Local Immunoglobulin E in the Nasal Mucosa: Clinical Implications

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Immunoglobulin E (IgE) can be highly elevated in the airway mucosa independently of IgE serum levels and atopic status. Mostly, systemic markers are assessed to investigate inflammation in airway disease for research or clinical practice. A more accurate but more cumbersome approach to determine inflammation at the target organ would be to evaluate markers locally. We review evidence for local production of IgE in allergic rhinitis (AR) and chronic rhinosinusitis with nasal polyps (CRSsNP). Diagnostic and therapeutic consequences in clinical practice are discussed. We describe that the airway mucosa has the intrinsic capability to produce IgE. Moreover, not only do IgE-positive B cells reside within the mucosa, but all tools are present locally for affinity maturation by somatic hypermutation (SHM), clonal expansion, and class switch recombination to IgE. Recognizing local IgE in the absence of systemic IgE has diagnostic and therapeutic consequences. Therefore, we emphasize the importance of local IgE in patients with a history of AR or CRSsNP.

Key Words: local IgE; allergic rhinitis; local allergic rhinitis; chronic rhinosinusitis with nasal polyps; diagnostics; treatment

INTRODUCTION

Immunoglobulin E (IgE) is a major contributing factor in multiple airway diseases, including allergic rhinitis (AR) and chronic rhinosinusitis with nasal polyps (CRSsNP). However, measuring IgE by classical systemic tests fails to give an adequate idea of local IgE in the target organ, the nose.¹² In this review, we summarize the evidence of local production of IgE in sinonasal diseases, and clinical implications will be discussed.

Diagnostic tools in rhinitis may not be sufficient to differentiate between allergic, non-allergic, and local allergic rhinitis (LAR), as local IgE normally is not measured.

In chronic rhinosinusitis, different disease subgroups exist³ with their inherent pathomechanisms, and it is challenging to find good markers to further categorize the nasal-polyposis population. It would be especially interesting to determine the endotype in which IgE, whether or not systemic, is crucial in the pathogenesis. The response to targeted therapy as anti-IgE can be predicted.

Classical pathway for development of IgE-positive B cells

Mature naïve B cells encounter antigen processed and presented by dendritic cells in peripheral lymphoid organs. They become activated after interaction with T cells specific for an incoming antigen. After activation on the boundary between B-cell follicles and T-cell zones, the B cells have 2 options. They migrate to the follicle, proliferate, and form germinal centers, or they migrate to an extra-follicular region, proliferate, and differentiate into short-lived plasma cells. B cells in the germinal center undergo antibody affinity maturation by means of somatic hypermutation (SHM), clonal expansion, and class switch recombination to IgE. Recognizing local IgE in the absence of systemic IgE has diagnostic and therapeutic consequences. Therefore, we emphasize the importance of local IgE in patients with a history of AR or CRSsNP.

SHM and CSR are necessary to create enormous diversity found in the antibody and T-cell receptor repertoires required for an effective immune response. As mentioned before, these reactions generally take place within germinal centers, which are typically located in secondary lymphoid tissues, such as tonsil tissue, lymph nodes, and the spleen. SHM is a modification of the genome sequence in somatic cells by substitution of a single base in variable regions of Immunoglobulin (Ig) genes in B cells. CSR in the Ig heavy chain gene locus of the constant region is necessary to class switch from IgM, IgG, or IgA to IgE, resulting in B cells expressing IgE. Both SHM and CSR are initi-

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ated by activation-induced cytidine deaminase (AID), and thus this molecule can be used as a marker for these processes. When a mature B cell alters its receptor in response to antigenic stimulation, this is called receptor revision (RR), initiated by recombination-activating gene products (RAG1 and RAG2).

Signals from T helper cells are mandatory for CSR to IgE+ cells, namely interleukin (IL) 4 and IL13, and the ligation between CD40 on B cells and CD40ligand on T cells. After binding of the promoter Igς, the production of germline transcripts (ςGLT) is initiated. This precedes IgE class switching and recombination of the heavy chain by AID. The outcome is a mature ε chain mRNA and a circular fragment of DNA that is looped out, known as a ‘switch circle’. Ego switch circles can be used as a marker of ongoing CSR in B cells.

Isotype-switched B cells that leave the germinal center reaction become either memory B cells or long-lived plasma cells. Memory B cells divide, whereas long-lived plasma cells do not self-renew. Memory B cells secrete little Ig, but rapidly provide antigen-specific antibody-secreting plasma cells upon antigen recall. Long-lived plasma cells provide long-term maintenance of antigen-specific antibody titers. This is likely the case for IgE as well. Both memory cells and long-lived plasma cells are the cellular source of IgE memory, and they ensure humoral memory. In a murine model, the results of the group of Erazo suggest that IgE+ cells that exit the germinal center reaction preferentially develop into plasma cells. The differentiation of B cells into plasma cells is directed by B cell-activating factor of the TNF-family (BAFF) and B-lymphocyte-induced maturation protein (BLIMP).

AR and non-AR

IgE in AR and non-AR

Rhinitis is traditionally categorized as allergic, infectious, or non-infectious non-allergic rhinitis (NINAR). NINAR is diagnosed by exclusion, meaning that this category includes a heterogeneous group of rhinitis patients with a poorly defined pathogenesis. Mostly, no etiology is found, and this subgroup is known as idiopathic rhinitis.

IgE is one of the most important markers of allergy, as it is the major contributing factor in most types of allergic disease. Traditionally, the distinction between allergic and non-allergic rhinitis is based on skin prick tests and serum IgE analysis. However, in a subgroup of patients with a typical history suggestive of allergy, these tests are negative, but measurement of local IgE can show elevated levels. This observation suggests that a local form of rhinitis may exist, known as ‘LAR’.

AR and Systemic IgE

Respiratory mucosa is a site of IgE induction during allergic airway inflammation. The group of Ecker-Dornal investigated the role of blood-derived plasma cells and B cells in the production of allergen-specific IgE. In a series of cell separation experiments performed by negative depletion and positive selection, they found that the majority of circulating specific IgE antibodies is not derived from IgE-producing cells in the blood. This result suggests that the production at the target organ is the probable source. Serum IgE in AR might thus be a result of spill over rather than the reverse; nearly all of the serum IgE may be derived from mucosal sites.

LAR

Conventionally, an immune response is primed in germinal centers within lymph nodes, whereas the effector function of antibodies is peripheral, e.g. mucosal. In AR, however, all events may occur peripherally. Localized IgE-mediated inflammation may be suspected in a small subgroup of the idiopathic rhinitis group with negative skin prick testing, where allergen-specific IgE is measurable in nasal secretions of patients. Also, studies have proven that the localized cellular pathogenesis is analogous to that in patients with AR, suggesting that these ‘non-allergic’ subjects are in fact allergic.

Evidence of Local IgE production

In AR, IgE-positive B cells would reside in the nasal mucosa, as local IgE production has been perceived in vitro and in vivo. The possibility that IgE is produced locally in the target organ was first proposed after an experiment in children with AR to house dust mite and asthma. IgE was measured in the serum and in nasal secretions in steady state, and kinetic studies were added. It was seen that after re-exposure, local IgE increased more rapidly than serum IgE. In another experiment, levels of specific IgE measured in nasal secretions exceed levels measured in serum of patients diagnosed with AR to ceder pollens. Furthermore, in cases where no serum specific IgE is measurable, elevated levels are found in nasal secretions. This was demonstrated for HDM-specific IgE and for specific IgE to grass/olive pollens in non-allergic patients with positive nasal provocation tests.

Kleinjan et al. provided evidence for local production of specific IgE, as IgE-positive mast cells, plasma cells, and B cells are found in inferior turbinate tissue from patients with AR. The presence of long-lived plasma cells in the mucosa of subjects with AR was also suggested by Smurthwaite et al. who demonstrated persistent IgE synthesis between seasons in explants of hay fever patients. Ex vivo stimulation of tissue obtained from allergic patients leads to an increase in IgE, meaning that all is available locally to produce IgE. Cameron et al. demonstrated elevated levels of IL4 in AR, meaning the nasal mucosa in AR is a favorable environment for CSR. Hence, local production and release of IL4 and IL13 by T cells and mast cells may regulate the local IgE production. These cells also carry the CD40 ligand necessary for DNA recombination and IgE synthesis. Regarding functionality, Pawankar et al. demonstrated up-regulation of FcεRI on mast cells in response to locally pro-
duced specific IgE. Subsequently to the increased IgE cross-linking, more mast cells are activated and degranulate, promoting allergic reactions.

As previously discussed, evidence points at the local synthesis and secretion of IgE. Does CSR to IgE+ B cells and affinity maturation occur in distant lymphoid tissues before migrating to the target organ or do they first migrate and only then class switch? In human nasal mucosa of allergic individuals, GC formation has not yet been demonstrated,

Although SH and CSR to IgE have been described. In an ex vivo experimental setting with nasal mucosa from grass pollen-sensitized subjects, class switching is found upon allergen stimulation. The group of Coker detected local bias to the VH5 germline gene family, while no VH5 overexpression was detected in the blood. Furthermore, AID is continuously expressed, representing a fundamental aberration in the mucosa of AR patients. Next to AR, local IgE formation is also present in CRSwNP. IgE in nasal polyps is polyclonal, whereas IgE in AR is monoclonal or oligoclonal. In contrast to AR, germinal center formation has been documented in nasal polyps,

autoimmune diseases, and lower airways. This information illustrates that the nasal mucosa has the intrinsic capability of affinity maturation by SHM, clonal expansion, and CSR to IgE.

Pathomechanisms underlying AR

To understand the role of IgE in allergic and ‘non-allergic’ rhinitis, knowledge of pathomechanisms is essential. Allergic inflammation typically comprises an early and a late phase organized by structural epithelial cells, resident mast cells, and infiltrating eosinophils/basophils/T cells. Cytokines released from mast cells and T cells mediate local IgE production by B cells.

1) The role of epithelial cells

Epithelial cells are not merely functional as a barrier, but upon activation they can also release immunomodulatory substances that regulate Th2 cytokine response, including eicosanoids, endopeptidases, cytokines (thymic stromal lymphopoietin [TSLP], IL25, and IL33), and chemokines. Epithelial cells can be activated by an IgE-mediated mechanism.

2) Immediate response

The activation of mast cells is crucial in the immediate response, and activation by antigen cross-linking of IgE is a well-known mechanism. Sensitized mast cells have both high and low affinity receptors for IgE on their surface, the abg2 tetramer FcRI and the ag2 trimer FcRII, respectively. The latter is also termed CD23 receptor and found on a broad range of cells. These receptors bind IgE, and upon cross-linking the mast cell degranulates and releases different mediators like histamine, tryptase, newly synthesized lipid mediators, cytokines, and chemokines. Histamine, leukotrienes, and prostaglandins cause clinical symptoms, such as sneezing, running nose, and nasal obstruction.

Basophils are phenotypically similar to mast cells, as they are granulocytes containing histamine and expressing FcεRI. Thus, upon cross-linking with specific IgE, they release histamine. However, basophils can migrate to lymphoid tissues, while mast cells cannot. It is suggested that effector functions of basophils are heterogeneous according to the eliciting factor. IL3-elicited basophils would be highly responsive to IgE, conversely to TSLP-elicited basophils that would be non-responsive to IgE.

3) Late phase response

The release of cytokines and chemokines by mast cells in the immediate phase response, which starts within minutes and lasts for 2-3 hours, is important for the subsequent late phase response. The late phase response starts 4-6 hours after stimulation and lasts for 18-24 hours. An infiltrate containing T helper 2 (Th2) cells, eosinophils, and basophils is characteristic of this phase. Th2 cells are important for the production of key cytokines, including IL4, IL5, IL9, and IL13. These cytokines are essential for antibody class switching, regulation of local and systemic IgE synthesis, recruitment of eosinophils, basophils, and mast cells, and survival of eosinophils. It is mostly assumed that eosinophils differentiate within the bone marrow in response to eosinophilopoietins. In AR, however, a subset of eosinophils would differentiate within the nasal mucosa in a highly IL5-dependent manner. Langerhans cells are dendritic cells of the skin and mucosa. They are professional antigen-presenting cells (APCs), which process aeroallergens deposited on the mucosa and subsequently present the antigens to T cells. Th2 cells release their mediators upon recognition of antigen presented by APCs. The Th2 cytokines IL4, IL13, and CD40L induce selective somatic recombination of Ig heavy chain regions in B cells before maturation into IgE-producing plasma cells.

Pathomechanisms underlying non-AR

In ‘non-allergic’ rhinitis, no systemic markers of allergy can be detected. These patients could however still be allergic in the case of LAR, although most cases are truly non-allergic. Here, a broad differential diagnosis has to be made. Endonasal infections are a common cause of non-allergic rhinitis, and complaints can persist long after resolution of the infection. Non-specific exogenous irritants, such as tobacco smoke, pollution, dust, smog, perfumes, solvents, and occupational irritants, could also trigger mucosal inflammation. Furthermore, rhinitis can be elicited by foods and beverages, namely ‘gustatory rhinitis.’ Changes in hormones due to pregnancy or hypothyroidism, for example, could result in rhinitis. In the elderly, watery rhinitis or geriatric rhinitis is often seen. Non-IgE-mediated hypersensitivity can also explain the discrepancy between IgE testing results and...
clinical findings, for example, T-cell mediated delayed type hypersensitivity and activation of mast cells by Ig free light chains (FLCs).33,34

Evidence increases for the possibility that a T cell-mediated inflammatory reaction to common antigens sustains asthma and AR, especially when atopic dermatitis is identified in the history.34 Here again, negative skin prick test (SPT) and RAST can be expected.

Ig FLC could serve as an alternative for IgE in eliciting inflammation in allergic disorders, including food allergy, atopic dermatitis, rhinitis, and asthma.35 FLCs can cause immediate hypersensitivity through mast cell activation, and they are found in both atopic and non-atopic rhinitis.36 Moreover, Powe et al.35 demonstrated a link between FLC immunoglobulins and mast cells in the nasal mucosa, which suggests an association between Ig FLC and mast cell-mediated nasal hyperreactivity. The role of FLC in AR is not clear, and further studies are necessary to investigate the function of FLC in local hypersensitivity.

Clinical implication: diagnostic workup of AR and non-AR

Currently, the diagnostic workup of AR must consist of a history of the patient, with most important differentiation between intermittent and persistent symptoms and between mild and moderate to severe AR following the ARIA guidelines. In addition, clinical examination along with anterior rhinoscopy and nasoendoscopy is required. At last, allergen extract-based IgE tests are indispensable, such as SPT and the measurement of specific IgE antibodies in serum. Though these conventional tests are expected to be negative in cases where IgE is only situated in the nasal mucosa and those where rhinitis is induced by non-IgE-mediated mechanisms. These patients are classified as ‘non-allergic,’ although additional testing for local IgE may reveal a local allergic response in a subgroup of patients. In these cases where history is truly suggestive of AR and currently used tests are negative, it has to be questioned whether measurement of systemic IgE levels is sufficient for the diagnostic workup.

Additional testing by means of nasal provocation, as proposed by Rondón et al.36 or measurement of local IgE in the nasal mucosa can be considered to reveal LAB. However, it is currently unclear how large this subgroup is; findings from our clinic/group point to rather small numbers.37 Methods to measure local mediators have not yet been standardized, and this will be discussed further in this review. To determine whether T cell-mediated, delayed hypersensitivity lies at the basis of the complaints, an Atopy Patch Test can be useful; however, the relevance of this test in AR has not yet been established.

Clinical implication: treatment of AR and non-AR

The correct diagnosis and knowledge of pathophysiology are essential in establishing the best treatment for a patient.

Mast cells are crucial in allergic inflammation, and different treatment options aim to block their effect. Traditionally, topical corticosteroids and antihistamines are prescribed.

Avoidance of allergens if possible is an important step, for instance, in AR to mold or cat allergens. In most cases, however, it seems that this is hardly effective or impossible, such as in the case of AR to grass pollen or house dust mite. Only when it is known to which allergen the patient is sensitized, avoidance of allergen is possible, and/or immunotherapy can be considered.

Recent diagnostic tests based on recombinant allergens, epitopes, and peptides allow more precise diagnosis of allergy. New immunotherapy strategies that would be more effective and safer than conventional allergy vaccines aim to suppress IgE-mediated inflammation. T-cell tolerance can be induced by administration of peptides containing T-cell epitopes, carrier-bound allergen peptides, or recombinant hypoallergens.

Targeted treatments using antibodies are developing. In allergic diseases, such as AR, asthma, food allergy, and allergic dermatitis, anti-IgE or omalizumab has been investigated. This humanized monoclonal antibody targets the Cε3 domain of IgE, reducing free IgE by forming a biological inert molecule. Furthermore, omalizumab down-regulates FcεRI on effector cells, and it would also decrease IgE synthesis by inducing an anergic state in IgE+ B cells.38 Omalizumab has a good clinical effect in allergic diseases38 that lasts for about 4 months, and it has a low incidence of side effects.39 This treatment is, in theory, an option in AR. However, the high cost-effectiveness ratios37 result in restriction of indications. Theoretically, rituximab or anti-CD20 could also be an option to inhibit IgE-producing cells.

To conclude, allergen avoidance (if possible), local corticoids, antihistaminic drugs, and leukotriene receptor antagonists are the first-line treatment options. Immunotherapy can be considered in certain cases where the conventional treatment is not sufficiently effective. In this way, a good control of symptoms is achievable and omalizumab or rituximab are not indicated for AR as such.

CRSwNP

Definition

The term chronic sinusitis covers a wide range of sinus diseases, in which the mucosa is inflamed with symptoms on most days lasting at least 12 weeks without complete resolution. In the pathophysiology, intrinsic (for example genetic) and extrinsic (for example microbial, environmental, and iatrogenic) factors may be involved, which can act locally or systemically.

Two phenotypes are currently distinguished, namely chronic rhinosinusitis without nasal polyposis (CRSsNP) and CRSwNP. However, some authors believe that different disease entities classified as chronic sinusitis are a continuum of the same disease. These phenotypes show considerable overlap in symptoms, but can be differentiated from each other with reasonable certainty by endoscopy.
Inflammation patterns

The definition ‘chronic sinusitis’ suggests a single clinical entity, but in reality it represents multiple overlapping entities with different inflammation patterns.

In CRSsNP, mostly Th1-skewed neutrophilic inflammation is found with elevated levels of interferon-γ (IFNγ) and transforming growth factor-β (TGFβ), whereas inflammation in CRSwNP is often Th2-skewed eosinophilic inflammation with elevated levels of IL5 and IgE. However, patients with CRSwNP also form a heterogeneous group with different endotypes, meaning that inflammation patterns and T helper subsets vary.

In the Western countries, the majority of nasal polyps is characterized by Th2-skewed inflammation, dominated by cytokines that promote production of IgE and eosinophilic inflammation. IL4 stimulates proliferation and differentiation in Th2 cells and mediates CSR to IgE by stimulating the synthesis of ε-germline gene transcript (GLT). IL5 is elevated in CRSwNP and remarkably increased in CRSwNP accompanied by non-allergic asthma and aspirin sensitivity. IL5 mediates eosinophil survival, differentiation, and recruitment to the nasal mucosa, and causes secretion of eosinophil cationic protein (ECP). ECP is an indicator of eosinophilic activation, and local ECP correlates with local IgE. The local inflammatory pattern could be related to the type of bacterial colonization. Colonization with Gram-positive bacteria, for example, *Staphylococcus aureus* (SA) is common in individuals with nasal polyps characterized by elevated IL5, ECP, and total IgE. In Asian populations, the inflammation in CRSwNP is frequently predominantly neutrophilic. This pattern is associated with greater Gram-negative bacterial colonization. Patients with nasal polyps and cystic fibrosis have Th1/Th17-skewed neutrophilic inflammation with highly increased expression of IL17 and IFNγ.

Biochemical markers that could be used to differentiate between CRSsNP and CRSwNP in Caucasians are ECP, IL5, and IgE. Markers to differentiate between CRSwNP and cystic fibrosis with nasal polyps also include IL8, IL17 and MPO.

IgE in CRSwNP

**Local IgE**

Total IgE is often highly increased in nasal polyp mucosa, and IgE specific to staphylococcal enterotoxins can be present independently of serum total/specific IgE. The presence of plasma cells in the mucosa insinuates local production of IgE, which is polyclonal and functional.

In atopic nasal polyp patients, local IgE production can be the effect of stimulation with allergens. However, local hyperimmunoglobulinemia is also present in non-atopic patients, meaning that elevated IgE levels result from other pathways as well. The cytokines IL25 and IL33 can induce IgE-mediated inflammation by stimulating a non-T cell source to produce IL4.

**Fig. 1.** Pathways resulting in local IgE production in CRSwNP. In the first part, dendritic cells of the skin and mucosa process aeroallergens deposited on the mucosa, and subsequently they present antigens to T cells. T helper 2 cells release their mediators upon recognition of antigens presented by antigen-presenting cells. The Th2 cytokines IL4, IL13, and CD40L induce selective somatic recombination of immunoglobulin heavy chain regions in B cells before maturation into IgE-producing plasma cells. IL5 stimulates eosinophil growth and differentiation. Alternatively, IgE is produced by stimulating innate lymphoid cells to release IL4, IL5, and IL13.
IL5, and IL13, namely innate lymphoid cells (ILC)\textsuperscript{59} (Fig. 1). A role of mast cells in enhancing eosinophilic inflammation in chronic rhinosinusitis is suggested. \textit{Ex vivo} experiments demonstrate that activation of IgE cross-linking in CRSwNP\textsuperscript{47} and FLCs present in nasal polyps could mediate local immune responses.\textsuperscript{50} FLC concentrations correlate with IL5, IL6, and local IgE. Furthermore, a decrease in local FLCs is seen after treatment with anti-IL5, presuming that IL5 creates an environment that favors FLC production.\textsuperscript{50} Next to IgE and FLC, locally produced IgA\textsuperscript{51} could also be involved in the activation of mast cells and eosinophils. The role of IgA is not clear, but elevated levels of IgA are often found in patients with chronic mucosal inflammation. In CRSwNP as well, elevated levels of IgA is found in tissue homogenates,\textsuperscript{51} although this could be explained by a decreased translocation of IgA from the epithelium of the lamina propria into nasal secretions.\textsuperscript{52}

\textbf{Germinal center reaction in nasal polyposis}

As mentioned earlier, it is generally believed that GC are situated in lymphoid tissues, meaning that affinity maturation by SHM and CSR takes place in these lymphoid tissues. The involvement of local IgE production has been investigated by comparing key markers of Th2 inflammation and GC reactions in nasal polyp tissue versus control tissue. We measured elevated levels of IL4, IL5, IFN-\gamma, mIgM, and local IgE, which all point at local CSR.

Enhanced differentiation of B cells into plasma cells in nasal polyps can be concluded from the increased number of plasma cells and ratio of plasma cells to B cells, together with the elevated BAFF levels. The up-regulation of RAG1/RAG2 points at local RR. The presence of switch circle transcripts points at ongoing local CSR. To conclude, our group was the first to reveal formation on GC-like structures in polyps with local RR, class switching to IgE, and B-cell differentiation into IgE-secreting plasma cells.\textsuperscript{5} Because T follicular helper (Tfh) cells and IL21 produced by these cells are important in regulating B cells in GC and lymphoid accumulations are found in nasal polyps, a new study was conducted. Here, IL21 and IL21- positive T helper cells were measured in nasal tissues of CRSwNP, CRSsNP, and controls. Elevated levels of IL21 and IL21-positive T helper cells were found in CRSwNP, and after stimulation with staphylococcal enterotoxin (SE) B during 24 hours a significant increase of IL21 in the CRSwNP was seen. In the same study, high expression of B lymphocyte-induced maturation protein-1 (Blimp-1) and B-cell lymphoma-6 (Bcl-6) was found. These molecules are antagonists with Bcl-6 being the regulator of Tfh cell differentiation and Blimp-1 suppressing generation of Tfh cells. The authors suggest that both mediators regulate differentiation of T and B cells in nasal polyp tissue. It is concluded that IL21 produced by Tfh cells stimulates GC formation in CRSwNP (unpublished data). The group of Mechtcheriakova\textsuperscript{49} used CRSwNP as a model to study AID-associated responses in diseases characterized by chronic inflammation. In nasal polyp tissue, they detected functional AID-positive ectopic follicular structures by real-time PCR analysis and immunostaining. No AID mRNA could be detected in the control CRSsNP tissue. The transcription of IgE and/or IgG was positively correlated with the expression of AID mRNA. These results indicate that CSR and SHM can take place locally in the airways at sites of chronic inflammation.

\textbf{SA-Specific IgE}

Local IgE in CRSwNP is polyclonal and associated with the presence of IgE antibodies to SEs.\textsuperscript{46,49} SA is one of the most frequently found bacteria in CRSwNP. Colonization with SA is significantly more prevalent in the population with nasal polyps compared to controls. Colonization is present in 27.3% of CRSsNP patients, 33% of control subjects, 60% of patients with CRSwNP and as high as 87% in the subgroup with asthma and aspirin intolerance, based on culture data.\textsuperscript{48} An increased colonization rate of SA was demonstrated in patients with CRSwNP (63.6%), but not in those with CRSsNP (27.3%) and control subjects (33%).\textsuperscript{55}

Immune response rates to enterotoxines released by SA follow the same trend, with SE-specific IgE present in 28% of CRSwNP and up to 80% in the subgroup with aspirin-exacerbated respiratory disease.\textsuperscript{49,56} SE contributes to nasal polyp formation in both atopic and non-atopic patients,\textsuperscript{46,57} since these enterotoxins have the ability to induce enormous activation of lymphocytes without undergoing processing by APCs. Consequently, severe eosinophilic inflammation arises, and polyclonal IgE and IgG/IgG4 are produced by B cells.\textsuperscript{57} Products of viral or bacterial microorganisms with this feature are identified as superantigens.

The activation of T- and B-cells by T-and B-cell superantigens, respectively, is nonspecific and polyclonal, and leads to massive cytokine release. This resultant polyclonal IgE saturates the receptors on local tissue mast cells, reducing the effect of specific IgE and the allergic response. Superantigens skew the inflammation toward Th2-mediated inflammation with increased levels of ECP, IL5, and IgE, or a pre-existing Th2 milieu might facilitate the persistence of SA. Moreover, macrophages are alternatively activated, causing deficient phagocytosis of SA in CRSwNP.\textsuperscript{57} Detection of IgE specific to SE has been demonstrated for tissue homogenates, but rarely in serum of subjects without comorbid asthma. Moreover, the levels of IgE specific to SEA and SEB detected in tissue homogenates correlate with levels of ECP and IL5, and thus eosinophilic inflammation.\textsuperscript{58}

\textbf{IgE}

Eosinophilic inflammation with elevated IgE found in nasal polyps suggests an atopic background, although atopy has no impact on tissue inflammatory patterns in Caucasian polyps. Epidemiological data support this notion, as CRSwNP is as prevalent in atopic patients as in non-atopic patients. Suh et al., in a recent study, found a higher prevalence of SA-specific IgE in CRSwNP patients compared to CRSsNP and control subjects. They also observed a higher prevalence of SA colonization in patients with CRSwNP compared to CRSsNP and control subjects. The presence of SA in CRSwNP is associated with higher levels of IgE-specific antibodies to SEs, indicating a possible role of SA as a triggering factor in eosinophilic inflammation.
本地IgE在鼻黏膜

由Sheahan等人在2010年发现，本地IgE在非过敏鼻窦炎患者的鼻息肉组织中以57%的频率出现，与强或弱的系统性过敏反应有关。在2010年，他们发现本地IgE在有或无哮喘的鼻窦炎患者中水平较高，与对照组和非过敏性鼻窦炎患者相比。

临床意义：诊断

诊断是不足的，如果仅仅基于临床和放射学评价，本地IgE可能用于鼻窦炎患者的诊断。

伴发哮喘

CRSwNP与哮喘和/或阿司匹林不耐受有关。CRSwNP和哮喘具有某些特征，常常共存，这使得认为它们可以共同表达同一类型的Th2介导的疾病。重要的是，鼻息肉组织和哮喘中本地IgE的水平很高，与哮喘的增加风险有关。在2010年，Sheahan等人发现本地IgE在非过敏性鼻窦炎患者中的水平较高，与对照组和CRSwNP相比。

临床意义：治疗

目前，CRSwNP的治疗主要包括药物治疗、鼻腔冲洗和口服皮质类固醇。然而，对于鼻息肉或鼻窦炎的长期复发，这些药物治疗的效果非常有限。

抗生素是治疗的选择，作为对微生物群的治疗。在实践中，对于鼻窦炎和鼻息肉的治疗，需要进行鼻窦手术。然而，CRSwNP是一种慢性疾病，鼻窦和鼻息肉的治疗需要长期的治疗。
good choices, because they have an anti-inflammatory action next to their antibiotic features.

Macrolides like erythromycin, clarithromycin, and roxithromycin affect neutrophils and eosinophils to reduce tissue damage by chronic bacterial colonization. The tetracycline antibiotic family has good tissue penetration in airway tissue, has a broad spectrum, and is especially effective against SA. The effect of doxycycline was examined by Van Zele et al. and had a significant, albeit more moderate, effect compared to that of oral glucocorticosteroids. The effect of 3-week doxycycline treatment with 100 mg/day was present for 12 weeks, whereas the effect of tapering methylprednisolone treatment over the same period only lasted for 8 weeks.

**New treatment options that target IgE directly or indirectly**

By the end of the last century, biological antibodies were developed, including anti-IL5, anti-IL4Rα, and anti-IgE. Unlike corticoids, these specific treatments are only effective in a selected population. Nevertheless, it might be an opportunity to predict responsiveness to a certain targeted treatment based on different T-cell patterns and cytokines in serum, in mucosa when surgery is mandatory, or in nasal secretions.

1) Direct inhibition of IgE

Considering Th2-skewed eosinophilic inflammation with marked local production of functional IgE antibodies, it can be assumed that treatment with anti-IgE is indicated for this endotype. Especially in patients with nasal polyps and comorbid asthma, treatment targeting IgE can be advocated, as high local IgE is frequently present in this subpopulation (Fig. 2). Indeed, this therapy has proven to be successful in CRSwNP and comorbid asthma with a substantial decrease in nasal polyp scores after 16 weeks in the omalizumab group, as compared to baseline. Upper and lower airway symptoms significantly improve by application of anti-IgE therapy in both atopic and nonatopic patients with nasal polyps. The independence of atopic status highlights the functionality of the locally produced IgE. This result is in contrast to that of a study conducted by Pinto in 2010. In this randomized controlled trial, the effect of omalizumab was determined in patients with chronic sinusitis, as compared to placebo. An improvement of sinus opacification in CT scans and the SNOT-20 questionnaire was found, but the difference was not significant. However, the population also included patients with CRSsNP, which may partially explain the limited benefit from omalizumab; furthermore, the patients were using intranasal corticosteroids, which were not tried to reduce. From both studies, we can conclude that patients have to be selected carefully when considering a treatment or a study with a targeted treatment.

2) Indirect inhibition IgE: Anti-IL4rα, Anti-TSLP, and Anti-IL33

IL4 is, next to IL13, an important component in airway inflammation, as the IL4/IL13/STAT6 pathway is crucial in Th2-mediated inflammation. Several IL4-antagonizing approaches have been proposed, including the soluble recombinant human IL4 receptor, IL4 mutein, humanized anti-IL4 monoclonal antibody, and human anti-IL4 receptor α monoclonal antibody. By targeting the α subunit of the IL4 receptor, both IL4 and IL13 are blocked; this could be more effective than targeting either one alone. Dupilumab is a fully human monoclonal antibody targeting the IL4 receptor α subunit. A randomized controlled trial has investigated the efficacy and safety of dupilumab in patients with persistent, moderate-to-severe asthma and a blood eosinophil count of at least 300 cells per microliter or a sputum eosinophil level of at least 3%. In view of the good clinical result in asthma, it would be interesting to investigate the efficacy in eosinophilic CRSwNP.

As mentioned before, TSLP is an epithelia derived cytokine that is important in promoting Th2-type inflammation. An increased expression of messenger RNA for TSLP and an increased number of TSLP-positive cells are demonstrated in nasal polyps. Moreover, there is a clear correlation between the number of TSLP-positive cells and the IgE level. It would be interesting to assess the effect of TSLP inhibition on the IgE level in CRSwNP. AMG 157 is a human antibody targeting TSLP, and thus theoretically this antibody could prevent need to same wole text IgE formation upstream. Similarly, next to TSLP, IL33, and IL25 represent a link between the epithelium and the Th2-dominated adaptive immunity in CRSwNP (Fig. 1). The IL33/ST2 pathway be a therapeutic target, and blockade by means of anti-IL33 or soluble ST2 could reduce inflammation in nasal polyps. To our knowledge, however, the impact of such blockade on local IgE in CRSwNP has not yet been investigated.

**CONCLUSIONS**

In many sinonasal diseases, IgE is crucial in the pathogenesis. Systemic markers to investigate inflammation in upper airway disease are sometimes not representative for local inflammation. IgE that is locally produced in the target organ could have diagnostic and therapeutic importance in AR and CRSwNP.

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