Parasitic lung diseases

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Abstract
Parasitic lung diseases are caused by a number of parasites as a result of transient passage in the lung or as a result of an immunologic reaction. The clinical presentation may be in the form of focal or cystic lesions, pleural effusion or diffuse pulmonary infiltrates. With increasing globalisation, it is important to consider parasitic infections in the differential diagnosis of lung diseases. This is particularly important since early identification and prompt therapy result in full cure of these conditions. In this review, we summarise the most common parasitic lung diseases.

Introduction
Parasitic lung diseases are selective diseases caused by a number of parasites. They may be related to the transient passage of the parasites in the lung or as a result of an immunological disease. Since parasites are usually a cause of disease among residents of developing countries, parasitic lung diseases are normally seen in these countries. However, globalisation, increased international travel and immigration may lead to the presentation of patients in well-developed countries as well [1]. In this review, we summarise the most common forms of parasitic lung diseases.

Pattern of parasitic lung disease
The most common causes of parasitic lung disease are listed in table 1. The clinical presentation of such patients may be in the form of focal or cystic lesions, coin-like, consolidation, pleural effusion, diffuse or transient pulmonary infiltrates, or alveolar or interstitial lung changes [2]. The most common routes of transmission of common pulmonary parasites are shown in figure 1. A wide range of parasites cause disease in humans and include nematodes, trematodes, cestodes and protozoa (table 1).

Parasitic eosinophilic lung disease
There are three types of parasitic eosinophilic lung disease, as shown in figure 2. These are Loeffler’s syndrome, tropical eosinophilia syndrome (TES) and visceral larva migrans.

Loeffler’s syndrome
The term Loeffler’s syndrome denotes two different syndromes: transient eosinophilic pulmonary infiltrate and eosinophilia with endocarditis or myositis, thus causing some confusion [3]. This confusion is compounded by the subsequent classifications of the causes of Loeffler’s syndrome: idiopathic, drugs or infections.
The initial syndrome was associated with pulmonary infiltrates and peripheral eosinophilia as a result of transbronchial migration of *Ascaris* species [3]. Additionally, the syndrome may result from a hypersensitivity reaction to the transient parasitic immigration into the lungs. It is also known as simple pulmonary eosinophilia as it lacks pulmonary symptoms or is only associated with mild symptoms [4]. It is characterised by a transient pulmonary infiltrate that spontaneously resolves within 2–4 weeks. Patients may have peripheral eosinophilia [2], severe bronchospasm, fatigue and fever [5]. It is more common in children than adults. The syndrome is associated with disruption of the respiratory epithelium, ciliastasis, mucus production and a significant release of platelet activating factor and leukotrienes, which in turn lead to bronchospasm and diffuse lung infiltrate. The major causes of Loeffler’s syndrome are *Ascaris lumbricoides*, *Ancylostoma duodenale*, *Necator americanus* and *Strongyloides stercoralis* [5]. A special form of Loeffler’s syndrome is caused by *S. stercoralis* in immunocompromised patients, such as those on steroid treatment with superinfection [6], and may be associated with Gram-negative bacteraemia.

Diagnosis is based on clinical suspicion as well as demonstration of the parasitic eggs of *Ascaris lumbricoides*, *Ancylostoma duodenale* or *N. americanus* in stool, or of the larva of *S. stercoralis* in stool, sputum or bronchoalveolar lavage [3]. Treatment is either mebendazole 100 mg orally twice daily for 3 days or albendazole (for children >2 years) 400 mg·day$^{-1}$ for 3 days [5]. For *S. stercoralis* with superinfection, the treatment is albendazole 400 mg orally twice daily for 7 days or ivermectin 200 $\mu$g·kg$^{-1}$ orally once daily for 1–2 days [6, 7].

**TES**

TES is a type of hypersensitivity reaction to a few parasitic agents. The reaction is usually directed towards microfilaria of *W. bancrofti* and *Brugia malayi*. Tropical pulmonary eosinophilia (TPE) was first described by Weingarten [8] in 1943 and was initially called “pseudotuberculosis with eosinophilia” [9]. Other names include filarial tropical pulmonary eosinophilia [10]. *W. bancrofti*, *B. malayi* and *Brugia timori* are the aetiologic agents of lymphatic filariasis and are transmitted to humans through different mosquitoes, such as Anopheles, Aedes, Culex, Mansonia and Ochlerotatus [11]. Lymphatic filariasis occurs mainly in the tropics and sub-tropics with highest transmission in Central and Latin America, Africa (west, coastal

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**TABLE 1** Summary of common parasitic lung diseases

| Nematodes | Trematodes | Cestodes | Protozoa |
|-----------|------------|----------|----------|
| *Ascaris lumbricoides*<br> *Ancylostoma duodenale* and *Necator americanus*<br>*Strongyloides stercoralis*<br>Tropical pulmonary eosinophilia: *Wuchereria bancrofti*<br>*Schistosoma haematobium* and *Schistosoma japonicum*<br>*Paragonimus westermani*<br>*Echinococcus granulosus* and *Echinococcus multilocularis*<br>*Amoebiasis: Entamoeba histolytica*<br>*Leishmaniasis (Ibaka): Leishmania donovani*<br>*Malaria: Plasmodium vivax, Plasmodium falciparum, Plasmodium malarii, Plasmodium ovale and Plasmodium knowlesi*<br>*Toxoplasmosis: Toxoplasma gondii*<br>*Babesiosis: Babesia microti* | | | |

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**FIGURE 1** Route of transmission of common lung parasites.
east and southern parts), India, Southeast Asia, Indonesia, Papua New Guinea and the western Pacific [11]. It was indicated that transmission increases with high rainfall and temperature, the growth of certain types of vegetation, and high population density and decreases at high altitudes [11].

TES occurs in only a small fraction (<0.5–1.0%) of patients infected with filaria and has a male predominance of 4:1 [1]. The disease occurs in Southeast Asia, China, India and Africa and may occur outside those geographic areas among immigrants or visitors [12]. The syndrome is slow in onset and occurs over many months. It is characterised by the occurrence of cough, dyspnoea and wheezing associated with systemic symptoms such as fever, malaise and weight loss. The exact pathophysiology of TES is not known, but it is thought that an increased IgE immune response to a filarial antigen known as G-glutamyl transpeptidase (gGT) may lead to a higher risk of TES [13]. The levels of gGT-specific IgE/IgG4 ratio were 45 and 107 in patients with TES and those who live in the tropics without TES [13]. In addition, acidic calcium-independent phospholipase A2 was found to be a master regulator of TPE pathogenesis [14].

The diagnosis of TES/TPE relies on a constellation of findings, which include being a resident of or coming from a filaria endemic area, the presence of blood eosinophilia, the absence of peripheral microfilariae, high antifilarial antibody titres and serum IgE level, as well as a clinical response to diethylcarbamazine (DEC) [13]. TPE may also mimic asthma but is associated with an eosinophilic count of >3×10⁹ cells and a high IgE level of >1000 IU·mL⁻¹ [15]. Diagnosis of TPS is based on the presence of IgG and IgM antifilaria antibodies.

One study found the following sensitivities: filarial antibody test 30%, radiological changes 45.5% and erythrocyte sedimentation rate 80% [16]. The authors proposed the following for the diagnosis of TPS: worsening cough at night, residence in a filarial endemic area, presence of an eosinophil count >3300 cells·mm⁻³ and a clinical and haematological response to DEC [16]. Another study showed high levels of filarial-specific IgG, IgM and IgE [17]. In a study of 12 patients, the main computed tomography (CT) scan finding was widespread ill-defined bronchocentric nodules [18]. In acute TPE, a mild-to-moderate obstructive pattern is usually observed, and chronic cases may show a restrictive pattern [1].

Therapy usually requires DEC for 3 weeks and is associated with a 20–40% failure rate in those with chronic disease [19], which may be related to inadequate treatment or advanced disease [1]. Residual airway inflammation and reduction of lung function after DEC therapy may be observed and can be treated with a course of corticosteroid or other potent anti-inflammatory agents [19, 20].

**Visceral larva migrans**

Pulmonary involvement in the case of visceral larva migrans is mainly associated with the migrating larva of *Toxocara canis*, but might also be caused by *Ascaris suum* or *Toxocara catis* [21]. Patients present with cough, wheezing, acute bronchiolitis, asthma and acute pneumonia with or without eosinophilia, and other symptoms related to allergic and inflammatory responses [22]. These symptoms are seen in about 80% of patients with visceral larva migrans [23]. Secondary chronic eosinophilic pneumonia may also develop [24]. Radiography may show non-specific calcified nodules suggesting old *Mycobacterium tuberculosis* or other nontuberculous infection (figure 1). The appearance on a CT scan could be multifocal subpleural nodules with halo or ground-glass opacities and ill-defined margins [21] (figure 3). In patients with pulmonary visceral larva migrans due to *A. suum*, a high-resolution CT scan showed multiple
nodules (57%) and nodular ground-glass opacities (29%) [25]. Similar findings were also reported in another study [26]. Diagnosis relies on serum enzyme-linked immunosorbent assay (ELISA) of antibodies directed towards the larva of these worms [27].

**Pulmonary schistosomiasis**

The burden of schistosomiasis is significant, with an estimated 200 million infected people globally, mainly in low- and middle-income countries such as Africa, parts of Asia and South America [28]. *Schistosoma* may cause an acute or a chronic pulmonary disease or the lung could be involved in Katayama fever [29]. Acute and chronic schistosomiasis results in nodular pulmonary lesions and the disease is likely related to eggs laid by adult worms during their migration in the chronic phase. In addition, the pulmonary nodules formed during Katayama fever (syndrome) are secondary to a systemic immune-allergic reaction [29]. The distinction between acute and chronic schistosomiasis is based on an arbitrary cut-off time frame of 10–12 weeks from the time of exposure [30]. Acute schistosomiasis (Katayama fever) usually occurs among nonimmune travellers and is associated with fever, malaise, fatigue, headache, cough, abdominal pain and urticarial rash [29, 30]. The disease is related to the migration, maturation and egg-production of schistosomal larva [31], and subsequent immunological reactions [31].

Radiographic manifestations may include distinct nodules, ground-glass opacities and mass-like lesions [32]. The diagnosis relies on finding schistosome eggs in stool, urine or tissue biopsy samples [32]. However, in Katayama fever the diagnosis can also be established by serum PCR if urine and stool samples are negative for *Schistosoma* eggs [33].

One of the manifestations of schistosomiasis is schistosomiasis-induced pulmonary arterial hypertension (SchPAH) [34]. SchPAH is considered a global disease due to the high prevalence of schistosomiasis [35–37], with a prevalence of 5% among those chronically infected with schistosomes [38] and 5–30% in patients with hepatosplenic schistosomiasis secondary to *Schistosoma mansoni* [39–41]. Another study showed that SchPAH results in 30.8% of all cases of pulmonary hypertension in endemic areas [42]. In a retrospective study from China, only 10 out of 18829 (0.053%) people with schistosomiasis had pulmonary hypertension diagnosed by echocardiography [43]. Other parasitic causes of pulmonary hypertension include *W. bancrofti, Clonorchis sinensis* in southeast Asia and *Echinococcus* (hydatid cysts) [44]. There are three possible explanations for the development of SchPAH: *Schistosoma* eggs obstructing pulmonary vasculature, schistosomal egg embolisation from portopulmonary hypertension and the development of inflammatory vasculopathy [38]. Pulmonary schistosomiasis is more common from *S. mansoni* and *Schistosoma japonicum*. It was found that *S. japonicum* caused milder SchPAH than that caused by *S. mansoni* in an animal model, which was thought to be due to the uniform versus clustered distribution of eggs laid by *S. mansoni* and *S. japonicum*, respectively [45]. In an experimental study, *S. japonicum* resulted in 35-fold smaller granulomas than those caused by *S. mansoni* due to attenuated type 2 inflammation [45].
The pathogenesis of SchPAH is thought to be related to angiogenesis. Thus, inhibition of angiogenesis by the use of bevacizumab was associated with less angiogenesis in a mouse model [46]. Current therapies for SchPAH rely on nitric oxide pathway inhibitors such as phosphodiesterase type 5 inhibitors (sildenafil and tadalafil), soluble guanylyl cyclase stimulators (riociguat), endothelin receptor antagonists (bosentan, ambrisentan and macitentan) and prostanoid analogues (epoprostenol, treprostinil, iloprost, and the nonprostanoid IP-receptor agonist selexipag), as fully described by Sibomana et al. [47]. Experience has shown that phosphodiesterase type 5 inhibitors improve symptoms and may also increase survival rates [48].

Other Schistosoma-associated pulmonary diseases include spontaneous pneumothorax [49], hydropneumothorax with pleural thickening [32] and nodular pulmonary lesions [50]. The latter can be acute in the case of travellers or chronic in residents of endemic areas [50]. A metabolically active pulmonary nodule with 18F-Fluorodeoxyglucose positron emission tomography/CT has been described in relation to schistosomiasis [51].

**Paragonimiasis-associated pleural effusion**

The most common species associated with paragonimiasis are *Paragonimus westermani*, *Paragonimus miyazakii*, *Paragonimus mexicanus* and *Paragonimus skrjabini* [52]. After oral ingestion, adult *Paragonimus* parasites pass through multiple organs or tissues and then later reach pulmonary tissues from the small intestine [52]. Paragonimiasis may be associated with cystic lesions and nodules, and 20% of patients may have asymptomatic lesions [3]. Pleural effusion had been described specifically with paragonimiasis and was reported in 40–70% of patients and associated with eosinophilia [52]. However, paragonimiasis may also cause nodules, infiltrates and cavities [52] (figure 4). The most common radiographic finding is pleural effusion (associated with massive eosinophil infiltration) [53]. Although the classic presentation of *Paragonimus* infection is cavitary pulmonary lesion, this finding was seen in only 5% of patients [53]. One study of chest CT scans showed nodular opacities (56.4%) and worm migration tracks (18.1%) [54]. A rare presentation of Loeffler’s syndrome was also described with *P. westermani* [55].

The diagnosis of paragonimiasis relies on detection of *Paragonimus* eggs in stools, sputum, bronchoalveolar lavage fluid or pleural effusion [53], but this is only seen in a minority (11.7%) of...
patients. The most widely used serology for the diagnosis of *Paragonimus* infection is indirect ELISA with a significant cross-reaction with schistosomiasis, fascioliasis and clonorchiasis [56]. Histopathology of tissue may exhibit chronic granulomatus inflammation with numerous parasitic eggs consistent with paragonimiasis (figure 5). The treatment relies on the use of biltricide orally at a dose of 25 mg·kg⁻¹ three times a day for 3 days [57]. A surgical approach is also considered in patients with severe pleural or pleuro-parenchymal pulmonary disease [58].

**Pulmonary amebiasis**

*Entamoeba histolytica* is the aetiologic agent of amebiasis and causes intestinal and extraintestinal manifestations including liver and pulmonary amebiasis. It was estimated that 7–20% of patients with liver amebiasis and 2–3% of those with invasive disease also have pleural effusion, lung abscesses or pleural empyema [59]. Amebiasis has also been found to cause empyema and vena cava obstruction [60, 61].
Pulmonary hydatid disease

The larvae of *Echinococcus* species, especially *Echinococcus granulosus*, are the aetiologic agent of hydatid disease [2]. The disease was reported in South America, the Mediterranean region, the Middle East, sub-Saharan Africa, Russia and China and its prevalence was estimated to be up to 79 cases per 100 000 people [62]. Pulmonary hydatid disease accounts for 20–30% of hydatid disease [63, 64].

The main presentation is incidental findings on radiography; however, symptoms such as cough, haemoptysis, chest pain or pneumothorax may occur [2]. Radiographic presentations of pulmonary hydatid disease differ between *E. granulosus* and *Echinococcus multilocularis*. Hydatid disease due to *E. granulosus* causes smooth margin, noncalcified nodules or masses when the cysts are not ruptured [65]. On the other hand, *E. multilocularis* causes alveolar echinococcosis and radiographically presents as lobulated nodules or masses and calcification [66]. The main radiographic characteristics of hydatid cysts are meniscus sign, crescent sign, Cumbo’s sign, onion peel sign, water lily sign and mass-within-a-cavity sign [67, 68]. One of the classical radiograph signs of hydatid cysts is the presence of a well-defined round radio-opacity in the lung with or without calcification (figure 6). Hydatid disease of the lung may become secondarily infected and on chest radiography may present with a cyst and an air fluid level (figure 7).

Treatment relies on surgical excision of the cyst with or without lung tissue resection [69]. In a systematic review, the use of uni-portal video-assisted thoracoscopic surgery was associated with an overall complication rate of 9.35% and no reported recurrence [69]. Generally, surgical removal of pulmonary hydatid cysts is associated with low morbidity and mortality [70]. One study of 12 patients with pulmonary hydatid disease showed a cure rate of 72% and there was no change in 18% patients using albendazole (800 mg·day\(^{-1}\)) for four courses with each course lasting 28 days followed by 14 days drug free [71]. The use of albendazole is not recommended pre-operatively due to thinning of the capsule but it is used...
post-operatively [64, 72] as surgical therapy and is associated with a high recurrence rate of 25% [73]. In one study, pre-operative albendazole therapy was associated with a reduction in the tensile strength values of the cuticular membrane of the hydatid cysts [72].

One study showed a recurrence rate of 18.7% for those who did not receive albendazole versus 4.16% for those who had pre- and/or post-operative albendazole in liver disease [74]. On the other hand, children especially those with small cysts, young cysts or cysts with daughter cysts may respond well to albendazole therapy without surgery [75]. Needle aspiration of hydatid cysts with antiparasitic therapy before and after the aspiration has also been advocated [76]. One of the therapeutic techniques for hydatid cysts is the use of the percutaneous aspiration of cyst contents, installation of scolicidal agent and re-aspiration (PAIR) technique. This technique had been used with great success with the caveat of the possibility of risks associated with PAIR of pulmonary cysts [77].

**Pulmonary dirofilariosis**

This parasitic disease is caused by a filarial nematode called *Dirofilaria immitis*. Mosquitoes transmit the parasites from dogs to humans [78]. Dirofilariosis is diagnosed most commonly in the east coast and southern parts of the United States, with infrequent cases elsewhere in the world [78–82]. A recent study from the Balkan Peninsula reported that two of 46 cases of dirofilariosis were pulmonary infections [80]. The disease is usually asymptomatic and manifests as a solitary pulmonary nodule that requires surgical excision [78, 79]. The disease does not have specific radiographic or symptomatology that differentiates it from other solitary pulmonary lesions [79].

**Pulmonary manifestations of malaria**

Malaria is caused by *Plasmodium falciparum*, *Plasmodium vivax*, *Plasmodium ovale* and *Plasmodium malariae* and is transmitted by the *Anopheles* mosquito [83]. The main symptoms of malaria are fever, chills, sweating, cytopenia and splenomegaly. The most frequent pulmonary manifestation of malaria is acute respiratory distress syndrome and it is one of the criteria for severe and complicated malaria [67]. Other pulmonary manifestations include pleural effusion, interstitial oedema and consolidation [84–86]. Transient cough was described in 36% of patients with *P. falciparum* infection and in 53% of those with *P. vivax* or *P. ovale* infections [87]. In addition, subclinical airflow obstruction was also demonstrated by spirometry as well as the occurrence of impaired gas exchange [87]. Other manifestations included bronchiolitis obliterans organising pneumonia [88]. The diagnosis is usually made by identifying trophozoites or other parasitic forms within the erythrocytes in a thin blood smear or parasites in a thick smear. Serologic and nucleic acid amplification tests are also available.

**Points for clinical practice**

- A number of parasites may cause pulmonary disease as a result of a transient passage in the lung or as a result of an immunological reaction.
- The clinical presentation may be in the form of focal or cystic lesions, pleural effusion or diffuse pulmonary infiltrates.
- Of particular interest are: Loeffler’s syndrome, TES and visceral larva migrans.
- Increasing globalisation may lead to the presentation of pulmonary lung disease in travellers and migrants.

**Conclusion**

The pulmonary involvement of different parasitic agents is well known globally. The most common manifestations involve radiographic abnormalities, with the possible occurrence of solitary pulmonary nodules. Other manifestations may reflect systemic involvements and it is important to recognise these symptoms for proper diagnosis and management, especially among travellers and migrants.

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