modell Foundation [14]. Chronic sinusitis is very frequent in PID patients [15], although this frequency varies with the type of study and PID, as shown in a systematic review where 10%-54% of patients with chronic sinusitis had PID [16]. We previously demonstrated that sinusitis was the most common ear, nose, and throat presentation in patients with primary antibody deficiency (PDD) [17]. The most frequent PID associated with sinusitis is humoral immunodeficiency. This disorder includes selective IgA deficiency (SlgAD), common variable immunodeficiency (CVID), and specific antibody deficiency (SAD), which is characterized by normal IgG levels but an impaired response to polysaccharide vaccines [17-21]. One study reported considerable differences in the incidence of sinusitis in PIDs, such as agammaglobulinemia (11.52%), CVID (9.4%), SAD (1.8%), IgG subclass deficiency (13.41%), SlgAD (6.58%), and hyper-IgM syndrome (0.7%) [19]. The manifestation of recurrent sinusitis in patients with immune dysregulation (especially PAD patients) demonstrates the importance of immunological testing in patients with recurrent sinusitis [22]. The Table shows some of the bacteria involved in sinusitis in PID patients.

Recurrent rhinosinusitis has a strong negative impact on quality of life [23]. Immunological tests in patients with recurrent sinusitis comprise measurement of antibody levels (IgG, IgA, and IgM), preimmunization and postimmunization specific antibody responses to tetanus toxoid and pneumococcal polysaccharide vaccines, CH50 testing, and determination of T-cell count and function (delayed hypersensitivity skin tests and flow cytometric enumeration of T cells). IgG subclasses should not be checked routinely when evaluating immunodeficiency, as the association between IgG subclass deficiency and recurrent sinusitis is controversial and the clinical significance of abnormal IgG subclasses in patients with recurrent infections is unclear [24]. Only a few studies have investigated the treatment of chronic sinusitis in patients with PIDs, although it seems that treatment involves both medical and surgical approaches, as occurs in immunocompetent individuals. In general, broad-spectrum antibiotics, saline nasal washes, anti-inflammatory agents, and immunoglobulin replacement therapy could reduce the frequency and severity of sinusitis in PID patients [18,25]. While the role of sinus surgery has not been well defined, it seems that PID patients experience similar benefits to immunocompetent populations in terms of symptoms and quality of life [26-29]. If the response to surgery of patients with PID is similar to that of the general population with chronic sinusitis, earlier surgery has been shown to decrease the harmful effects of chronic sinusitis [30].
Innate immunity deficiencies:
- Neutrophil defects:
- Macrophage defects:

Primary antibody deficiency:
- Streptococcus pneumoniae, Haemophilus influenzae
- Moraxella catarrhalis, Chlamydia trachomatis, Bordetella pertussis
- Pseudomonas aeruginosa, Neisseria meningitidis, Ureaplasma urealyticum,
  Mycobacterium hominis, Mycobacterium avium,
- Hemophagocytic lymphohistiocytosis (HLH): Herpesviridae family viruses
- Hyper-IgE syndrome:

Burkholderia pseudomallei, Serratia marcescens, Staphylococcus aureus,
- Scedosporium prolificans, Histoplasma capsulatum, Cryptosporidium parvum,

Combined immunodeficiencies:
- Mycobacterium hominis, Mycobacterium avium,
- Complement deficiencies:

Pseudomonas aeruginosa, Neisseria meningitidis, Ureaplasma urealyticum,
- Moraxella catarrhalis, Chlamydia trachomatis, Bordetella pertussis

Otitis media
- Primary antibody deficiency: Streptococcus pneumoniae and Haemophilus influenzae
- Complement deficiency: Streptococcus pneumoniae and Haemophilus influenzae

Complement deficiencies: Encapsulated bacteria and Neisseria meningitidis

Innate immunity deficiencies: Streptococcus pneumoniae, Haemophilus influenzae, and Neisseria meningitidis

Complement deficiencies: Pneumocystis jiroveci, Pseudomonas aeruginosa, attenuated vaccines (Bacillus Calmette–Guérin strain), Aspergillus fumigatus, Scedosporium prolificans, Histoplasma capsulatum, Cryptosporidium parvum, respiratory syncytial virus (RSV), adenovirus, parainfluenza 3, paramyxovirus, cytomegalovirus, Streptococcus pneumoniae, and Haemophilus influenzae
- Hyper-IgE syndrome: Staphylococcus aureus

Hemophagocytic lymphohistiocytosis (HLH): Herpesviridae family viruses

Congenital phagocytosis deficiency: Klebsiella pneumonia, Aerobacter aerogenes, Burkholderia pseudomallei, Serratia marcescens, Staphylococcus aureus, Pseudomonas aeruginosa, Aspergillus nidulans, Candida albicans, and Nocardia asteroides

Bronchiectasis
- Primary antibody deficiency: Streptococcus pneumoniae and Haemophilus influenzae
- Innate immunity deficiencies:
  - Neutrophil defects: Staphylococcus aureus, Nocardia, Aspergillus and Candida species
  - Macrophage defects: Mycobacteria, Histoplasma, Listeria and Salmonella species

Otitis Media

Otitis media is an infection or inflammation of the middle ear that may point to a variety of underlying problems. This disease affects both children and adults, although it is less common in adults owing to the anatomical development of the upper airways. Four or more serious ear infections in a year are 1 of the 10 warning signs of PID proposed by the Jeffrey Modell Foundation [14]. Recurrent otitis media is considered to be one of the most common infections in PID, although its frequency varies between the different forms of PIDs. Yarmohammadi et al [15] reported a high rate of otitis media in patients with PID (81 of 113 patients, 71.6%), which was significantly higher than in patients without immune defects. Owayed and Al-Herz [31] found the rate of otitis media to be lower (59 of 202 patients, 29.2%). In the third report of the Iranian registry, we reported 46 cases in 731 patients (6.3%) in whom the first presentation of PID was otitis media [3].

Conley and Howard [32] indicated that almost all of the patients who had X-linked agammaglobulinemia (XLA) after the first year of life had a history of recurrent otitis media. Other studies have also reported a high rate of otitis media in XLA patients [31-34]. In one study, we demonstrated that 81.1% of XLA patients manifested acute otitis media. Although this was the second most common infectious disease among Iranian XLA patients, chronic otitis media was the most common complication among chronic manifestations [35]. We found a lower rate of otitis media in CVID patients with hypogammaglobulinemia than in those with agammaglobulinemia [17,36]. This rate was higher in other studies [37-39], although this difference could be due to differences in the number of patients in each study (high in our study). Moreover, since B cells and secondary lymphoid organs are present in CVID patients, in contrast with patients with agammaglobulinemia, a lower rate of otitis media could be logical in CVID. Otitis media is more common in children with CVID than in adults, probably as a result of the anatomic characteristics of the upper airways in children [37]. Patients with a defect in polysaccharide antibody production (with or without IgG subclass deficiency) are very prone to otitis media [40,41]. Infection is more severe in patients with a defect in polysaccharide antibody production associated with IgG subclass deficiency and SlgAD [42-44]. IgG2 may protect against polysaccharides, because it has been indicated that patients with low IgG2 had recurrent episodes of otitis media despite exhibiting normal levels of IgA, IgM, IgG and IgG1 [45]. Sinopulmonary complications such as otitis media are more common in PAD than in combined immunodeficiency.

Table. Microorganisms Involved in Infectious Pulmonary Complications of Primary Immunodeficiency

<table>
<thead>
<tr>
<th>Pulmonary Manifestation</th>
<th>Germs in Primary Immunodeficiency</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sinusitis</td>
<td>Primary antibody deficiency: Streptococcus pneumoniae, Haemophilus influenzae and Neisseria meningitidis</td>
<td>[131], [132]</td>
</tr>
<tr>
<td></td>
<td>Complement deficiencies: Streptococcus pneumoniae, Haemophilus influenzae</td>
<td></td>
</tr>
<tr>
<td>Otitis media</td>
<td>Primary antibody deficiency: Streptococcus pneumoniae and Haemophilus influenzae</td>
<td>[33]</td>
</tr>
<tr>
<td></td>
<td>Complement deficiency: Streptococcus pneumoniae and Haemophilus influenzae</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Complement deficiencies: Encapsulated bacteria and Neisseria meningitidis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Innate immunity deficiencies: Streptococcus pneumoniae, Haemophilus influenzae, and Neisseria meningitidis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Combined immunodeficiencies: Pneumocystis jiroveci, Pseudomonas aeruginosa, attenuated vaccines (Bacillus Calmette–Guérin strain), Aspergillus fumigatus, Scedosporium prolificans, Histoplasma capsulatum, Cryptosporidium parvum, respiratory syncytial virus, adenovirus, parainfluenza 3, paramyxovirus, cytomegalovirus, Streptococcus pneumoniae, and Haemophilus influenzae</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Hyper-IgE syndrome: Staphylococcus aureus</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hemophagocytic lymphohistiocytosis (HLH): Herpesviridae family viruses</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Congenital phagocytosis deficiency: Klebsiella pneumonia, Aerobacter aerogenes, Burkholderia pseudomallei, Serratia marcescens, Staphylococcus aureus, Pseudomonas aeruginosa, Aspergillus nidulans, Candida albicans, and Nocardia asteroides</td>
<td></td>
</tr>
<tr>
<td>Pneumonia</td>
<td>Primary antibody deficiency: Streptococcus pneumoniae, Haemophilus influenzae, Moraxella catarrhalis, Chlamydia trachomatis, Bordetella pertussis, Pseudomonas aeruginosa, Neisseria meningitidis, Ureaplasma urealyticum, Mycobacterium hominis, Mycobacterium avium, adenovirus, and enterovirus</td>
<td>[133], [134], [135], [34], [49], [136], [34], [6], [8], [137], [78], [86], [138]</td>
</tr>
<tr>
<td></td>
<td>Complement deficiencies: Encapsulated bacteria and Neisseria meningitidis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Innate immunity deficiencies: Streptococcus pneumoniae, Haemophilus influenzae, and Neisseria meningitidis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Combined immunodeficiencies: Pneumocystis jiroveci, Pseudomonas aeruginosa, attenuated vaccines (Bacillus Calmette–Guérin strain), Aspergillus fumigatus, Scedosporium prolificans, Histoplasma capsulatum, Cryptosporidium parvum, respiratory syncytial virus, adenovirus, parainfluenza 3, paramyxovirus, cytomegalovirus, Streptococcus pneumoniae, and Haemophilus influenzae</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Hyper-IgE syndrome: Staphylococcus aureus</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hemophagocytic lymphohistiocytosis (HLH): Herpesviridae family viruses</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Congenital phagocytosis deficiency: Klebsiella pneumonia, Aerobacter aerogenes, Burkholderia pseudomallei, Serratia marcescens, Staphylococcus aureus, Pseudomonas aeruginosa, Aspergillus nidulans, Candida albicans, and Nocardia asteroides</td>
<td></td>
</tr>
<tr>
<td>Bronchiectasis</td>
<td>Primary antibody deficiency: Streptococcus pneumoniae and Haemophilus influenzae</td>
<td>[86], [87]</td>
</tr>
<tr>
<td></td>
<td>Innate immunity deficiencies:</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Neutrophil defects: Staphylococcus aureus, Nocardia, Aspergillus and Candida species</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Macrophage defects: Mycobacteria, Histoplasma, Listeria and Salmonella species</td>
<td></td>
</tr>
</tbody>
</table>
Lower Respiratory Tract

Angioedema is characterized by localized swelling that is generally asymmetric owing to increased vascular permeability with leakage of plasma. While this complication can involve all mucosal layers and subcutaneous tissues, in the respiratory tract, it can cause potentially life-threatening obstruction and asphyxiation. Several types of PID should be considered in patients with hereditary angioedema (HAE). Monogenic disorder of the complement system due to decreased level or dysfunction of C1 esterase inhibitor (C1INH) has been estimated to have a prevalence of 1 case per 50,000 individuals. It is important to distinguish between HAE and acquired angioedema in order to modify management and treatment of patients, which can differ for each group and should include purified or recombinant C1INH and bradykinin receptor antagonists.

Pneumonia

Pneumonia is an inflammatory condition of the lung that is most often caused by infection in patients with PID (opportunistic organisms such bacteria, viruses, fungi [Pneumocystis jiroveci]), although it can also be noninfectious. Pneumonia as a complication of the lower respiratory tract is usually multifactorial owing to immune dysregulation (eg, chronic inflammation, autoimmunity, and allergy), adverse effects of therapies, and malignancies [8]. Lower respiratory tract complications are examined in the following sections.

Acute bronchitis is characterized by cough due to acute inflammation of the trachea and large airways when there is no evidence of pneumonia [59]. Cough associated with acute bronchitis typically lasts about 2 to 3 weeks and is usually the chief complaint of patients referred to clinics [59]. Many patients with bronchitis fail to respond to appropriate treatment and are referred to physicians with chronic productive cough. This should therefore be differentiated from chronic bronchitis, which lasts about 2 to 3 weeks [25]. The clinical picture of PID is not distinctive, and symptoms may be difficult to detect at the clinical level. Approximately 70% to 80% of patients experience recurrent respiratory infections, including bronchitis [60].

As occurs in pneumonia (the most frequent type of PID) patients with antibody deficiencies are at increased risk of bronchitis. Karaulov et al [61] demonstrated that 54.4% of CVID patients presented bronchitis as the first
manifestation [61]. Plebani et al [62] reported that lower respiratory tract infections were most frequent in patients with XLA and that 78% of patients had bronchitis. In milder forms of humoral immunodeficiency, recurrent bronchitis manifests in IgG3 deficiency, the most common subclass deficiency (88.1%), and IgG4 deficiency (15.3%) [63]. IgG subclass deficiency is characterized by exacerbation-prone phenotypes, decline in lung function, and subsequent poor prognosis [63]. Although these patients did not manifest evidence of pneumonia or bronchitis, which was defined as ≥3 episodes of acute bronchitis per year [63]. In North American and European studies, antibiotic treatment has not been shown to benefit patients with acute bronchitis, and antibiotic treatment is not recommended for this condition in immunocompetent or immunocompromised patients in developed countries [64].

Bronchiectasis

Bronchiectasis is characterized by chronic and abnormal dilation of the airways and is caused by impaired clearance of various microorganisms and recurrent infection. Previous studies show that 26%-53% of cases of bronchiectasis have no known cause [65,66]. Chronic airway inflammation is supposed to be the primary cause of bronchiectasis, as observed in a condition that involves the airways, such as recurrent pulmonary infections, and autoimmune disease. PID is a common cause of recurrent bronchiectasis. Indeed, PID contributes to increased susceptibility to pulmonary infections and chronic inflammatory airways, which in turn lead to bronchiectasis. This condition is characterized by obstructive or mixed patterns in pulmonary function tests in patients with PID, is generally observed in the lower or middle lobes, and is predictive of a poor prognosis.

Most patients with hypogammaglobulinemia experience the mild type of bronchiectasis [67]. Patients with PADs are at a significantly increased risk for developing bronchiectasis [65,68,69], as we indicated in a study in which 37.5% of patients with bronchiectasis were diagnosed with defects in antibody-mediated immunity [70]. The exact prevalence of bronchiectasis in patients with PAD is unclear, although a systematic review has reported that bronchiectasis manifests in 17%-76% of cases, usually with a tubular/cylindrical pattern affecting the proximal bronchi [71]. More than 70% of CVID patients develop bronchiectasis and bronchial wall thickening [47,72], indicating that bronchiectasis is a well-recognized complication of CVID. The incidence of bronchiectasis in patients with XLA is almost 32%, although XLA has been associated with up to 3% of cases of childhood bronchiectasis [73-75] and is rare in adults. We indicated that the rate and severity of bronchiectasis were greater in CVID patients than in those with XLA, probably because of the earlier diagnosis and treatment of XLA patients [76]. Moreover, CVID patients had a greater likelihood of developing lung disease, possibly owing to delayed diagnosis and immune dysregulation, as compared with XLA patients [76]. Mild to severe vitamin D deficiency in CVID patients is considered to be associated with a risk of bronchiectasis. In the milder form of PAD (eg, SlgAD and IgG subclass deficiencies), the prevalence of bronchiectasis is much lower than that of XLA and CVID. IgG subclass deficiency, particularly that of IgG2, has been associated with bronchiectasis in children [77]. Given that the incidence of bronchiectasis in patients with IgG subclass deficiency varies widely (anywhere from 4% to nearly 50%), it is difficult to establish a correlation between bronchiectasis and IgG subclass deficiency [65,69,77-79]. In one study, SAD was reported in 58% of patients with idiopathic bronchiectasis [80]. However, the study was small, with no matched controls in which the immunological criteria used for SAD were queried [81]. Other larger series of adult patients with bronchiectasis show that the incidence of specific antibody deficiency varies from 4% to 11% [65,82]. Antibody responses to polysaccharide vaccines are variable and depend on age. Furthermore, up to 10% of the healthy population may be nonresponders [75,83]; consequently, it is difficult to evaluate SAD as a cause of bronchiectasis without further studies involving large numbers of bronchiectasis patients and matched controls.

The incidence of bronchiectasis in other rare PIDs such as hyper-IgE syndrome (<2.5% in children and very rare in adults) [74,84,85], phagocyte defects (<1%-10% in children and <1% in adults) [86,87], and transporter antigen peptide deficiency (rare in children and very rare in adults) [87,88] is low. The microorganisms associated with bronchiectasis in patients with PID are shown in the Table.

High-resolution computed tomography (HRCT) is considered a reliable test for assessing bronchiectasis in patients with PID [89-91]. HRCT should be used in all patients with chronic chest symptoms to monitor disease progression. Findings such as bronchial dilatation, signet ring sign (large airway involvement), tree-in-bud sign (small airway involvement), mucopulent plugs, bronchial wall thickening, lack of tapering, and bronchi visible closer than 2 cm to the pleural surface are considered characteristic of bronchiectasis [92]. Since we and others demonstrated that patients with DNA repair defects and some CVID patients show increased radiation sensitivity [93-95], a radiation-free alternative to CT scan or chest X-ray could be magnetic resonance imaging to assess pulmonary changes and alterations [96]. Recognizing the cause of bronchiectasis may improve management and prognosis, eg, initiation of immunoglobulin replacement in PAD patients, which may prevent the progression of irreversible lung damage. While patients with bronchiectasis require higher doses to achieve a satisfactory trough level, the intravenous route is more commonly recommended for administration of immunoglobulin.

Interstitial Lung Disease

Interstitial lung disease (ILD) comprises a group of chronic inflammatory diseases that are major complications of PIDs. ILD is symptomatic in later stages and is associated with pulmonary fibrosis. In advanced stages, patients can develop pulmonary hypertension, cor pulmonale, and respiratory failure. In PID patients with a recurrent respiratory infection, ILD is more frequent and much more prevalent than expected in the general population, especially in antibody-deficient individuals. IgG subclass deficiency and SlgAD,
respective, are the most and least common immune disorders associated with ILD. However, some specific PIDs (e.g., ataxia telangiectasia and chronic granulomatous disease) have been associated with greater susceptibility to ILD [98]. They rarely occur in childhood but represent a large number of conditions involving the parenchyma of the lung including the alveoli, the alveolar epithelium, the capillary endothelium, and the spaces between those structures, as well as perivascular and lymphatic tissue. These disorders are associated with considerable rates of morbidity and mortality. ILD comprises nonmalignant disorders that are generally not caused by identified infectious agents. Moreover, there is little consensus regarding management of most types of ILD.

Many types of PID carry an increased risk of systemic autoimmune disorders that involve respiratory interstitial tissue. Autoimmunity-associated ILD may be more responsive to treatment than idiopathic forms, particularly with targeted therapy using rituximab or anti–tumor necrosis factor (TNF) agents. The main PIDs that manifest with systemic autoimmune disorders are PAD (associated with rheumatoid arthritis, systemic lupus erythematosus, polymyositis, vasculitis, Sjögren syndrome, and scleroderma), phagocytosis defects (antiphospholipid syndrome), and complement deficiencies (systemic lupus erythematosus and vasculitis).

A decrease in the diffusing capacity of the lungs for carbon monoxide could be an early sign of progression to a restrictive lung disease or ILD and should be monitored using additional spirometry testing [99]. In imaging, lung ultrasound is reported to be a sensitive tool for detection of ILD, and combining chest x-ray and lung ultrasound for further evaluation provides various complementary features. The combination of both approaches could reduce the need for HRCT [100]. However, HRCT is the diagnostic choice for early detection and confirmation of suspected ILD, better evaluation of the extent and distribution of disease, and identification of coexisting complications [101]. Antifibrotic agents, especially pirfenidone and nintedanib, can be used in patients with ILD [10].

Granulomatous-lymphocytic interstitial lung disease (GLILD) is defined as the pulmonary manifestation of a multisystem disease and is an umbrella term encompassing lymphocytic intestinal pneumonia, follicular bronchiolitis, pulmonary nodular lymphoid hyperplasia, and reactive lymphoid infiltrates [102,103]. GLILD is frequently accompanied by diffuse autoimmune cytopenia, adenopathy, splenomegaly, and extrapulmonary granulomatous disease, which mainly affects the lymph nodes, spleen, liver, and gastrointestinal tract. The etiology of noncaseating granuloma formation remains unknown, although an increase in levels of TNF, human herpesvirus 8, Epstein-Barr virus, and cytomegalovirus has been proposed [104,105]. No studies report GLILD in XLA, suggesting that T-cell dysfunction is the probable pathogenic mechanism. Hypomorphic mutations in recombination-activating gene 1 (RAG1), haploinsufficiency of cytotoxic T lymphocyte antigen-4 (CTLA4), and deficiency in lipopolysaccharide responsive beige-like anchor protein (LRBA) has been described in patients with GLILD [106].

Of all the ILDs associated with PID, GLILD is the most common, the most widely investigated, and the most closely associated with poor clinical outcomes. The incidence of granulomas in CVID has been estimated to be between 5% and 20% in various cohorts [107-109]. In patients with PID, GLILD manifests with the gradual development of dyspnea on exertion and cough or may be asymptomatic. The presence of GLILD in PID patients points to a poorer prognosis and increases the prevalence of lymphoproliferative disorders [110]. The finding of interstitial fibrosis (with or without architectural remodeling) requires additional study for its effect on prognosis [111]. The typical imaging pattern of ILD in PAD is a generalized diffuse reticular change and consolidation, with or without a ground-glass appearance, predominantly in the lower lobe.

In a recent study [112], positron emission tomography (PET)-CT scanning using fludeoxyglucose was used to assess and monitor the response to treatment in CVID patients with GLILD. The authors found a widespread and high level of metabolic activity in the lungs and lymph nodes, thus suggesting the potential utility of this imaging modality in this subset of patients. In order to decrease the delay in diagnosis, increase early detection, and prevent disease progression, it has been recommended to screen all symptomatic and asymptomatic CVID patients for lung disease [113]. The choice of treatment should be based on the presence of symptoms, history of an inciting medication, extent of extrapulmonary involvement, and careful assessment of the histopathologic grade of the lesion. Immunoglobulin replacement therapy seems to be an effective and satisfying strategy and is also the oldest modality for prevention of progression of GLILD in various forms of PID. However, corticosteroids are the first-line approach in several studies and have been maintained until pulmonary function tests and radiology findings show an improvement in the patient’s condition. In some studies, combination therapies are indicated for patients with GLILD in order to achieve a complete response, because statistics show that although many patients respond well to high-dose corticosteroids, this modality is not effective in some cases [114].

Organizing Pneumonia

Organizing pneumonia (OP), formerly known as bronchiolitis obliterans OP (BOOP), is a relatively rare presentation in PID. It involves a nonspecific response of the bronchial epithelium to infections (particularly those caused by Mycoplasma species), inflammation (autoimmunity, post-HSCT), and fibrosis. OP is clinically characterized by concentric luminal narrowing leading to subacute illness with dyspnea and cough and resting hypoxemia in advanced stages of fibrosis, with a mixed obstructive and restrictive pattern in pulmonary function tests. OP has been reported in CID and PAD patients [72,115,116]. Rao et al [111] evaluated CVID patients and reported OP in 87.5%, mostly in mild forms of the disease. Pulmonary function tests reveal a mild-to-moderate restrictive pattern. We previously reported OP in childhood as a complication of PID due to an LRBA mutation and suggested that mutations in this gene could lead to a variety of immunodeficiencies, ranging from immunoglobulin deficiency to low B-cell count [117].

The diffusing capacity of the lung for carbon dioxide is usually reduced in patients with fibrosis. The chest CT scan shows a typical pattern of bilateral patchy alveolar infiltration. Inspiratory and expiratory HRCT images have been recommended to identify air trapping downstream from the obstruction and show scattered bilateral alveolar opacities.
and ground-glass consolidation in the presence of cryptogenic OP. The histological diagnosis can be made by transbronchial lung biopsy, although open lung biopsy might sometimes be required, as the disease may go undiagnosed in small specimens. Histologically, OP is characterized by Masson bodies, which are spherical proliferations of loose collagen-embedded fibroblasts and myofibroblasts set in a pale, myxoid stroma located within airspaces and the adjacent interstitium. Analysis of bronchoalveolar lavage fluid typically reveals foamy macrophages and an increase in counts of all cell types, mainly lymphocytes and plasma cells in the interstitium.

**Pulmonary Adenopathies and Malignancies**

Localized hilar and/or mediastinal adenopathies can be seen in the context of lower respiratory tract infections, although they are classically associated with either lymphoproliferative disorders or malignancy. GLILD (usually presenting in CVID), chronic granulomatous disease, and immune dysregulation (including hemophagocytic lymphohistiocytosis and autoimmune lymphoproliferative syndrome) are common lymphoproliferative PIDs associated with pulmonary adenopathies. Primary lymphoma (extranodal marginal zone B-cell lymphoma of mucosa-associated lymphoid tissue due to increased genetic vulnerability to Epstein-Barr virus–induced complications) [120,121] and secondary lung metastatic malignancy (in PAD and CID patients with DNA repair system defects, NK-cell defects, and cytotoxic T-cell defects) may present initially with thoracic adenopathy [122-124]. Infiltration of the lung with metastases and subsequent restrictive disease is more common than primary lung tumors.

Pulmonary adenopathies and malignancies could lead indirectly to pulmonary arterial hypertension and superior vena cava syndrome. Malignancy is the second most common cause of death in PID (6%-10% worldwide).

**Hyperreactive Airway Diseases**

Patients with PAD, especially CVID and SlgAD (15%-45% of patients), may have a concomitant history of asthma and other types of hyperreactive airway diseases, although it can be difficult to differentiate between these atopic complications in severe pulmonary conditions [50,125,126]. Both asthma and small hyperreactive airway diseases can cause dyspnea, wheezing, cough, and sputum production. Detection of both conditions is confirmed by the presence of a reversible obstructive pattern in pulmonary function tests and a positive result in the methacholine provocation test.

**Pulmonary Dysgenesis**

Velocardiofacial syndrome (also known as DiGeorge syndrome, affecting 10% of patients) is a hereditary congenital disease that could be associated with immunodeficiency if it involves impaired thymic development due to 22q11.2 deletion. However, in some affected individuals, pulmonary dysgenesis and other structural airway abnormalities may be present in addition to abnormal development of the pharyngeal arch. On the other hand, in cardiac defects, pulmonary artery anomalies such as pulmonary atresia and absent pulmonary valve syndrome can affect lung function [127].

**Pulmonary Complications of PID Treatment**

Pulmonary adverse effects associated with immunoglobulin therapy can be classed as immediate reactions (flu-like symptoms, hypersensitivity reaction, and transfusion-related acute lung injury) and delayed reactions (mainly due to thromboembolic events) [128].

Pulmonary graft-versus-host disease (GVHD) is one of the respiratory complications that can be observed in PID patients with severe CID-induced lack of maternal T-cell engraftment or in other types of PID after HSCT [129]. Pulmonary GVHD can be acute (rare, occurring around 5 months after transplantation with perihilar or diffuse interstitial fibrosis, pulmonary cysts, and pulmonary nodules) or chronic (usually presents as BOOP) [130].

**Conclusions**

The respiratory system is the most frequent site of the clinical manifestations in PID and associated complications are often the first warning sign of PID. Pulmonary complications must be recognized and diagnosed accurately in patients with PID, since they determine prognosis. Since pulmonary complications could be considered a significant cause of morbidity and mortality in PID patients, appropriate awareness of these manifestations is essential, especially for the pulmonologist.

**Funding**

The authors declare that no funding was received for the present study.

**Conflicts of Interest**

The authors declare that they have no conflicts of interest.

**References**


Pulmonary Complications in PID


Manuscript received March 13, 2017; accepted for publication April 24, 2017.

Asghar Aghamohammadi

Children’s Medical Center Hospital
62 Qarib St., Keshavarz Blvd.
Tehran14194, Iran
E-mail: aghamohammadi@sina.tums.ac.ir