Bilateral anterior ischaemic optic neuropathy associated with optic disc drusen and systemic hypotension

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SUMMARY We report a case of bilateral anterior ischaemic optic neuropathy in a 23-year-old woman which was probably attributable to optic disc drusen and systemic hypotension related to peritoneal dialysis for renal failure.

Non-arteritic anterior ischaemic optic neuropathy (AION) appears to be the result of obstruction of axoplasmic flow owing to ischaemia, leading to a cascading effect via further small vessel compromise in the setting of underlying disc crowding. It has been reported in patients, some relatively young, with drusen of the optic nerve head. We report on a young woman with systemic hypotension who had optic disc drusen and developed bilateral AION while on home peritoneal dialysis.

Case report

A 23-year-old white female noticed on awakening the painless loss of inferotemporal vision in the left eye in late February 1986. On 14 March her visual acuity was 20/20 on the right and 20/25 +3, with marked relative afferent papillary defect and marked constriction of visual field on the left. There were drusen of both optic discs, with swelling of the left. A CT scan of the head and orbits was considered normal.

On 22 May her visual acuity was 20/20 in each eye without relative afferent papillary defect. The right disc pale and elevated, the left flat and mildly atrophic. In early June she awoke with marked diminution of vision in the right eye and was referred to our Neuro-ophthalmology Clinic on 20 June.

She had been on haemodialysis for chronic renal failure following post-streptococcal glomerulonephritis until 1984 and on peritoneal dialysis thereafter. She was chronically hypotensive, with blood pressure 52/38 and 62/38 mm Hg at clinic visits in January and February 1986. An oral contraceptive (Ortho-Novum, norethisterone with ethinyloestradiol) taken for approximately four years was discontinued when visual symptoms first developed. Her visual acuity was finger counting at 2–3 feet (60–90 cm), with hand motion only superotemporally in the right eye, and 20/20 +3 with the field concentrically constricted on tangent screen in the left eye. Colour plates (AO-HRR): 0/6 OD, 6/6 OS.

External, motility, and slit-lamp examinations gave normal results. Tension by applanation was 16 in each eye. Her pupils were sluggish, with a marked right relative afferent defect. There were bilateral disc drusen, with right more than left optic atrophy, and diffuse thinning of the nerve fibre layer (Figs. 1 and 2). Calcification was evident in both optic nerve heads on review of her orbital CT scan (Fig. 3).

Discussion

Various factors may have contributed to apparent bilateral consecutive AION in our patient, with optic disc oedema preceding visual loss by weeks in the second eye, as may occur. She had optic disc drusen in both eyes, which have been associated with transient visual obscurations and permanent visual loss, including AION. Preceding transient visual...
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Fig. 1 Appearance of right optic disc on 20 June 1986.

Fig. 2 Appearance of left optic disc on 20 June 1986.

Fig. 3 Orbital CT scan in March 1986 showing bilateral optic nerve head calcification.

loss and recurrent episodes of AION have been reported. Hypotension has been suggested, in addition to increase of intraocular pressure, as a basis for the frequent finding of visual loss in AION, as in our patient, on awakening. Impairment of autoregulation of optic nerve perfusion may have been an additional factor in our patient, who was chronically hypotensive. This was postulated by Taylor et al. in cases of AION following sudden reduction in systemic blood pressure in chronically hypertensive children with disc swelling. Drusen, through mass effect, may have raised the intraneural tissue pressure, leading to greater susceptibility to ischaemia. The crowded discs reported with drusen, as in our patient and others, may have been another factor. Increased circulating vasoconstricting agents may exacerbate hypoperfusion in the setting of hypotension in posthaemorrhagic AION and in AION following cardiopulmonary bypass. Peritoneal dialysis may have contributed to her AION, as suggested by the association with haemodialysis together with hypotension and intraocular pressure elevation in that setting. Oral contraceptives, reported to be associated with various forms of vaso-occlusive disease (including possibly AION) even months after their cessation, seem unlikely to be of significance in our patient, though not entirely excluded.

References


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