Bony Orbital Surgery for Graves’ Ophthalmopathy

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Abstract

The majority of Graves’ ophthalmopathy, or thyroid eye disease, can be managed medically; however, in refractory or severe cases, surgical intervention with orbital decompression may be necessary. The majority of the published literature is retrospective in nature, and there is no standardized approach to orbital decompression. The purpose of this review is to evaluate the various surgical approaches and techniques for orbital decompression. Outcomes are ultimately dependent on individual patient factors, surgical approach, and surgeon experience.

Keywords

► orbital decompression
► thyroid eye disease
► Graves’ ophthalmopathy

Graves’ ophthalmopathy, or thyroid eye disease (TED), is an autoimmune condition which can affect up to 50% of patients with Graves’ disease, and causes both cosmetic disfigurement and functional deficits. It can lead to exophthalmos, eyelid retraction and exposure keratopathy, orbital pain and pressure, diplopia, and even vision loss secondary to compressive optic neuropathy with significant impact on quality of life. Important consideration in management of patients with TED is medical management of the hyperthyroidism and comorbid cigarette dependence. Cigarette smoking has been found in many studies to increase the risk of developing TED. Patients with Graves’ disease should be counseled about tobacco cessation in regards to both the risk of developing TED and the other well-known negative health effects of cigarette smoking. Patients with TED should also be informed that tobacco cessation may decrease the progression of the disease. The first-line treatment of Graves’ disease is antithyroid drugs, while radioactive iodine and total thyroidectomy are reserved for refractory cases. It is recommended that patients who are treated with radioactive iodine also undergo prophylactic oral steroid treatment to help mitigate the risk of TED development. However, in patients presenting with TED, thyroidectomy and radioactive iodine have not been shown to affect the natural course of TED. Initial treatment during the acute phase of TED includes supportive medical management, and typically includes topical lubrication, head elevation, corticosteroids, and orbital radiotherapy. If patients develop acute vision-threatening orbital compression or enter remission and seek to restore normal globe position, surgical intervention is considered. The purpose of this manuscript is to review the role of bony orbital decompression in the management of TED.

Pathophysiology of Thyroid Eye Disease

The pathophysiology of TED is not fully understood; however, it is speculated to be caused by autoantibodies to the orbital connective tissue. Thyroid stimulating hormone receptor autoantibodies, also known as long-acting-thyroid stimulators (LATS), have been shown to be involved in the etiology of Graves’ disease and are thought to be responsible for the development of hyperthyroidism. A correlation between severity of elevation of these LATS and the development of TED has been described. There are two phases to the disease: an active phase and quiescent phase. During the active phase, lymphocytes infiltrate the orbital tissue with subsequent fibroblast activation, release of proinflammatory mediators, and subsequent fibrosis.

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cytokines, and deposition of glycosaminoglycans. There is increased production of hyaluronan, which accumulates within the orbit leading to edema and orbital soft-tissue volume expansion. The orbital fibroblasts may then differentiate into adipocytes or myofibroblasts and may explain the heterogeneity in clinical presentation. Ultimately, fibrosis of the extraocular muscles and fat hypertrophy occurs. The active phase waxes and wanes for many years, and is followed by a quiescent phase with no further progression of disease. Rarely, patients may experience flare-ups of their disease burden with further inflammation.

**Indications for Surgical Decompression**

In the majority of patients with TED, conservative management is sufficient to alleviate symptoms and surgical intervention is avoided; however, approximately 20% of patients will undergo surgical intervention. Indications for urgent orbital decompression include visual loss from compressive optic neuropathy or corneal exposure. Delayed decompression usually occurs after a euthyroid state is achieved and an interval of 6 months has passed assuring surgical intervention during the quiescent phase of disease. Indications for delayed decompression include asymmetric or severe proptosis, moderate-to-severe extraocular muscle hypertrophy, diplopia, ocular pain, and ocular hypertension. Orbital decompression should take place prior to any strabismus and/or eyelid surgery.

**Surgical Approaches to Bony Decompression of the Orbit**

Bony orbital decompression is performed in the operating room under general anesthesia. Intravenous antibiotics and corticosteroids are administered preoperatively, and depending on the surgical approach, surgical navigation can be set up at this time. After the induction of general anesthesia, local anesthesia based on surgeon’s preference is administered at the planned incisions. There are four bony boundaries of the orbit; however, only three are routinely decompressed: the orbital floor, medial wall, and lateral wall. Depending on the degree of decompression needed, a single wall or a combination of walls can be decompressed. Most commonly, the medial orbital wall is addressed first, followed by the orbital floor. If additional decompression is needed, then the lateral wall is decompressed. The patient is reassessed by ophthalmology, and if further decompression is needed, albeit rarely used, the orbital roof is decompressed. The surgical approach for each wall is discussed below.

**Medial Orbital Wall**

The medial orbital wall can be approached in one of two ways: transorbital or transnasal. The transorbital approaches include a transcutaneous approach using a Lynch incision and a transcaruncular approach; however, the transnasal endoscopic approach has become the most widely utilized approach to the medial orbital wall due to its many advantages. The transnasal endoscopic approach has garnered more attention recently as it provides a direct approach to the medial wall with the advantage of no external incisions. This approach is typically performed by otolaryngologists with experience in endoscopic anterior skull base surgery. In addition, the introduction of surgical navigation technology in endoscopic endonasal surgery provides enhanced efficacy and safety. The nasal mucosa is decongested, and local anesthetic is infiltrated. After medialization of the middle turbinate, uncinectomy is performed. A total ethmoidectomy is completed to identify the lamina papyracea and create space for the orbital contents after decompression. Orbital decompression is performed by removal of the entire lamina papyracea with incision into the periorbita such that the orbital contents are herniated into the ethmoid sinuses. The periorbita is incised with a sickle knife beginning posteriorly and extending the incision anteriorly. The endoscopic transnasal approach provides better visualization and safer access to the posterior medial wall for

![Fig. 1](https://example.com/fig1.png)

**Fig. 1** Endoscopic approach to the medial orbital wall. (A) After ethmoidectomy, the lamina papyracea is removed to expose the periorbita. The periorbita is incised. (B) After incision of the periorbita, the orbital fat is prolapsed into the middle meatus. Care is taken to prevent injury to the medial rectus.
decompression. If needed, the optic nerve can be decompressed via this technique. If the optic nerve is to be decompressed, a wide sphenoidotomy is performed and the fovea ethmoidalis is identified posteriorly. Within the sphenoid, the carotid artery and optic nerve are identified. The thick bony overlying the junction between the orbital apex and the sphenoid sinus, also known as the optic tubercle, is then removed. This bone is thick and requires the use of a diamond drill. The bone overlying the optic nerve within the sphenoid sinus is then removed, and this bone is thinner and can be removed with a Freer elevator. Once the bone has been removed off the orbital apex and optic nerve, the optic nerve sheath is then incised keeping in mind the ophthalmic artery is found in the posteroinferior quadrant of the nerve. Complications of the transnasal endoscopic approach are low, and are minimized by intimate endoscopic anatomical knowledge and experience with the technique.\textsuperscript{15} Care is taken to avoid injury to the anterior and posterior ethmoid bundle which can retract into the orbit and cause a retrobulbar hematoma. Additionally, during incision into the periorbita, the medial rectus is identified as injury to this structure can cause diplopia.

**Orbital Floor**

The orbital floor can be approached transorbitally using a transconjunctival incision and transantrally using the Caldwell–Luc approach.

The transconjunctival approach utilizes an incision in the palpebral conjunctiva of the lower eyelid inferior to the tarsus with two potential routes: retroseptal or preseptal. The retroseptal approach provides direct access to the infraorbital rim and after incision of the periorbita, exposure of the orbital floor is performed in usual fashion. The preseptal approach begins with eversion of the lower eyelid to identify the tarsal plate. The incision is made below the tarsus and the preseptal space is entered. With direct visualization of the septum, blunt dissection is carried down to the infraorbital rim, and the orbital floor is exposed as previously described. The transconjunctival approach can also be combined with a lateral canthotomy with inferior cantholysis, termed the “swinging eyelid” approach, to visualize both the orbital floor and lateral orbital wall, and has been described extensively for orbital decompression.\textsuperscript{16–18}

The transantral approach avoids a lower eyelid incision, and affords widest exposure to the orbital floor (\textsuperscript{\textbullet} Fig. 2).\textsuperscript{19} The maxillary sinus is accessed via a Caldwell–Luc approach. An incision is made in the gingivobuccal sulcus above the canine fossa and carried down to the anterior maxillary wall. The periosteum is elevated and the infraorbital nerve is identified. After entering the maxillary sinus, the mucosa is stripped and the orbital floor can be removed. The peri-orbita is incised away from the infraorbital nerve and allows the periorbital fat to herniate into the maxillary sinus. The anterior ethmoid air cells can also be removed to expose the medial orbital wall through this approach. This approach includes a risk of paresthesia and anesthesia due to injury to the infraorbital nerve, oroantral fistula, and devitalization of maxillary teeth.

Regardless of approach, decompression of the orbital floor is limited to removal of the bone medial to the infraorbital nerve, which limits the risk of paresthesia and anesthesia of the cheek and upper lip. In addition, a strut of bone is left anteriorly to minimize the risk of hypoglobus.

**Lateral Wall**

The lateral wall is typically approached by transcutaneous incisions, including the lateral eyelid crease and lateral transconjunctival incisions with inferior cantholysis (\textsuperscript{\textbullet} Fig. 3). The “swinging eyelid” approach may also be utilized if the orbital floor is to be decompressed as well. The lateral one-third of an upper blepharoplasty incision is made and dissection down to the lateral orbital rim is performed. Cantholysis is then
performed, and with a cutting burr, the lateral orbital rim is excised and placed into saline. The lateral periorbita is bluntly dissected away from the lateral wall and the orbital contents are retracted medially. The orbit is expanded laterally by drilling the lateral orbital wall until the temporalis muscle is exposed. The dissection is taken posteriorly through the greater wing of the sphenoid bone into the marrow space with or without exposure of the dura. After adequate bone removal, the periorbita is incised above and below the lateral rectus to allow the orbital contents to herniate into the expanded bony space. The lateral orbital rim is then replaced, secured with miniplates, and canthoplasty is performed to resuspend the lateral canthal tendon. Complications of this approach include cerebrospinal fluid leak secondary to dural violation, anesthesia secondary to injury to the zygomaticotemporal and zygomaticofacial neurovascular bundles, and oscillopsia.

**Orbital Roof**

A superior blepharoplasty or sub-brow incision is used to approach the orbital roof (►Fig. 4). Soft tissue dissection is then carried forth to the superior orbital rim, taking care to preserve the supraorbital neurovascular bundle. A subperi orbital dissection is then carried forth along the orbital roof, and the greater wing of the sphenoid and frontal bone posterior to the frontal sinus can be removed with Freer or number nine periosteal elevator. The dura is identified and preserved during this procedure. Like the other approaches, the periorbita is incised and orbital fat can herniate the expanded bone region.

**Outcomes**

The primary goal of delayed orbital decompression for TED is improvement in proptosis. The choice in which wall to decompress is both individualized to the patient and dependent on surgeon comfort with technique. Likewise, the degree of decompression is dependent on surgical technique, and can be enhanced with the addition of orbital fat decompression. Single-wall decompression rarely occurs in the contemporary management of TED due to the limited reduction in proptosis compared with two-wall or three-wall decompression. If a single-wall approach is chosen, some have advocated for a deep lateral wall decompression which can reduce proptosis by 4 to 6 mm.20–22 Jefferis et al compared both their personal experience with orbital decompression and a literature review on the different approaches. Their study reported a 4.2 mm mean reduction with single-wall and between 2.9 and 7.6 mm mean reduction for a combination of two- or three-wall decompression.23 The literature review revealed a mean single-wall decompression of 3.6 to 4.8 mm and a mean two- or three-wall decompression of 2.5 to 8.0 mm. When a two-wall
approach is chosen, surgeons may decompress the medial wall and orbital floor, termed an “inferomedial decompression,” or a “balanced decompression” involving the medial and lateral orbital walls. Thapa et al showed more effective proptosis reduction with the inferomedial decompression alone.\textsuperscript{24} If inferomedial decompression is performed, the inferomedial orbital strut should be preserved to maintain the position of the eye.\textsuperscript{25} However, many oculoplastic surgeons have moved away from inferomedial decompression in favor of the “balanced decompression” approach.\textsuperscript{26}

In recent years, there has been an increased focus on the effect of therapeutic interventions on patient-reported outcome measures. TED has a significant impact on patient quality of life, and Wickwar et al have shown approximately 26% of patients with TED having signs of clinical depression.\textsuperscript{27} A disease-specific quality-of-life questionnaire has been developed for TED called the Graves’ Ophthalmopathy Quality of Life (GO-QOL) scale. With regard to orbital decompression, two studies have shown a significant impact on visual function and appearance.\textsuperscript{28,29}

**Special Considerations**

**Balanced Decompression**
The “balanced decompression” involves removal of the lateral wall and medial wall while maintaining an intact orbital floor, and has become a favored technique among oculoplastic surgeons.\textsuperscript{26} The purpose is to reduce the asymmetric shift in orbital contents and thus reduce the rate of complications such as hypoglobus, new-onset diplopia, and muscle imbalance.\textsuperscript{30} Studies have shown a reduction in proptosis of approximately 4 to 6 mm.\textsuperscript{10,31,32} Additional decompression can be achieved with removal of fat.

Unal et al reported an effective reduction in proptosis with a balanced decompression with a reduced incidence of postoperative diplopia when compared with a three-wall decompression.\textsuperscript{33} However, Goldberg et al reported a lower incidence of strabismus with a lateral orbital-wall decompression compared with a balanced medial and lateral orbital-wall decompression.\textsuperscript{34} The conflicting results suggest that outcomes are likely surgeon-specific and technique-dependent.

**Two-Wall versus Three-Wall Decompression**

With various combinations of decompression options and preponderance of retrospective studies, there remains debate about the best approach for reducing proptosis among TED patients. Many authors report a significant improvement in proptosis reduction of approximately 7 to 8 mm with a three-wall decompression.\textsuperscript{35,36} However, three-wall decompression is more often associated with postoperative complications such as new-onset or worsening diplopia.\textsuperscript{33,36,37} Given the potential for increased incidence of diplopia when compared with two-wall decompression, three-wall decompression is typically reserved for patients with severe proptosis.

**Future Directions**

In the past, there was no specific Food and Drug Administration (FDA)-approved medical therapy for TED. Initial therapy was directed toward symptomatic management using corticosteroids, and when refractory, patients proceeded with orbital decompression. In January 2020, a monoclonal antibody to insulin-like growth factor I receptor, teprotumumab, received FDA approval specifically for the treatment of TED. In a phase 3 multicenter randomized control trial, Douglas et al showed reduction in proptosis of 2.82 mm in patients with active TED treated with teprotumumab compared with placebo as well as a significant improvement in GO-QOL.\textsuperscript{38} The introduction of this novel therapy will likely result in a major shift in the medical and surgical management of TED with fewer patients necessitating surgical intervention in the future.

**Conclusion**

TED refractory to medical management can be managed surgically with orbital decompression resulting in improved cosmesis and potential return to baseline function. There are multiple surgical techniques for orbital decompression, and the approach can be individualized based on patient characteristics and surgeon experience. Intimate knowledge of orbital anatomy and careful surgical technique can minimize complications.

**Note**
The work herein does not necessarily represent the views of the United States government, Department of Defense or its affiliates.

**Conflict of Interest**
None declared.

**References**

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