Acute dacyrocystitis complicated by orbital cellulitis and loss of vision: A case report and review of the literature

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A B S T R A C T

INTRODUCTION: Acute dacyrocystitis usually presents as a pre-septal cellulitis since the lacrimal sac lies anterior to the orbit septum. Orbital cellulitis secondary to acute dacyrocystitis is very rare but has been reported resulting in abscess formation with risk of visual loss. Typically, orbital cellulitis responds well to systemic antibiotic therapy and surgical drainage without permanent loss of vision [1]. We describe a case of otherwise healthy adult who presented to our academic institution with complete visual loss following orbital cellulitis and abscess formation secondary to acute dacyrocystitis. Our case has been prepared and reported in line with the SCARE criteria in: “The SCARE Statement: Consensus-based surgical case report guidelines”. International Journal of Surgery 2016; 34:180–186. The authors further stress that they have no financial disclosures related to their recommendations [1].

1. Introduction

Acute dacyrocystitis commonly causes pre-septal cellulitis as the lacrimal sac lies anterior to the orbital septum. Due to natural anatomic barriers to the orbit, orbital cellulitis secondary to acute dacyrocystitis is very rare but has been reported resulting in abscess formation with risk of visual loss. Typically, orbital cellulitis responds well to systemic antibiotic therapy and surgical drainage without permanent loss of vision [1]. We describe a case of otherwise healthy adult who presented to our academic institution with complete visual loss following orbital cellulitis and abscess formation secondary to acute dacyrocystitis. Our case has been prepared and reported in line with the SCARE criteria in: “The SCARE Statement: Consensus-based surgical case report guidelines”. International Journal of Surgery 2016; 34:180–186. The authors further stress that they have no financial disclosures related to their recommendations [1].

2. The case

A 35-year old male presented with pain, redness and eyelid swelling of the right eye for 3 days. His condition was rapidly progressing with proptosis, restricted eye movement and decreased vision in the right eye over the following day. He had no associated history of fever. He was otherwise healthy and had no history of trauma. Prior to his presentation, the right eye was completely normal with normal vision and eye movement. He had visited another hospital 2-days prior to his presentation to us but did not have any symptoms suggestive of cellulitis at that time. He was diagnosed with acute dacyrocystitis and pre-septal cellulitis on the right side, was given oral antibiotics and discharged home with a follow up appointment in 3 weeks. Despite compliance with the oral antibiotics, his symptoms continued to worsen until he lost the vision in his right eye, which brought him to our emergency room seeking urgent medical advice.

On evaluation, his vital signs were stable. The clinical examination showed right upper and lower eyelid erythema and edema, which was tense on attempted retraction, and right eye proptosis with complete ophthalmoplegia (Fig. 1A & B). The visual acuity of the right eye was no light perception and of the left eye was 20/20. The intraocular pressure of the right eye was 50 mmHg while it was 16 mmHg in the left eye. The right pupil was fixed, mid-dilated, and nonreactive to light. The slit lamp examination of the right eye showed conjunctival chemosis and signs of exposure keratopathy.
Fundus examination of the right eye showed hyperemic disc and choroidal folds. The slit lamp examination and fundus examination of the left eye were unremarkable.

A computed tomography (CT) of the right orbit with contrast showed dilatation of the right lacrimal sac as well as the ipsilateral nasolacrimal duct. The lacrimal sac was showing rim enhancement extending into the extra-orbital post septal region. There was a large extraconal non-enhancing collection measuring 4.7 cm (anteroposterior) × 2.0 cm (transverse) × 2.1 cm (cranio-caudal) seen adjacent to the right medial rectus muscle and causing mass effect on the globe (Fig. 1C & D). The findings were in favor of right-sided dacryocystitis associated with orbital cellulitis and extraconal orbital abscess.

He was immediately started on intravenous ceftriaxone and clindamycin and underwent urgent surgical drainage of the abscess. Intraoperatively the abscess was drained from the lacrimal sac and medial part of the orbit (Fig. 2A) and a swab was taken from the purulent material for microbiology. Posterior rupture of the lacrimal sac into the orbit was expected. Irrigation was done using normal saline and cefazolin. Irrigation of the lacrimal drainage system showed the reflux coming through the drain which confirmed the nasolacrimal duct obstruction (Fig. 2B).

Postoperatively, proptosis had markedly reduced as well as the eyelid edema and erythema. Improvement in motility was also noted and conjunctival chemosis became much less. His visual acuity remained at no light perception. Culture results were positive for staphylococcus epidermidis which was sensitive to ceftriaxone and clindamycin. Intravenous dexamethasone was administered on the third postoperative day with marked improvement in the extraocular motility and periorbital swelling. He was discharged on the seventh postoperative day on oral amoxicillin/clavulanic acid and was followed up as an outpatient. Three months later, he underwent right external dacrtyocystorhinostomy and intubation without complications. He was followed up regularly with a plan for stent removal 4 months after the surgery. He did not regain vision and developed right optic nerve atrophy (Fig. 2C & D).

3. Discussion

Dacryocystitis is an infection of the lacrimal sac due to nasolacrimal duct obstruction. Acute dacryocystitis can present as a pre-septal cellulitis, which generally involves soft tissue in the pre-septal area. Orbital cellulitis is a vision-threatening infectious process involving the ocular adnexal structures posterior to the orbital septum. Orbital cellulitis can rarely occur secondary to acute dacryocystitis and typically responds well to systemic antibiotic and surgical drainage without permanent visual loss [2]. The attachment of orbital septum to the lacrimal crest prevents the spread of infection to the posterior orbit. In addition, other anatomical structures such as the lacrimal fascia, the posterior limb of the medial canthal ligament, and deep heads of the pre-tarsal and pre-septal orbicularis muscles also act as a barrier to posterior extension. Orbital abscess secondary to dacryocystitis generally occurs in the medial and inferior aspects of the globe because of the anteroinferior location of the lacrimal sac. The anterior and inferior location of the lacrimal sac in relation to the globe can result in a channel of communication between the medial and inferior rectus muscles directly to the intraconal space, which can result in rapid loss of vision necessitating urgent surgical intervention [3].

There are 7 cases of visual loss following orbital cellulitis secondary to acute dacryocystitis reported in the literature. Summary of the cases (including ours) is shown in Table 1. Kikkawa et al.
Fig. 2. A: Significant purulent material was drained from the lacrimal sac and medial part of the orbit. B: Irrigation of the lacrimal drainage system showed the reflux coming through the wound drain which confirmed the nasolacrimal duct obstruction. C: Color fundus photo of the right eye showing temporal optic disc pallor. D: External photograph of the patient which was taken in the last follow up visit.

Table 1
Summary of all 8 cases of acute dacryocystitis complicated by orbital cellulitis and loss of vision including our present case.

| Author/year       | Patient age/gender | VA on presentation/final VA | Location of orbital abscess/side | Previous dacryocystitis | Mechanism of vision loss | Organisms |
|-------------------|--------------------|-----------------------------|----------------------------------|--------------------------|--------------------------|-----------|
| Kikkawa/2002, [3] | 38 years/ F 64 years/ F | NLP/NLP                     | Extranodal/ left Intracanal/ right | Yes                      | Optic nerve compromise | Staphylococcus aureus, Nonenterococcal group D streptococci, streptococcus viridans, streptococcus intermedius. Gram-positive diplococci. NS. |
| Maheshwari/2009, [2] | 50 years/ F 65 years/ F | NLP/NLP                      | Intracanal/ right Extranodal/ left | Yes                      | Optic nerve compromise | Optic nerve compromise |
| Coşkun/2011, [5]  | 45 years/ F        | LP/NLP                       | Extranodal/ left                 | No                       | Central retinal artery occlusion | Coagulase–negative staphylococcus. |
| Wladis/2016, [7]   | 66 years/ F        | NLP/NLP                       | NS                               | NS                       | Ophthalmic artery occlusion | Streptococcus pyogenes.       |
| Pfeiffe/2016, [14] | 50 years/ F        | NLP/NLP                       | Extranodal/ left                 | No                       | Optic nerve compromise | Staphylococcus epidermidis. |
| Alsalamah * 2018  | 35 years/ M        | NLP/NLP                       | Extranodal/ right                | No                       | Optic nerve compromise | Staphylococcus epidermidis. |

VA = visual acuity, F = female, M = male, NLP = no light perception, LP = light perception, NS = not specified.
* Present case.
found that a prior episode of dacryocystitis is a risk factor for orbital extension by distension of the lacrimal sac and disruption of its posterior barriers. These barriers weaken from distension increasing the likelihood of posterior rupture and spread [3]. Although our patient had no history of dacryocystitis, chronic distention of the lacrimal sac may have led to posterior rupture and orbital extension. Other theories included spread of infection to ethmoid sinus via lamina papyracea, with subsequent extension from the sinus to the orbit [4]. Other causes include hematogenous spread from other systemic causes and, in some cases a primary orbital cellulitis that can extend into the lacrimal sac before the presence of dacryocystitis. Immunosuppression can also be a contributing factor [5].

The microorganisms found in dacryocystitis-induced orbital cellulitis are similar to the microorganisms found in uncomplicated dacryocystitis [6]. Intraoperative cultures revealed that Gram-positive bacteria are the most common pathogens [7]. The most frequently isolated microorganisms are Staphylococcus species (Staphylococcus aureus and coagulase-negative Staphylococcus) and Streptococcus species [8,9]. These microorganisms are also found in paranasal sinus infections [10]. The mechanism of vision loss is thought to be due to optic nerve compression and ischemia as in our patient or secondary to the elevation of intraocular pressure by the mass effect of the abscess, which results in central retinal artery obstruction or ophthalmic artery obstruction [5,7]. Thrombophlebitis in the valve-less orbital veins and optic neuritis caused by invasion of organism may also be an explanation [10,11]. Prompt diagnosis and emergency treatment is very essential in orbital infections. A delay in treatment can result in serious complications like vision loss, cavernous sinus thrombosis, meningitis, frontal abscess, and even death [9,12,13]. If an abscess is present, incision and drainage to prevent rupture of the lacrimal sac and posterior extension into the orbit is highly recommended, with dacryocystorhinostomy being the definitive treatment [14].

In conclusion, acute dacryocystitis should be considered as one of the causes of orbital cellulitis and abscess formation. Patients with orbital cellulitis and/or orbital abscess should be questioned about symptoms of lacrimal drainage obstruction. We recommend that patients with acute dacryocystitis should be carefully monitored with very close follow up and should be warned about symptoms and signs of pre-septal and orbital cellulitis, which can rapidly progress and cause severe visual sequelae. Prompt recognition and appropriate treatment are essential to prevent visual loss.

Conflicts of interest

The authors have no conflict of interest related to this case report.

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Ethical Approval

Case reports do not require ethical approval at our institution.

Consent

General informed written consent has been obtained from the patient. The General informed written consent includes patient’s approval for anonymous use of relevant clinical and surgical information in an anonymous way.

Author contribution

Abrar K. Alsalamah: Review of chart, literature review and drafting of the case report

Hind M. Alkatan: Critical review and correction of the manuscript and the corresponding author.

Yasser H. Al-Fakya: Surgical care of the patient and manuscript review

Registration of Research Studies

No Registration is needed for Case reports.

Guarantor

Hind Mana Alkatan, MD

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