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From Handlebar to Enucleation: Management and Prosthetic Outcome of a Severe Traumatic Globe Luxation

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ABSTRACT

Background: Traumatic globe luxation (TGL) is a rare, severe ocular emergency involving the complete displacement of the eyeball from the orbit. It presents a profound clinical challenge, demanding a rapid and accurate assessment of complex prognostic factors to guide the difficult decision between globe salvage and primary enucleation. **Case presentation:** A 33-year-old male presented 18 hours after a motorcycle handlebar strike to his left orbit. The examination revealed a left globe luxation with No Light Perception (NLP) vision, a total afferent pupillary defect, and complete ophthalmoplegia. Computed tomography confirmed a closed-globe injury with a superior orbital rim fracture and a large retrobulbar hematoma, but could not delineate soft tissue integrity. Surgical exploration revealed two critical, paradoxical findings: an anatomically intact optic nerve despite its functional death, and a catastrophic avulsion of five of the six extraocular muscles. The medial, lateral, and inferior recti, along with both oblique muscles, were detached, while the superior rectus muscle was uniquely spared. **Conclusion:** Based on the catastrophic loss of vascular supply from the avulsed muscles, which rendered the globe biologically non-viable, a primary enucleation was performed. This case suggests that in TGL, the integrity of the extraocular musculature is a paramount prognostic indicator, potentially superseding the anatomical status of the optic nerve in determining globe viability. It highlights the necessity of intraoperative exploration to definitively assess the extent of injury and illustrates a scenario where primary enucleation is not a treatment failure, but a definitive, rehabilitation-focused therapeutic strategy.

1. Introduction

Among the myriad injuries in the spectrum of orbital trauma, traumatic globe luxation (TGL) represents a uniquely devastating event. Defined as the complete, forceful displacement of the eyeball from the protective confines of the bony orbit, TGL is an absolute ophthalmic emergency that challenges the core principles of clinical assessment and surgical decision-making.¹ Its incidence is low, a fortunate consequence of the substantial force required to overcome the globe's robust anatomical tethers. Epidemiological data, while sparse due to the condition's rarity, consistently identify high-velocity

blunt force trauma as the primary etiology, with motor vehicle accidents—particularly involving unrestrained or un-helmeted individuals—physical assaults, and high-impact sports injuries being the most common culprits. The significant kinetic energy involved means that TGL is frequently a component of a larger constellation of polytrauma, often accompanied by life-threatening intracranial hemorrhage or severe craniomaxillofacial fractures that demand immediate, multidisciplinary attention.²

The pathophysiology of TGL is a study in extreme orbital dynamics.³ The orbit, a conical bony cavity with a volume of approximately 30 cm³, provides a

rigid container for the 7.5 cm³ globe and its surrounding soft tissues. Luxation occurs when a sudden, massive increase in intraorbital pressure exceeds the tensile strength of the globe's retaining structures. This pressure surge can be generated through several biomechanical mechanisms.⁴ A "lever effect" can occur when an elongated object, such as a tree branch or a tool, enters the orbit medially and pries the globe forward. A "wedge effect" describes a similar outcome from a laterally-directed force. However, in the context of blunt trauma, the most common mechanism involves the transmission of a powerful shockwave through the orbital rim. This force can cause a concomitant fracture and, more critically, can rupture orbital vessels, leading to the rapid formation of a retrobulbar hematoma. This expanding hematoma acts as a hydraulic piston within the confined orbital space, propelling the globe forward. Once the globe's equator passes anterior to the orbital aperture, a vicious cycle often ensues: intense pain triggers a powerful blepharospasm of the orbicularis oculi muscle, which clamps down behind the displaced globe, trapping it externally and strangulating its vascular supply.⁵

The clinical management of TGL is fraught with complexity, as there is no universally accepted treatment algorithm.⁶ The attending surgeon is faced with a series of critical decisions that must be made rapidly, often with incomplete information. The initial visual acuity is the single most important prognostic indicator for function. A finding of No Light Perception (NLP), especially when coupled with a total afferent pupillary defect, suggests a catastrophic and likely irreversible injury to the optic nerve or retina. The structural status of the globe (open vs. closed) and the optic nerve (avulsed, transected, or stretched) are paramount. An open globe injury or a complete optic nerve transection often necessitates primary enucleation.⁷ The integrity of the extraocular muscles (EOMs) is a crucial, yet often underappreciated, factor. The EOMs, particularly the four rectus muscles, are not merely responsible for motility; they are the primary conduits for the anterior ciliary arteries,

which provide the majority of the blood supply to the anterior segment of the eye. The "generally accepted rule" in ophthalmic trauma, a principle derived from decades of clinical experience and case series rather than large-scale trials, posits that the avulsion of more than two, and certainly more than three, rectus muscles leads to irreversible anterior segment ischemia.⁸ This condition results in a cascade of complications, including corneal opacification, intractable glaucoma, and chronic pain, ultimately leading to a shrunken, non-functional eye (phthisis bulbi). This principle forms a critical, though sometimes debated, cornerstone of the decision-making process. It forces the surgeon to weigh the desire to preserve the globe for cosmetic reasons against the certainty of creating a painful, dying organ.⁹

The novelty of this case report is multifold. It presents a rare constellation of findings—a functionally dead but anatomically intact optic nerve, a closed-globe injury, and a near-total EOM avulsion—that created a unique and challenging clinical paradox. Furthermore, the specific pattern of injury, with the unique sparing of a single rectus muscle, provides a rare opportunity to analyze the intricate biomechanics of focused orbital trauma.¹⁰ Therefore, the aim of this study is to provide a detailed, scientifically rigorous analysis of this case, moving beyond a simple narrative to explore the underlying pathophysiology, critically examine the clinical decision-making algorithm, and use these unique findings to refine our understanding of globe viability in the context of severe trauma. We aim to illustrate a scenario where intraoperative findings are paramount and to suggest that a management strategy focused on definitive, early rehabilitation can yield a superior outcome.

2. Case Presentation

A 33-year-old male, not wearing a helmet, was brought to our tertiary care emergency department 18 hours following a single-vehicle motorcycle accident. He reported that while traveling uphill, he lost control

of his vehicle and was struck with significant force by the left handlebar directly in the left orbital region. The patient reported immediate, severe pain and a complete and persistent loss of vision in his left eye. He had not received any prior medical attention. A standard Advanced Trauma Life Support (ATLS) primary survey was unremarkable for other life-threatening injuries, and his tachycardia (110 bpm) and hypertension (150/90 mmHg) were attributed to acute pain and distress. The ophthalmic examination revealed a stark contrast between the two eyes. The right eye was entirely normal, with a visual acuity of 20/20 and an intraocular pressure (IOP) of 14 mmHg. The left eye, however, presented with a devastating

injury profile. Visual acuity was definitively confirmed as No Light Perception. The globe was completely luxated anteriorly, wedged between severely edematous and ecchymotic eyelids. Due to the globe's position and the patient's distress, quantitative exophthalmometry was not feasible, and IOP was not measurable. The constellation of findings, detailed in Figure 1, painted a grim picture of a functionally dead eye with severe adnexal trauma. The 18-hour delay in presentation likely exacerbated the exposure-related keratopathy and may have contributed to the ischemic conjunctival necrosis, though the primary cause of the necrosis was likely the vascular strangulation from the initial injury.

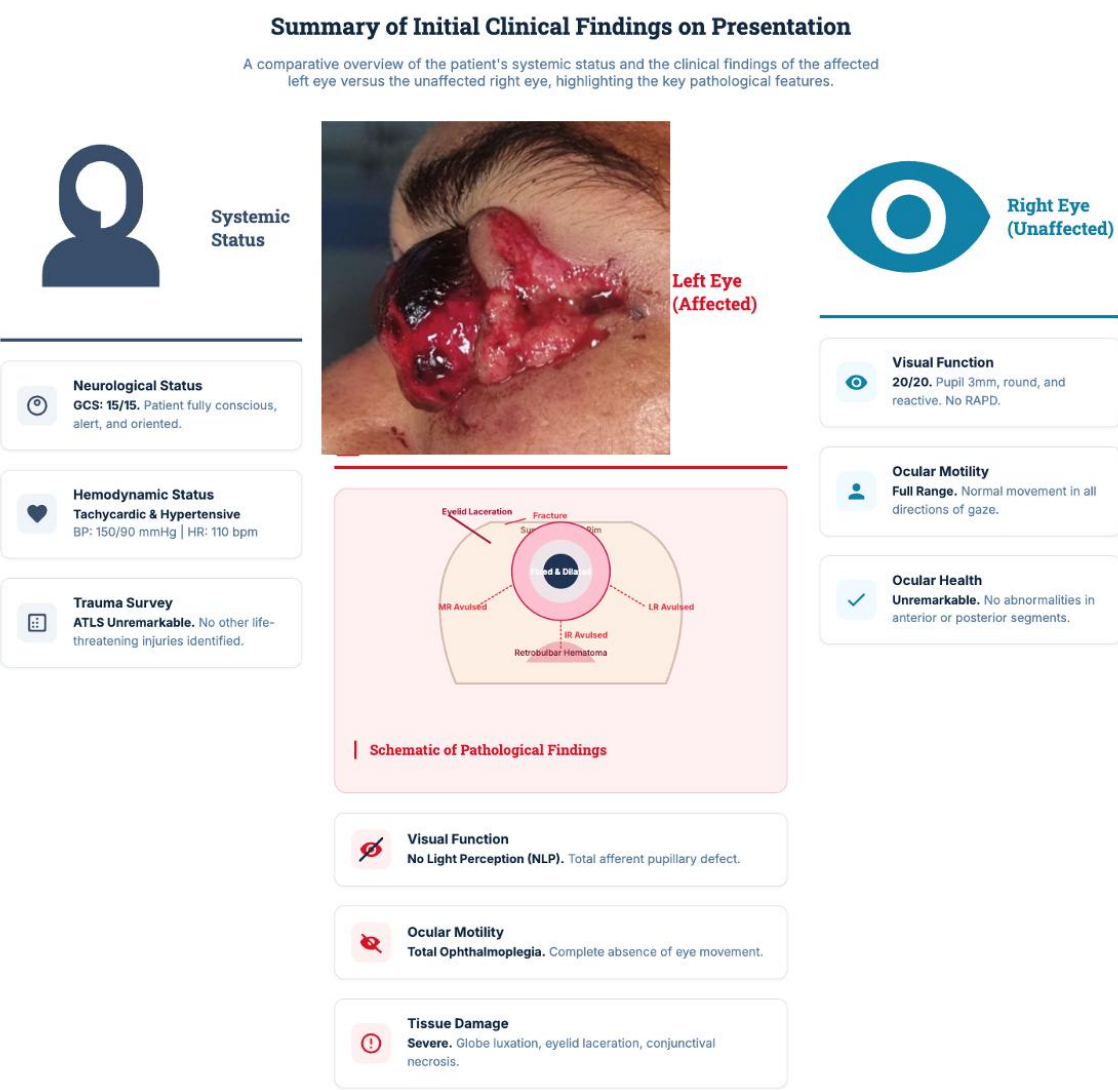


Figure 1. Summary of initial clinical findings on presentation.

CT findings (Figure 2) showed a complex pattern of injury that was critical for surgical planning while simultaneously underscoring the limitations of radiological imaging in acute, severe trauma. The axial and 3D reconstruction views provided a clear anatomical map of the damage. The first crucial finding, as detailed in the summary panel, was the absence of any acute intracranial pathology, which definitively ruled out an immediate neurosurgical emergency and allowed the clinical focus to shift entirely to the devastating orbital injury. The scan was pivotal in confirming a closed-globe injury. The integrity of the globe's contour, with no signs of rupture or a "flat tire" sign, was a key finding that kept the possibility of globe preservation on the table, at least from a structural standpoint. The imaging also clearly delineated the forces at play: a non-displaced linear fracture of the superior orbital rim confirmed the point of high-energy impact, while a large, hyperdense retrobulbar hematoma was identified posterior to the globe. This hematoma was the radiological correlate of the severe clinical proptosis, confirming significant internal bleeding and explaining the immense hydraulic pressure that propelled the globe forward out of its socket. However, the most important finding from the CT scan was, paradoxically, what it could not reveal. The final panel, Diagnostic Limitation, highlights that the extensive retrobulbar hematoma created a significant artifact that completely obscured the fine anatomical details of the optic nerve and the extraocular muscles. This diagnostic ambiguity is visualized in the central schematic, where the nerve and muscles are shown as indistinct, dashed lines disappearing into the hematoma. This limitation was critical: it rendered the CT scan unable to answer the most vital questions regarding the globe's biological viability. The integrity of these crucial structures could not be confirmed or denied by imaging alone, making surgical exploration not just a therapeutic step, but an essential and unavoidable diagnostic necessity to determine the true extent of the damage and dictate the definitive course of management.

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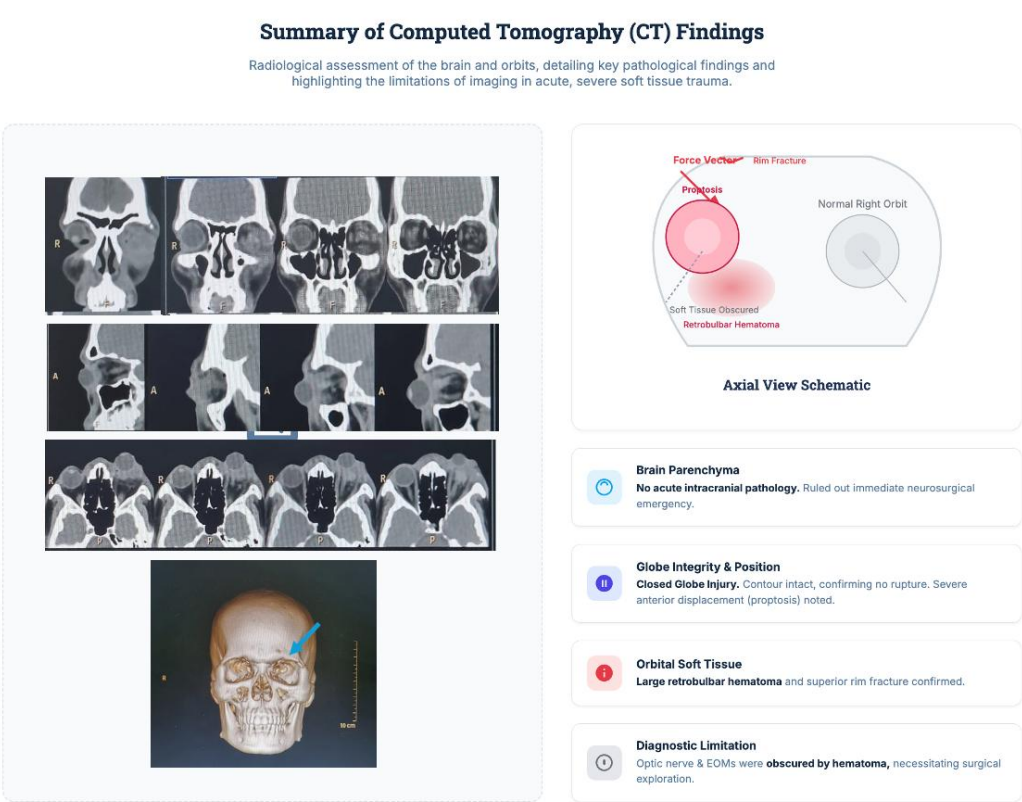


Figure 2. Summary of computed tomography (CT) findings.

After extensive discussion with the patient and his family regarding the NLP vision and the exceedingly poor prognosis, informed consent was obtained for surgical exploration with a high likelihood of primary enucleation. The patient was taken to the operating theater under general anesthesia. The intraoperative findings, detailed in Figure 3, The figure tell a story of two starkly different realities coexisting within the same traumatized orbit: one of unexpected structural preservation and another of catastrophic, irreparable damage. The right panel methodically outlines the surgical progression, beginning with the essential first step of exposure & debridement. This initial phase was not merely preparatory but diagnostic, allowing for the removal of necrotic tissue and providing a clear view of the underlying globe and its adnexal structures. It was during the second step, the Critical Finding of Muscle Avulsion, that the true, devastating extent of the injury was revealed. As depicted in the central schematic on the left, systematic exploration confirmed the complete avulsion—a tearing away from their scleral insertions—of the medial, lateral, and inferior rectus muscles, as well as both oblique muscles. The red color-coding and dashed lines in the schematic vividly illustrate this catastrophic detachment, signifying a profound loss of both motility and, more importantly, vascular supply. This discovery was immediately contrasted by the paradoxical finding of intact structures. Amidst the widespread destruction, the surgical team identified an anatomically intact optic nerve and a single, securely attached superior rectus muscle. These structures, highlighted in green, represent a significant clinical paradox. The intact optic nerve, despite being functionally dead as determined by pre-operative examination, could have created a moment of surgical hesitation. The sparing of the superior rectus muscle, likely due to anatomical shielding from the specific injury vector, stands as a unique and scientifically intriguing detail of the case. However, it was the fourth and final step, the Definitive Decision for Enucleation, that resolved this paradox. The decision was not based on the non-functional optic

nerve but was a direct and necessary consequence of the catastrophic muscle damage. As the panel explains, the loss of three of the four rectus muscles confirmed that the globe was biologically non-viable due to irreversible anterior segment ischemia. The avulsion had severed the majority of the anterior ciliary arteries, cutting off the blood supply to the front of the eye. This finding represented a point of no return, making primary enucleation the only logical and humane course of action to prevent the inevitable, painful sequelae of a dying eye.

The patient's post-operative course was uneventful. He was managed with a 3-day course of intravenous antibiotics followed by a 7-day course of oral antibiotics. His pain was well-controlled, and he was discharged on the third post-operative day. The subsequent rehabilitation pathway, detailed in Table 4, was seamless. Figure 4 showed a comprehensive timeline of the post-operative management and rehabilitation pathway, visually chronicling the patient's successful journey from a state of acute surgical trauma to complete cosmetic and functional restoration. This figure is not merely a sequence of events; it is a testament to a well-executed, multi-stage clinical strategy that prioritizes patient comfort, tissue healing, and psychosocial well-being, demonstrating that the surgical intervention of enucleation is only the first step in a longer, equally important process of holistic care. The timeline begins with the crucial phase of Immediate Post-operative Care, spanning the first three days. This period, managed in a hospital setting, was focused on mitigating the primary risks of infection and pain. The administration of intravenous antibiotics provided systemic protection against potential pathogens introduced during the trauma or surgery, while analgesics ensured the patient's comfort. The use of a pressure bandage over the anophthalmic socket served a dual purpose: it helped to control post-operative edema and hemorrhage, and it maintained the position of the internal conformer, which is essential for preserving the shape of the conjunctival fornices—the deep pockets of the socket that will

eventually support the prosthesis. The next milestone, at two weeks, was Wound Healing & Suture Removal. The transition to oral antibiotics upon discharge continued the prophylactic coverage as the patient recovered at home. This follow-up appointment was a critical checkpoint. The finding of "excellent wound healing" confirmed that the initial surgical reconstruction of the eyelids and canthus was successful and that there were no signs of infection or wound dehiscence. The removal of sutures at this stage signifies that the tissues have regained sufficient tensile strength, paving the way for the next phase of rehabilitation. The well-formed socket observed at this point is a direct result of the meticulous surgical closure and the proper use of the conformer. The Prosthetic Fitting Process, initiated at week four, marks the shift from surgical healing to active rehabilitation. The referral to a specialized ocularist is a key step. The creation of an impression mold is a highly precise procedure, akin to taking a dental impression, that captures the unique topography of

the patient's anophthalmic socket. This custom approach is what separates a truly functional prosthesis from a simple "glass eye." It ensures that the final device will fit intimately against the orbital tissues, allowing for optimal comfort, preventing socket discharge, and, crucially, enabling the transmission of movement from the underlying orbital implant and tissues to the prosthesis, which provides a more natural, dynamic appearance. The final panel, Successful Rehabilitation at week six, represents the culmination of this entire process. The fitting of the final, custom-fabricated, hand-painted prosthesis is the moment of transformation. The report of excellent comfort and motility, along with a satisfactory cosmetic outcome, validates the entire preceding management strategy. The patient's high satisfaction is the ultimate measure of success, signifying not only the restoration of facial symmetry but also the profound psychosocial benefit of mitigating a visible disfigurement, allowing the patient to reintegrate into their life with confidence.

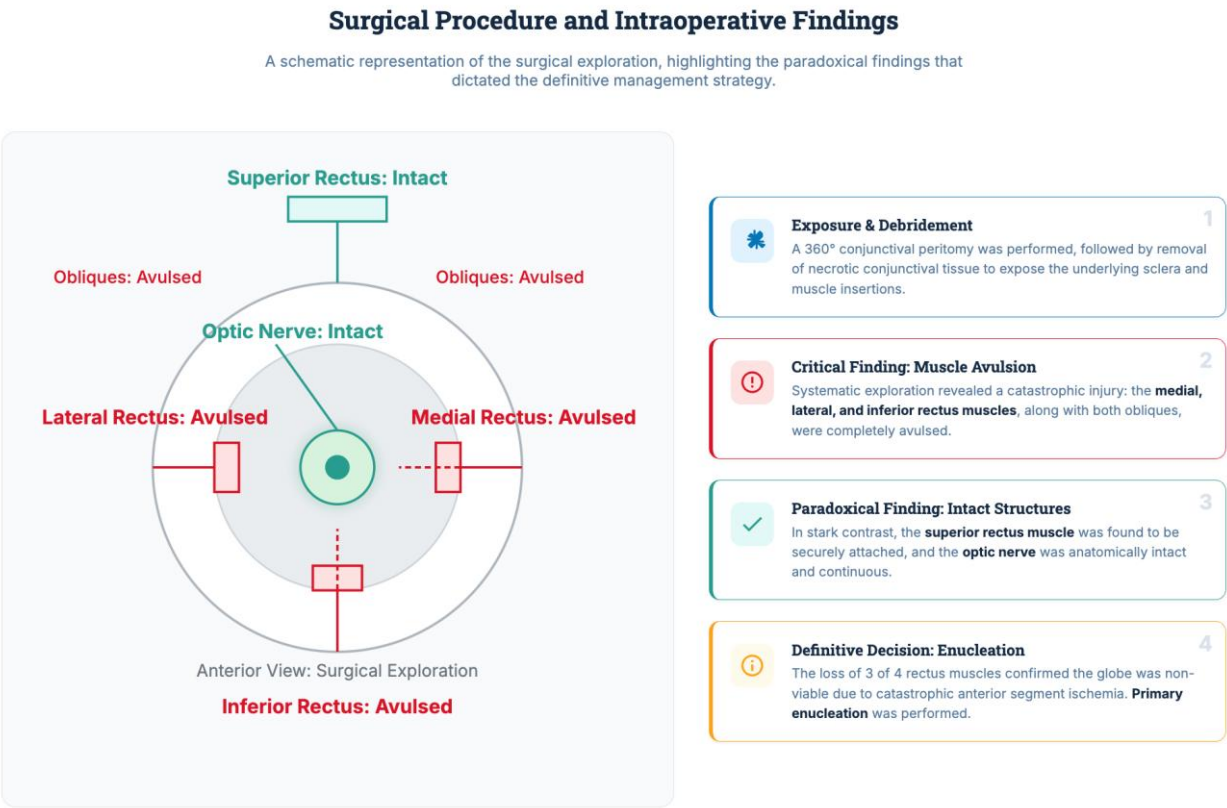


Figure 3. Surgical procedure and intraoperative findings.

Post-operative Management and Rehabilitation

A timeline illustrating the key milestones in the patient's recovery, from immediate post-surgical care to the final successful prosthetic fitting.

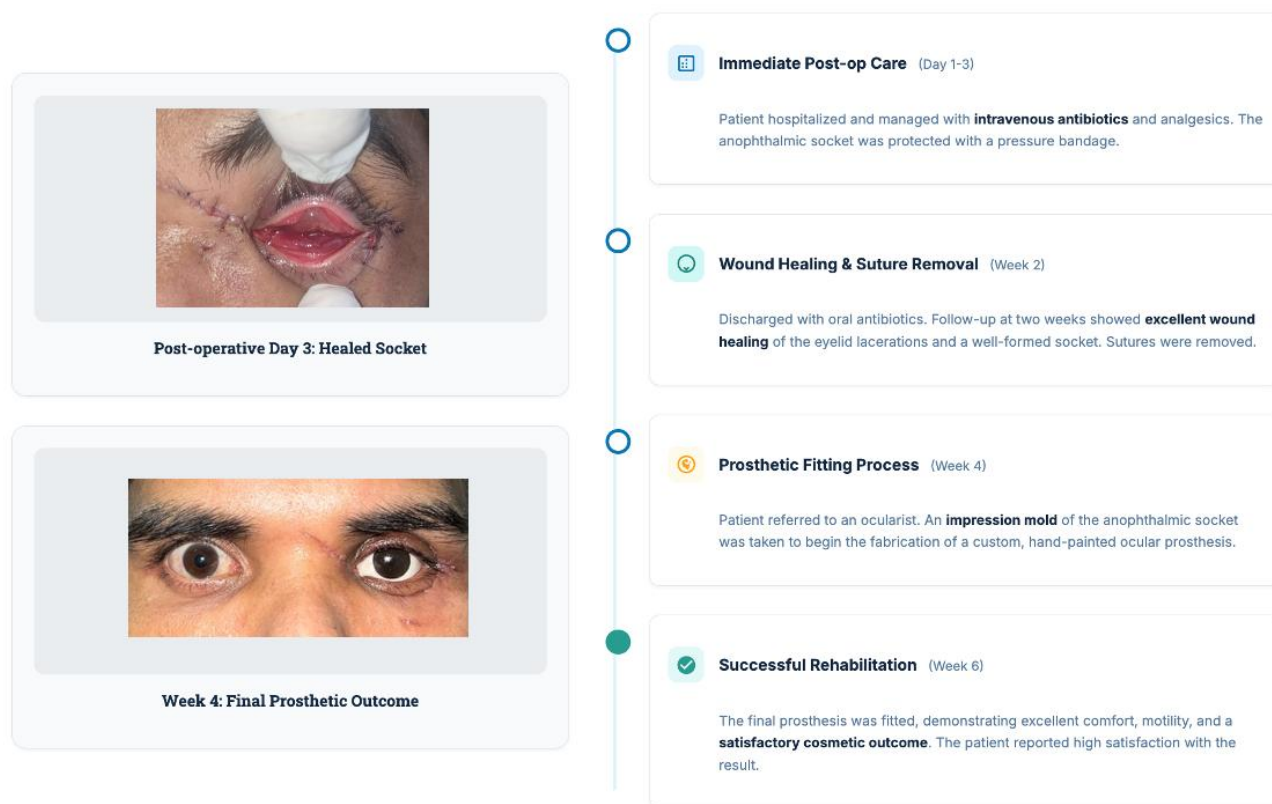


Figure 4. Post-operative management and rehabilitation.

3. Discussion

The management of this case of severe traumatic globe luxation offers a profound opportunity to move beyond a simple narrative and engage in a deep, critical analysis of the biomechanics, pathophysiology, and clinical decision-making that define modern ophthalmic trauma care. The unique constellation of findings forces a confrontation with established clinical axioms and highlights the indispensable role of intraoperative exploration in resolving the ambiguities left by pre-operative imaging. Understanding the totality of the damage begins with a granular analysis of the injury mechanism.¹¹ A motorcycle handlebar is not a simple blunt object; it is a tool that concentrates immense kinetic energy along a narrow, cylindrical line of contact. Unlike a diffuse

impact that might be absorbed by the zygomatic and maxillary buttresses, the focused force of the handlebar strike on the superolateral orbital rim acted as both a battering ram and a hydraulic plunger.¹² The "coup" impact delivered the energy necessary to fracture the superior orbital rim. Simultaneously, this force was transmitted through the orbital tissues, creating a high-pressure shockwave that propagated posteriorly and medially. This "contrecoup" wave was likely responsible for the shearing of orbital vessels and the subsequent retrobulbar hematoma. However, the most scientifically intriguing finding of this case is the pattern of muscle injury: the avulsion of all EOMs except the superior rectus. This is not a random occurrence; it is a direct consequence of the injury's vector and the unique anatomy of the orbit, and it

demands a specific analysis. The impact occurred superolaterally. The force vector would have been directed inferiorly and medially. The superior rectus muscle, originating from the annulus of Zinn and coursing directly beneath the orbital roof and the levator palpebrae superioris muscle, was likely shielded from the primary shockwave by the very bone that was fractured. The energy would have been transmitted around and below it, directly impacting the medial, inferior, and lateral recti. The oblique muscles, with their more complex paths and delicate trochlear and scleral insertions, would have been highly susceptible to the rotational and shearing forces generated by the globe's violent anterior displacement.¹³ The clean avulsion of the muscles at their scleral insertions, rather than mid-belly ruptures, suggests that the primary mechanism of injury was a rapid, extreme hyper-rotation and anterior displacement of the globe. This placed the tendinous insertions, the points of fixation, under sudden, intolerable tensile stress, causing them to tear away from the sclera. The superior rectus, being located at the pivot point of this superolateral impact, may have experienced less of this rotational shear and more of a direct compressive force against the orbital roof, to which it is more resilient.¹⁴ This specific pattern of injury underscores a critical point: the damage in TGL is not a uniform explosion but a highly vectored event. A detailed understanding of the impact's location and nature can help predict the likely pattern of internal damage and should be a key component of the clinical assessment.¹⁵

The core of this case lies in the concurrent and devastating injuries to two separate biological systems: the neural pathway (axonal injury) and the vascular pathway (anterior segment ischemia). Understanding the distinction between these is fundamental to justifying the surgical management. The finding of no light perception (NLP) vision in the presence of an anatomically intact optic nerve is a classic, if tragic, clinical paradox. This was not a compressive optic neuropathy that might have been partially reversible; the absence of optic nerve sheath

distension on CT and the sheer velocity of the trauma point overwhelmingly to a diagnosis of severe traumatic optic neuropathy (TON) secondary to axonal shear. During the luxation event, the optic nerve was subjected to extreme stretching and rotational forces. While the tough outer dural sheath of the nerve can withstand this, the millions of delicate, microscopic axons within cannot. They were stretched beyond their elastic limit and sheared apart. This initiates an immediate and irreversible cascade at the cellular level: disruption of the axonal cytoskeleton, failure of axoplasmic transport, mitochondrial dysfunction, and the initiation of apoptosis (programmed cell death). This is why the afferent pupillary pathway was completely obliterated, resulting in a fixed, non-reactive pupil. The nerve was, for all functional purposes, dead at the moment of impact. The intraoperative finding of its anatomical continuity, while noted, had no bearing on the visual prognosis. This is the central and most educational aspect of the case. The concept of "biological viability" in this context is defined by the integrity of the globe's blood supply, specifically to the anterior segment.¹⁶ The cornea, iris, and ciliary body are metabolically active tissues that receive their nourishment primarily from the seven anterior ciliary arteries, which travel with the four rectus muscles (two with each rectus, except the lateral rectus, which has one). The avulsion of the medial, inferior, and lateral rectus muscles severed five of these seven critical arteries.¹⁷ This represents a massive, acute ischemic insult, cutting off over 70% of the blood flow to the front of the eye. Had salvage been attempted, the subsequent pathophysiological cascade is predictable and horrifying. The immediate result would be severe anterior segment ischemia. Within days to weeks, the patient would develop: Severe, Intractable Pain: From ciliary body inflammation and ischemia; Corneal Decompensation: The corneal endothelium would fail, leading to diffuse, permanent stromal edema and opacification; Intraocular Inflammation: Chronic, severe uveitis would ensue; Neovascular Glaucoma: The ischemic retina and iris would release vasoproliferative factors

(like VEGF), causing abnormal blood vessels to grow over the iris (rubeosis iridis) and into the drainage angle, leading to an aggressive and painful form of glaucoma that is notoriously difficult to treat; Phthisis Bulbi: Ultimately, this cycle of ischemia,

inflammation, and high pressure would cause the internal structures of the eye to atrophy and disorganize. The globe would shrink, becoming a small, disfigured, and chronically painful phthisical eye.¹⁸

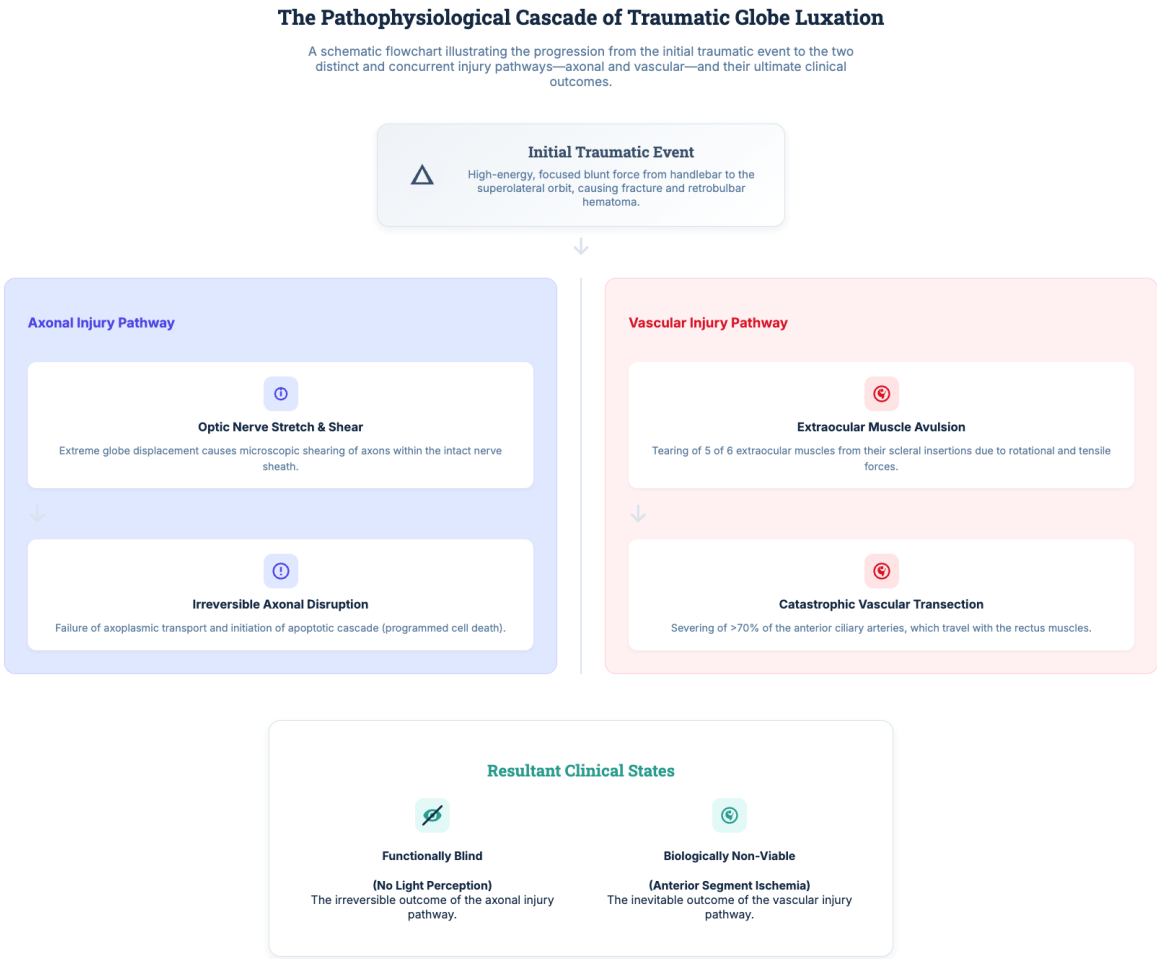


Figure 5. Pathophysiological cascade of traumatic globe luxation.

Figure 5 shows a meticulously detailed schematic flowchart that serves as the scientific and conceptual cornerstone of this entire case report. It masterfully deconstructs a single, violent moment of trauma into its constituent, parallel, and devastating pathophysiological cascades. This figure is not merely an illustration; it is a visual argument that logically and inexorably leads to the clinical conclusion that was reached. It provides a profound and granular

understanding of how a high-energy, focused blunt force injury can simultaneously trigger two distinct and independent pathways of organ death—one neural and one vascular—which together render the globe both functionally blind and biologically non-viable, making the decision for primary enucleation not just a reasonable choice, but a therapeutic imperative. The flowchart begins with the Initial Traumatic Event, the genesis of all subsequent

pathology. The schematic correctly identifies this not as a diffuse impact, but as a "high-energy, focused blunt force" from the motorcycle handlebar. This specificity is crucial. The focused nature of the impact explains why the energy was not dissipated across the broader facial skeleton but was instead concentrated on the superolateral orbital rim. This concentration of force acted as both a battering ram, causing the linear orbital fracture, and, more importantly, as a hydraulic plunger. The impact created a sudden, massive spike in intraorbital pressure, which propagated through the non-compressible orbital fat and soft tissues. This is the primary mechanical event that initiated the two distinct injury pathways depicted. The schematic correctly identifies the immediate consequences: a retrobulbar hematoma from torn vessels and the application of extreme shear and stretch forces to all the delicate structures within the orbit. From this single, violent origin point, the pathophysiology bifurcates into two separate, tragic narratives. The first branch of the flowchart, the Axonal Injury Pathway, details the destruction of the eye's neural connection to the brain. This pathway addresses the critical clinical paradox of why an eye with an anatomically intact optic nerve could present with no light perception (NLP) vision. The first step, optic nerve stretch & shear, is the key. The violent, instantaneous anterior displacement of the globe subjected the optic nerve to a degree of stretching that far exceeded its physiological tolerance. While the tough, fibrous outer sheath of the nerve—the dura—is remarkably resilient and can withstand significant stretching without tearing, the millions of delicate, microscopic axons within it cannot. These axons, the individual nerve fibers that transmit visual information, were subjected to an intolerable shear force. They were stretched to their breaking point and microscopically torn, even as the larger nerve structure remained grossly intact. This is a crucial distinction between macroscopic anatomical integrity and microscopic functional integrity. The second step in this pathway, Irreversible Axonal Disruption, describes the inevitable cellular and molecular consequences of this initial shear

injury. The tearing of the axons triggers a cascade of events known as Wallerian degeneration. Axoplasmic transport—the vital process by which nutrients and proteins are moved from the neuron's cell body in the retina down the length of the axon—comes to an immediate halt. Without this lifeline, the distal portions of the axons begin to die. This is not a slow, degenerative process but a rapid, programmed cell death, or apoptosis. The mitochondria within the axons fail, the cellular scaffolding collapses, and the nerve fibers are irreversibly damaged.¹⁸ This is the biological basis for the immediate and permanent NLP vision. The afferent pupillary pathway, which relies on these same axons, is completely obliterated, resulting in the fixed, dilated pupil observed clinically. This pathway, therefore, leads directly and inexorably to the first of the two final clinical states: the eye was Functionally Blind. This outcome was sealed at the very moment of impact, and no amount of subsequent surgical intervention could have restored the flow of information through these millions of severed neural connections. The second, parallel branch of the flowchart, the Vascular Injury Pathway, details the destruction of the globe's own biological life support system. This pathway is independent of the neural injury but equally catastrophic. It begins with the extraocular muscle avulsion. The same rotational and tensile forces that stretched the optic nerve also acted upon the six extraocular muscles that tether the globe within the orbit. The schematic correctly identifies that five of these six muscles were torn away from their insertions on the sclera. This is not just a loss of motility; it is a critical vascular event. The next step, catastrophic vascular transection, explains why. The four rectus muscles are not merely tendons for movement; they are the primary conduits for the anterior ciliary arteries, the vessels that provide the vast majority of the blood supply to the entire anterior segment of the eye—the cornea, the iris, and the ciliary body. By avulsing the medial, inferior, and lateral rectus muscles, the trauma had severed more than 70% of this vital arterial supply. This is akin to severing the major arteries to a limb; the organ

downstream is immediately rendered ischemic. This event triggered a predictable and devastating cascade of Anterior Segment Ischemia. Had the globe been salvaged, the consequences of this ischemia would have been swift and agonizing. The corneal endothelium, which relies on a constant supply of oxygen and nutrients to power its pumps that keep the cornea clear, would have failed within days, leading to diffuse, permanent corneal edema and opacification. The iris and ciliary body would have become inflamed and necrotic, causing severe, deep, boring pain (uveitis). In response to the profound ischemia, the eye would have released a flood of vasoproliferative factors, such as VEGF, triggering the growth of abnormal new blood vessels on the iris (rubeosis iridis) and in the drainage angle of the eye. This would have inevitably led to an aggressive, intractable, and exquisitely painful form of neovascular glaucoma. Over a period of months, this relentless cycle of ischemia, inflammation, and high pressure would have caused the internal structures of the eye to atrophy and disorganize, leading to the globe shrinking into a small, disfigured, and chronically painful ball known as phthisis bulbi. This vascular pathway, therefore, leads directly to the second of the two final clinical states: the eye was Biologically Non-Viable. It had lost the blood supply necessary to sustain its own life as an organ. Finally, the flowchart converges on the resultant clinical states, which represent the synthesis of these two independent pathways. The axonal pathway resulted in a functionally blind eye. The vascular pathway resulted in a biologically non-viable eye. The presence of either one of these conditions alone would have presented a profound clinical challenge. The concurrent presence of both provided an unequivocal and compelling mandate for the surgical course of action. The eye was not only without any hope of vision, but it was also guaranteed to become a source of chronic, severe pain and disfigurement.¹⁹ The flowchart, therefore, brilliantly illustrates that the decision for primary enucleation was not a choice to sacrifice a potentially salvageable organ. It was the logical, evidence-based,

and humane response to an organ that had, for all practical purposes, already died twice over—once neurologically, and once vascularly.

The decision to perform a primary enucleation was based on the definitive intraoperative finding of multi-muscle avulsion. To strengthen the justification for this choice, it is instructive to play devil's advocate and critically analyze the hypothetical alternative: attempting globe salvage. A salvage procedure would have involved a heroic and complex attempt to microsurgically reattach the three avulsed rectus muscles to the sclera. Such a procedure is technically fraught with difficulty, as the avulsed muscle stumps are often retracted deep into the orbital fat.²⁰ Even if reattachment were technically possible, it would have been medically futile. The anterior ciliary arteries within those muscles were severed at the moment of avulsion. Reattaching the muscle tendon to the sclera does not re-establish this lost blood flow. The anterior segment would have remained profoundly ischemic. The surgeon would have been repositioning a dying organ back into the orbit. This "best-case scenario" for salvage would have initiated the painful cascade of phthisis bulbi described above. The patient would have been subjected to months or years of suffering, requiring multiple subsequent procedures—glaucoma surgeries, pain management injections, and ultimately, a secondary enucleation or evisceration. This secondary enucleation would have been performed in a scarred, contracted socket, making subsequent prosthetic fitting more difficult.

Therefore, the decision for primary enucleation was not a choice of "giving up." It was a proactive, evidence-based, and humane intervention. It was a definitive treatment to prevent a predictable and miserable clinical course. It acknowledged the limitations of modern medicine and prioritized the patient's long-term quality of life and comfort over the sentimental preservation of a non-functional, non-viable anatomical structure. This case strongly suggests that the clinical algorithm for TGL must include a critical checkpoint: upon intraoperative confirmation of the avulsion of three or more rectus

muscles, the surgical goal should pivot from salvage to primary enucleation. The final chapter of this case is the successful prosthetic rehabilitation, a process that highlights the synergy between surgical technique and ocularist artistry. A primary enucleation that preserves the conjunctival fornices and orbital volume provides the ideal foundation for a custom prosthesis. The ocularist's work is a crucial component of the patient's recovery, transforming a disfiguring injury into a state of restored normalcy. The custom-molded, hand-painted prosthesis restores facial symmetry, supports eyelid architecture, and, by moving in concert with the remaining orbital tissues, provides a lifelike appearance that is essential for psychosocial recovery. This case also opens doors to translational research questions. The inability of pre-operative CT to delineate the muscle and nerve damage highlights a critical technological gap. This generates a clear need for the development of advanced, acute imaging modalities.

4. Conclusion

This case of a severe handlebar-induced traumatic globe luxation illustrates a complex clinical scenario where the ultimate surgical decision was dictated by a rare and devastating pattern of injury. It strongly suggests that the paradigm for managing TGL should include a critical assessment of extraocular muscle integrity as a primary determinant of globe viability, a factor that can, in certain circumstances, outweigh the anatomical status of the optic nerve. The unique sparing of the superior rectus muscle provides a fascinating insight into the biomechanics of focused orbital trauma. The findings generate the hypothesis that in such injuries, catastrophic muscle avulsion may be a more decisive pathology than previously emphasized. Ultimately, this report details a scenario where primary enucleation should not be viewed as a treatment failure, but rather as a successful, patient-centered, and definitive therapeutic strategy. It is a choice that preempts inevitable suffering and provides the most direct and effective pathway toward complete functional and cosmetic rehabilitation.

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