

Content available at: https://www.ipinnovative.com/open-access-journals

Indian Journal of Clinical and Experimental Ophthalmology

ONNI DUBLICATION

Journal homepage: www.ijceo.org

Case Series

Optic nerve head: A diagnostic Saga!

Jyoti Bhatt1*

¹Regional Institute Ophthalmology, Sitapur, Uttar Pradesh, India



ARTICLE INFO

Article history: Received 19-01-2024 Accepted 18-04-2024 Available online 30-09-2024

Keywords:
Optic neuropathy
Optic nerve head edema
Optic disc hyperemia
Optic disc cupping

ABSTRACT

Optic nerve head assessment can be a window to diagnosing a plethora of diseases. It can provide us with a hint into glaucoma as well as many non glaucomatous optic neuropathies associated with neurological conditions like space occupying lesion, raised intracranial tension, demyelination, & systemic diseases like uncontrolled hypertension, diabetes, vascular auto- dysregulation, as well as toxic dietary influences & deficiencies, which can have serious consequences in terms of not just a threat to vision, but also systemic morbidity & mortality.

This is an Open Access (OA) journal, and articles are distributed under the terms of the Creative Commons Attribution-NonCommercial-ShareAlike 4.0 License, which allows others to remix, tweak, and build upon the work non-commercially, as long as appropriate credit is given and the new creations are licensed under the identical terms.

For reprints contact: reprint@ipinnovative.com

1. Methodology & Case Description

This is a retrospective case series of ten patients who presented at a tertiary eye care, with clinical features suggestive of an Optic neuropathy. The patients were evaluated & diagnosed on the basis of clinical judgement & investigations.

2. Case 1

A 43-year-old female patient presented with complaint of dimness of vision left eye since 1 month. BCVa RE-6/6, LE- 6/60, RE-disc edema along nasal margin, LE temporal pallor. Visual fields - LE unequivocal & RE total loss of right half, respecting vertical meridian. Suspecting Foster Kennedy syndrome, patient was further investigated & found to have a bilateral frontal space occupying lesion, for which she underwent bifrontal craniectomy with supraorbital osteotomy & excision of SOL which was adherent to optic chiasma. Biopsy confirmed olfactory groove sarcoidosis.

E-mail address: drjyotismailbox@gmail.com (J. Bhatt).

3. Case 2

61 year old female, diabetic & hypertensive, presented with complain of decreasing vision left eye since 10 days. BCVa, RE-6/9 & LE- 6/36p, RAPD in LE, IOP 19 mmHg BE, .75 & .85 cupping RE & LE respectively with prominent disc pallor. LE showed very advanced visual field defect, but RE field clinched the diagnosis of neurological fields. Investigations revealed pituitary microadenoma compressing left optic nerve & crossing fibres of right eye along optic chiasma, for which she underwent surgery with a successful removal of the tumor.

4. Case 3

A 53 year old male, hypertensive, presented with complaint of decreased vision. BE had shallow anterior chamber, occludable angles, BCVa RE-6/12p, LE-6/9, IOP 18 & 19 mmHg respectively & CCT of 490 microns BE, RE C:D-. 7 & LE C:D-.5 prominent temporal pallor of both optic nerve heads, left homonymous hemianopia. Field defects appeared to break through to the opposite side with a questionable superior arcuate form. He gave a history of left hemeperesis 4 years back with CT brain showing

^{*} Corresponding author.

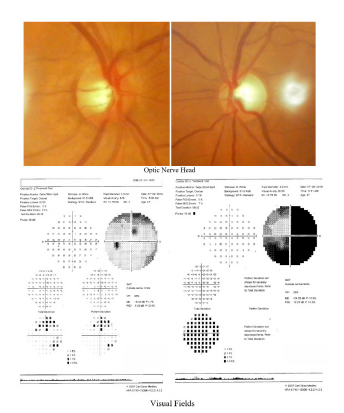


Figure 1: Case 1: Olfactory groove sarcoidosis

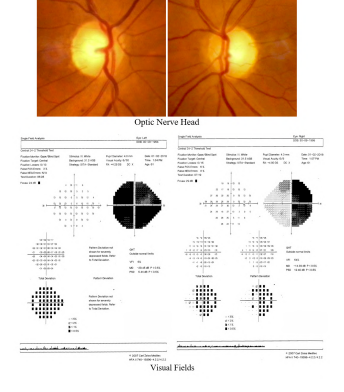


Figure 2: Case 2: Pituitary microadenoma

retrochiasmatic right hemorrhagic infarct.

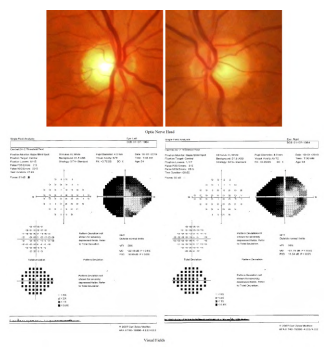


Figure 3: Case 3: Retrochiasmatic right hemorrhagic infarct

5. Case 4

A 24-year-old female patient presented with complaint of dimness of vision both eyes since 1 year, tinnitus & some hearing loss. BCVa 6/60 OU, IOP 17 mmHg, normal ODs with .3 cupping, open angles. She had been diagnosed as a malingerer elsewhere, in view of all normal findings, including a normal color vision test. We did Visual field test, which revealed a central, para central field defect. Suspecting some form of optic neuropathy, a VEP was advised, which showed bilateral prolonged P-100 latencies, confirming an optic neuropathy of a non-glaucomatous type, for which she is being further investigated.

6. Case 5

A 53 year old male patient presented with complaints of progressive dimness of vision both eyes & headache since few months , but a sudden & gross dimness of vision RE , followed soon in LE 20 days back. His Vn both eyes was CF 1 Feet, IOP-15 & 11 mmHg RE, LE respectively, open angles & C:D .75 RE, .3 LE , with disc pallor, almost total color vision defect ,marked field defects both eyes. After an intensive questioning & history taking, he revealed heavy alcohol intake daily since many years. A diagnosis of toxic optic neuropathy/atrophy, was made.

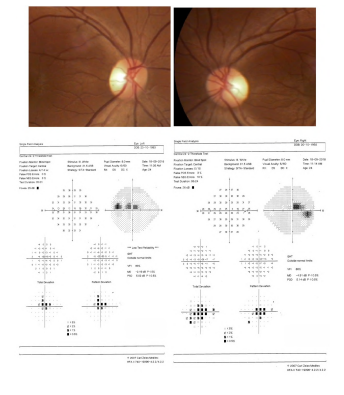


Figure 4: Case 4: Non glaucomatous optic neuropathy

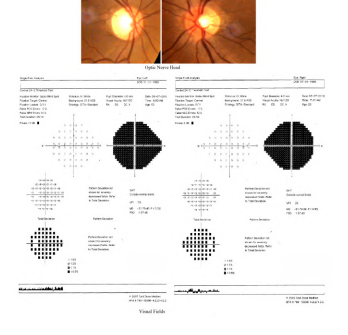


Figure 5: Case 5: Toxic optic neuropathy/atrophy

7. Case 6

A 50 year old diabetic patient presented with complaint of discharge in left eye. His BCVA BE was 6/5, N-6, LE showed RAPD, IOP OU-19/ 20 mmHG respectively, RE optic disc normal, LE showed disc edema. Color vision BE- normal, Visual fields RE- Normal, LE showed early/ relative inferior altitudinal field defect, uncontrolled blood sugar levels. A diagnosis of non arteritic anterior ischemic optic neuropathy [NA-AION] was made.

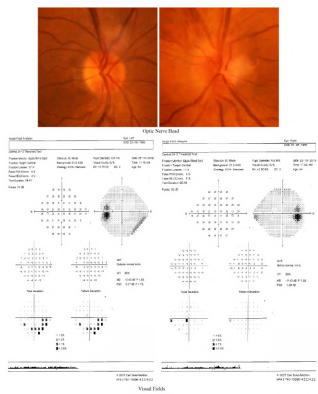


Figure 6: Case 6: Non arteriticanterior ischemic optic neuropathy

8. Case 7

A 55 year old male patient presented with complains of sudden & gross dimness of vision RE followed by some dimness of vision LE since 1 year. BCVA RE 6/60, LE 6/18,IOP- 17 mmHg .RE- Disc gliosis with pallor, LE disc temporal pallor, with both eyes cupping not clear. RE showed gross field defect, LE altitudinal field defect. ESR -22,borderline hyperlipidemia, CRP normal, other features like jaw claudication were absent & patient denied temporal artery biopsy . A diagnosis of Arteritic anterior ischemic optic neuropathy was made [A-AION].

9. Case 8

A 42 year-old female patient came for a regular glass prescription. BCVa 6/6, N/6, uneventful ocular exam, except

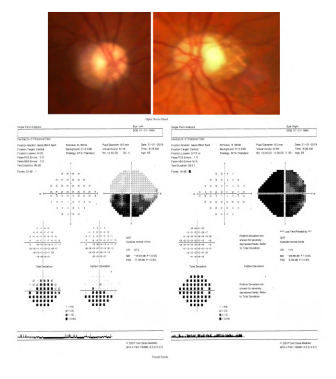


Figure 7: Case 7: Arteritic anterior ischemic optic neuropathy

a bilateral disc edema. On questioning she told that she was a patient of Benign intracranial hypertension under long term diuretic treatment. Visual fields showed a classic enlargement of blind spot with peripheral constriction of field.

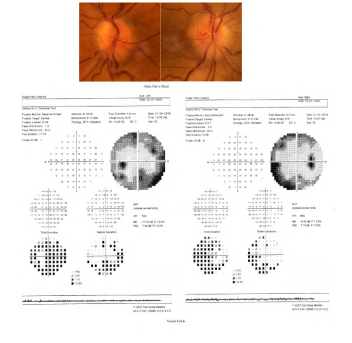


Figure 8: Case 8: Papilledema benign intracranial hypertension

10. Case 9

A 49 year old female patient presented with complain of pain right eye with decreased vision since few days. BCVA RE 2/60, RAPD, total color vision loss, & disc hyperemic & edematous. LE normal with BCVA 6/6. Visual fields LE-normal, while RE showed a gross irregular field defect, Borderline neutrophilia. MRI brain & spinal chord did not show MS plaques. Optic neuritis was diagnosed & treated with high dose systemic steroids, with vision & fields returning to normal.

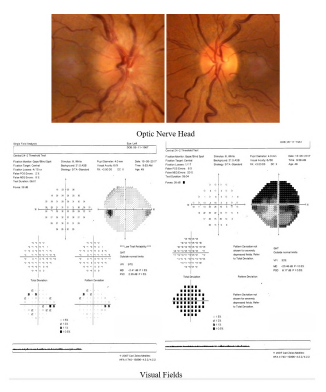


Figure 9: Case 9: Optic neuritis

11. Case 10

A 40-year-old male patient presented with complaint of gradual & gross dimness of vision since last 15 years. Vision BE 1/60, IOP 18 mmHg, defective color vision. Optic discs showed disc pallor & peripapillary telangiectasis & Visual fields showed irregular constriction. MRI plain & contrast Brain & orbit - T2 hyperintense signal seen in optic nerve & chiasma. VEP & CSF analysis - normal. Suspecting Leber's hereditary optic neuropathy, a genetic study was conducted for mitochondrial gene variation evaluation, which showed a homoplasmic missense variation in the MT-ND4 gene, mutations of which have been reported to be associated with LHON.

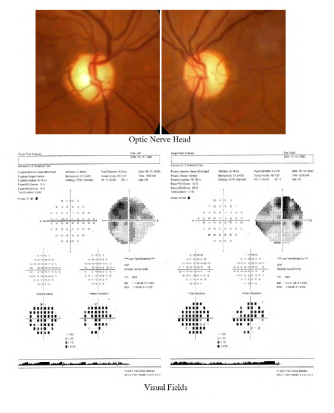


Figure 10: Case 10: Leber's hereditary optic neuropathy

12. Discussion

In case 1 a timely diagnosis with an optimum surgical intervention saved the patient from systemic & ocular morbidity, with an excellent visual recovery in LE from 6/60 pre-op to 6/5 post -op. Case 2 was being treated for glaucoma elsewhere. Resection of tumor unfortunately lead to collateral damage, with loss of vision in the better seeing eye, which could have been avoided / mitigated with a timely diagnosis & intervention. In Case 3 neurological diagnosis was clear but in view of the occludable angles, thin CCT, antiglaucoma medication was also started, as disc & field picture was inconclusive in making a confirmed diagnosis of a concurrent glaucoma. In cases 1-3 Visual field analysis confirmed a neurological pathology. 1 Case 4 - Its important to thoroughly investigate a patient before labeling them as malingerer to avoid missing a potentially vision/ life threatening pathology. ² Case 5 Toxic optic neuropathy is often under /late diagnosed, with patients presenting with painless, progressive, bilateral symmetrical visual loss, papillomacular bundle damage, central or cecocentral scotoma, and reduced color vision, where both toxic and nutritional factors play a synergistic role. ³ Case 6 & 7 – In NA- AION there is transient nonperfusion / hypoperfusion of ONH circulation in posterior ciliary arteries [PCAs], seen more often in microangiopathy associated with hypertension & Diabetes mellitus, hence usually less severe & extensive ONH damage than A-AION,

in which there is thrombotic occlusion of PCAs. Almost half of the N-AION patients can have almost normal Vision & about 40% show spontaneous visual improvement, 4 in sharp contrast to A-AION, where no such recovery is seen.⁵ A relative altitudinal defect with RAPD, unilateral disease, good vision & a small cup in other eye were helpful diagnostic criteria for N-AION in case 6. A massive visual loss with no recovery, bilateral involvement, extensive field defect in RE with an absolute inferior altitudinal field defect in LE helped clinch the diagnosis of A-AION in case 7. Absence of systemic sypmtoms like jaw claudication does not rule out the disease. Temporal artery biopsy should be done for confirming the diagnosis. 4-9 History taking & ONH examination, with relevant investigations are crucial, as infrequent or a single episode of transient obscuration of vision, a feature of papilledema, may not be registered by the patients. 10 Correlating complaints with clinical picture seen on ONH can confirm a diagnosis of anterior typical optic neuritis. The optic neuritis treatment trial (ONTT) showed that IV methylprednisolone followed by a tapering course of oral prednisone accelerated visual recovery by a few weeks, &, that the presence of asymptomatic white matter lesions on the MRI scan is the strongest predictor for MS. 11,12 LHON presents with simultaneous or sequential painless loss of vision with either an acute or sub-acute onset & defects in color vision. 12

13. Conclusion

A thorough detailed examination of Optic Nerve Head can give insight into a plethora of ocular & systemic diseases. A timely diagnosis, followed by appropriate treatment can thus avoid major ocular & systemic morbidities.

14. Source of Funding

None.

15. Conflict of Interest

None.

References

- Kedar S, Ghate D, Corbett JJ. Visual fields in neuro-ophthalmology. *Indian J Ophthalmol*. 2011;59(2):103–9.
- Dias DT, Ushida M, Battistella R, Dorairaj S, Prata TS. Neurophthalmological conditions mimicking glaucomatous optic neuropathy: analysis of the most common causes of misdiagnosis. BMC Ophthalmol. 2017;17(1):1.
- Sharma P, Sharma R. Toxic optic neuropathy. *Indian J Ophthalmol*. 2011;59(2):137–41.
- Hayreh SS, Zimmerman MB. Nonarteritic anterior ischemic optic neuropathy: natural history of visual outcome. *Ophthalmology*. 2008;115(2):298–305.
- Newman NJ. The Ischemic Optic Neuropathy Decompression Trial. Arch Ophthalmol. 2007;125(11):1568–70.
- Hayreh SS. Anterior ischemic optic neuropathy. V. Optic disc edema an early sign. Arch Ophthalmol. 1981;99(6):1030–40.

- 7. Hayreh SS, Zimmerman B. Management of giant cell arteritis. Our 27-year clinical study: new light on old controversies. *Ophthalmologica* . 2003;217(4):239–59.
- 8. Hayreh SS, Zimmerman B. Visual field abnormalities in nonarteritic anterior ischemic optic neuropathy: their pattern and prevalence at initial examination. *Arch Ophthalmol*. 2005;123(11):1554–62.
- 9. Hayreh SS. Ischemic optic neuropathy. *Prog Retin Eye Res*. 2009;28(1):34–62.
- Wall M, Hart WM, Burde RM. Visual field defects in idiopathic intracranial hypertension (pseudotumor cerebri). Am J Ophthalmol. 1983;96(5):654–69.
- 11. Guier CP, Stokkermans TJ. Optic Neuritis (Internet). Treasure Island (FL): StatPearls Publishing; 2023.

 Newman NJ. The Optic Neuritis Treatment Trial. Ophthalmology. 2020;127(4S):172–3.

Author biography

Jyoti Bhatt, Associate Professor

Cite this article: Bhatt J. Optic nerve head: A diagnostic Saga!. *Indian J Clin Exp Ophthalmol* 2024;10(3):603-608.