Allergic Fungal Rhinosinusitis

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ABSTRACT

Allergic fungal rhinosinusitis (AFRS) has always remained a topic of discussion at all rhinology meets. Despite so much of literature available, the nature of this disease, its diagnosis, pathogenesis, classification and appropriate management continue to generate debate and controversy even after three decades of research and investigation. AFRS is an endemic disease in North and South India. In spite of this, there has been no optimal management protocol for this disease being followed in India yet. To overcome this, a national panel was conducted on AFRS at the ENT Surgical Update 2011, held at Postgraduate Institute of Medical Education and Research, Chandigarh with experts from all over the country so that a consensus can be achieved regarding the workup and management of AFRS.

Keywords: Rhinosinusitis, Allergic fungal sinusitis, Management protocol.


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INTRODUCTION

Allergic fungal rhinosinusitis (AFRS) was recognized for the first time in 1976 when in some patients thick, dark, inspissated mucus was noticed filling the paranasal sinuses, similar both grossly and microscopically to that seen in the bronchial passages of patients with allergic bronchopulmonary aspergillosis (ABPA).1-3 It is invariably associated with nasal polyposis and the presence of allergic fungal mucin. Allergic fungal mucin is thick and tenacious macroscopically having a brown or greenish black color. This mucus consists of onion-skin laminations of necrotic eosinophils in various stages of degeneration, occasional small hexagonal crystals of lysophospholipase (Charcot–Leyden crystals) and scanty fungal hyphae on histology. It is a nontissue invasive fungal process, representing an allergic/hypersensitivity response to the presence of extramucosal fungi within the sinus cavity in which a strong IgE-mediated hypersensitivity to fungal elements drives the inflammatory process. Aspergillus and the dematiaceous fungi are most commonly found in AFRS mucus.4-6 The preferred terminology for this condition is now ‘allergic fungal rhinosinusitis’ or AFRS though once it was called ‘allergic’ Aspergillus sinusitis. There are a lot of variations seen in patients with similar clinical presentation as AFRS. It has been noted that in some cases, the eosinophilic mucin from the sinuses does not have identifiable fungal elements.6,7 Ferguson described an AFRS-like condition with slightly different clinical features and proposed the term ‘eosinophilic mucin rhinosinusitis’ (EMRS) to describe cases in which fungus was not identified histologically.8

On the other hand some patients with clinical features of AFRS may have demonstrable fungus within their eosinophilic mucin, but no allergy.9 There have been cases isolated with no allergy; no fungi in mucin but still have eosinophilic mucin. Ponikau et al postulated that most, if not all, CRS was a hypersensitivity response to fungi and that fungi could be universally cultured from nasal secretions also further clouded the distinction between AFRS and AFRS-like CRS.10 The nature of this disease, its diagnosis, pathogenesis, classification and appropriate management continue to generate debate and controversy even after three decades of research and investigation.

PATHOGENESIS

AFRS was described as distinct clinical entity by Millar in 19813 and Katzenstein2 in 1983 characterized by atopy, sinonasal polyposis and the presence of allergic mucin. It was thought that the fungus in the sinuses would have potential for tissue invasion so extensive surgical debridement followed by the use of systemic antifungal agents was the treatment of choice those days. The realization that AFRS may represent an immunological response to presentation of a fungal antigen within a susceptible host leads to the discontinuation of such treatment. As in ABPA, an atopic host is exposed to fungi through normal nasal respiration, thus providing an initial antigenic stimulus. Gel and Coombs type I (IgE) and III (immune complex) mediated reactions trigger an intense eosinophilic inflammatory response. Any anatomic obstruction, such as septal deviation or turbinate hypertrophy, accentuates the already inflamed mucosa which in turn leads to obstruction of sinus ostia resulting in stasis of mucus within the sinuses. This, in turn, creates an ideal environment for further proliferation of the fungus, thus increasing the antigenic exposure. This sets up a vicious cycle and produces a lot of allergic mucin.

This immunological aspect in the pathogenesis was supported by studies by Manning and Holman.11 In their
first study on eight patients, culture-positive bipolaris AFRS were prospectively compared with 10 controls with chronic rhinosinusitis. All eight patients with AFRS had positive skin test reactions to bipolaris antigen, as well as significant levels of bipolaris specific IgE and IgG by in vitro testing. Eight of the 10 control demonstrated negative results to both skin and serologic testing, suggesting that the presence of allergy to fungus as being important in the pathogenesis of AFRS.

In another study by them they showed that serum IgE and IgG levels to multiple fungi was found that in patients with AFRS and AFRS-like group were prospectively compared with 10 controls with chronic rhinosinusitis. Immuno-histochemical analysis for eosinophilic mediators (major basic protein and eosinophil derived neurotoxin) and a neutrophil-derived mediator (neutrophil elastase) was performed to compare the underlying inflammatory processes within each cohort. Inflammatory mediators derived from eosinophils predominated over neutrophil-derived mediators in the AFRS group, whereas significant differences were not present within the control group. The relative predominance of eosinophil-derived inflammatory mediators as compared to neutrophil-derived inflammatory mediators further support the association between non-infectious (i.e. immunologically mediated) inflammation and AFRS, and helps to differentiate this disease from other forms of chronic rhinosinusitis. This is further supported by the study by Stewart and Hunsaker in which they analyzed fungal-specific serum IgE and IgG levels in nonatopic controls, allergic rhinitis patients, non-AFRS polyp patients, AFRS-like patients and AFRS patients. It was found that in patients with AFRS and AFRS-like group there was elevated serum levels of IgE and IgG to multiple fungi.

The concept of eosinophilic activation associated with AFRS was further supported by Feger et al who studied eosinophilic cationic protein (ECP) levels in the serum and mucin of patients with AFRS. No differences in serum ECP were detected between patients with AFRS and controls, but ECP levels were significantly higher in mucin from patients with AFRS as compared to controls.

These studies by Manning et al and Feger et al while supporting that AFRS represents an immunologically-mediated disorder rather than an early stage of invasive fungal disease fuelled further speculation regarding the pathophysiology of AFRS.

In 1999 a ‘unifying hypothesis’ of CRS was proposed by Ponikau et al. They used a specially designed culture technique, and found that 93% of 101 consecutive patients with CRS demonstrated the presence of fungi obtained from nasal lavage and surgically obtained specimen. In contrast to prior AFRS studies, conventional IgE-mediated allergy to fungi was not consistently observed. It was therefore proposed that virtually all cases of chronic rhinosinusitis were associated with sensitization to colonizing fungi. It was further suggested that the term ‘allergic fungal rhinosinusitis’ be replaced with ‘eosinophilic fungal rhinosinusitis’ (EFRS). These findings have led to their belief that IgE-mediated inflammation is not crucial to the development of AFRS, and that eosinophilic chemotaxis and activation may result from a T lymphocyte-mediated inflammatory cascade. One potential problem with the common etiology that was proposed by the authors was the fact that fungi were also observed in 100% of normal control subjects.

In a study on humoral immune response in patients with EMCRS including AFRS Pant et al found that patients with AFRS had increases in fungal-specific IgE and total IgE but these were no different from a control group with allergic rhinitis. Though there was a poor correlation between fungal species present in the eosinophilic mucin of AFRS patients and the specific fungal allergy (42%) but elevated fungal-specific IgG3 was a distinguishing serologic feature that separated EMCRS and AFRS patients from those with fungal allergic rhinitis and other forms of CRS. Moreover, serum IgE levels could be used to distinguish EMCRS from AFRS. Another clinically important distinguishing feature of AFRS is type I hypersensitivity. Therefore, type 1 hypersensitivity to fungal antigens, as assessed by specific allergy tests, helps to distinguish AFRS from other forms of EMCRS and has implications for treatment.

Recently in 2009 Luong et al found that peripheral blood mononuclear cells from AFRS patients are stimulated by fungal antigens to secrete TH2-type cytokines.

In spite of all these studies supporting humoral immune factors, the underlying pathophysiology in AFRS remains steeped in controversy. Although it appears clear that the eosinophil plays an important role in the development of both AFRS and some forms of chronic rhinosinusitis, the factors that ultimately trigger eosinophilic inflammation remain in question.

In summary, it can be said that initiation of the inflammatory reaction leading to AFRS is a multifactorial event, governed by IgE-mediated sensitivity (atopy), humoral expression, exposure to specific fungi and aberration of local mucosal defence mechanisms.

Epidemiology and Clinical Presentation

It is estimated that approximately 5 to 10% of those patients with chronic rhinosinusitis actually carry a diagnosis of AFRS. It is most common among adolescents and young adults (mean age; 21.9 year). The incidence of AFRS...
appears to be impacted by geographic factors. On review of literature it was found that the majority of sites reported cases of AFRS to be located in temperate regions with relatively high humidity. In United States most cases are found in the southern central region of the country along the Mississippi basin. In India, fungal sinusitis is maximally reported from North India and South India. Initially, *Aspergillus* was believed to be the causative organism in AFRS, but dematiaceous fungi were most commonly found in AFRS mucus, in different studies conducted in the United States. In other parts of the world, *Aspergillus* is still found to be a common isolate in cases of AFRS and nonallergic eosinophilic fungal sinusitis. Identification of the specific fungal organism is important only for making a diagnosis, it does not have any prognostic value nor does it make any difference in the planning of the treatment protocol.

AFRS occurs in adolescents and young adults who often have asthma that is exacerbated by their sinusitis. All patients are immunocompetent and have a strong history of atopy. All have nasal polyps and chronic sinusitis. There is no increased aspirin sensitivity despite the association with asthma and nasal polyps.

The development of nasal airway obstruction is gradual and the patient is unaware of its presence and presents only when there is complete nasal obstruction or may develop more serious symptoms like headaches and visual disturbances. Facial dysmorphism if present is also often so slow that its identification escapes notice of the patient and family members. The slow accumulation of allergic fungal mucin imparts unique and predictable characteristics to the disease. Allergic fungal mucin is sequestered within involve paranasal sinus cavities. As its quantity increases, the involved paranasal sinus begins to resemble and behave in a way consistent with a mucocele (sometimes referred to as a fungal mucocele). With time, bony remodeling and decalcification may occur, causing the disease to mimic ‘invasion’ into adjacent anatomic spaces. The location of bone destruction seems to be determined simply by the location of the disease, and this destruction often gives rise to exophthalmos, facial dysmorphism or intracranial extension without tissue invasion.

In cases of allergic fungal sinusitis, the mechanism of vision loss has thus, far been assumed to be secondary to direct compression or direct inflammation of the optic nerve. Gupta et al hypothesized that there may be aberrant anatomical pathways present in the region of the optic canal could have been responsible for direct inflammation of the nerve in the absence of obvious bony erosion apart from mechanical compression of the optic nerve, a local immunological reaction to fungal antigens might be responsible for the visual loss seen in allergic fungal sinusitis.

A patient with AFRS classically is a young (mean age is 22 years), immunocompetent patient with unilateral or asymmetric involvement of the paranasal sinuses, a history of atopy, nasal casts and polyposis, and a lack of significant pain. Nasal casts are green to black rubbery formed elements made of allergic mucin. The presentation may be dramatic, with a significant number of patients presenting with proptosis, telecanthus or gross facial dysmoria. The diagnosis for AFRS is usually derived from the clinical, radiological, microbiological and histopathological information combined together. There has been a lot of debate regarding the diagnostic criteria for AFRS.

A number of different attempts at establishment of diagnostic criteria have been proposed, most of which focus upon some combination of the radiologic, immunologic, clinical and histologic manifestations of the disease. Allphin et al felt that the combination of opacified paranasal sinuses and radiography, characteristic histologic findings of allergic mucin, and laboratory evidence of allergy was sufficient to differentiate AFRS from other forms of rhinosinusitis. Loury and Schaefer proposed a more specific set of diagnostic criteria, which included eosinophilia, immediate skin reactivity or serum IgE antibodies to fungal antigen, elevated total IgE level, nasal mucosal edema or polyposis, histopathologic findings of allergic mucin containing noninvasive fungal hyphae, and characteristic radiological findings.

In 1994, Bent and Kuhn published their diagnostic criteria centered on the histologic, radiographic and immunologic characteristics of the disease and they still remain the standard and most widely accepted worldwide.

According to Bent and Kuhn patients must meet all the major criteria for diagnosis, while the minor criteria support the diagnosis. The major criteria include a history of type I hypersensitivity by history, skin testing or *in vitro* testing; nasal polyposis; characteristic computed tomography (CT) scan findings; the presence of eosinophilic mucin without invasion and a positive fungal stain of sinus contents removed at the time of surgery. The minor criteria include a history of asthma, unilateral predominance of disease, radiographic evidence of bone erosion, fungal cultures, presence of Charcot-Leyden crystals in surgical specimens and serum eosinophilia (Table 1).

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<th>Major</th>
<th>Minor</th>
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<td>Type I hypersensitivity</td>
<td>Asthma</td>
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<tr>
<td>Nasal polyposis</td>
<td>Unilateral disease</td>
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<tr>
<td>Characteristic CT findings</td>
<td>Bone erosion</td>
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<tr>
<td>Eosinophilic mucin without invasion</td>
<td>Fungal cultures</td>
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<tr>
<td>Positive fungal stain</td>
<td>Charcot-Leyden crystals</td>
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<td>Serum eosinophilia</td>
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Table 1: Bent and Kuhn’s diagnostic criteria
Histology of Allergic Fungal Mucin

Microscopic picture on hematoxylin and eosin (H&E) staining of mucin is an important diagnostic tool and would show typical inflammatory infiltrate composed of eosinophils, eosinophilic breakdown products or Charcot-Leyden crystals, lymphocytes and plasma cells. The mucosa will be hypertrophic and hyperplastic but should not have evidence of necrosis, giant cells, granulomas or invasion into surrounding structures. It is important to note that examination of the unique allergic fungal mucin itself, and not the surrounding mucosa, is the most reliable indicator of disease. Grossly, this thick, highly viscous, variably colored mucin has been described as being similar to peanut butter or axle grease. H&E staining accentuates the mucin and cellular components of allergic fungal mucin. Using this stain, background mucin will often take on a chondroid appearance, whereas eosinophils and Charcot-Leyden crystals are heavily stained and become easily detected. Fungi fail to stain with H&E, however, and may be implicated only by their resulting negative image against an otherwise stained background. Given that fungal hyphae are frequently rare, scattered, and fragmented within allergic mucin, identification is extremely difficult unless specific histological stains are used. Fungal elements are recognized for their unique ability to absorb silver. This property is the basis for various silver stains, such as Grocott’s or Gomori’s methenamine silver (GMS) stain, which turn fungi black or dark brown. The use of a fungal stain complements the findings of initial H&E stain and is extremely important in the identification of fungi.

Fungi commonly identified in the electron microscopy are from the dematiaceous family and Aspergillus species. Dematiaceous fungi include the genera Alternaria, Bipolaris, Cladosporium, Curvularia, Drechslera and Helminthosporium. These fungi are darkly pigmented due to dihydroxynaphthalene melanin in the cell walls of the hyphae or conidia.

Fungal hyphae do not invade tissue: The presence of fungal tissue invasion has been considered incompatible with a diagnosis if AFRS. Many patients with polypoïd CRS and eosinophilic mucin lack other important clinical characteristics of AFRS: Demonstrable fungi and fungal allergy. These patients should not be classified as having AFRS. In a study Dhiwakar et al pointed out that the combination of nasal polyps, CT scan hyperattenuation and elevated titers of anti-Aspergillus IgE have high predictive value for AFRS, though considered in isolation they are not specific. There is a lot of overlap that exists between AFRS, EMCRS and CRS from other causes, but Bent and Kuhn criteria can still distinguish between these.

Immunologic Findings

The original reports describing AFS focused on the fungus Aspergillus. Miller et al described immediate cutaneous reactivity to Aspergillus fumigatus antigen in all five patients in their original case series. Katzenstein et al found specific IgE and immunoglobulin G (IgG) to Aspergillus in their series. The total IgE level has served as a useful tool to follow the clinical activity of allergic bronchopulmonary aspergillosis. Based on similar IgE behavior associated with recurrence of AFRS, total IgE levels have been proposed as a useful indicator of AFRS clinical activity. Total IgE values are generally elevated in AFRS, often to more than 1,000 U/ml elevated in all patients with AFRS and corresponded with the results of fungal cultures. In contrast, levels of fungal-specific IgE were not elevated within the control group of patients with chronic rhinosinusitis. Moreover, patients with AFRS appear to demonstrate a broad sensitivity to a number of fungal and nonfungal antigens. Mabry et al have reported their experience, which indicates that patients with AFRS are allergic to multiple fungal antigens, as well as many typical nonfungal antigen. Patients with AFRS generally demonstrate positive skin test and in vitro (RAST) responses for both to fungal and nonfungal antigens. Manning et al first demonstrated the sensitivity of RAST, who compared 16 patients with histologically confirmed AFRS with a control group with chronic rhinosinusitis. Levels of fungal-specific IgE were uniformly elevated in patients with AFRS and corresponded with the results of fungal cultures.

Morpeth et al in their review on AFS literature noted the following immunologic findings: 48% with positive skin tests to fungi, 44% with elevated total IgE, 40% with peripheral eosinophilia, 33% with elevated specific IgG, and 28% with serum precipitins to fungal antigens. Schubert and Goetz found that 67% of their patients had an elevated total IgE (greater than 199 IU/ml). The mean total IgE for their patients was 668 IU/ml. All patients have positive skin tests for aeroallergens with specific IgE to their presumably causal mold and two-thirds of patients having elevated specific IgG to molds.

Non-specific inflammatory findings in the surgical debris removed from AFS patients include elevated levels of eosinophilic cationic protein and major basic protein. This indicates eosinophilic degranulation.

Ponikau et al postulated that there is a non-specific eosinophilic response in these patients to the presence of fungal elements in the nose and sinuses. We need further studies to confirm these findings, because if the inflammatory process in AFS is actually driven mostly by a non-specific eosinophilic reaction rather than by specific IgE, then steroids will continue to be the primary medical therapy.
and immunotherapy with fungal antigen extracts will have no role. If the etiology of AFS is a specific IgE-mediated immunologic response then appropriate immunotherapy may be initiated and would be efficacious.

**Fungal Culture**

Fungal cultures provide only supportive evidence helpful in the diagnosis and subsequent treatment of AFRS, but must be interpreted with caution and the diagnosis of AFRS is not established or eliminated based on the results of these cultures as yield is variable from 64 to 100%. A diagnosis of AFRS in the presence of a negative fungal culture is possible but there may be a situation when a positive fungal culture fails to confirm the diagnosis of AFRS, because it may merely represent the presence of saprophytic fungal growth.

A panel of international experts in 2004 convened to establish working diagnosis for chronic rhinosinusitis which included AFRS. The impact of IgE-mediated sensitivity to fungus in AFRS was acknowledged by the panel and they proposed diagnostic criteria based upon the combination of histologic, immunologic, clinical and radiologic factors (Table 2).

**RADIOLOGICAL FEATURES**

AFRS has characteristic features on CT scan or MRI considered extremely important for diagnosis. CT scan shows multiple opacified sinuses with central hyperattenuation, sinus mucocele formation and erosion bone. Ghegan et al showed that 56% of AFRS cases presented with radiographic evidence of skull base erosion or intraorbital extension, whereas similar findings were only noticed in 5% of other cases of chronic sinusitis. Zinreich et al have described the CT and MRI findings in patients with AFRS. There is characteristic serpiginous attenuation in CT scan particularly in bone window. Ferromagnetic elements produced by fungi are believed to be the cause of this heterogeneous destruction. The combination of hyperattenuation and bony erosion on CT scan and type I hypersensitivity may be considered as preoperative predictors of AFRS (Fig. 1).

MRI is complimentary to CT scan and done in cases with intraorbital or intracranial extension of disease on CT scan. On MRI, there is a central low signal on T1 and T2 imaging by sinuses that correspond with areas of eosinophilic mucin (Fig. 2). Peripheral high-signal intensity

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**Table 2: Diagnostic criteria for AFRS (2004)**

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<th>Symptoms</th>
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<td>Requires ≥ one of the following:</td>
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<td>• Anterior and/or posterior nasal drainage</td>
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<td>• Nasal obstruction</td>
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<td>• Decreased sense of smell</td>
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<td>• Facial pain-pressure-fullness</td>
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<tr>
<th>Objective findings</th>
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<tr>
<td>Requires all of the following:</td>
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<tr>
<td>• Presence of allergic mucin (fungal hyphae with degranulating eosinophils on histopathology)</td>
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<tr>
<td>• Evidence of fungal-specific IgE</td>
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<tr>
<td>• No histologic evidence of invasive fungal disease</td>
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<table>
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<tr>
<th>Radiologic findings</th>
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<tr>
<td>Highly recommended:</td>
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<tr>
<td>• Sinus CT demonstrating</td>
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<tr>
<td>• Bone erosion</td>
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<tr>
<td>• Sinus expansion</td>
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<tr>
<td>• Extension of disease into adjacent anatomic areas</td>
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<tr>
<th>Other diagnostic measures</th>
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<tr>
<td>Possible, but not required:</td>
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<tr>
<td>• Fungal culture</td>
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<tr>
<td>• Total serum IgE</td>
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<td>• Imaging by more than one technique</td>
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**Fig. 1:** CT scans showing characteristic hyperattenuation seen in AFRS
corresponds with inflamed mucosa.\textsuperscript{37-39} MRI demonstrates hypointense regions surrounded by mucosal inflammation in T2 weighted images. Manning et al in a series of 10 cases of AFRS, demonstrated that hypointense central T1 signal, central T2 signal void, and the presence of increased peripheral T1/T2 enhancement was highly specific for AFRS as compared with other forms of fungal sinusitis (invasive fungal sinusitis and fungal ball) and mucocele.\textsuperscript{38}

CT findings of sinus expansion with central areas of irregular high attenuation should prompt suspicion for AFRS. Bony erosion noted is likely due to pressure resorption but may be due to the effect of inflammatory mediators. Although the CT scan and MRI findings in AFRS are considered important in diagnosis, definitive diagnosis requires histologic verification and other clinical information (Figs 3 and 4).

**Treatment**

With the better understanding and recognition of the disease process, the treatment of AFRS has also evolved from aggressive radical surgery and toxic antifungal medications to conservative endoscopic surgery and adjunctive medical therapy directed at suppressing inflammation and reducing the burden of fungal antigen in the nose. The recognition of AFRS now, to be a noninvasive, immunologically mediated hypersensitivity to fungal antigens has lead to the treatment options to be considered accordingly.

The treatment of AFRS is surgical extirpation of the allergic mucin and fungal muck and providing drainage and aeration of the sinuses. External radical approaches with removal of the sinus mucosa were done earlier\textsuperscript{40,41} but with the advent of nasal endoscopes, most cases are amenable to tissue preserving endoscopic approach.\textsuperscript{28} External surgeries are not necessary except in rare circumstances and obliterator procedures should be avoided. Nasal polyposis is inherent to AFRS and can range from subtle to extensive, distortion of local anatomy and loss of useful surgical landmarks which increases the risks of complications during surgery. Image guidance can be an important tool for...

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**Fig. 2:** T2 weighted MRI showing demonstrates hypointense regions surrounded by mucosal inflammation

**Fig. 3:** CT scan showing hetrodense shadow in sphenoid sinuses suggestive of AFRS

**Fig. 4:** CT scan showing hetrodense shadow in frontal sinus suggestive of AFRS
facilitating a complete surgery as missing diseased cells with eosinophilic mucin would decrease the effectiveness of medical treatment and increase the chances of recurrence.\(^{42}\)

In cases of recurrence, if the medical management fails to clear an exacerbation, eosinophilic mucin accumulation and polyposis, then surgical treatment is warranted which would need a more aggressive surgery and wide sinusotomies. Regardless of the completeness of surgical exenteration, recurrences are common. Hence, the need for medical adjuncts following surgery is mandatory.

Medical treatment for AFRS is essential to obtain long-term symptom control, retard polyp formation and delay or prevent revision surgery. Different forms of medical therapies ranging from immunotherapy to systemic steroids have been used for treating AFRS though randomized blinded clinical trials are lacking. Systemic anti-inflammatory agents are usually required in the treatment of AFRS and appear to be the most effective medical therapy. Systemic steroids are at least transiently effective for the treatment of polypoid rhinosinusitis as shown by a few placebo controlled, randomized trials\(^{43,44}\) so may be used as an adjunct after surgery. The side effect profile of systemic corticosteroids warrants careful consideration when they are used in a long-term approach to control AFRS. Therefore, as a general rule, systemic steroids are best confined to the perioperative period and for use in short bursts to suppress recurrent polyps and address acute exacerbations of disease. Topical intranasal steroids play an important role in the long-term medical management of AFRS. Topical delivery avoids or minimizes most of the acute and chronic long-term toxicities of corticosteroids, yet is successful in maintaining control of inflammation for prolonged periods. Topical steroids have been shown to be effective in the treatment of nasal polyp disease in AFRS\(^{45}\) and in higher doses than normal.\(^ {46}\) Budesonide has been tried in the form of atomized spray as it would deliver a larger total dose of steroid compared with conventional steroid nasal sprays.\(^ {47}\) Unfortunately, although topical intranasal steroids appear to be effective, they are often not sufficient to completely eliminate the use of systemic steroids.

The recognized toxicity of repeated courses of systemic steroids has led to a search for nonsteroid treatment alternatives. Macrolides\(^ {30}\) and leukotriene receptor antagonists or synthesis inhibitors have also been tried for polypoid CRS because of their safety and possible steroid-sparing effect though it lacks effective control.\(^ {48}\)

Systemic antifungal therapy for AFRS was initially proposed to control the theoretical potential for progression to invasive forms of fungal sinusitis. The early use of amphotericin B yielded to the use of less toxic agents, such as ketoconazole, itraconazole and fluconazole, but the poor \textit{in vivo} activity of these agents against dematiaceous fungi was soon discovered.\(^ {11}\) The role of antifungals had been controversial and one view is that these are too expensive and toxic for routine use, but some studies however, reported good result with the use of systemic itraconazole therapy.\(^ {49-51}\) None of these studies however, gave any evidence that antifungals decrease reliance on systemic steroids. Moreover, the efficacy of agents such as itraconazole may not be due to a reduced fungal burden in the nose, but rather due to the anti-inflammatory properties of the molecule or its inhibition of prednisone metabolism. Antifungal treatments are sometimes employed for AFRS with the aim of decreasing the fungal exposure within the sinonasal cavities.\(^ {46,52}\) Like systemic antifungals, there have been no trials of topical antifungals specifically for AFRS.

Randomized, controlled trials have failed to show a significant therapeutic benefit of topical antifungal (amphotericin) for the treatment of chronic polypoid rhinosinusitis.\(^ {53}\) Despite the purported fungal cause of AFRS, antifungal therapies need further investigation to establish their efficacy before their use is widely adopted.

Immunotherapy is another treatment modality that holds potential as an effective treatment option for AFRS. The anti-inflammatory effect of specific allergen immunotherapy has the potential to decrease reliance on systemic steroids in the treatment of AFRS or may reduce the need for revision surgery. Although the relative importance of type I hypersensitivity in AFRS continues to be debated, the rationale for immunotherapy is that AFRS is at least partially a result of allergen-specific IgE-mediated inflammation. The evidence to support the use of immunotherapy was incidental\(^ {54,55}\) In a nonrandomized not double blinded study on comparing AFRS patients treated with and without immunotherapy with an average 33 months of follow-up, Folker et al showed that the immunotherapy treated patients had better endoscopic mucosal appearance, lower CRS survey scores, required fewer courses of oral steroids (0 vs 2), and showed less reliance on nasal steroids (27 vs 73%). At present therapy for AFRS is directed toward reducing inflammation and reducing fungal antigen exposure.\(^5\) Significant uncertainty about the ideal treatment approach will persist as long as high-level evidence from randomized, controlled trials is lacking.

**Follow-up**

Follow-up of the patients can be done according to the following criteria:

a. \textbf{Visual analog scale:} Clinical symptoms like nasal obstruction, olfactory dysfunction, rhinorrhea, facial pain, sneezing, headache and visual symptoms are evaluated with a visual analog scale (VAS) ranging from 0 (no symptoms) to 10 (maximum symptoms). The severity of each symptoms collected from the results of
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Endoscopic physical findings: Objective assessment by rigid nasal endoscopy following surgery can be done by a standard endoscopic staging system described by Kupferberg et al:

- Stage 0: No evidence of disease
- Stage 1: Mucosal edema ± allergic mucin
- Stage 2: Polypoid mucosa ± allergic mucin
- Stage 3: Polyps with allergic mucin

Objective CT score- can be done based on Lund Mackay staging system that attributes points based on sinomucosal disease, opacification and obstruction:

- 0 point: No abnormality
- 1 point: Partial opacification
- 2 points: Total opacification

Points are calculated by adding for each sinus and on both sides.

Interpretation of Lund and Mackay Staging System

- 0 to 4: Normal
- 5 to 9: Minimal
- 10 to 14: Moderate
- 15 to 24: Severe

Recurrence of Disease

The potential for AFRS recidivism is well-recognized and ranges from 10 to 100% depending on the length of follow-up. To emphasize the importance of long-term surveillance, Bent et al pointed out that in their experience the often-dramatic initial response to surgical therapy was eventually replaced by recurrence of AFRS in the absence of ongoing therapy. Similarly Kupferberg et al followed the appearance of sinonasal mucosa of 24 patients treated with combined medical and surgical therapy for AFRS. Nineteen of 24 eventually developed recurrence of disease after discontinuation of systemic steroids, but they observed that endoscopic evidence of disease generally preceded return of subjective symptoms. AFRS recidivism appears to be influenced by long-term postoperative therapy.

Indian perspective: Consensus arrived by a national panel on AFRS at the ENT Surgical Update 2011, held at Postgraduate Institute of Medical Education and Research, Chandigarh.

Preoperative Workup and Preparation

Imaging: CT scan of nose, paranasal sinuses with orbits, axial, coronal and sagittal sections is manadatory in all patients to be taken up for surgery. Patients having intraorbital or intracranial extension of disease on CT scan to be subjected to a MRI. CT findings of sinus expansion with central areas of irregular high attenuation should prompt suspicion for AFS.

Workup for general anesthesia: All routine hematological and biochemical investigations necessary to obtain fitness for surgery to be done.

Preoperative steroids: Oral prednisolone in a dose of 0.5 mg/kg body weight to be started 2 weeks preoperatively. Oral deflazacort to be preferred in diabetic patients in a dose of 0.5 mg/kg.

Surgical Management

Consensus was reached that endoscopic sinus surgery to be preferred, reserving the open approach for rare circumstances. The role of surgery in patients of AFRS is to remove all the obstructing inspissated allergic mucin and to address the diseased mucosa establishing a permanent aeration of the sinuses. Early surgery warranted in all cases especially in all patients especially having intraorbital or intracranial extension of disease.

Postoperative Management

The experts were of the opinion that oral prednisolone or deflazacort in a dose of 0.5 mg/kg body weight should be used for 3 to 6 months. Topical steroidal sprays to be continued for 6 months and then as and when required. In the topical steroids, fluticasone furoate has a better compliance than mometasone or budesonide. The role of systemic antifungals in patients with intracranial and intraorbital extension of disease was debated. These are being used but because of lack of scientific evidence, all the panelists were of the opinion that further studies are required before any conclusion can be drawn. This protocol would be reviewed again after one year. Saline douches to be continued preferably lifelong. Further, it was deliberated that leukotriene receptor antagonists are not effective. Thirty percent of the experts were of the opinion that these are not effective and 30% said they are effective while the rest 40% had not used them. Role of immunotherapy in India has yet not been established.

Follow-up

Postoperative endoscopic suction and clearance to be done weekly for 1 month bimonthly for 3 months, once a month for 6 months and then 3 monthly for 5 to 6 years. It was further proposed to get postoperative CT scan after 6 weeks and then as and when required.

Recurrent Disease

The first line of treatment for recurrence is medical therapy in the form of oral and topical steroids. If the patient fails
medical therapy, revision endoscopic surgery to be done after adequate imaging. Patient to be labeled as disease-free, if there is no radiological evidence of disease on CT scan for 5 years.

DISCUSSION

Fungus is ubiquitous, present in all our surroundings and the air we inhale. Most healthy people do not react to the presence of fungus due to a functioning immune system. However, in rare instances, fungus may cause inflammation in the nose and the sinuses. Fungal sinusitis can manifest in various forms, differing in pathology, symptoms, course, severity and the treatment required.

A simplified classification of fungal sinusitis is as follows:

A. Noninvasive fungal sinusitis
   i. Fungus ball
   ii. Allergic fungal sinusitis
   iii. Nonallergic fungal sinusitis

B. Invasive fungal sinusitis
   i. Acute invasive fungal sinusitis
   ii. Chronic invasive fungal sinusitis
   iii. Granulomatous invasive fungal sinusitis.

We would be restricting ourselves to AFRS. AFRS is a distinct form of chronic polypoid rhinosinusitis characterized by accumulation of eosinophilic mucin with fungal hyphae in the sinuses, type 1 hypersensitivity to fungi, and a propensity for mucocele formation and bone erosion (Figs 5 and 6).

As mentioned earlier, approximately 5 to 10% of those patients with chronic rhinosinusitis carry a diagnosis of AFRS. It is most common among adolescents and young adults (mean age, 21.9 year). The incidence of AFRS appears to be impacted by geographic factors. In India, AFRS is reported from North and South India because of hot and humid conditions.

Patients with AFRS usually present with rhinosinusitis symptoms lasting months or years and they may not seek medical attention until complete nasal obstruction, headaches, visual disturbances or facial distortion develop. Proptosis or telecanthus are frequently seen at presentation, especially in younger patients. (Figs 7 to 9).
Gupta et al in a study compared AFRS in adults and children and found that clinical profile of both the groups was the same except for higher incidence of proptosis in pediatric cases (60% as compared to 20% in adults) and high chances of having coexistent asthma and rhinobronchial allergy in adults (50% compared to 1% in children). There was higher incidence of facial deformities in the children compared to adults in their study with proptosis in 60%, telecanthus in 40% and nasal pyramid widening in 18% children and 20, 9 and 0% in adults respectively. McClay has reported an incidence of 42% facial deformities in children compared to 10% in adults. Gupta et al found a higher incidence of facial deformities, proptosis, intraorbital/intracranial extension and a higher rate of recurrence in children, with an earlier presentation, therefore, suggesting a more aggressive nature of AFS in children than adults mandating an early diagnosis, proper management and regular follow-up in these cases.

On radiology the involvement of the sinuses has been found to be asymmetrical in 70% of the children compared to 30% in adults and similar findings have been found by other authors. The involvement of orbit was seen more commonly with the children (68%) compared to adults (33%) Similarly, the intracranial extension was seen more often in children than adults. It is in accordance with the findings by McClay et al. Overall bony erosion was seen in 88% of the children compared to 36% in adults. Various studies quote a range of 20 to 90% incidence of bone erosion. McClay reported an incidence of 25% bone erosion in children compared with 23% in adults.

The treatment of AFRS is surgical extirpation of the allergic mucin and fungal muck and providing drainage and aeration of the sinuses with endoscopic sinus surgery being preferred over external approaches (Figs 10 and 11).

Preoperative steroids have been proved beneficial in obtaining a better operative field as well as having a better disease control.
Kennedy et al conducted a prospective study to evaluate the effect of preoperative high dose systemic corticosteroids and radiographic and endoscopic appearance of AFRS. They reported a marked reduction of Lund-Mackay scores following high dose steroids. They advocated minimally invasive endoscopic sinus surgery following therapy with steroids. Furthermore, a short course of preoperative systemic corticosteroids will shrink polyps and decrease bleeding during surgery. Kinsella et al who reviewed a series of 15 patients who underwent variable postoperative medical therapy for AFRS found that of the seven patients who had no recurrence at 6 months follow-up, four had been administered oral steroid for 2 to 4 weeks postoperatively. None of the eight patients with recurrent disease had received oral steroid. Similar encouraging results were found by DeShazo and Swain. Both groups of researchers while recommending the use of oral steroids in the postoperative period based on anecdotal evidence highlighted the need for prospective, controlled studies. Schubert and Goetz further studied the role of systemic corticosteroids in the postoperative management AFRS, demonstrating a significant increase in the time to revision sinus surgery in those patients with AFRS who received prolonged courses of postoperative corticosteroids. Postoperative corticosteroid therapy in this study ranged from 2 to 12 months, with improved outcomes recorded among those patients who were placed on longer courses of therapy. Kupferberg et al in a review of 28 patients with AFRS stressed the role of systemic corticosteroids after endoscopic sinus surgery. Patients who received corticosteroids postoperatively had less recurrence compared to the patients who did not receive steroids. They recommended oral prednisolone at a dose of 40 mg/day for 4 days, 30 mg/day for 4 days and 20 mg/day for 1 month. The dose was then adjusted to the lowest possible dose at which the patient could be maintained at stage 0. Kuhn and Javer followed a protocol lasting about 7 months with prednisolone being administered in a tapering dose starting at 40 mg/day and maintained at 0.1 mg/kg day for the last 2 months of treatment. The long duration of therapy was essentially to maintain a recurrence free state. The use of concomitant high-dose inhaled steroid was also emphasized. However, long-term treatment with systemic corticosteroids increases the risk of both acute and long-term morbidity particularly in children. The effect of corticosteroids in growth and bone development are extremely important in the pediatric population. Studies have shown that long-term systemic steroid usage can cause developmental growth delay and bone demineralization. In a study by Chesney et al children with childhood glomerular disease receiving prednisone were 5.3% shorter and had 16.7% more demineralization than matched control subjects. Shegren et al experimentally proved that alternate day dosing lessened the effect of corticosteroids but still cause premature closure of epiphyseal plates. Linear growth may be inhibited by a little as 5 mg of prednisone per day or 30 mg every other day. Schubert et al reported no adverse effects among their series of 67 patients with AFRS treated for up to 1 year with systemic corticosteroids, but long-term follow-up for this form of therapy is lacking. In view of these side effects, the panel recommended steroids in the form of prednisolone to be started 2 weeks preoperatively and 3 to 6 months postoperatively in a dose of 0.5 mg/kg body weight rather than 1 mg/kg body weight. Topical intranasal steroids play an important role in the long-term medical management of AFRS. Topical delivery avoids or minimizes most of the acute and chronic
long-term toxicities of corticosteroids, yet is successful in maintaining control of inflammation for prolonged periods.

Macrolides and leukotriene receptor antagonists or synthesis inhibitors have also been tried for polypoid CRS because of their safety and possible steroid-sparing effect though it lacks effective control.

Systemic antifungal therapy for AFRS was initially proposed to control the theoretical potential for progression to invasive forms of fungal sinusitis. The early use of amphotericin B yielded to the use of less toxic agents, such as ketoconazole, itraconazole and fluconazole, but the poor in vivo activity of these agents against dematiaceous fungi was soon discovered. Systemic antifungals are too expensive and toxic for routine use, but some studies however, reported good result with the use of systemic itraconazole therapy probably because of their anti-inflammatory properties. Randomized, controlled trials have failed to show a significant therapeutic benefit of topical antifungal (amphotericin) for the treatment of chronic polypoid rhinosinusitis.

In accordance with the literature, the panel concluded that leukotriene receptor antagonists are not effective and antifungal irrigation has no role while systemic antifungals have any role except in repeated recurrences. The advantages and usefulness of saline douches was emphatically stressed by the panel and preferably to be continued lifelong.

Immunotherapy may be an effective treatment option for AFRS as it is at least partially a result of allergen-specific IgE-mediated inflammation. Although subcutaneous immunotherapy has clearly demonstrated efficacy in allergic rhinitis and asthma randomized, controlled trials that examine the efficacy of immunotherapy specifically for AFRS are lacking. There are chances of worsening of disease due to introduction of foreign extraneous fungal antigens and damage to tissues due to type III Gel and Coomb’s occasionally. Moreover, there is lack of substantial high-level evidence from randomized, controlled trials, a consensus was reached at the meeting that immunotherapy in the current form has no role in AFRS.

In cases of recurrence, if the medical management fails to clear an exacerbation, eosinophilic mucin accumulation and polyposis, then surgical treatment is warranted which would need a more aggressive surgery and wide sinusotomies (Figs 12A and B).

The panel categorically emphasized the importance of long-term surveillance in cases of AFRS and a follow-up schedule by nasal endoscopic examination and CT scan was suggested to look for recurrence and recidivism. AFRS recidivism appears to be influenced by long-term postoperative therapy. Schubert et al reported the long-term clinical outcome of 67 patients following initial surgical therapy for AFRS. Patients treated with at least 2 months of oral corticosteroids were compared with those who received no corticosteroids. At 1 year after initial surgery, patients treated with oral corticosteroid were significantly less likely to have experienced recurrent AFRS. It is important to realize that AFRS recidivism remains high despite appropriate postoperative medical therapy. Nasal endoscopy at regular intervals is the best way to monitor the activity of disease, and patients should be encouraged to return early for any symptom exacerbation.

**KEY POINTS**

Indian perspective: Consensus arrived by a national panel on AFRS at the ENT surgical update 2011, held at PGI, Chandigarh.
1. CT scan of nose and PNS with axial, coronal and sagittal sections mandatory for endoscopic sinus surgery.
2. Role of preoperative steroids: Oral prednisolone to be started in normal patients and deflazacort in diabetics in a dose of 0.5 mg/kg body weight preoperatively.
REFERENCES


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