Current Insights into the Pathogenesis of Graves’ Ophthalmopathy

Abstract

Environmental, genetic, and immune factors are at play in the development of the variable clinical manifestations of Graves’ ophthalmopathy (GO). Among the environmental contributions, smoking is the risk factor most consistently linked to the development or worsening of the disease. The close temporal relationship between the diagnoses of Graves’ hyperthyroidism and GO have long suggested that these 2 autoimmune conditions may share pathophysiologic features. The finding that the thyrotropin receptor (TSHR) is expressed in orbital fibroblasts, the target cells in GO, supported the notion of a common autoantigen. Both cellular and humoral immunity directed against TSHR expressed on orbital fibroblasts likely initiate the disease process. Activation of helper T cells recognizing TSHR peptides and ligation of TSHR by TRAb lead to the secretion of inflammatory cytokines and chemokines, and enhanced hyaluronic acid (HA) production and adipogenesis. The resulting connective tissue remodeling results in varying degrees extraocular muscle enlargement and orbital fat expansion. A subset of orbital fibroblasts express CD34, are bone-marrow derived, and circulate as fibrocytes that infiltrate connective tissues at sites of injury or inflammation. As these express high levels of TSHR and are capable of producing copious cytokines and chemokines, they may represent an orbital fibroblast population that plays a central role in GO development. In addition to TSHR, orbital fibroblasts from patients with GO express high levels of IGF-1R. Recent studies suggest that these receptors engage in cross-talk induced by TSHR ligation to synergistically enhance TSHR signaling, HA production, and the secretion of inflammatory mediators.

Introduction

Graves’ ophthalmopathy (GO), also known as Graves’ orbitopathy or thyroid eye disease, is an inflammatory autoimmune disorder [1]. While the majority of patients with this condition experience only mild eye irritation and redness, some 3–5% suffer from more severe disease. Disease manifestations vary include redness and swelling of the conjunctivae and lids, forward protrusion of the globes (proptosis), ocular pain, debilitating double vision, and even sight loss due to compressive optic neuropathy or breakdown of the cornea [2]. From a mechanistic standpoint, these features derive largely from progressive enlargement of the orbital adipose tissues and extra-ocular muscles within the confines of the bony orbit. As the orbital pressure rises, proptosis may develop and venous drainage may be compromised with impairment of inflammatory mediator efflux. Diplopia in early disease appears to result from active inflammation within the enlarged and edematous muscles. When encountered in late, inactive disease, diplopia likely reflects muscle fibrosis. This chapter will review current insights into the environmental, genetic, and pathobiological mechanisms responsible for orbital tissue remodeling in GO.

Risk Factors for the Development of GO

Patients with GO are more likely to be women by a 2:1 ratio while men with Graves’ disease appear to be at higher risk for the development of severe GO [3]. Racial differences in the prevalence of GO have also been noted with Asians having a lower likelihood of developing the disease than do Caucasians [4]. Genetic studies have identified candidate genes associated with the development of...
Graves’ hyperthyroidism including the immune regulatory genes, human leukocyte antigen (HLA)-DRB1-Arg74, cytotoxic T lymphocyte antigen (CTLA)-4, PTPN22, and CD40, and the thyroid-related genes thyroglobulin and thyrotropin receptor (TSHR) [5]. However, none of these genes contribute more than a 4-fold increase in risk, are essential for disease development or have been clearly shown to confer separate risk for GO [6]. Overall these findings suggest that Graves’ disease has a complex genetic basis with multiple susceptibility alleles that, in combination with epigenetic, post-translational, and environmental factors, predispose to the various disease phenotypes, including the presence of GO [7, 8].

Smoking is the risk factor that has most consistently been linked to the development or worsening of GO [9]. It was the primary risk factor identified for GO outcome with an odds ratio among smokers vs. nonsmokers of 5.2 in a recent trial of patients with newly diagnosed Graves’ hyperthyroidism treated with either radioactive iodine or antithyroid drugs [10]. Upon post-hoc analysis with a better outcome definition, smoking was found to impart an even higher odds ratio of 9.8. Overall more than 40% of the smokers either developed GO or experienced deterioration in the disease, a rate almost double that seen in nonsmokers. The GO risk relative to active smoking is proportional to the number of cigarettes smoked per day, and former smokers have significantly lower risk than current smokers [11]. The potential impact of secondhand smoke was suggested in a retrospective, questionnaire-based study demonstrating a higher rate of childhood GO in the European countries having the highest prevalence of adult smoking [12]. While in vitro studies have shown that both cigarette smoke extract [13] and hypoxia [14] increase hyaluronic acid (HA) production by orbital fibroblasts and that cigarette smoke extract synergizes with interleukin (IL)-1 [13] to enhance adipogenesis in these cells, mechanisms underlying the deleterious impact of smoking in GO (as well as in other autoimmune diseases) are unclear. Clearly important insights into GO pathogenesis lie in better understanding of this association.

Multiple studies have concluded that radioactive iodine (RAI) treatment of Graves’ hyperthyroidism negatively impacts the eyes [15, 16] and that smoking may represent an additive risk factor in that setting [10]. The association of GO with RAI may be in part explained by the elevated levels of autoantibodies directed against the thyrotropin receptor (TRAb) seen immediately following RAI therapy that remain above pre-RAI levels for approximately a year’s time, then gradually decline over the ensuing several years [17]. In contrast, treatment with antithyroid medication or thyroidectomy is generally followed by a gradual decrease, then disappearance of TRAb in 70–80% of patients after 18 months. While the risk of GO development or progression was found in one randomized controlled trial to be 15% for RAI (compared with 3% for antithyroid drug therapy) [15] other studies reported higher risk for RAI at 33% (compared with 10% for antithyroid drugs and 16% for thyroidectomy) [14], and 39% for RAI (compared with 21% for antithyroid drugs) [10]. Risk factors for RAI-induced activation of GO include male sex, advanced age, and shorter time from onset of GO and post-RAI hypothyroidism [18, 19]. The majority of patients developing GO after RAI therapy have mild and transient disease requiring no treatment and glucocorticoids administered concurrently with RAI therapy with tapering over 6–8 weeks may prevent the development or worsening of mild GO [9, 20, 21]. However, as these studies included only patients with no GO at the time of RAI or only mild disease, the risk of disease worsening and the potential impact of corticosteroid administration in patients with more than mild disease is unclear. Recent practice guidelines suggest that hyperthyroidism in patients with severe GO be treated with either antithyroid drugs or thyroidectomy [22]. Finally, both hyper- and hypothyroidism have been independently shown to confer increased risk for development or deterioration of GO [9]. Accordingly, initiating early thyroxine replacement to prevent hypothyroidism following RAI therapy or close monitoring of thyroid hormone levels in patients treated with antithyroid medication is advised.

### Cellular Functions and Interactions

The autoimmune nature of GO is evidenced by the presence within the orbit of a perivascular and diffuse infiltration with mononuclear cells, including predominately CD4+ T cells with occasional populations of CD8+ cells, B cells, and macrophages [1, 23]. CD4+ T cells were primarily γδT cells, while αβT helper cells were rare [24]. Activation of infiltrating T cells in early disease initiates the secretion of primarily Th1-type cytokines [including interferon (IFN)-γ, tumor necrosis factor (TNF)-α, interleukin (IL)-1β, IL-2] whereas in later disease Th2 responses (IL-4, IL-5, IL-10) predominate [25]. Activated T cells additionally produce leukoregulin, a cytokine capable of potent stimulation of hyaluronic acid (HA) or hyaluronan synthesis by GO fibroblasts [21, 26].

The enlarged extraocular muscles (predominately the inferior and medial recti) and expanded orbital adipose tissue volume are a consequence of HA accumulation in the muscles and connective tissues and the development of new fat cells (adipogenesis) within the orbital fat [1]. In addition, as HA is a profoundly hydrophilic glycosaminoglycan, the orbital tissues retain water and become very edematous. Evidence from several laboratories suggests that fibroblasts investing the extraocular muscle fibers and residing within the orbital connective tissues are the autoimmune target cells in GO [27–30]. Early studies showed that orbital-infiltrating CD8+ T cells recognize orbital fibroblasts and not eye muscle extracts, and that they respond by proliferation via major histocompatibility complex (MHC) class II and CD40 signaling [28]. Orbital fibroblasts may be classified based on the presence or absence of the cell surface glycoprotein CD90, also known as thymocyte antigen-1 ( Thy-1) [31]. Fibroblasts expressing this antigen are abundant in the extraocular muscles. They are capable of copious HA production and can differentiate into myoblasts when treated with transforming growth factor (TGF)-β. In contrast, fibroblasts found within the orbital connective tissues are primarily Thy-1 negative and characteristically undergo adipogenesis under appropriate conditions.

Fibrocytes are CD34+ cells that derive from monocyte and B cell lineages within the bone marrow [32]. These cells circulate as peripheral blood mononuclear cells, infiltrating connective tissues at sites of injury or inflammation in response to chemokines [32]. Recent studies from the laboratory of Dr. Terry J. Smith have suggested that fibrocytes express high levels of thyrotropin receptor (TSHR), produce copious inflammatory cytokines, and may similarly infiltrate orbital tissues in GO [33, 34]. If confirmed by other groups, these cells may represent a subset of orbital fibroblasts playing an important role in the orbital tissue remodeling characteristic of GO.
The Role of TSH Receptor

A close temporal relationship exists between the onset of Graves’ hyperthyroidism and the onset of clinically apparent GO; regardless of which occurs first, the other develops in 80% of patients within 18 months [35]. GO can be diagnosed in 25–50% of patients with Graves’ hyperthyroidism. However, when sensitive imaging techniques are used, evidence of ocular involvement is detectable in the majority of patients [9]. These clinical observations suggest that Graves’ hyperthyroidism and GO share etiologic factors. That said, it is notable that GO is seen on occasion in patients with no thyroid dysfunction. However, using adequately sensitive assays, autoantibodies directed against the thyrotropin receptor (TRAb) can be detected in essentially every patient with GO [36]. Further, levels of TRAb correlate with the severity and clinical activity of the disease [37,38] and high TRAb levels in early GO predict a poor prognosis [39]. As autoantibodies directed against TSHR on thyroid follicular cells stimulate the production of thyroid hormone in Graves’ hyperthyroidism, it was postulated early on that immunoreactivity against TSHR in the orbit may play a role in GO pathogenesis.

Insight into the link between the thyroid and the orbit was gained by the demonstration that orbital fibroblasts express TSHR [40,41]. Studies from several laboratories are in general agreement that TSHR mRNA and protein are present in both GO and normal orbital tissues and fibroblast cultures, and that significantly higher levels of the receptor are detectable in GO cells [42–44]. Further, a positive correlation has been demonstrated between TSHR mRNA levels in individual GO orbital adipose tissue specimens and the activity of the patient’s disease, suggesting a role for TSHR in the development of the clinical disease [45]. TSHR signaling pathways in orbital fibroblasts are similar to those found in thyrocytes [46], including activation of both adenyl cyclase/cAMP and phosphatidylinositol 3-kinase (PI3K)/pAkt cascades [47]. In order to investigate whether TSHR activation plays a role in adipogenesis in GO orbital fibroblasts, Zhang and colleagues introduced an activating mutant TSHR into these cells and demonstrated an increase in adipocyte differentiation, as shown by 2 to 8-fold elevations in levels of early to intermediate adipocyte markers [48]. Our laboratory found increased expression of late adipocyte genes (adiponectin and leptin) and accumulation of lipid in GO orbital fibroblasts cultured in the presence of a stimulatory monoclonal TRAb (termed M22) or bovine TSH. This was inhibited by co-treatment with the PI3-kinase inhibitor LY294002, suggesting that PI3K/pAkt signaling is involved in adipogenesis in these cells [49].

As HA accumulation is another histologic feature of GO orbital tissues, several groups have studied the impact of TSHR activation on HA synthesis in orbital fibroblasts. Zhang and colleagues treated cells with bovine TSH or with 2 different monoclonal TRAb and showed increased HA production in normal orbital fibroblast cultures, but not in similarly treated GO fibroblasts [48]. However, when this same group transfected GO orbital fibroblasts with an activating mutant TSHR, they demonstrated the induction of hyaluronan synthases 1 and 2 and elevated HA production compared with control transfected cells. In our studies of GO orbital fibroblasts treated with either bovine TSH or a potent stimulatory TRAb (termed M22), we found elevated levels of cAMP, pAkt and HA [50]. We also determined that these effects could be abrogated by co-treatment with a small molecule TSHR antagonist, termed C-1 [51], further implicating activation of this receptor in the cellular changes characteristic of GO.

Perhaps the strongest evidence to date that TSHR is the primary autoantigen in GO lies in a recently developed animal model. The protocol, developed by Moshkelgosha and colleagues, involves genetic immunization of inbred mice with human TSHR ectoderm plasmid using close field electroporation [52]. The animals develop TSHR stimulating or blocking antibodies and either hyperthyroidism or hypothyroidism. In addition, following prolonged immunization, inflammatory cell infiltrates consisting of CD3+CD4+ T cells, macrophages and mast cells can be demonstrated in the extraocular muscles in the majority of animals. Outward signs of orbital involvement include inflammation of the eyelids and conjunctivae. Expansion of the orbital adipose tissues is evident in about 10% of animals and magnetic resonance imaging (MRI) reveals hypertrophy of extraocular muscles and proptosis in most. While useful animal models of Graves’ hyperthyroidism have long been available [53], this is the first demonstrating convincing eye changes similar to those seen in human GO. It is likely that the chronic stimulation of TSHR accomplished in this model, as well as the unique electroporation protocol, allowed for induction of GO in these animals.

The Potential Role of IGF-1 Receptor

The insulin-like growth factor-1 receptor (IGF-1R) has emerged as potentially involved in GO pathogenesis via interactions with TSHR [26,54]. Indeed IGF-1R is known to participate in crosstalk with several other receptors and signaling pathways [55]. This receptor, as well as the structurally related insulin receptor (IR), contains 2 α- and 2 β-chains that form a heterotetrameric structure linked by covalent disulfide bridges [56,57]. Binding of either IGF-1 or insulin initiates dimerization, autophosphorylation, and activation of the receptor complex. Phosphorylation of tyrosine substrates follows and leads to binding and activation of the regulatory subunit of PI3K with subsequent stimulation of PI3K activity. The resulting elevated phosphatidylinositol 3,4,5-trisphosphate levels function to phosphorylate and activate Akt at the cell membrane. The PI3K/Akt pathway has been shown to mediate many different cellular functions, including cell survival [58] and the promotion of adipogenesis [59].

Early studies demonstrated IGF-1 binding in orbital cells from patients with GO and not in normal orbital cells [60,61], and later studies showed expression of IGF-1R to be greater in GO orbital fibroblasts than in normal orbital cells [62]. Smith and colleagues treated GO orbital fibroblasts with IGF-1, human recombinant TSH (hrTSH), or immunoglobulins from patients with Graves’ disease (GD-IgG) and studied both HA synthesis [63] and T cell chemotactant expression [62]. While they found GD-IgG to be a potent inducer of both HA synthesis and chemokine expression, hrTSH had no effect in these cells. They additionally showed neutralization of GD-IgG by a potent IGF-1R blocker (termed 1H-7) or by transfection of fibroblasts with a dominant negative mutant IGF-1R [64]. They concluded that HA synthesis and chemokine expression in these cells might be augmented by autoantibodies specifically targeting IGF-1R and that ligation of this receptor might mediate some TSH-induced signaling. Our laboratory performed studies in which GO orbital fibroblasts treated with bovine TSH (potentially more potent than hrTSH), M22 or IGF-1, with or without co-treatment with 1H7. We found that cAMP production, phosphorylation of Akt and HA synthesis stimulated by each of these agents can be abrogated by co-treatment with 1H7, suggesting that HA syn-
thesis induced by TSHR activation in these cells may involve IGF-1R signaling.

Moshkelgosha and colleagues used their animal model of GO (described above) to address the potential role of IGF-1R as an autoantigen in GO [52]. However, rather than immunizing animals with TSHR cDNA, they used IGF-1Rα plasmid. They found that while these mice generated high levels of IGF-1Rα antibody, they developed no apparent ocular or other pathology. However, some of the animals immunized with TSHR A subunit alone developed low-titer IGF-1Rα antibodies shortly after immunization. How these secondary antibodies are produced and whether they might impact TSHR signaling in orbital fibroblasts is unknown.

Recent studies by Krieger and colleagues suggest that IGF-1R and TSHR increase the production of HA by GO orbital fibroblasts in a synergistic fashion [65]. These investigators showed progressively higher levels of HA production by TSH-treated cells co-treated with increasing IGF-1 doses. In addition, the dose-dependent stimulation of HA secretion by M22 was shown to be biphasic with the higher potency phase mediated in part via IGF-1R signaling. These data provide evidence for the existence of cross-talk between TSHR and IGF-1R in GO orbital fibroblasts induced by TSHR activation. That these receptors might form physical complexes has been suggested in studies demonstrating immunoprecipitation of both receptors from GO orbital fibroblast or thyrocyte preparations using specific monoclonal antibodies directed against either receptor and showing their co-localization in the perinuclear and cytoplasmic compartments of these cells [60].

Despite attempts by several laboratories, antibodies that specifically target IGF-1R have not been found to be more prevalent or elevated in GO than in normal sera, making it unlikely that such antibodies play a role in GO pathogenesis [66]. However, it is possible that TSHR signaling pathways within the GO orbit are influenced by locally produced IGF-1 acting in an autocrine or paracrine fashion and/or that downstream signaling pathways common to both TSHR and IGF-1R are engaged following TSHR activation [54].

**Therapeutic Implications**

Evidence presented in this review concerning TSHR expression and function in GO orbital cells and cross-talk between this receptor and IGF-1R suggests potential therapeutic avenues involving receptor antagonists. Small molecular ligands that inhibit stimulated as well as constitutive TSHR signaling have been developed via molecular modeling and functional experiments [67]. These allosteric modulators sit within the transmembrane helices of TSHR and prevent contact with intracellular residues essential for agonist activity. As such, their action does not involve competition with extracellular TSH or blockade of TRAb binding sites. Neumann and colleagues found TRAb-stimulated thyroid function in female mice to be reduced following treatment with a small molecule TSHR antagonist [68]. Further, as discussed above, a particular TSHR antagonist, termed C-1, inhibits TSH- and TRAb-mediated HA secretion in GO orbital cells [47]. These early studies suggest that small molecule TSHR antagonists might be useful in the treatment of Graves’ hyperthyroidism and ophthalmopathy, and could potentially be used to prevent the development of GO in patients with Graves’ disease. Because these compounds can be produced in large quantities and are not degraded in the GI tract, oral administration may be possible. Another antireceptor approach, the blocking of IGF-1R with teprotumumab, is currently under investigation in a multi-center trial (NCT01868997). Based on current concepts of GO pathogenesis, combined anti-TSHR and anti-IGF-1 therapy might in future prove particularly effective.

Multiple studies implicate both Th-1–type and macrophage-derived cytokines in early GO pathogenesis [1, 26]. Therefore, mononuclear antibodies that target proinflammatory cytokines and chemokines might be of therapeutic benefit in the disease. Specifically, biological agents that block IL-1 receptor (anakinra), IL-6 (tocilizumab), TGF-β (Iberelimumab, GC1008), or TNF-α (infliximab, adalimumab, or etanercept) are attractive as potential therapy. Case reports have suggested that rituximab, a monoclonal antibody directed against B cell surface CD20, might be useful therapy for patients with active GO. However, 2 recently completed randomized controlled trials showed disparate results, indicating that the agent is not uniformly effective [69, 70]. Because GO duration was shorter in the study showing benefit (mean duration 4 months vs. 12 months), it is possible that a subgroup of patients with short disease duration might benefit from the therapy. Other approaches might include modulation of co-stimulatory pathways with agents directed against cytotoxic T-lymphocyte-associated protein (CTLA)-4 (abatacept) or reduction of inflammation and potentially orbital adipogenesis with novel selective peroxisome proliferator-activated receptor (PPAR)-γ antagonists.

**Summary**

Environmental, genetic and immune factors are at play in the development of the variable clinical manifestations of GO. Both cellular and humeral immunity directed against TSHR expressed on orbital fibroblasts likely initiate the disease process (Fig. 1).
Activation of helper T cells recognizing TSHR peptides presented by CD40+ cells and ligation of TSHR by TRAb lead to the secretion of inflammatory cytokines and chemokines by activated T cells and orbital fibroblasts, and enhanced HA production and adipogenesis in the latter. The resulting connective tissue remodeling results in varying degrees extracellular enlargement and orbital fat expansion. A subset of orbital fibroblasts express CD34, bone-marrow derived, and circulate as fibrocytes that infiltrate connective tissues at sites of injury or inflammation. As these cells express high levels of TSHR and are blasts express CD34, are bone-marrow derived, and circulate as enlargement and orbital fat expansion. A subset of orbital fibroblasts express CD34, bone-marrow derived, and circulate as inflammation. Curr Opin Pharmacol 2012; 12: 491–496

Conflict of Interest ▼

The authors declare no conflict of interest.

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