Multimodality imaging of the orbit

Pradipta C Hande, Inder Talwar
Department of Radiodiagnosis and Imaging, INHS Asvini, Colaba, †Department of Radiodiagnosis, Bombay Hospital Institute of Medical Sciences, Marine Lines, Mumbai, India

Correspondence: Dr. Pradipta C Hande, Department of Radiodiagnosis and Imaging, INHS Asvini, Colaba, Mumbai - 400 005, India.
E-mail: pradipta.hande@gmail.com

Abstract
The role of imaging is well established in the evaluation of orbital diseases. Ultrasonography, Computed tomography and Magnetic resonance imaging are complementary modalities, which allow direct visualization of regional anatomy, accurate localization and help to characterize lesions to make a reliable radiological diagnosis. The purpose of this pictorial essay is to highlight the imaging features of commonly encountered pathologies which involve the orbit.

Key words: Imaging; magnetic resonance imaging; multidetector computed tomography; orbit

Introduction
The orbit is the site of a large number of pathologies of diverse etiologies, and imaging has to be tailored to the symptoms and clinical findings. The commonest clinical indications for orbital imaging are proptosis/exophthalmos, diminished vision, enophthalmos, diplopia, leucocoria, pain, tumor and trauma. It is also used for evaluation in craniofacial developmental anomalies and epiphora.

Relevant anatomy
Good knowledge of detailed anatomy of the orbits and its contents is essential to evaluate orbital pathologies. The orbits are pyramid shaped bony cavities with the base anteriorly (orbital opening). The rigid bony orbital walls are formed by seven bones namely frontal, sphenoid, ethmoid, lacrimal, maxilla, zygoma, and the palatine. The walls of the orbit are lined by the periosteum (periorbita) with potential extraperiosteal and subperiosteal spaces.

The orbits are closely related anatomically to the adjacent paranasal sinuses, pterygopalatine fossa, and cerebral contents making them vulnerable to extension of disease processes involving these structures.

Imaging considerations
The inherent soft tissue contrast of the fluid-filled globe, retrobulbar contents within the fat and bony walls of the orbit is separated from the ethmoidal air cells by a very thin orbital plate of ethmoid (lamina papyracea). The walls of the orbit are vulnerable to fractures due to head injuries and facial trauma.

The main contents of the orbit include globe/eyeball, seven extraocular muscles (EOM), blood vessels, cranial nerves (II with meninges, III, IV, V, VI), sympathetic and parasympathetic nerves, lacrimal gland and apparatus surrounded by fat and connective tissue. The optic canal, superior and inferior orbital fissure are apertures that transmit vital nerves and vessels to and from the orbit and brain at the orbital apex. The tendinous ring (annulus) of Zinn is the common origin of the EOM including the–medial rectus (MR), lateral rectus (LR), superior rectus (SR) except inferior rectus (IR). The four recti attach anteriorly onto the globe forming a muscle cone tapering towards the apex, which divides the orbit into intraconal, conal, extraconal compartments and are well identified on imaging. This descriptive approach is useful in localizing the lesions of the orbit to a specific compartment, thereby helping in the diagnosis and further planning the surgery. The orbital walls lined by the periosteum (periorbita) have potential extraperiosteal and subperiosteal spaces.
orbits delineates the orbital anatomy well [Figures 1 and 2]. Ultrasonography USG (B-scan) remains a popular and easily available modality for examination of ocular abnormalities[4] and is a cost effective diagnostic tool without harmful effects of ionizing radiation to the lens. 7.5-13 MHz transducers give high resolution images for visualization of the anterior and posterior segments of the eyeball. It has the added advantage of real time imaging and allows dynamic scanning synchronized with ocular movements while performing the scan. The role of USG is well established for vitreal, choroidal, and retinal abnormalities[4] and also in cases of ocular trauma. Foreign bodies (FB) within the globe, that are not radio-opaque or suspected ferromagnetic metallic FB, can be visualized easily and localized with USG. The use of Doppler has enhanced its utility in evaluation of the vessels and also helps in assessing vascularity of lesions. The main limitation of USG for orbits is its inability to image the bony architecture, lack of detailed visualization of orbital apex and for intracranial extension of various pathologies.[1]

Computed tomography (CT) and Magnetic resonance (MR) are used widely in imaging of the orbits and each has its advantages and pitfalls. The protocols for CT/(MR) should be planned as per clinical indication. Intravenous (IV) contrast in recommended doses should be used as required to evaluate enhancement.

Routine CT exams are obtained parallel to the orbitomeatal line (OML) with 3 mm contiguous axial sections, with thinner slices up to 1-2 mm, when required. Direct 2-3 mm thin coronal sections perpendicular to OML may be required. Multidetector CT (MDCT) has the advantage of rapid acquisition of volumetric dataset (isotropic imaging) allowing excellent multiplanar reconstruction (MPR) images. Non-ionic iodinated contrast (300 mg/ml) is used in appropriate doses to assess the tissue enhancement patterns in many orbital conditions.

Radiation dose to the lens (at least 50 mGy or more) needs to be remembered. Proptosis can be objectively assessed on axial CT[1]. Normally, at least one third of the eyeball lies posterior to the line joining lateral orbital margins in axial scans [Figure 1A, blue line].[1] MR images obtained using head coil on a high field strength magnet (1.5-3.0 T) enables adequate visualization at orbital apex and beyond. Short scan times with turbo/fast spin echo (SE) sequences, T1W, T2W without and with fat suppression (FS) techniques like short tau inversion recovery (STIR) images. Thin (3 mm or less) slice thickness with acquisition in at least two scan planes, small field of view (FOV), high resolution matrix (256 x 256) and T2W imaging of the brain for evaluation of the entire visual pathways. Gadolinium (Gd) enhancement is seen as high intensity in post-contrast FS T1W images. Additional sequences with Gradient echo images (GRE) can be acquired depending on the indication.

Inflammatory diseases
Infections/cellulitis
Bacterial infections are amongst the commonest pathologies constituting nearly 60% of primary orbital pathologies,[1,5] majority of which originate from the paranasal sinuses. It is the most common cause of proptosis in children.[6] CT and MRI can demonstrate the various stages of orbital cellulitis[7] [Figure 3A and B]: Edema, (not a stage of orbital cellulitis) phlegmon and abscess formation [Figure 4]. Complications like cavernous sinus thromboses and intracranial extension[7] can be detected early.

Orbital fungal infections like mucormycosis or aspergillosis may follow an aggressive course in patients with diabetes
Hande and Talwar: Multimodality imaging of orbit

mellitus or an immunocompromised status\[1\] sometimes resulting in intracranial extension [Figure 5].

**Idiopathic non-granulomatous inflammatory syndrome:**
Patients with Inflammatory orbital syndrome (IOS) (also called as orbital pseudotumour) present with acute or sub acute pain\[1\] and swelling associated with restricted eye movements and may sometimes be bilateral.\[8\] There is usually no local or systemic cause found.\[8, 9\] The common types of IOS are:

- Anterior orbital group which is characterized by uveal-scleral thickening and inflammation at optic nerve junction with enhancement on CT/MR. It needs to be differentiated from orbital cellulitis or leukemic infiltration.\[9\]
- Diffuse type represents a tumefactive or infiltrative process, involving retrobulbar space without bony erosion. A differential diagnosis of lymphoma should be considered.\[9\]
- Myositic type of IOS is an inflammation of the extra ocular muscles. CT/MRI show enlarged muscles with shaggy margins, typically involving the tendons up to the insertion with enhancement and obliteration of peripheral surgical fat planes, usually affecting the superior group [Figure 6] and medial rectus and should be differentiated from thyroid myopathy.\[9\]
- Lacrimal gland inflammation\[1\] has a similar imaging appearances sarcoid or lymphoma of the lacrimal gland.
- Orbital apex inflammation\[1\] infiltrating posteriorly into optic nerve sheath complex (ONSC) or posterior EOM involvement.
- Tolosa-Hunt syndrome\[1\], which is characterized by painful external ophthalmoplegia [Figure 7].

**Parasitic infestations**
Parasitic infestations are common in the tropical countries and may be related to certain occupations.

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**Figure 3A:** Orbital cellulitis, left. Axial post-contrast computed tomography sections (a and b) show eyelid edema (asterisk), enhancing thickened sclera, extraocular muscles (arrow) with sinusitis causing proptosis. Coronal (c), volume rendered (d) images shows boggy collection (phlegmon) (curved arrow) with displacement of the globe

**Figure 4 (A-C):** T2W coronal magnetic resonance images (A), post-contrast T1W FS axial, coronal images (B and C) shows left orbital cellulitis with subperiosteal abscess (asterisk) with bilateral pansinusitis (arrow)

**Figure 3B:** Magnetic resonance T2W axial images (a and b) show left orbital cellulitis diffusely involving preseptal and postseptal tissues with myositis (curved arrow) extending upto the orbital apex (asterisk) and associated sinusitis (arrow)

**Figure 5 (A-C):** Mucormycosis in human immunodeficiency virus (HIV) positive patient with left proptosis. Coronal T1WI (A) show hypointense areas (curved arrow) within medial aspect of hyperintense orbital fat. Signal void within ethmoid air cells (apparent aerated sinus) is misleading. Post-contrast T1WI FS (B and C) show enhancement of medial rectus (arrow) and enhancing sinonasal mucosa (asterisk) suggesting inflammation
and environmental conditions. They are endemic in regions with poor hygiene and sanitation as human infestation occurs due to ingestion of ova in food or water. Larval forms may be found as cysts within orbital tissues which may be detected by imaging. [Figure 8] Cysticercosis may involve EOM and can be associated with neurocysticercosis [Figure 9]. The differentials on imaging are other granulomatous orbital lesions [Figure 10] e.g., tuberculomas. Toxocara infestation may cause inflammation (endophthalmitis) when they die, which can present with leucocoria in children.

Graves’ dysthyroid ophthalmopathy/graves’ orbitopathy (GO)
Graves ophthalmopathy is one of the commonest causes of exophthalmos in adults and is almost always bilateral although asymmetric involvement may be seen in few. It is presumed to be an autoimmune condition with a female preponderance consisting of a triad of goiter with hyperthyroidism, infiltrative ophthalmopathy, and dermopathy. CT/MRI can be used to assess and follow-up the changes in response to steroid and/or immunomodulatory therapy. EOM are enlarged with sparing of tendons. The muscle margins are sharply defined with preserved fat planes. There is increased volume of retro-orbital fat with bulging of orbital septum anteriorly [Figure 11]. Involvement of the inferior rectus precedes involvement of the superior and the medial rectus.[9] In cases with isolated lateral rectus involvement, other differentials need to be considered. CT may show low density within the muscles due to deposition of lymphocytes and mucopolysaccharides. MRI reveals T2 hyper intensity due to infiltration and interstitial edema of involved EOM in active disease which can be easily demonstrated in T2W inversion recovery sequences (TIRM) with FS.[10] The cross-sectional area of the most inflamed EOM muscle belly can be measured on 3 mm coronal T2W IR fat suppressed sequences EOM enlargement can be easily measured as its maximum radial diameter of the belly of the involved muscle in coronal sections.[11] This can be used for objective follow-up to assess response to therapy.

Figure 6 (A-D): Orbital pseudotumor. Axial and coronal T2W (A and C) images show minimal hyperintensity of globe with uveo-scleral thickening (curved arrow) (Tenon’s fasciitis), inflammation (arrow) of muscle and surrounding orbital fat with enhancement (asterisk) in post-contrast T1WI FS (B and D)

Figure 7 (A, B): Tolosa-Hunt syndrome subtype. T1W FS post-contrast gadolinium coronal (A), axial (B) shows enhancing inflammatory infiltration in the left orbital apex (arrow) extending into the cavernous sinus (curved arrow)

Figure 8 (A, B): (A, B) Swelling medial aspect of left orbit. High resolution USG left orbit shows linear, serpiginous contents within extraocular cystic lesion (arrow) with distal enhancement medial to globe. Realtime slow movements of these structures were seen during scanning. Live adult filarial worms were seen on surgery.

Figure 9 (A, B): Cysticercosis. Magnetic resonance T2W sagittal sections show multiple hyperintense cysticerci with hypointense eccentric scolecides in the orbit within (A) optic nerve (arrow) and inferior rectus muscle (curved arrow) and (B) superior rectus muscle (curved arrow). Note heavy infestation of brain and other soft tissues.
with steroids. In chronic disease, atrophy of EOM may be seen with fat replacement which can be detected on T1W images without FS.

There may be associated compressive optic neuropathy probably as a result of pressure at the orbital apex by increased volume of orbital contents, overcrowding and stretching of the nerve due to proptosis.\cite{10} High-resolution volume acquisition (T1W-3D) with curved multiplanar reformatting can be used to measure the optic nerve diameter along its entire length.\cite{11} Decompression surgery may prevent irreversible optic nerve atrophy in early stage of disease.

**Orbital trauma**

CT is the modality of choice in evaluation of orbital trauma.\cite{1} Bony fractures and displacements, small comminuted fragments [Figure 12], and radio-opaque foreign body (FB) are best seen on CT. Volume rendering techniques (VRT) with MDCT allow excellent 3D reconstructions that help surgeons to visualize the fractures in relation to other anatomic structures and evaluate treatment plans\cite{12} [Figure 13]. Blowout fractures classically involve the floor with sparing of the orbital rim. The orbital contents including the orbital fat, inferior rectus and the inferior oblique muscles may herniated through the floor of the orbit into the maxillary antrum due to increased intra orbital pressure.\cite{13} [Figure 14A]. Soft tissue injury, inflammation, muscle herniation, and entrapment [Figure 14B] are well visualized with CT/MRI.
on axial and coronal sections. Associated vascular and nerve injury, and intracranial hematomas[13] are better evaluated with MRI [Figure 15]. However, MR is contraindicated if there is suspicion of intraocular ferromagnetic FB.

**Tumors**
The orbit may be involved by tumors which are primary masses arising from the various orbital structures.

**Lymphoid neoplasms**
These range from the benign reactive lymphoid hyperplasia to malignant lymphomas and are more commonly seen in adults and very rarely in children.[8] All such patients should undergo evaluation to rule out a systemic lymphoproliferative disorder.[3] Malignant lymphoma, either arise in and are limited to the orbit or extend into the orbit arising from the sino-nasal cavities.[6] Imaging is not specific in differentiating them from pseudotumors. CT and MR reveal enhancing, solid soft tissue masses in the retrobulbar region [Figure 16] without bony destruction, except rarely in very aggressive malignant variants.[13] Lacrimal gland lymphoma shows enhancement and enlargement with associated displacement of the globe medially.[14] MR is more sensitive in visualizing early infiltration of ocular structures.

**Leukemias**
Orbital involvement is due to infiltration of soft tissue or orbital bone, usually in Acute lymphoblastic leukemia (ALL) (up to 75%)[15] and 20% in Acute myeloid leukemia (AML) (Chloromas).[15] CT/MRI may reveal a sub-periosteal lesion [Figure 17] involving the lateral wall of the orbit, which may extend into the temporal fossa or a mass on the medial wall involving ethmoid sinuses extending through the cribriform plate into the anterior cranial fossa.[1] Meningeal infiltration and intracranial extension can be detected early in post-contrast scans.

Vascular lesions represent a large group of vascular malformations involving the orbit.[16] MRI has the advantage over CT as it is very sensitive in detecting blood and blood products and vessels with flow.

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**Figure 14 (A, B):** T1W coronal image (A) shows classic blowout fracture of floor of right orbit with herniated contents into the roof of maxillary antrum (arrow). Axial computed tomography (B) shows herniation of medial rectus into adjacent ethmoid sinus (curved arrow) with fracture of right lamina papyracea.

**Figure 15 (A, B):** Subperiosteal hematoma in orbital trauma. Axial (A), coronal (B) T1WI shows subacute hyperintense hematoma right orbit with mass effect (arrow). Associated hemosinus within right maxillary sinus (asterisk) and subdural hematoma (curved arrow).

**Figure 16 (A, B):** Lymphoma left orbit in a case of unilateral proptosis. Axial magnetic resonance shows diffusely infiltrating lesion involving left retrobulbar region (asterisk), appearing hypointense to orbital fat on T1WI (A), hypointense on T2WI (B) with preseptal soft tissue thickening.

**Figure 17 (A, B):** Orbital leukemic deposits in patient with ALL. Axial Magnetic resonance short tau inversion recovery (A) shows bilateral subperiosteal lesions (asterisk) extending superior and laterally up to the lacrimal glands, hypointense to orbital fat extending posteriorly on sagittal T1WI (B).
Cavernous hemangioma (Venous angioma) is the commonest vascular tumor in adults presenting as a well-defined progressively increasing mass. It is usually seen on CT as a smoothly marginated, rounded enhancing soft tissue density intracranal mass that may have calcification [Figure 18]. MR T1W shows a hypointense mass, hyperintense on T2W with variable contrast enhancement [Figure 19]. Differentials of other intracranal masses like meningiomas, schwannomas or hemangiopericytoma need to be considered.[11,16]

Capillary hemangiomas present in infants within first year of life and then tend to diminish in size from the time of presentation due to gradual involution. They usually have definite arterial supply from either external/internal carotid artery,[16] and are composed of thin small vascular spaces due to capillary proliferation with no distinct capsule. CT/MR may show extraconal masses along the superior aspect of the orbit that may extend intracranially. These masses show intense enhancement in the early phase with early washout of the contrast in delayed phases.[11] On MR, they are usually hypointense on T1W images and have a heterogeneous appearance on T2 weighted images with evidence of flow voids within the lesion [Figure 20].

Other vascular lesions include lymphangioma [Figure 21], veno-lymphatic, and arteriovenous malformations.[1,16] Orbital varix present usually with intermittent proptosis and is a common cause of spontaneous orbital hemorrhage.[17]

Imaging performed in prone position can demonstrate a dilated venous channel which is usually hyperintense on both T1W and T2WI. Carotid-cavernous fistula causes pulsating exophthalmos, with engorgement of superior ophthalmic vein (SOV) which can be very well visualized on CT and MRI [Figure 22]. However, in the late stages there can be stenosis due to thrombosis of the SOV.

Optic nerve sheath complex
MR is the modality of choice for imaging of the entire visual pathway from its origin at the posterior point of the globe to the visual cortex.[1] Intraorbital optic nerve (ON) is seen well on CT due to the attenuation differences with surrounding hypodense retrobulbar fat. Intraorbital part of ON is hypointense on T1WI within hyperintense fat, isointense on T2WI with surrounding subarachnoid cerebrospinal fluid (CSF) within the meningeal sheath.
is hyperintense. The intracanalicular portion is not well visualized on CT due to the dense cortical bone at the apex and is displayed very well on MR. The intracranial ON, optic chiasma, and tracts are well visualized on coronal and sagittal MR and appear isointense to cerebral white matter. The optic radiation is within the white matter of temporoparietal and occipital lobes extending up to the calcarine cortex (primary visual cortex) flanked by the easily identified calcarine sulcus on the medial aspect of the occipital lobe.

Enlargement of ONSC can be due to primary or secondary tumors involving the ON or due to inflammatory/infiltrative processes.

Optic nerve glioma

Intraorbital gliomas appear early within first decade of life, presenting with painless proptosis, usually with associated loss of vision. CT/MR shows marked fusiform enlargement of ON with kinking or buckling. Calcification is usually absent. Bilateral lesions are seen in Neurofibromatosis (NF) Type 1, thus entire visual pathway should be evaluated. MR is more sensitive and 3mm thin sections are recommended which can avoid missing small lesions in the intracanalicular portion. Usually, it appears hypointense to orbital fat on T1WI (isointense to cerebral cortex), it is hyperintense to cortex, white matter and fat on T2WI. Post-contrast enhancement is common except in areas with cystic changes, [19-21] [Figure 23]. Sometimes, the tumor grows circumferential around the ON into the perineural space to mimic a perioptic meningioma [19] [Figure 24].

Optic nerve sheath meningioma

These are usually benign tumours arising from the meningo endothelial cells with a higher incidence in females. They commonly occur in the 4th-5th decade, however if they are found in children, a possibility of Neurofibromatoses type II needs to be kept in mind. The patterns usually seen on imaging are as an eccentric localized mass on one side of the optic nerve or circumferential along the length as thickening or fusiform enlargement of the ONSC. [19]
Intratumoral calcification is common and is well detected on CT. It may cause bony hyperostosis and optic canal enlargement\cite{19,20} [Figure 25]. It shows decreased signal intensity on both T1W and T2W MRI. Post-contrast enhancement is marked [Figure 26] and “tramtrack” type in circumferential type of lesions.\cite{1} Diagnosing intracanalicular meningiomas is a challenge\cite{22} and Gd-enhanced MRI with FS helps in detection of tiny lesions, especially at the orbital apex.\cite{23} In children, they tend to be more aggressive and may infiltrate into other orbital structures.

Optic neuritis
Acute inflammation causes enlargement of ON due to infections by micro-organisms causing papillitis, or following viral infections due to immune mechanisms (parainfections).\cite{1} Non-infective granulomatous diseases like sarcoidosis, systemic autoimmune diseases like systemic lupus erythematosus (SLE) may involve the ON.\cite{24} Optic neuritis is an early manifestation of multiple sclerosis (MS) which is seen well on MRI as thickening with focal hyperintense plaques on T2WI and better seen on STIR images\cite{25} [Figure 27]. FS T1WI pre-and post-contrast images are useful in optic neuritis and demonstrates localized or diffuse enhancement within the nerve [Figure 28]. Brain and spine MRI is usually required in combination while evaluating patients of demyelinating diseases such as MS.

The orbit may be secondarily involved due to invasion of malignant tumors from the surrounding structures, most commonly malignancy from sino-nasal cavities [Figure 29] or due to metastatic deposits from known or unknown primary.\cite{1} The most common primary cancers in adults are breast, prostate gland, and lung cancer. In about 10% of patients who present with ophthalmic symptoms due to secondary deposits, the primary site remains obscure in spite of detailed systemic evaluation. In the pediatric age group, neuroblastoma is one of the commonest primary tumors that metastasise to the orbits.

Diffusion weighted imaging (DWI) MRI with apparent
diffusion coefficient (ADC) values and ratios are being increasingly used to analyze molecular diffusion at the cellular level in head and neck imaging, especially in cancers.\[^{[26]}\]

**Ocular tumors**

*Retinoblastoma*

Retinoblastoma is the commonest intraocular tumor of childhood (usually presenting below 5 years) and is thought to be congenital in origin.\[^{[1]}\] It is a malignant tumor arising from the neuroectodermal cells of the retina and must be differentiated from other benign lesions for early management and better prognosis. Clinically, leucocoria\[^{[27]}\] is associated with proptosis, which may be bilateral. The main patterns of the lesion are exophytic, endophytic or diffuse type.\[^{[1]}\] The intraocular mass with calcification and areas of necrosis, associated retinal detachments and vitreous involvement (endophytic type) can be well visualized on USG. CT and MRI are very useful in determining retrobulbar and extraocular spread, intracranial involvement, and exclude trilateral retinoblastoma with pineal gland tumors when bilateral disease is detected. CT is very sensitive as >90% of tumors show evidence of calcification\[^{[1]}\] (Figure 30). On MR, they are hyperintense to normal vitreous on T1WI, moderate to markedly hypointense on T2WI depending on the calcification (Figure 31), demonstrated well with contrast administration and thin sections (1.5-3 mm) with fat-suppression. It is important to differentiate the tumor from persistent hyperplastic primary vitreous (PHPV) (Figure 32), retinopathies of prematurity (ROP), Coats’ disease (primary retinal telangiectases), organized subretinal hemorrhage, endophthalmitis, and other intraocular masses.\[^{[26]}\]

*Malignant melanoma*

These are highly malignant tumors arising from the uvea (choroid, ciliary body, iris), that tend to metastasize hematogenously to liver, lung, bone, kidney, and brain. Imaging must focus to search for metastatic spread which guides management. CT reveals a hyperdense, well-marginated, typically enhancing mass and should to be differentiated from other primary and secondary uveal

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**Figure 28:** Optic Neuritis in multiple sclerosis. Axial T1WFS image showing enhancement of segment of left intraorbital (curved arrow) optic nerve extending posteriorly involving the intracanalicular portion (arrow)

**Figure 29 (A, B):** Orbital involvement in sinonasal malignancy. Coronal computed tomography (A) shows expansile enhancing mass (asterisk) in sinonasal cavity right causing erosion of adjacent orbital bony walls (B) and extension into right orbit (arrow)

**Figure 30 (A, B):** Bilateral retinoblastomas. Axial computed tomography (CT) scans (A) showing bilateral intraocular soft tissue density mass lesions (curved arrow) with specks of calcifications (arrow). Follow-up CT (B) shows chunky calcifications with shrunken globe (phthisis bulbi) left (arrow)

**Figure 31 (A, B):** Retinoblastoma. Magnetic resonance axial T1WI (A) shows a left intraocular mass (arrow) hyperintense to vitreous and markedly hypointense to vitreous (curved arrow) on axial T2WI (B)
and choroidal tumors. On MR, the lesions are of moderately high signal on T1W (paramagnetic properties of melanin or hemorrhage) [Figure 33], and are hypointense on T2W. Gd contrast is used for evaluation of ON and retrobulbar extension.[27]

**Lacrimal gland and fossa lesions**

Lacrimal gland is located in the lacrimal fossa in the superolateral, extraconal compartment adjacent to superior and lateral rectus tendons. Inflammatory diseases are usually acute, may be a part of spectrum of orbital pseudotumor or related to trauma.[28] Chronic dacrtyoadenitis may be associated with connective tissue disorders, non-infective granulomatous diseases (Sarcoidosis, Wegener's), Mikulicz syndrome or other non-specific infiltrative conditions.[21] CT/MR demonstrates unilateral or bilateral diffuse enlargement of the gland with marked contrast enhancement [Figure 34], and associated adjacent Superior rectus / Lateral rectus myositis or tendinitis.

Lymphomatous processes may be unilateral or bilateral, ranging from benign infiltration to malignant lymphomas, often causing diffuse enlargement. Tumors of epithelial origin represent half the masses involving the lacrimal gland; 50% of these are benign pleomorphic adenomas.[28,29] Bony changes in the fossa may be seen on CT/MRI associated with indentation of the globe and distortion of the muscle cone [Figure 35A and B].

**Miscellaneous conditions**

Post-transplantation lymphoproliferative disorder (PTLD) Seen in 2-3% patients, within first year after organ transplantation, due to uncontrolled lymphocytic proliferation response.[30] Orbit is a common site for this lesion, seen as soft tissue mass in lacrimal region, may have contrast enhancement, and at times aggressive as opposed to lymphoma [Figure 36A and B].

Craniofacial abnormalities and craniosynostosis affects the development of the calvarium and facial bones which usually result in secondary changes of cranial fossae, skull base, and cranial vault and orbits. There may be other congenital developmental abnormalities associated.
Tumor-like conditions of the craniofacial bones like Fibrous dysplasia (FD)\cite{31} with extensive involvement of the maxilla, mandible, and skull base may lead to anatomic deformities of the orbit and encroachment of the neurovascular canals\cite{1} [Figure 37]. CT is the gold standard for imaging and 3D reformats are very useful in pre-operative assessment and post-surgical follow-up.\cite{32} MR is the preferred modality for imaging of associated brain and spinal abnormalities in the evaluation of various craniofacial syndromes.

Figure 35A: Contrast enhanced CT images (CECT) show enhancing mass lesion of the right lacrimal gland (asterisk) with mass effect (a) and remodeling of adjacent bone (curved arrow) (b)

Figure 35B: Lacrimal gland pleomorphic adenoma. Magnetic resonance T1W image (a) shows enlarged globular right lacrimal gland (white arrow head), hypointense with enhancement in post-contrast gadolinium enhanced images (b-d) (white arrows) suggestive of benign tumor

Figure 36A: Post-bone marrow transplant in a patient of acute leukemia with left proptosis. Magnetic resonance T1WI axial (a), sagittal (d) shows extracranial space infiltration on left (small on right), hypointense to orbital fat (white arrow), short tau inversion recovery coronal (b), axial (c) shows hypointense lesions involving lacrimal glands extending superiorly and displacing the left globe

Figure 36B: Lesion on magnetic resonance axial short tau inversion recovery (a), post-contrast T1W FS (b) shows no enhancement (black arrow), with true restriction of diffusion appearing bright on diffusion weighted imaging axial (c) with increasing B values (0,500,1000) and correspondingly dark on apparent diffusion coefficient map (white arrows)

Figure 37 (A-C): (A-C) Craniofacial fibrous dysplasia with left proptosis. Computed tomography MPR images show expansion of diploic space with ground-glass appearance (asterisk) involving hemicraniofacial bones on left. Note obliteration of sinuses, encroachment of orbit with overcrowding of orbital contents
Conclusion

Multimodality approach for imaging of orbits helps in narrowing the differentials in establishing a confident radiologic diagnosis. CT is the modality of choice for bony detail and calcifications, whereas MR is superior for evaluation of the visual pathways, the globe and soft tissues. High resolution realtime USG, MDCT with valuable post-processing techniques and high field strength MRI are very useful in visualization of and characterization of orbital lesions. MRI DWI, MR spectroscopy, perfusion studies, Positron Emission Tomography (PET)-CT/MR has increasingly become important in the recent years, in evolving towards functional imaging.

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