Bronchiolitis in adult: A review

Alcibey Alvarado* and Isabel Arce

1Internal Medicine and Neumology, Clínica de Diagnóstico Médico, San José, Costa Rica
2Medicine and General Surgery, Medicine School, University of Costa Rica. San José, Costa Rica

Abstract

Few diseases have a greater effect on the health of young children than viral respiratory illness of lower tract. This does not happen in adults, possibly due to the different states of activation and expression of the innate immune response and acquired immune responses in these two age groups. Bronchiolitis is a general term used to describe non-specific inflammatory injury that primarily affects the small airways and generally limits the extent to interstice. In the adult clinic, conventional and high-resolution radiology and respiratory functional studies are suggestive of the diagnosis but the etiology usually requires tissue. For this reason, in this work, although there are clinical and radiological classifications, we will use the histologic classification. The goal is a simple, concise and updated monograph issue discussing the different types of adult bronchiolitis, pathophysiology, diagnosis and current therapeutic options.

Introduction

Bronchiolitis is a disease of the small airways, which are defined as airways with less than 2 mm in diameter and no cartilage [1]. Membranous bronchioles have full fibromuscular wall but this wall is very thin [2]. Bronchioles derived from the tertiary bronchus and form a transition area between the major airways and lung parenchyma and fundamentally have a centrilobular location. For its abundant structure (occupy an important area of the airway), its respiratory mechanics (without cartilage and small diameter), the transport of gases to and from the alveoli and their defense mechanism, are particularly vulnerable to infectious, inhaled, vascular, pharmacological and immunological injury [3]. The primary symptoms of bronchiolitis are coughing and dyspnea and during the respiratory auscultation crackles and/or wheezing can be perceived. Respiratory function tests revealed a non-reversible obstructive defect, hyperinflation and generally decreased diffusion (DLCO). Some jobs just used as functional obstructive criteria the reduced FEV1, FVC and FEV1/FVC [2]. There is subtle evidence of early detection of small airway dysfunction such as closing volume, closing capacity, dynamic compliance, the isoflow-volume and forced oscillation technique (FOT) [4,5]. FOT is a convenient tool for measuring respiratory mechanics (resistance and reactance) by applying external oscillatory pressure during tidal breathing [6]. The detail in these tests is generally used in research and does not always have work routine diagnosis of respiratory diseases, particularly in third world countries. The flow-volume curve has greater availability in resource-limited countries [14,15]. The picture is not so clear in adults with acute bronchiolitis. In adults, it has been reported in patients with Mycoplasma pneumoniae, RSV, measles, influenza, pertussis, parainfluenza, and adenovirus [16]. The clinical presentation is poorly

Correspondence to: Alcibey Alvarado, Internal Medicine and Neumology, Clínica de Diagnóstico Médico, San José, Costa Rica, Tel: 50622237134; 50622566439; 50688783325; 50687351858; Fax: 50622216754; E-mail: alcialvagonza@yahoo.com.mx

Key words: small airways, inflammation, bronchiolitis

Received: January 14, 2017; Accepted: February 03, 2017; Published: February 06, 2017
defined unlike children. Often, a disease of the upper respiratory tract precedes the onset of symptoms of lower respiratory tract of cough, exertional dyspnea, fever, tachypnea, and wheezing. Generally, if no risk factors, is self-limited [1].

RSV is now recognized as a significant problem in certain populations of adults, including the elderly patient, people with cardiopulmonary disease, and immunocompromised host. The impact of RSV in older adult may be similar to influenza not pandemic. In addition, RSV causes 2-5% of community-acquired pneumonia in adults [17]. In long-term care facilities RSV is predictable cause of respiratory disease, infecting 5-10% of residents per year, with rates of pneumonia 10-20% and death in 2-5%, but the percentage of these patients with bronchiolitis is not known. In one series, CT showed thickening of bronchial walls, centrilobular nodules and / or pattern of “tree-in-bud” [18]. Occasionally, this acute syndrome has been reported to progress to bronchiolitis obliterans. They have recently reported other viruses such as EV D68 (enterovirus) causing bronchiolitis in adults, associated with low levels of IgG3 and bronchial hyperreactivity up to 5-7 years after bronchiolitis in infancy associated with polymorphism of TLR4 (Toll-like receptors) and deterioration of lung function in children with polymorphism the TLR7 [8,19].

**Proliferative bronchiolitis**

It is a distinctive histopathological pattern in which characteristic intra-luminal fibrotic buds, called Masson bodies, extend beyond alveolar ducts to alveoli. When proliferative bronchiolitis is associated with extension of inflammatory cells into the more distal pulmonary parenchyma, the process is called organizing pneumonia. It is not uncommon finding foamy macrophages in the alveolar spaces [20]. This histopathological finding is associated with many pulmonary disorders. Proliferative bronchiolitis is particularly extensive or prominent in patients with cryptogenic organizing pneumonia (COP), previously called idiopathic bronchiolitis obliterans organizing pneumonia (BOOP). COP is one of seven idiopathic interstitial pneumonias and results from injury to the alveolar epithelium, although the bronchioles are usually committed due to its close proximity [21]. It can also be found as a minor finding in other interstitial diseases, such as chronic eosinophilic pneumonia, hypersensitivity pneumonitis, idiopathic pulmonary fibrosis and nonspecific interstitial pneumonia. It may also occur by exposure to nitrogen dioxide. HCRT show patchy airspace consolidation, “ground-glass” opacity and small nodular opacities in the periphery and in lower lung areas. When the proliferative bronchiolitis is the predominant histological finding, the next step is to determine whether it is cryptogenic organizing pneumonia or secondary. This bronchiolitis typically responds to glucocorticoids; the specific regimen depends on the clinical changes and the underlying cause [22].

**Follicular bronchiolitis**

This bronchiolitis (FB) is a polyclonal hyperplasia of bronchiolar associated lymphoid tissue (BALT) produced by antigen stimulation, with reactive germinal centers distributed along the bronchioles and bronchi to a lesser extent, and associated with low interstitial disease. Well-structured lymphoid follicles are located between the bronchioles and pulmonary arteries and hyperplasia following compress the light of the bronchioles causing obstruction or complete obliteration [23]. This entity is part of the “lymphoproliferative pulmonary diseases” (LPDs) and is associated with connective tissue diseases (rheumatoid arthritis, Sjögren’s syndrome), interstitial lung diseases (ILDs), disorders of mixed collagen-vascular, obstructive diseases of airways and states immunodeficiency including HIV and common variable immunodeficiency (CVID) secondary FB [10,24]. When there is no such association it’s classified as primary FB. LPDs, in turn, include three groups of pathologies: 1) reactive/non-neoplastic lymphoid lesions that are subdivided based on the pattern of lung involvement in lymphoid nodular hyperplasia (LNH) (focal commitment); FB (peribronchiolar commitment); lymphoid interstitial pneumonia (LIP) (diffuse involvement with lung cysts) 2) lymphoproliferative malignant parenchymal lesions that are subdivided into primary and secondary. Primary (0.5% of all primary lung neoplasms) include lymphomas of extranodal marginal zone of origin MALT (lymphoma MALT), diffuse large B-cell lymphoma (DLBCL) and lymphomatoid granulomatosis (LYG). The second ones include Non-Hodgkin lymphoma (NHL) and Hodgkin lymphoma (HL). [3] immunocompromised host lymphoproliferative disorders, including lymphoma related to acquired immune deficiency syndrome (AIDS) (ARL) and post-transplant lymphoproliferative disorder (PTLD) [25]. Pulmonary function tests in patients with FB are generally nonspecific and may be normal, restrictive, obstructive or mixed pattern of airflow limitation.

CT cardinal changes are the small centrilobular nodules associated, sometimes to peribronchiolar nodules and “ground-glass” areas. One of the most distinctive patterns that have been associated with bronchiolar disease is “tree-in-bud” pattern, representing impaction with secondary bronchiolar exudate material infection or inflammation [26]. The presence of peribronchial inflammation and formation of peribronchiolar lymphoid follicles may appear as a “tree-in-bud” on HRCT, because the lymphoid follicles become heavily concentrated in the area adjacent to the bronchioles gap and fade away from the interstitium furthest from the airway. This pattern on HRCT has been described as “cotton-in-bud”. The dilated and thick-walled bronchioles give the appearance that pathologically correlates with follicular bronchiolar obstruction [26]. Histological diagnosis requires two fundamental changes: the presence of lymphoid follicles well formed on the walls of the bronchioles and secondary obstruction or complete obliteration of the bronchiolar light. Distribution is mainly in the interlobular septa and low in the alveolar septa, whereas the diffuse involvement of the latter is considered the key feature of the LIP [27]. Yet, many pathologists today describe these two entities as a continuum of reactive, lymphoid lung diseases, where in many cases distinction is arbitrary. FB associated secondary histological changes are organizing pneumonia, obstructive pneumonia, and infiltrates of intraluminal bronchiolar neutrophils [28]. Immunohistochemistry is essential to rule out malignancy. In the absence of primary immunodeficiency states, FB usually reveals positive for CD79a and CD20 on B cells, predominantly within lymphoid peribronchiolar aggregates, and CD3 positive in T cells, predominantly in the alveolar interstice when there is overlap with LIP [29]. Clinicopathological and radiological correlation is essential for differential diagnosis.

In FB, the basic operation is to treat the underlying disease. In patients with CVID, intravenous gamma globulin may reduce the frequency and severity of lung infections. Rituximab and azathiorpne combined have recently demonstrated radiographic improvement and lung function in these patients [30]. When FB is associated with interstitial disease, treatment usually involves immunosuppressive therapy. The FB associated with HIV generally improves with the initiation of anti-retroviral therapy [24]. In the idiopathic form, the anecdotal use of steroids has reported improvement in clinical symptoms and radiological abnormalities [31]. However, there are...
Respiratory bronchiolitis

Respiratory bronchiolitis (RB) is a well-recognized histological lesion found in the lungs of many young smokers, but is usually asymptomatic. It is recognized by the presence of clusters of tan-pigmented macrophages in the respiratory bronchioles. These intraluminal macrophages are accompanied by a patchy submucosal and peribronchiolar infiltrate of lymphocytes and histiocytes. Peribronchiolar fibrosis extends in a stellate pattern into contiguous alveolar walls [4]. The inflammatory profile in normal smokers is very similar to that of patients with chronic obstructive pulmonary disease (COPD), but less prominent. The concept that emerges is that there is an amplified response in patients with COPD [34]. The RB may be responsible for subtle functional alterations in young smokers and may precede the development of emphysema in genetically predisposed patients. This becomes very important because the RB can be resolved after stopping smoking and this information could be relevant in the anti-smoking campaigns. When the RB is more severe, symptoms usually translate a greater interstitial involvement and then is called respiratory bronchiolitis-associated interstitial lung disease (RB-ILD). On HRCT, RB-ILD is associated with diffuse or patchy “ground-glass” opacities, fine nodules, and air trapping, predominantly in the upper lobes.

Airway-centered interstitial fibrosis

Airway-centered interstitial fibrosis (ACIF) is a disease that has been recognized relatively recent and particularly affects women between 40-60 years of age [35]. Also, known as idiopathic bronchiolocentric interstitial pneumonia or chronic bronchiolitis with fibrosis, is a form of bronchiolitis in which the key histopathological finding is a centrilobular and bronchiolocentric inflammatory infiltrate with peribronchiolar fibrosis and an absence of granulomas [36]. Prominent epithelial hyperplasia compromising the adjacent alveolar septa is usually present; squamous metaplasia, goblet cell metaplasia and necrosis have been described. In clinical groups, it was thought to have hypersensitivity pneumonitis (HP) characteristics but not specific antigens were identified. Many patients reported to have been smokers. Others had chronic silent microaspiration, hypersensitivity or toxic reactions, turning it difficult to segregate the contribution of each factor [37]. The clinical picture is of a chronic nonproductive cough, progressing faster than HP, no specific interstitial pneumonia (NSIP) or RB-ILD, the other diagnoses have a similar presentation. The lung function tests can show obstructive or restrictive patterns. HRCT scans of ACIF show a combination of “ground-glass” opacities, traction bronchiectasis, and bronchial wall thickening. This latter finding is characteristic of most advanced peribronchiolar fibrosis [20]. Unlike HP, the percentage of lymphocytes in the bronchoalveolar lavage (BAL) is less than 40%. The diagnosis is based on lung biopsy. The treatment is not known. Some patients improve with inhaled or systemic glucocorticoids, but the disease progresses generally in at least half of the patients reported, although ACIF has better prognosis than idiopathic pulmonary fibrosis [35].

Diffuse aspiration bronchiolitis

This bronchiolitis is a poorly defined and occult entity. It’s found most often in young or middle-aged patients with identifiable risk factors such as gastroesophageal reflux disease (GERD), drug abuse and dysphagia [38]. A retrospective study of 20 patients found that cough, sputum production, dyspnea, and fever in patients with a history of recurrent pneumonia that did not respond to antimicrobial therapy were the most common manifestations [39]. A high index of suspicion is required to make the diagnosis. Chest CT findings are micro-nodules and opacities “tree-in-bud”. The management of these patients should focus on the prevention of recurrent aspiration and treatment of GERD.

Diffuse panbronchiolitis

Diffuse panbronchiolitis (DPB) is a distinct clinicopathologic syndrome that primarily affects Japanese, Korean, Chinese and Thai middle age men and rarely reported outside South-East Asia. It affects the lower and upper respiratory tract, leading to progressive evolution bronchiectasis, recurrent infections and generally sinusitis [9]. Histological injury DPB is focused on the respiratory bronchioles, and consists of a transmural infiltrate of lymphocytes, plasma cells and distinctive lipid-laden “foamy” macrophages, with extension to peribronchiolar tissue. In advanced stages, there’s obstruction and constriction of lumen, proliferation of lymphoid follicles and secondary ectasia of terminal bronchioles. The functional pattern is essentially obstructive.

The etiology is unknown, although it appears to be a genetic predisposition. For example, the HLA-B54 is associated with PBD in Japanese while HLA-A11 in Korean [40]. It has been suggested that in any position between HLA-B and HLA-A, a mutation of a suspicious gene susceptible to the disease in a single ancestral chromosome containing HLA-B54 and HLA-A11 occurred. Then it is possible that a series of genetic recombination events around the locus (location on a chromosome) of the disease could have resulted in the association of the disease with HLA-B54 in Japanese and HLA-A11 in Korean. Additionally, the diseases caused by HLA genes identified in the region of susceptibility to PBD have been investigated. One of these, “bare lymphocyte syndrome I” (BLS I) has a number of similarities with PBD, including chronic sinusitis, inflammation and bronchiolar nodules, and H. influenzae and Pseudomonas aeruginosa in those affected. As the PBD, the BLS I respond to treatment with erythromycin, showing resolution of symptoms. The similarity between the two entities, the result to the same type of treatment, and the fact that the gene responsible for BLS I was causing the PBD, narrows the field for a responsible gene within the PBD area [41]. The radiological image consists of small nodular shadows to 2 mm in diameter, diffusely scattered in both lungs that can be invisible by hyperinflation of alveoli and the sign of “tree-in-bud” [42]. Effective treatment is based on macrolides such as erythromycin and roxithromycin and for extended time and at low doses, mainly for its immunomodulatory effect [43]. While it is true that the histological diagnosis is desirable, with the appropriate facilities, compatible physiological and radiological changes and the geographical location, lung biopsy is usually unnecessary and a course of macrolides is adequate. The prognosis is good. The survival rate at 10 years of PBD is about 90%. The disease has no known cure.

Obliterants bronchiolitis

Also, known as obliterative bronchiolitis (OB), it is a syndrome associated with small airways damage caused by a spectrum of exposure
to drugs, inhalation, or infection and also to transplantation of the lung and hematopoietic cells (HSCT) [16]. The clinical syndrome is typically characterized by dyspnea, reverse-and-flow airway limitation with bronchodilators, and normal chest x-ray or pulmonary hyperinflation. The pathogenesis is not clear. Injury to the bronchial epithelium seems to initiate the process that may involve the alveoli adjacent to the small pathways. As in many other inflammatory processes in the human economy, repair may result in complete recovery or excessive proliferation of granulation tissue that causes obstruction or obliteration of the bronchiolar lumen (constrictive bronchiolitis) [44].

There is a wide range of agents and diseases that can cause the obliterating event. Inhalation of nitrogen dioxide, nitrogen mustard, ammonia, welding vapors, ash, food flavoring fumes (diacetyl and 2,3-pentanedione), SRV infection, adenovirus, or Mycoplasma pneumoniae, busulfan, gold or penicillamine treatment, and exposure to combustion air wells in the desert in soldiers, have been described as causes [45,46]. Rheumatoid arthritis and other rheumatic diseases can lead to OB [7]. In addition, OB is a respiratory manifestation of graft vs host disease (GVHD) in patients undergoing lung transplantation or HSCT. It occurs in 6% of allogeneic HSCT and in this group of patient’s survival is only 13% to 5 years [47]. Club cells (originally called Clara cells) promote the regeneration of epithelial airway and may be reduced in number or eliminated as a result of epithelial injury. Certain polymorphisms of genes of the immune system favor this phenomenon which represents a rejection of the allo-graft towards the host and is an expression of GVHD [2]. The condition typically develops within the first 2 years of transplantation, although it can occur several years later.

In patients with lung transplantation there is an added complication of microvascular insufficiency in the small airways of the transplanted lung, which presumably occurs because blood supply to the bronchial arteries is interrupted during transplantation. This could lead to defective repair if subsequent immune or non-immune injury occurs [48,49]. Compared with normal lungs, transplanted lungs are more susceptible to allo-immune immunological insult and airway insult, and they have limited resilience. It is therefore not surprising that OB affects longer-term lung transplant survivors, with the likelihood of remaining free from disease (OB) less than 30% at 10-year term [50,51].

Common immunosuppressive regimens have also been implicated in the pathogenesis of OB. They include calcineurin inhibitors (cyclosporin, tacrolimus and everolimus), inhibitors of purine synthesis (azathioprine or mycophenolate mofetil) and glucocorticoids [52-54]. Treatments for OB after HSCT or lung transplant have several controversial aspects. First, they are based on small and retrospective series. Second, the studies evaluated responses to treatments in which the primary focus was GVHD and not specifically OB. Third, studies are difficult to interpret because the severity of OB varied widely among the patients studied. Possibly the OB has different clinical phenotypes, which condition different responses to therapy. Azithromycin has improved lung function in 50% of patients transplanted with OB and also survival [55,56]. The mechanism of action appears to be a decrease in airway neutrophilia and related cytokine activation [57]. FAM (fluticasone, azithromycin and montelukast) have been used for OB post-transplantation. Lung transplantation in end stage OB in HSCT or in a first lung transplant is an accepted therapy in carefully selected patients [2]. As with other types of bronchiolitis, OB can be seen in inflammatory bowel disease. Pemphigus vulgaris is a life-threatening autoimmune disease of the skin and mucous membranes caused by autoimmune IgG against desmoglein 3 (Dsg3), a cell-to-cell adhesion molecule of keratinocytes [58]. Rarely, OB has been reported associated with paraneoplastic pemphigus, sometimes occurring prior to the appearance of the underlying neoplasia. On rare occasions, OB is idiopathic or cryptogenic [59]. OB should be considered in patients presenting with insidious onset of dyspnea and cough, especially when the symptoms and signs do not follow a typical pattern of asthma or COPD or if the patient has a predisposing exposure or condition; for example, symptoms of viral infection, exposure to toxic vapors, history of organ transplantation, or concomitant connective tissue disease. OB must enter the differential diagnosis of irreversible obstruction to airflow or associated with reduction in the transfer of gases in nonsmokers. Pulmonary function tests may initially be normal or show an obstructive pattern with gas trapping and no reversibility with inhaled bronchodilators [46]. Given the vast number of bronchioles, of which a large number will be narrowed or clogged, a significant loss of function occurs, which explains the occasional restrictive disorder and the reduction of DLCO [60].

In many cases of OB, conventional radiography is normal or shows only thickening of the bronchial wall or hyperinflation (without flattening of diaphragms or hyperlucent areas suggestive of emphysema). HRCT is the technique that best identifies the findings consistent with OB, particularly if images are obtained in inspiration and expiration. Trapping of expiratory gas (diffuse or mosaic), thickening of bronchial walls (linear opacities that bifurcate in “Y” or “V”), centrilobular nodules and opacities in “ground glass” are the most frequent patterns [11]. Except for a mosaic pattern of attenuation, which is highly suggestive of OB, it is often difficult to distinguish between severe asthma and bronchiolitis. Bronchoscopy and BAL are non-specific and only help to rule out other processes such as anatomic stenosis or endobronchial tumor, sarcoidosis, or hypersensitivity pneumonitis. Transbronchial biopsy offers conflicting results in post-transplantation OB. Definitive diagnosis usually requires a thoracoscopic or open lung biopsy. The findings are basically: infiltration of Τ lymphocytes around the small airways, bronchiolar smooth muscle hypertrophy, and bronchiectasis with mucus stasis, distortion, fibrosis, and total scarring of the bronchioles [16].

**Bronchiolitis and asthma**

Severe early-onset bronchiolitis is associated with an increased risk of asthma, especially after rhinovirus or RSV bronchiolitis, and an increased risk of asthma may persist into early adulthood [61,62]. An undefined aspect is up to which point early bronchiolitis in infancy damages normal lung development and predisposes to subsequent bronchospasm or even where certain infants have a preexisting aberration of immune response or airway function that predisposes them to severe bronchiolitis and recurrent bronchospasm [63]. Some work suggests the possibility that premorbid lung function may be abnormal among infants who have bronchiolitis in the first year of life [64]. A genetic predisposition to severe early bronchiolitis in life and the subsequent development of asthma is suggested by reports of associations between polymorphism in genes involved in the innate immune response and allergic responses, surfactant proteins and inflammatory cytokines [65]. From another perspective, Stein et al, presented recent evidence suggesting that the difference in the incidence of asthma between Old Order Amish versus the genetically related Hutterites may be due to variations in lifestyle that generates differences in the environmental microbiome (specifically domestic dust) [66]. Recent data reveal that the intestine-lung axis, particularly fecal microbiotic, may be a preventive factor against the development of viral respiratory tract infection [67]. It is conceivable that the asthma associated with viral respiratory tract disease could be
prevented by alteration of the lung-intestine microbiotic. In the study by Ege et al., the protective effect against asthma in children exposed to a farm environment could be due to the influence of microbiotic on helper T cells type 1 and 2 [68]. All this indicates that there is a complex interaction between genetic and environmental factors in the development of asthma.

Differential diagnosis

In adults, asthma differs from bronchiolitis due to the presence of an obstruction to reversible airflow in spirometry. However, severe asthma may be associated with an irreversible limitation of airflow. A reduction in DLCO is uncommon in asthma, but common in bronchiolitis. The perfusion mosaic is very rare in asthma, but it is found in at least 50% of patients with bronchiolitis obliterans. Some patients with severe asthma have bronchiolitis obliterans with hypereosinophilic syndrome. These patients have poor response to inhaled steroids, peripheral eosinophilia (> 1000 cells / μl serum), more than 25% of eosinophils in BAL. In HRCT, there are more changes in bronchiolitis than in asthma, and lung biopsy shows prominent infiltration of bronchiolar wall with eosinophils. Some authors classify this entity as hypereosinophilic bronchiolitis obliterans [69]. Sarcoïdosis may have a similar presentation in terms of cough and dyspnoea, and occasionally airway sarcoid is associated with airflow limitation. More frequently, sarcoïdosis is associated with a restrictive pattern and decreased DLCO and unusually lung function may be normal. Lung biopsy will show well-formed non-caseous granulomas that make the diagnosis of sarcoïdosis more likely.

Bronchocentric granulomatosis predominantly affects the bronchi but peribronchial inflammation may be present. The presence of fungal hyphae strongly supports the diagnosis.

Diffuse idiopathic pulmonary neuroendocrine cell hyperplasia (DIPNECH) is a neuroendocrine proliferation associated with a carcinoid tumour in which neuroendocrine cells are confined to the bronchial and bronchiolar epithelium. It is seen predominantly in women with cough and dyspnoea, airflow limitation in spirometry and multiple small nodules (4-10 mm) in HRCT [70,71]. In one series, 44% of patients with DIPNECH had constrictive bronchiolitis [70]. Serum levels of chromogranin A may be elevated. Diagnosis requires lung transbronchial or surgical biopsy. The pulmonary neuroendocrine cells (PNE-C) are specialized epithelial cells, located through the respiratory tract from the trachea to terminal airways, and synthesize, store and release various amines and peptides. Among them, serotonin, calcitonin, chromogranin A (CgA) and gastrin-releasing peptide (GRP) [72]. DIPNECH is a form of preinvasive lesion that is located within the spectrum of neuroendocrine cell neoplasms and is associated with OB [73].

Conclusions

1) Bronchiolitis is a better-known event in the pediatric population than in the adult population.

2) In the adult population in many cases represents a pathophysiological response to various types of injury, monomorphic and nonspecific and only in some cases a well-established nosological entity.

3) In most patient’s tissue is required to make the diagnosis and classification.

4) Even with histology, an adequate clinical, functional, radiological and histological correlation is required, since the findings may be common to several etiologies.

5) Treatment depends on the underlying cause.

6) More primary and clinical research is required to better delineate groups, pathophysiology, and to define diagnostic and prognostic markers.

7) This information will be basic to establishing prospective treatment protocols.

Acknowledgement

This work was carried out in collaboration between both authors. Authors AA and IA contributed equally in the planning, data collection, data analysis, writing and critical review. Both authors read and approved the final manuscript.

References


activation of interferon signaling networks. Pediatr Infect Dis 32: e68-e76. [Crossref]