

OPEN ACCESS

SUBMITTED 03 September 2025

ACCEPTED 02 October 2025

PUBLISHED 01 November 2025

VOLUME Vol.06 Issue11 2025

COPYRIGHT

© 2025 Original content from this work may be used under the terms of the creative commons attributes 4.0 License.

Longitudinal Retinal Imaging In Stage IV-A Retinopathy Of Prematurity: A Case Series On Disease Progression And Management

Dr. Rohan M. Sharma

Department of Pediatric Ophthalmology and Strabismus, Aravind Eye Hospital, Madurai, India

Dr. Priya K. Singh

Neonatal Intensive Care Unit, Department of Pediatrics, All India Institute of Medical Sciences (AIIMS), New Delhi, India

Abstract: Background: Retinopathy of Prematurity (ROP) is a leading cause of childhood blindness, with advanced stages like Stage IV-A presenting significant management challenges. While wide-field retinal imaging has revolutionized ROP screening, its utility in monitoring the detailed evolution of the disease, particularly in advanced stages, is not extensively documented. This study aims to illustrate the longitudinal progression of Stage IV-A ROP using serial retinal imaging.

Methods: This retrospective case series was conducted at a tertiary neonatal care center. Premature infants diagnosed with Stage IV-A ROP who underwent serial examinations with a wide-field infant retinal camera were included. A comprehensive review of their medical records and sequential retinal images was performed. The evolution of key pathological features, including the extent of retinal detachment, vascular abnormalities, and ridge characteristics, was documented and correlated with clinical management decisions.

Results: The case series details the clinical and imaging journey of infants with Stage IV-A ROP. The longitudinal images provided a dynamic view of the disease, revealing subtle yet critical changes in the retinal detachment and vascular activity that influenced the timing and choice of interventions, including laser photocoagulation and vitreoretinal surgery. The imaging data served as an objective tool for monitoring

treatment response and guiding subsequent management steps.

Conclusion: Serial wide-field retinal imaging is an indispensable tool for the nuanced management of Stage IV-A ROP. It provides crucial, objective insights into disease dynamics that surpass traditional examinations, facilitating more precise and timely clinical decision-making. The integration of such technology into standard care protocols is vital for improving outcomes in this vulnerable population.

Keywords: Retinopathy of Prematurity, Retinal Imaging, Stage IV-A ROP, Wide-Field Imaging, Case Series, Preterm Infants, Retinal Detachment.

Introduction:

1.1 Background on Retinopathy of Prematurity (ROP)

The global increase in the survival rates of premature infants represents one of the most significant triumphs of modern medicine. However, this success has brought with it a corresponding rise in the incidence of morbidities associated with preterm birth. The World Health Organization (WHO) estimates that 15 million babies are born preterm every year, a figure that continues to rise, making complications from prematurity the leading cause of death among children under five years of age [1]. Among the most devastating of these complications is Retinopathy of Prematurity (ROP), a vasoproliferative disorder of the developing retina that stands as a primary cause of preventable childhood blindness worldwide.

The pathophysiology of ROP is intrinsically linked to the delicate process of retinal vascularization, which normally proceeds from the optic nerve towards the retinal periphery and is not complete until full-term gestation. In a premature infant, this process is abruptly interrupted. The peripheral retina, now existing in a relatively hyperoxic extrauterine environment compared to the hypoxic intrauterine state, experiences a downregulation of vascular endothelial growth factor (VEGF), leading to a cessation of normal vessel growth (Phase I). As the infant matures and the avascular retina becomes increasingly metabolic, a state of pathological hypoxia ensues, triggering an uncontrolled overproduction of VEGF and other angiogenic factors. This leads to neovascularization, the hallmark of ROP (Phase II), where fragile, abnormal blood vessels proliferate at the junction of the vascular and avascular retina. These vessels are prone to leakage and hemorrhage, leading to the formation of fibrous scar tissue that can contract and pull on the retina, causing tractional retinal

detachment and severe vision loss.

The severity of ROP is classified according to an international system based on its location (Zone I, II, or III), its stage (Stages 1 through 5), and the presence of "plus disease," which refers to the venous dilation and arteriolar tortuosity of posterior pole vessels, indicating significant disease activity. Stage I is characterized by a faint demarcation line, progressing to a more defined ridge in Stage II, and extraretinal fibrovascular proliferation in Stage III. Stage IV marks the onset of a partial retinal detachment, which is subdivided into Stage IV-A (extrafoveal detachment) and Stage IV-B (foveal detachment). Stage V represents a total, funnel-shaped retinal detachment, often resulting in irreversible blindness. The management of ROP is critically time-sensitive; treatment with laser photocoagulation or anti-VEGF injections is typically indicated for severe Stage III disease to prevent progression to retinal detachment. Once a detachment occurs, particularly in Stage IV-A, the management becomes significantly more complex, often requiring advanced vitreoretinal surgery [3].

1.2 The Challenge of ROP Screening and Diagnosis in India

While ROP is a global problem, its burden is disproportionately felt in middle-income countries like India, which are experiencing a "third epidemic" of ROP. This epidemic is characterized by a high number of at-risk infants due to improving neonatal care, but concurrently, a healthcare infrastructure that is often ill-equipped to provide the systematic screening and treatment required. The debate surrounding the need for India-specific ROP screening guidelines is ongoing, driven by the observation that ROP can occur in larger, more mature infants in India compared to their counterparts in high-income nations [2]. Factors such as inconsistent oxygen monitoring, sepsis, and fluctuating clinical stability contribute to a different risk profile, necessitating screening criteria that may need to be broader than those recommended by Western bodies [2, 6].

Compounding this issue are the significant socioeconomic and logistical barriers to effective screening. The cost of equipment, particularly traditional indirect ophthalmoscopes and advanced imaging systems, can be prohibitive for many institutions [7]. Furthermore, there is a severe shortage of ophthalmologists trained and willing to perform ROP screening, especially in rural and semi-urban areas where a large proportion of at-risk infants are born. This disparity in access to care means that many infants are either never screened or are screened too late for effective intervention.

Even when screening is available, the diagnosis of ROP, especially the determination of plus disease, is notoriously subjective. Studies have demonstrated only moderate agreement among experienced pediatric ophthalmologists in diagnosing plus and pre-plus disease based on clinical examination alone [8]. This variability is a critical issue, as the decision to initiate potentially vision-saving treatment often hinges on this very assessment. The subjective nature of the diagnosis can lead to both under-treatment, risking disease progression, and over-treatment, exposing fragile infants to unnecessary procedural risks.

1.3 Advancements in Retinal Imaging for ROP

In response to these challenges, the field of pediatric ophthalmology has witnessed transformative advancements in retinal imaging technology. Digital retinal imaging, particularly with wide-field cameras, has emerged as a powerful tool to objectify and enhance the screening, diagnosis, and monitoring of ROP. These systems can capture high-resolution images of the infant retina, providing a detailed and permanent record of the retinal vasculature that can be reviewed, compared over time, and shared remotely.

A significant breakthrough in making this technology more accessible has been the development of novel, low-cost, wide-field infant retinal cameras. For instance, the "Neo" camera, developed in India, provides a portable and affordable solution for capturing high-quality retinal images, making it suitable for use in diverse clinical settings, including neonatal intensive care units (NICUs) with limited resources [4]. The safety and efficacy of such devices have been validated for use in premature infants, paving the way for wider adoption [4].

The integration of such imaging technology with telemedicine platforms has further amplified its impact. Programs like the Karnataka Internet Assisted Diagnosis for Retinopathy of Prematurity (KIDROP) in India have demonstrated the profound effectiveness of a telemedicine-based service delivery model [5]. In this model, trained technicians in remote centers capture retinal images, which are then transmitted to a central team of ROP experts for diagnosis and management advice. This approach has successfully bridged the gap between the availability of at-risk infants and the scarcity of expert ophthalmologists, democratizing access to high-quality care and significantly improving the reach of ROP screening services [9].

1.4 Literature Gap and Study Rationale

The existing body of literature provides extensive

coverage of ROP screening guidelines [6], surgical techniques and outcomes [3], and the impact of innovative service delivery models [9]. However, there remains a relative scarcity of detailed, longitudinal reports that document the temporal evolution of advanced ROP using modern imaging modalities. Stage IV-A ROP represents a critical juncture in the disease course, where the retina has begun to detach but the macula is still attached, presenting a window of opportunity for intervention to preserve central vision. The management decisions at this stage are complex and rely heavily on a nuanced understanding of the disease's trajectory.

While single-point imaging is invaluable for diagnosis, serial imaging offers the potential to visualize the dynamic nature of the disease—the subtle shifts in traction, the waxing and waning of vascular activity, and the early response to treatment. Documenting these changes is essential for building a more comprehensive understanding of the pathophysiology of retinal detachment in ROP. Therefore, this case series aims to fill this gap by documenting and analyzing the evolution of Stage IV-A ROP through a series of sequential wide-field retinal images. By presenting a detailed account of disease progression and its correlation with clinical management, we seek to highlight the indispensable utility of this technology in navigating the complexities of advanced ROP and optimizing patient care.

2.0 Methods

2.1 Study Design and Patient Selection

This study was conducted as a retrospective case series. The clinical data and retinal images were sourced from the ROP database of a tertiary-level neonatal care unit and ophthalmology center, renowned for its high volume of premature infant care and its integrated ROP screening program. The study protocol was reviewed and approved by the Institutional Review Board and Ethics Committee. Given the retrospective nature of the analysis of de-identified data, the requirement for individual parental consent for this specific study was waived. All procedures and examinations described were part of the standard of care provided to the infants, and initial consent for screening and potential treatment had been obtained from the parents or legal guardians at the time of care.

Infants were included in this case series if they met the following criteria: (1) a confirmed diagnosis of Stage IV-A Retinopathy of Prematurity in at least one eye, based on expert clinical examination and supported by wide-field retinal imaging; (2) born at or before 34 weeks of gestational age or with a birth weight of 2000 grams or less; (3) had undergone a minimum of three wide-field retinal imaging sessions, including a baseline image at

the time of Stage IV-A diagnosis and subsequent follow-up images; and (4) had a complete medical record available for review. Exclusion criteria were: (1) infants with Stage IV-B or Stage V ROP at the time of initial diagnosis; (2) presence of other significant congenital ocular abnormalities that could confound the retinal findings, such as persistent fetal vasculature or congenital cataracts; and (3) incomplete or poor-quality imaging records that precluded detailed longitudinal analysis.

2.2 Screening and Diagnostic Protocol

All infants admitted to the NICU who met the institutional ROP screening criteria—typically infants with a birth weight ≤ 1700 grams or a gestational age of ≤ 34 weeks, as well as larger, more mature infants with a history of clinical instability—were enrolled in the screening program. This protocol is broadly aligned with national and international recommendations [2, 6]. The first screening examination was performed at 4 weeks of chronological age or at 31 weeks of postmenstrual age, whichever was later. Subsequent examinations were scheduled based on the findings, with the frequency ranging from every few days to every two weeks, as determined by the examining pediatric ophthalmologist.

All examinations were performed by an experienced pediatric ophthalmologist. The diagnosis of ROP stage, zone, and extent, as well as the presence or absence of plus disease, was made using binocular indirect ophthalmoscopy with a 28D lens after pupillary dilation with a combination of topical phenylephrine (2.5%) and tropicamide (0.5%). The diagnosis of Stage IV-A ROP was confirmed when a partial, extrafoveal tractional or rhegmatogenous retinal detachment was identified.

2.3 Imaging Technique

Digital retinal imaging was performed using a wide-field portable infant retinal camera (e.g., Neo, RetCam). All imaging sessions were conducted at the infant's bedside in the NICU to minimize transport and handling stress. Prior to imaging, the infant's pupils were dilated as per the screening protocol. A single drop of topical proparacaine hydrochloride (0.5%) was instilled for local anesthesia. A sterile, disposable lid speculum was used to gently open the eyelids, and a water-based gel was applied to the tip of the camera lens to serve as a coupling medium and ensure clear image acquisition.

The imaging technician or ophthalmologist captured a series of images of each eye, focusing on obtaining a clear view of the posterior pole, the optic disc, the macula, and the retinal periphery in all quadrants. The

goal was to comprehensively document the entire extent of the vascularized and avascular retina, the junctional ridge, and the area of retinal detachment. Specific attention was paid to documenting the features of plus disease, including the tortuosity and dilation of the major retinal vessels. All imaging procedures were performed with continuous monitoring of the infant's vital signs, and safety measures were in place to manage any potential adverse events, such as bradycardia or desaturation, although none were noted in the selected cases. The schedule of imaging for the cases included in this series was determined by the clinical need for close monitoring, with sessions typically performed at baseline diagnosis and repeated at intervals of 1-2 weeks or more frequently if rapid progression was suspected.

2.4 Data Collection and Analysis

A comprehensive review of the electronic medical records and the digital imaging database was conducted for each selected case. The following data points were systematically extracted: maternal demographics, infant's gestational age at birth, birth weight, sex, Apgar scores, and significant neonatal comorbidities (e.g., respiratory distress syndrome, sepsis, intraventricular hemorrhage). The complete ROP-related timeline was documented, including the age at first screening, the postmenstrual age at diagnosis of each ROP stage, and the details of any interventions performed (e.g., laser photocoagulation, anti-VEGF injections, vitreoretinal surgery).

The longitudinal series of retinal images for each case was analyzed qualitatively by a panel of two senior pediatric ophthalmologists. The analysis focused on identifying and describing the temporal evolution of specific pathological features. These included: (1) the location, height, and circumferential extent of the retinal detachment; (2) the appearance of the fibrovascular ridge, including its height and the presence of tractional elements; (3) the degree of vascular tortuosity and dilation (plus disease); (4) the presence of any retinal hemorrhages or exudates; and (5) the appearance of the retina post-intervention, noting any signs of reattachment or persistent detachment. The analysis was descriptive, aiming to create a detailed narrative of the disease's progression as visualized through the sequence of images.

3.0 Results

The demographic and clinical characteristics of the three infants included in this case series are summarized in Table 1. Each case presented a unique trajectory of Stage IV-A ROP, highlighting the heterogeneity of the disease and the value of individualized monitoring.

Table 1: Summary of Patient Demographics and Clinical Characteristics

Feature	Case 1	Case 2	Case 3
Gestational Age (weeks)	28	26	25
Birth Weight (grams)	1100	850	720
Sex	Male	Female	Male
Major Comorbidity	Respiratory Distress	Necrotizing Enterocolitis	Sepsis
PMA at Stage IV-A Dx	37 weeks	37 weeks	38 weeks
Initial Treatment	Laser Photocoagulation	Intravitreal Bevacizumab	Laser Photocoagulation
Follow-up Intervention	Observation	Vitrectomy	Intravitreal Bevacizumab + Vitrectomy
Final Anatomical Outcome	Spontaneous Reattachment	Successful Reattachment	Successful Reattachment
<i>PMA = Postmenstrual Age; Dx = Diagnosis</i>			

3.1 Case Presentations

Case 1: Controlled Progression Following Laser Photocoagulation

Infant A was a male neonate born at 28 weeks of gestational age with a birth weight of 1100 grams. His neonatal course was complicated by respiratory distress syndrome requiring mechanical ventilation for two weeks. ROP screening was initiated at 32 weeks postmenstrual age (PMA). He was diagnosed with Stage II ROP in Zone II with pre-plus disease. By 35 weeks PMA, the disease had rapidly progressed to Stage III in Zone II with marked plus disease, for which

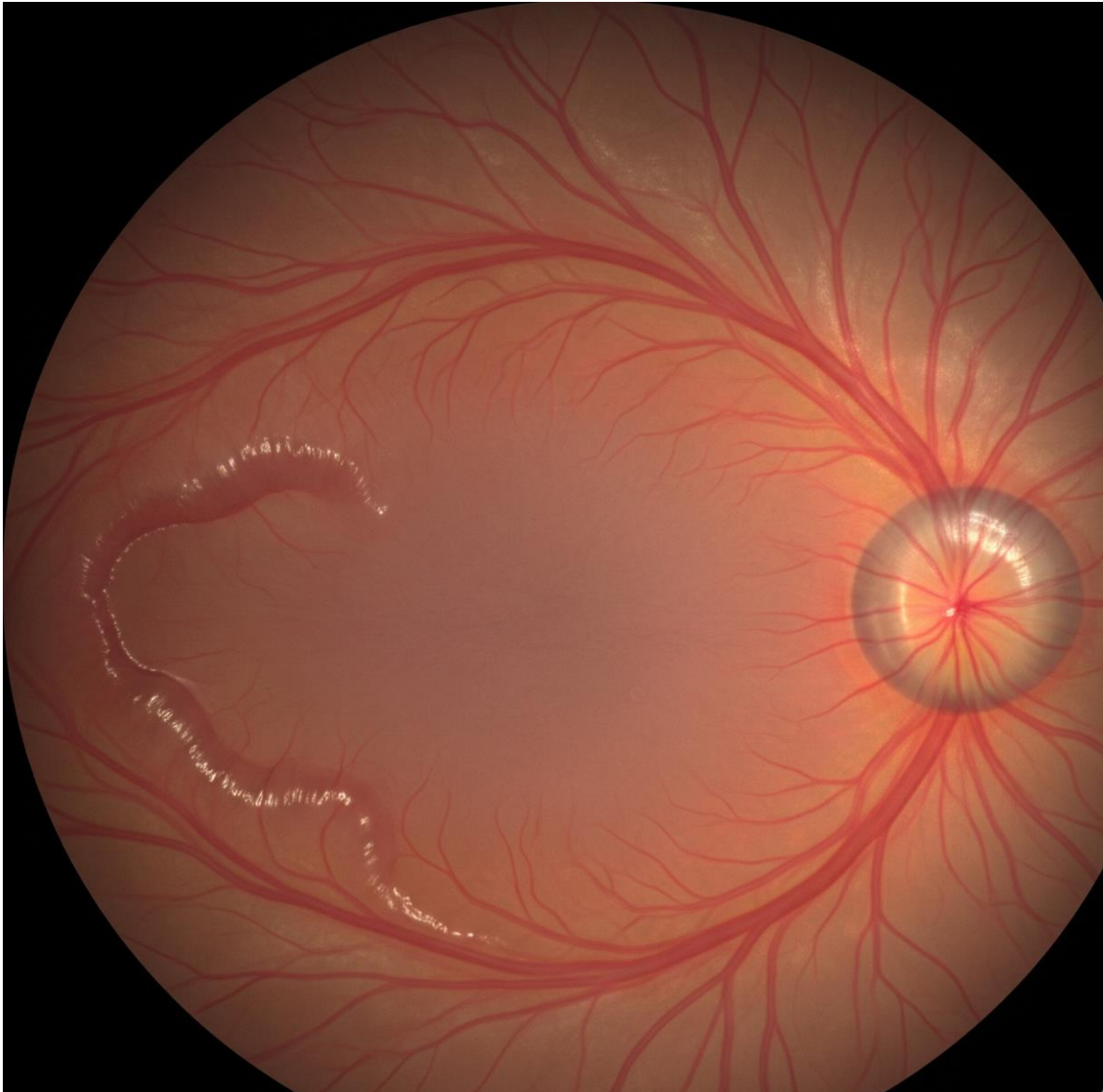
he underwent confluent laser photocoagulation to the avascular retina in both eyes.

Despite laser treatment, weekly follow-up imaging revealed the development of a shallow, sickle-shaped tractional retinal detachment in the temporal periphery of the right eye at 37 weeks PMA, consistent with a diagnosis of Stage IV-A ROP. The baseline wide-field image at this time (Figure 1A) clearly delineated a well-defined fibrovascular ridge with visible tractional forces pulling the retina anteriorly over an area of approximately four clock hours. The posterior pole vessels remained tortuous, though slightly less

engorged than before the laser treatment.

Serial imaging was performed weekly. The images at 38 and 39 weeks PMA showed a remarkable stability in the detachment. The tractional forces did not appear to increase, and the height of the detachment remained unchanged. Importantly, the fovea remained clearly visible and flat. The plus disease continued to slowly regress, with vessels at the posterior pole appearing less tortuous in each subsequent image. Based on this documented stability on serial imaging, a decision was made to continue with close observation

rather than proceeding immediately to vitreoretinal surgery. By 42 weeks PMA, imaging showed early signs of spontaneous reattachment at the posterior border of the detachment (Figure 1B). The infant was discharged with a plan for continued outpatient follow-up. This case illustrated the utility of serial imaging in confidently managing a case of Stage IV-A ROP non-surgically, avoiding the significant risks of surgery in a stable situation.



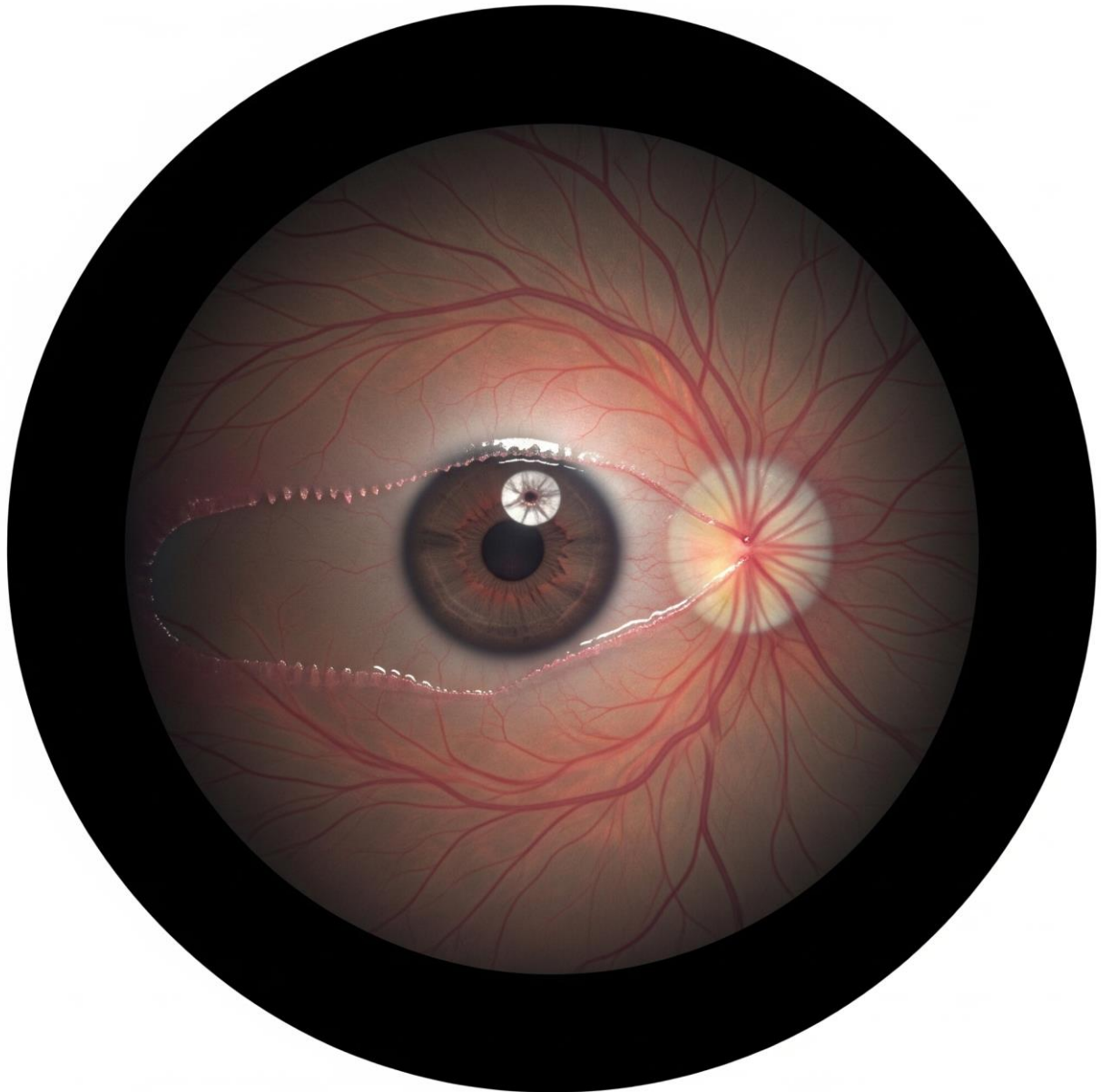


Figure 1: Case 1 - Right Eye

- (A) Wide-field retinal image at 37 weeks PMA at the time of Stage IV-A ROP diagnosis. The image shows a shallow, temporal, extrafoveal retinal detachment (white arrows) with a prominent fibrovascular ridge. Laser scars from prior treatment are visible in the periphery.
- (B) Follow-up image at 42 weeks PMA. The detachment has become significantly flatter with evidence of early spontaneous reattachment. The plus disease has resolved, and posterior pole vessels appear calm.

Case 2: Rapid Progression Necessitating Urgent Surgical Intervention

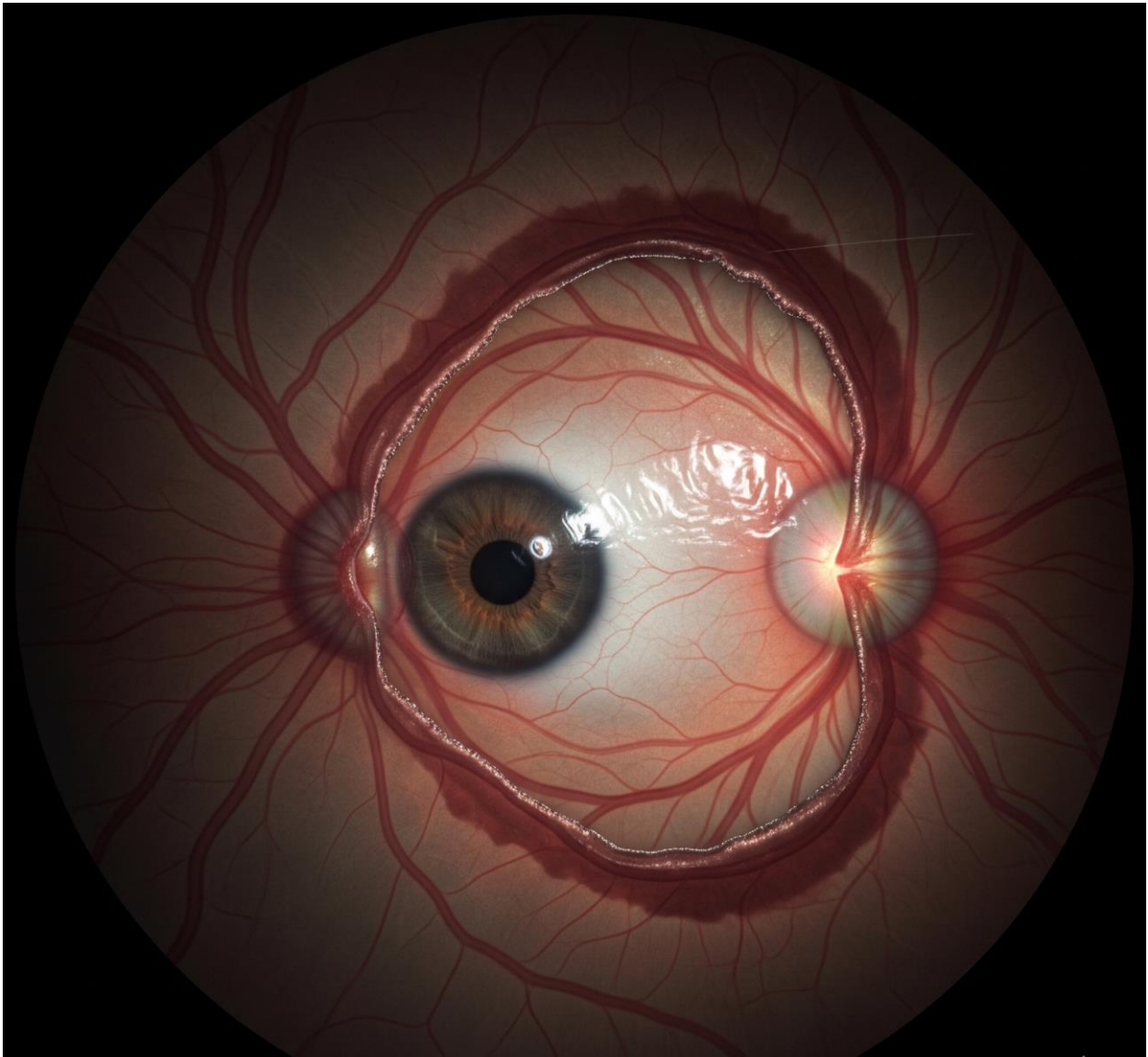
Infant B, a female born at 26 weeks gestation with a birth weight of 850 grams, had a tumultuous neonatal course with necrotizing enterocolitis and culture-

positive sepsis. She was first screened at 31 weeks PMA and was found to have aggressive posterior ROP (AP-ROP) in Zone I. She received an intravitreal injection of bevacizumab (anti-VEGF) in both eyes. The initial response was favorable, with regression of the neovascularization.

However, at 36 weeks PMA, follow-up imaging detected a reactivation of the disease. The right eye developed a broad, elevated ridge with extensive fibrovascular proliferation. Within a week, at 37 weeks PMA, the condition progressed to Stage IV-A ROP. The baseline image (Figure 2A) showed a circumferential tractional detachment extending from the 2 o'clock to the 8 o'clock position, creating a prominent fold that appeared to be exerting traction towards the fovea. The vessels were severely dilated and tortuous, indicative of florid plus disease.

Given the aggressive nature and posterior location, the decision for close monitoring with imaging every 3-4 days was made. The image taken just four days later (Figure 2B) showed a dramatic worsening. The detachment had increased in height, and the tractional fold was now visibly closer to the foveal center. The rapid progression documented by the images prompted an urgent consultation with the vitreoretinal surgery team. The detailed images were invaluable for surgical planning. The infant underwent a 25-gauge

pars plana lensectomy and vitrectomy [3]. Post-operative imaging at one week showed that the retina was successfully reattached, although some peripheral scarring was noted. This case highlights how high-frequency serial imaging can be critical in identifying rapidly progressing disease, enabling timely surgical intervention to save vision.



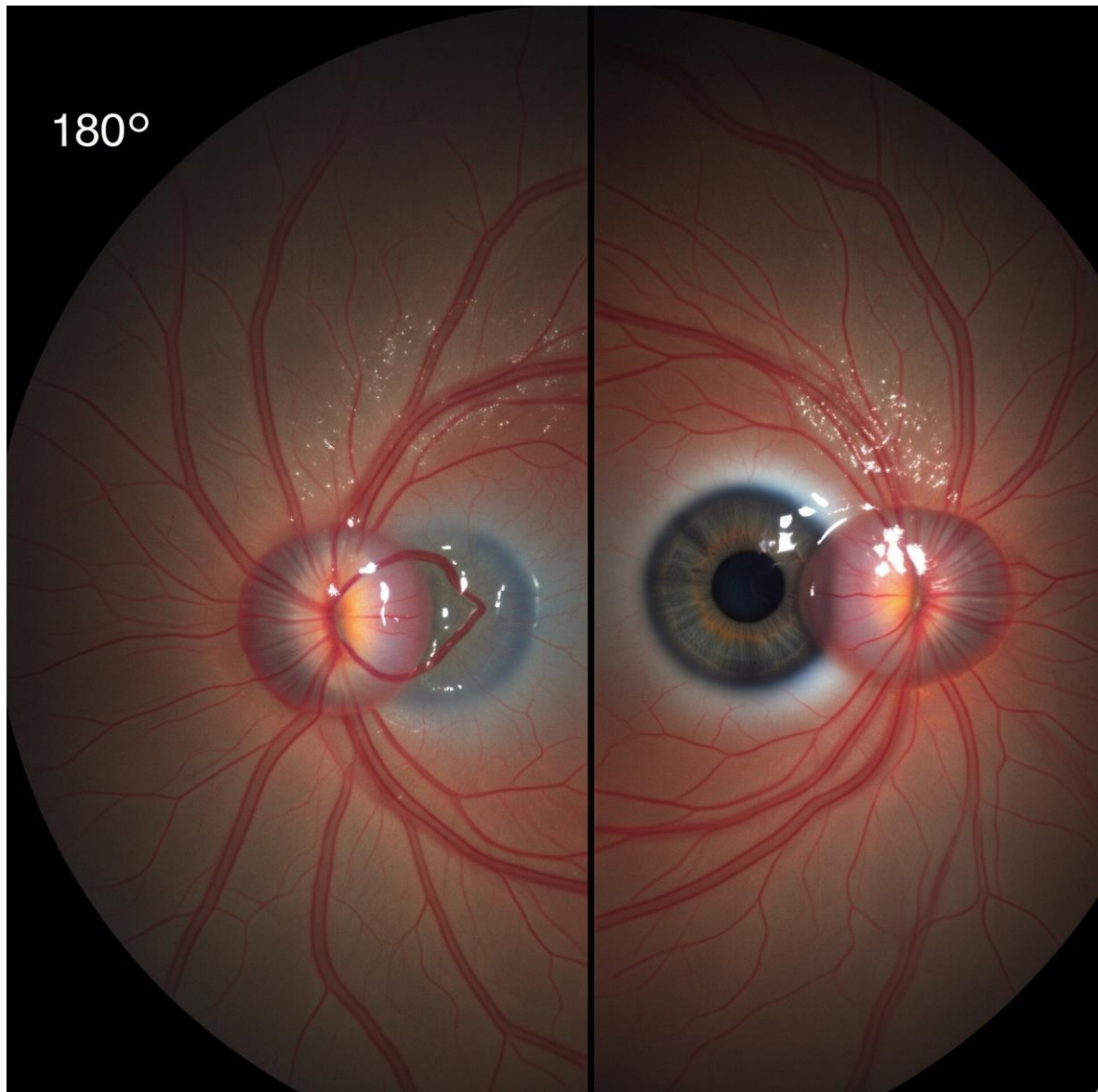


Figure 2: Case 2 - Right Eye

- (A) Image at 37 weeks PMA showing Stage IV-A ROP with a large, circumferential tractional detachment and a prominent retinal fold. Severe plus disease is evident with significant vascular tortuosity and dilation.
- (B) Image taken four days later demonstrating rapid progression. The height of the detachment has increased (white arrows), and the tractional fold is now closer to the macula, indicating worsening of the condition and prompting surgical intervention.

Case 3: Complex Evolution and the Role of Imaging in Guiding Combined Therapy

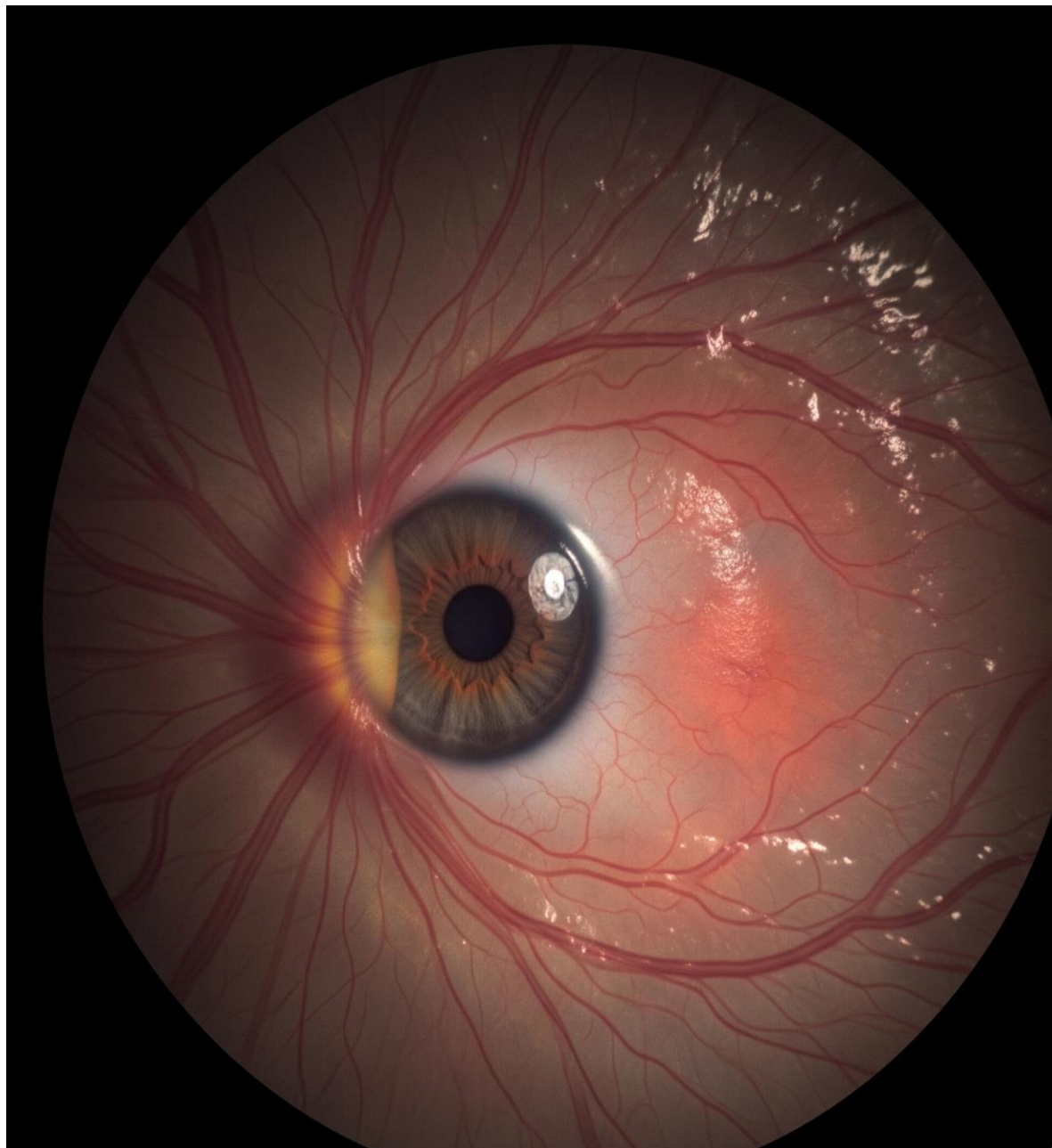
Infant C was an extremely preterm male, born at 25 weeks gestation with a birth weight of 720 grams. He was diagnosed with Stage III ROP in Zone I with plus disease at 34 weeks PMA and was treated with laser photocoagulation. Follow-up was challenging due to persistent media haze from poor pupillary dilation. At

38 weeks PMA, when a clear view was finally obtained with wide-field imaging, a diagnosis of Stage IV-A ROP was made in the left eye. The baseline image (Figure 3A) revealed a multi-quadrant, bullous-appearing retinal detachment in the periphery, with significant exudation and subretinal fluid, a feature less commonly described. The plus disease was severe.

The clinical team decided on a combined therapeutic approach. An intravitreal bevacizumab injection was administered to reduce the vascular activity and exudation, with a plan for subsequent surgery. Serial imaging post-injection was instrumental. The image at one week post-injection (Figure 3B) showed a marked reduction in the plus disease and a partial collapse of the bullous detachment, suggesting a decrease in the exudative component. However, the underlying tractional elements became more apparent as the exudation resolved. The fibrovascular ridge appeared more organized and taut.

This evolution, clearly captured on the image series, informed the timing and strategy for surgery. The surgeons, armed with images showing the location of the primary tractional forces, performed a vitrectomy two weeks post-injection. Post-operative imaging confirmed successful anatomical reattachment of the

retina. This case demonstrates the power of longitudinal imaging in dissecting complex disease pathophysiology (tractional vs. exudative components) and guiding a multi-modal therapeutic strategy.



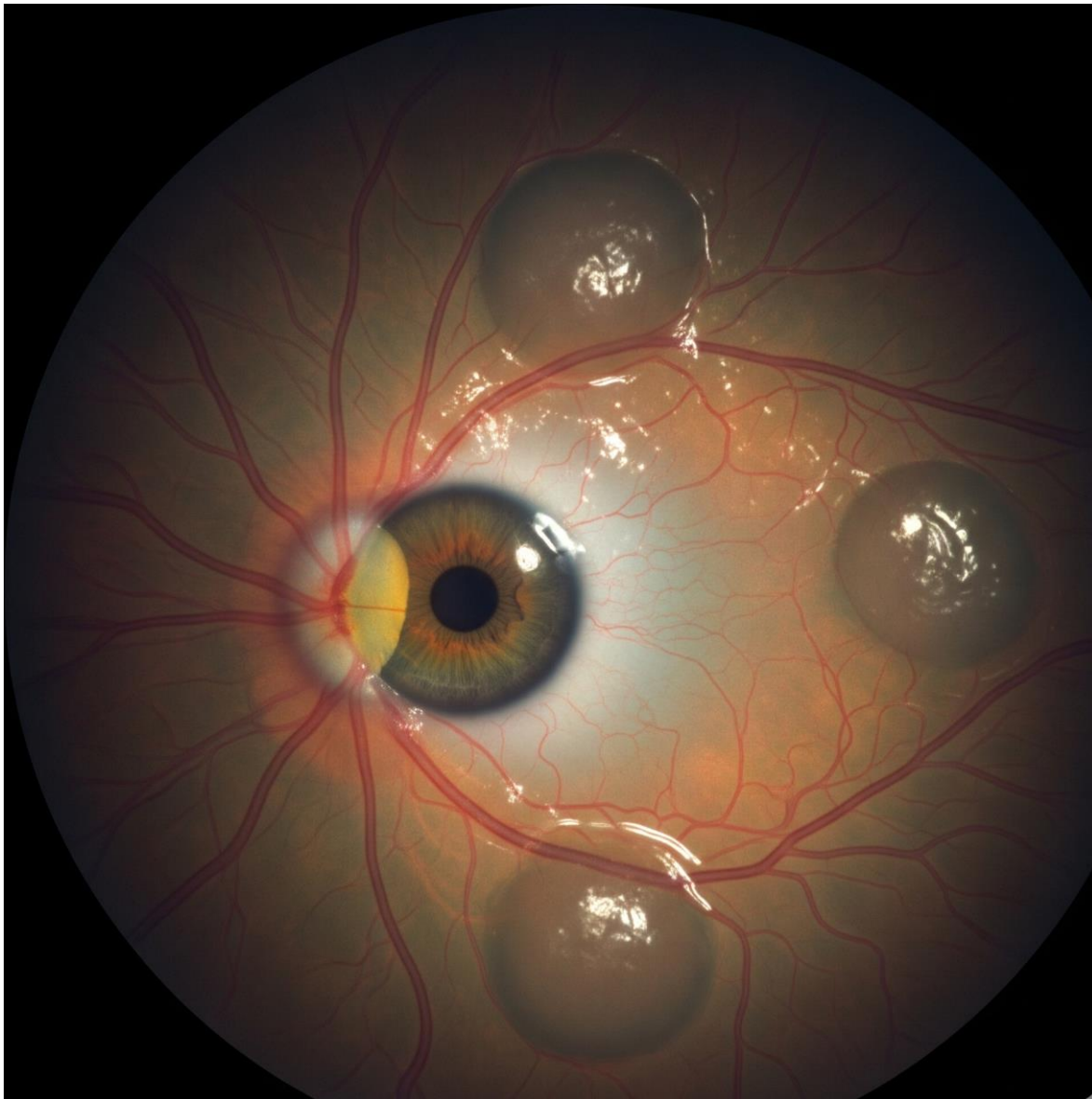


Figure 3: Case 3 - Left Eye

- (A) Baseline image at 38 weeks PMA showing a complex Stage IV-A ROP with a bullous, exudative-appearing detachment in the periphery (white arrows). Marked plus disease is visible at the posterior pole.
- (B) Image taken one week after an intravitreal bevacizumab injection. There is a significant reduction in the exudative component of the detachment and a decrease in vascular engorgement. The underlying tractional forces on the ridge have now become more clearly defined.

3.2 Summary of Cross-Case Findings

Across this series of cases, several key patterns emerged. First and foremost, the use of serial wide-field retinal imaging provided a dynamic and detailed narrative of Stage IV-A ROP progression that was far more informative than what could be gleaned from intermittent clinical examinations alone. The objective, high-resolution documentation allowed for precise tracking of the detachment's configuration, height, and

proximity to the fovea. Secondly, the technology was invaluable in monitoring the response to treatment. In Case 1, it documented stability, allowing for a conservative approach. In contrast, in Case 2, it captured rapid progression, triggering an urgent surgical response. Thirdly, the images facilitated a deeper understanding of the disease's character in individual patients. As seen in Case 3, imaging helped differentiate between exudative and tractional components, which had direct implications for the therapeutic plan. In all cases, the images served as a critical tool for communication and planning among the ophthalmology, neonatology, and surgical teams, fostering a collaborative and well-informed approach to managing these complex cases.

4.0 Discussion

4.1 Principal Findings and Interpretation

The case series presented in this paper offers a granular, longitudinal view of the clinical course of Stage IV-A Retinopathy of Prematurity, a critical juncture in the

disease where the potential for preserving vision still exists but is acutely threatened. The overarching finding from our analysis is the profound heterogeneity in the progression of this disease stage. Our cases demonstrate that a single diagnostic label of "Stage IV-A ROP" encompasses a wide spectrum of clinical behaviors, ranging from relative stability amenable to conservative observation (Case 1), to explosive progression demanding immediate surgical intervention (Case 2), and complex pathophysiological presentations that benefit from a carefully sequenced, multi-modal therapeutic approach (Case 3).

This variability highlights the inherent limitations of a management strategy based solely on single-point clinical examinations. The decision to observe or to subject a fragile, premature infant to the significant physiological stress and risks of vitreoretinal surgery [3] is one of the most challenging in pediatric ophthalmology. The findings from this series strongly suggest that incorporating serial wide-field retinal imaging into the management protocol provides a vital, objective data stream that can powerfully inform this decision-making process. By documenting the rate and character of change over time, serial imaging allows for a more dynamic and personalized risk assessment. The documented stability in Case 1, for example, provided the clinical confidence to defer surgery, whereas the rapid worsening captured in Case 2 presented an unequivocal mandate for urgent intervention. This ability to tailor management based on the individual disease trajectory, rather than a static diagnosis, represents a significant step towards optimizing outcomes and practicing a more precise form of medicine for these vulnerable infants.

4.2 The Transformative Role of Technology in Modern ROP Management

The insights gleaned from our cases are inextricably linked to the technological advancements that made them possible. The shift from traditional indirect ophthalmoscopy to wide-field digital retinal imaging is not merely an incremental improvement; it is a paradigm shift in the way ROP is diagnosed, monitored, and managed.

From Subjective Sketch to Objective Record

For decades, the gold standard for documenting ROP findings was a hand-drawn sketch in the patient's chart, supplemented by clinical notes. While valuable in the hands of a skilled expert, this method is inherently subjective, operator-dependent, and lacks the detail to capture subtle but critical changes over time. The quality of documentation varies, and the interpretation of another clinician's drawing can be ambiguous. In contrast, wide-field digital imaging

provides a high-resolution, objective, and permanent photographic record. This has several profound advantages. First, it allows for direct, side-by-side comparison of retinal images over time, making the assessment of progression or regression far more precise. Changes in the height of a detachment, the tortuosity of a vessel, or the extent of a fibrovascular ridge can be clearly visualized and even quantified, removing a significant degree of subjectivity.

This objectivity is particularly crucial in the assessment of plus disease, a key indicator of disease activity and a primary trigger for treatment. The diagnosis of plus disease has been shown to have only moderate inter-observer agreement among experts, even in the era of standardized photographic comparison sets [8]. The subtle gradations between "no plus," "pre-plus," and "plus" disease can be difficult to discern and communicate. A digital image freezes the retina in time, allowing for careful, unhurried examination of vascular characteristics. This static image can be magnified, and its contrast can be adjusted, aiding in the identification of early signs that might be missed during a brief examination of a moving infant. As a result, digital imaging provides a more reliable and reproducible basis for the diagnosis and longitudinal tracking of plus disease, leading to more consistent and appropriate treatment decisions.

Democratizing Expertise through Accessible Technology and Telemedicine

Perhaps the most significant impact of modern imaging technology is its ability to democratize access to expert ROP care. The global burden of ROP is highest in regions where the number of trained ophthalmologists is lowest [2]. This "expertise gap" is a primary barrier to preventing childhood blindness. Two parallel technological developments are directly addressing this challenge: the creation of affordable imaging devices and the implementation of telemedicine networks.

The development of innovative, low-cost, and portable wide-field cameras, such as the Indian-developed "Neo" camera, is a pivotal achievement [4]. These devices dramatically lower the financial barrier to entry for many hospitals and clinics in low- and middle-income countries, which previously found the cost of legacy imaging systems prohibitive [7]. Their portability allows imaging to be performed at the bedside in any NICU, eliminating the need for a dedicated imaging suite and the risks associated with transporting fragile infants. By making high-quality imaging feasible in a wider range of settings, these devices are the essential first step in building a scalable ROP screening infrastructure.

This infrastructure's true power is unlocked when imaging devices are connected to telemedicine

platforms. Models like the Karnataka Internet Assisted Diagnosis for Retinopathy of Prematurity (KIDROP) program have provided a blueprint for how to effectively leverage technology to bridge the expertise gap [5, 9]. In the KIDROP model, trained technicians, who are not ophthalmologists, are deployed to district-level NICUs to capture retinal images. These images are then uploaded to a central server where a team of ROP experts can review them remotely and provide an immediate diagnosis and management plan. This "store-and-forward" tele-ophthalmology approach is a force multiplier; it allows a small team of experts to serve a vast geographic area and a large population of at-risk infants. It effectively uncouples the diagnostic act from the physical location of the expert, ensuring that an infant born in a remote, underserved area can receive the same level of diagnostic expertise as one born in a major metropolitan center [9]. This model not only improves access to care but also ensures a standardized quality of diagnosis, directly addressing the challenge of diagnostic variability [8].

4.3 Expanding Clinical, Educational, and Communicative Implications

The integration of routine imaging into ROP management has implications that extend far beyond initial diagnosis and monitoring. It fundamentally changes how clinicians plan treatments, how trainees learn, and how medical teams communicate with families.

Enhancing Surgical Precision and Therapeutic Strategy

As illustrated in our case series, particularly in Cases 2 and 3, wide-field images are invaluable tools for pre-operative planning. A detailed clinical note might describe a Stage IV-A detachment, but a high-resolution image reveals its precise topography. The surgeon can study the exact location and orientation of tractional forces, identify the points of most significant vitreoretinal adhesion, and visualize the relationship of the detachment to key structures like the ora serrata and the lens. This detailed anatomical map allows the surgeon to anticipate challenges and devise a more precise and effective surgical strategy, potentially improving anatomical success rates and reducing operative time [3]. Furthermore, as seen in Case 3, imaging can help untangle complex presentations and guide a multi-modal approach. The ability to visualize the differential response of exudative and tractional components to an anti-VEGF injection provided critical information that optimized the timing and goals of the subsequent vitrectomy.

Revolutionizing Medical Education and Training

The creation of large, annotated digital imaging

archives represents a revolutionary resource for medical education. ROP is a disease of subtle signs and rapid changes, and developing expertise traditionally requires years of direct clinical exposure under mentorship. A digital archive allows residents and fellows to review thousands of cases, spanning the entire spectrum of the disease from mild to severe. They can follow individual cases longitudinally, observing how Stage I progresses to Stage III, how plus disease evolves, and how the retina responds to laser or anti-VEGF therapy. This virtual, asynchronous learning experience accelerates the development of pattern recognition skills and provides a standardized, high-quality educational foundation that is not dependent on the specific case mix at their institution during their training period. It allows for the creation of teaching modules, self-assessment quizzes, and certification programs that can elevate the standard of ROP training globally.

Fostering Communication and Empowering Families

Finally, the communicative power of a clear image cannot be overstated. Explaining the concept of an abnormal blood vessel or a detached retina to a worried parent can be difficult with words alone. Presenting them with an actual image of their child's eye makes the condition immediately tangible and understandable. Clinicians can use the images to point out the specific areas of concern, explain the rationale for a proposed treatment, and later, show the post-treatment results. This visual communication fosters a deeper understanding, builds trust between the medical team and the family, and can significantly improve adherence to complex and often arduous follow-up schedules. Empowering parents with knowledge about their child's condition makes them active partners in the care process, which is essential for achieving the best possible long-term outcomes.

4.4 Limitations of the Study

It is essential to acknowledge the inherent limitations of this study. As a case series, the primary limitation is the small number of patients and the absence of a control group. Therefore, the observations made are descriptive and cannot be used to establish causality or determine the incidence of different progression patterns in the broader population of infants with Stage IV-A ROP. The findings may not be generalizable to all infants, as the course of ROP can be influenced by a multitude of systemic factors and variations in clinical care.

The retrospective nature of the study is another limitation. The data and images were collected as part of routine clinical care, not for a research protocol. This could introduce a potential for selection bias, as cases

with more complete and interesting imaging series may have been more likely to be included. Furthermore, the timing of imaging was based on clinical judgment rather than a predetermined schedule, which may have influenced the observed patterns. A prospective study with a larger cohort and a standardized imaging protocol would be required to validate these preliminary findings.

4.5 Future Directions and Conclusion

This case series opens the door for several avenues of future research. Large-scale, prospective, multicenter studies are needed to systematically document the natural history and post-treatment course of Stage IV-A ROP using standardized imaging protocols. Such studies could help identify specific imaging biomarkers that are predictive of rapid progression, which would be immensely valuable for clinical decision-making. There is also significant potential for the application of artificial intelligence (AI) and machine learning algorithms to this area. AI models could be trained on large datasets of sequential retinal images to automatically quantify changes in vascular tortuosity or detachment area over time, providing an objective and powerful decision support tool for clinicians and potentially predicting disease progression before it is apparent to the human eye.

In conclusion, this case series reinforces and highlights the indispensable role of longitudinal wide-field retinal imaging in the modern management of Stage IV-A Retinopathy of Prematurity. This technology provides a window into the dynamic pathophysiology of the disease, transforming a subjective and often uncertain clinical picture into an objective, trackable narrative. By enabling a more nuanced understanding of disease progression and treatment response, serial imaging empowers clinicians to make more informed, timely, and personalized decisions. The integration of this technology into the standard of care—leveraged through accessible devices and scalable telemedicine networks—is not merely an enhancement but a fundamental and necessary evolution in our global fight against ROP-related blindness.

References

1. WHO Fact-sheets Preterm birth. Available from: <https://www.who.int/news-room/fact-sheets/detail/preterm-birth>
2. Honavar, Santosh G. Do we need India-specific retinopathy of prematurity screening guidelines?. Indian Journal of Ophthalmology 67(6): p 711-716, June 2019. https://doi.org/10.4103/ijo.IJO_973_19
3. Sinha R, Talawar P, Ramachandran R, Azad R,

Mohan VK. Perioperative management and post-operative course in preterm infants undergoing vitreo-retinal surgery for retinopathy of prematurity: A retrospective study. J Anaesthesiol Clin Pharmacol 2014; 30: 258-62. <https://doi.org/10.4103/0970-9185.130050>

4. Vinekar A, Rao SV, Murthy S, Jayadev C, Dogra MR, Verma A, Shetty B. A Novel, Low-Cost, Wide-Field, Infant Retinal Camera, "Neo": Technical and Safety Report for the Use on Premature Infants. Transl Vis Sci Technol. 2019 Mar 8; 8(2): 2. <https://doi.org/10.1167/tvst.8.2.2>
5. Karnataka Internet Assisted Diagnosis for Retinopathy of Prematurity (KIDROP) Available at <https://www.narayananeethralaya.org/pediatric-ophthalmology/kidrop/>
6. Fierson WM. Screening Examination of Premature Infants for Retinopathy of Prematurity. Pediatrics. 2018 Dec; 142(6): e20183061. <https://doi.org/10.1542/peds.2018-3061>
7. Moshfeghi, D. and Capone, A. (2008) Do cost concerns limit screening for retinopathy of prematurity? Retinal Physician. Available at: <https://www.retinalphysician.com/issues/2008/september-2008/do-cost-concerns-limit-screening-for-retinopathy-o> (Accessed: 05 May 2024).
8. Wallace D, Quinn G, Freedman S, Chiang M. Agreement among pediatric ophthalmologists in diagnosing plus and pre-plus disease in retinopathy of prematurity. J AAPOS. 2008; 12: 352-356. <https://doi.org/10.1016/j.jaapos.2007.11.022>
9. Vinekar A, Bhende P. Innovations in technology and service delivery to improve Retinopathy of Prematurity care. Community Eye Health. 2018; 31(101): S20-S22. PMID: 30275664; PMCID: PMC6157804