When he was 28 years old, suffered what in Augustus D’Este, grandson of King George III., even before von Gräfe, in December 1822, Sir Sir John Nettleship, which is here recounted.

Optic neuritis, often named retrobulbar neuritis or optic papillitis, is one of the commonest symptoms of multiple sclerosis (MS). At some stage it affects over 50% of MS patients. An early description in 1864 was by Albrecht von Gräfe [Graefe] (1828-1870). Modern techniques made diagnosis easier and more precise, but the early descriptions and much-argued concepts are seldom discussed. A thinning of the serial ganglion cell layer and inner plexiform layer is the pathogenesis of acute optic neuritis.

Ancient references to optic nerve dysfunction as a mechanism for loss of vision are found in Arabic texts of the ninth century.2 Possibly the first textbook of ophthalmology was written by Hunayn ibn Ishaq, (808-873) a Nestorian Christian and chief physician to the Caliph al-Mutawakkil. Like Galen, he believed that the optic nerve was hollow to transmit psychic pneuma* that flowed from the brain; the lens was the organ of vision. This he deduced by shutting one eye, whereupon the pupil of the other became enlarged to allow the escape of diverted punea. When the closed eye was opened, the enlarged pupil contracted to normal size.2 He described three different forms of ocular paralysis; those involving sight alone, those involving eye movements alone, and those involving both; but he failed to separate optic neuritis from other eye diseases.

The vision has ceased or diminished without our finding any change in the pupil and there is heaviness in the head and particularly its deep part and the parts surrounding the orbit. We know that the affection is caused by abundant moisture, which has run to the optic nerve...

Even before von Gräfe, in December 1822, Sir Augustus D’Este, grandson of King George III., when he was 28 years old, suffered what in retrospect was an attack of retrobulbar neuritis, though its nature was not realised at the time. In successive years, he noted progressive weakness, numbness, difficulty in walking, painful spasms and depression — all typical of MS. Although he was aged 54 when he died, no formal diagnosis was made, but ‘the meticulous notes in his diary justify a posthumous diagnosis’.

In 1823, George Frick (1793-1870)4 in the first American textbook (Figure 1) of ophthalmology (1823) had described varieties of amaurosis that included but did not demarcate optic neuritis:

“The terms amaurosis, gutta serena, suffusion nigra, &c are applied to a species of blindness which is produced by some immediate affection of the optic nerve or its expansion into the retina ... Amaurosis may take place suddenly or slowly and be transient, permanent or intermittent.” (Pp. 138-141. 1826 edition)

But before the ophthalmoscope, Frick was unable clearly to distinguish optic neuritis from uveitis, glaucoma, orbital tumours, and other eye and systemic disorders. Shrewdly, he had noted severe pain in the orbit before visual loss and abnormal pupillary responses to light. He related:

Amaurosis from whatever cause ... is generally characterized by a very dilated pupil which is not affected by any degree of light which is made to fall upon the retina ... [The patient] is obliged to turn his head to render them [objects] distinct. (p. 142)

The invention of the ophthalmoscope by Hermann von Helmholtz (1821–1894); in 1845, allowed optic neuritis to be separated from many other ocular disorders. So valuable was the ophthalmoscope that by 1871, Thomas Nettleship (1845-1913) in 1884 described its principal features.5 Nettleship acknowledged that both Leber and Jonathan Hutchinson had previously described cases of optic neuritis. However, Leber included instances of tobacco amblyopia and Hutchinson included several other pathologies. Nettleship’s comprehensive account in 1884 emphasised pain on eye movement, abnormal disk appearances and he stressed the impaired colour vision. Eleven of his 16 patients had a central scotoma. He accurately characterised its features:

Failure of sight limited to one eye, often accompanied by neuralgic pain about the temple and orbit and by pain in moving the eye; many recover but permanent damage and total blindness may ensue; there is at first little, sometimes no, ophthalmoscopic change, but the disc often becomes more or less atrophic in a few weeks... The defect in vision is often described at first as a “gaze” or a “yellow mist” or a “dark patch” or a “spot” which covers the object looked at and gives an unnatural colour, the hand looking, for example, as if covered by a brown glove.”

Although he identified all the salient features of optic neuritis, he did not mention a relationship to other relapsing and remitting neurologic symptoms, characteristic of MS. In the 19th century, optic neuritis was often used as a descriptive term for papilloedema. Its most common cause was widely said to be a brain tumour. It was also recognised as a discrete disease of the optic nerves,* though its aetiology was often uncertain.

After Nettleship’s seminal description, Thomas Buzzard (1831-1919)† (Figure 2.) in 1893 reported five patients with a history of disseminated sclerosis, who had episodes of visual failure with recovery consistent with optic neuritis. He was one of the first to recognise optic atrophy as a feature of disseminated (multiple) sclerosis.

And in this seminal text, he recognised the salient features of optic neuritis and ‘atrophic amaurosis’ and the frequent confusion with ischaemic optic neuropathy.7 1860 von Graefe ((1828–1870),8 and more meticulously, Edward Nettleship (1845-1913) in 1884 described its principal features.9 Nettleship acknowledged that both Leber and Jonathan Hutchinson had previously described cases of optic neuritis. However, Leber included instances of tobacco amblyopia and Hutchinson included several other pathologies. Nettleship’s comprehensive account in 1884 emphasised pain on eye movement, abnormal disk appearances and he stressed the impaired colour vision. Eleven of his 16 patients had a central scotoma. He accurately characterised its features:

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Neuromyelitis optica (Devic’s disease)

In 1870, (Sir) Thomas Clifford Allbutt, then Physician to several infirmaries at Leeds, first reported the association between myelitis and an optic nerve disorder.13 He briefly described a case of myelitis followed by optic neuritis approximately three months later. Earlier, EO Hocken had reported a patient with spinal cord inflammation and amaurosis in 1841.14 CM. Durrant described a probable case in 1850.15 Wilhelm Erb in 1879 described a man who developed recurrent optic neuritis succeeded by subacute myelitis.16 Dreschfeld in 1882 described the first pathologically examined case17 and showed inflammation in both the spinal cord and optic nerves; the brain was normal. In 1894, Eugène Devic (1853–1927)18 presented his case at the First Congress of Internal Medicine in Lyon, and with Gault summarised 16 patients with loss of vision, who within weeks developed spastic paraparesis. Devic’s telling question, “Why such a peculiar localisation?” remains unanswered. Neuromyelitis optica (NMO) has well known similarities to multiple sclerosis, but despite the finding of NMOIgG (Aquaporin 4) in about two-thirds of cases of NMO, whether it is an Aquaporin 4 + NMO autoimmune astrocytopathy, a disease sui generis, or an MS variant remains arguable.19

In typical optic neuritis, visual function improves spontaneously over four to six weeks, and within 12 months 93% have acuity of at least 20/40. High-dose corticosteroids may hasten recovery, but have little effect on long-term visual outcome. The cumulative probability of developing MS by 15 years is 50%. White matter plaques on the first magnetic resonance image increase that risk to 72%.

Edward Nettleship (1845-1913)

Of the many who contributed to the descriptions of optic neuritis it is Nettleship whose comprehensive writings first clearly delineated the disorder. Born on 3 March 1845 in Kettering, Northamptonshire, he attended Kettering grammar school. Intending at first to become a farmer, he entered King’s College, London, and the Royal Veterinary College, and was admitted a Licentiate of the Society of Apothecaries and received the diploma of the Royal College of Veterinary Surgeons in 1867. He was appointed Professor of Veterinary Surgery at the Royal Agricultural College but a year later returned to the London Hospital, as dresser and assistant to (Sir) Jonathan Hutchinson.20 Nettleship became his firm friend, and most distinguished pupil. He qualified in surgery (F.R.C.S.), from the London Hospital and the Blackfriars Hospital for Skin Diseases. To specialise in ophthalmology, in 1868, he studied at Moorfields Eye Hospital under Hutchinson and Waren.21 He was appointed Curator of the Museum and Librarian.

On 22 January 1869 he married Elizabeth Endacott Whiteway from Devon; they had no children. At Moorfields he began his researches into eye diseases,22 but left for the post of medical superintendent at the Ophthalmic School at Bow (1873–4), working with impoverished children suffering from eye infections. He was then appointed ophthalmologist at the South London Ophthalmic Hospital, St Thomas’s Hospital (1878–95), and surgeon at Moorfields (1882-98). In 1888 he became Dean of the Medical School. He served and became President of the Ophthalmological Society, and advised the Board of Trade on Sight Tests for the Mercantile Marine.

Nettleship acquired a considerable reputation as an eye surgeon and teacher. Sir John Parsons described him as the most scientific teacher of his time. He removed a cataract from William Ewart Gladstone, and attended Queen Victoria for the same condition, but advised against surgery.

Whilst working at St Thomas’s and Moorfields Eye hospital, he wrote the definitive text: The Student’s Guide to Diseases of the Eye. (1884 edn)