Variations in appearance of the normal optic nerve head

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The vast majority of patients examined in primary eye care have normal, healthy eyes. To discriminate between ocular disease and the normal eye, it is essential to know the many presentations that a normal eye can make and a collection of photographs of these normal variations at the optic nerve head is presented to supplement the information provided in atlases of ocular disease. The following is adapted from Clinical Procedures in Primary Eye Care (2007, 3rd edition) and its associated website (http://evolve.elsevier.com/Elliott/eyecare/).

Size and shape
The optic nerve head or disc comes in a variety of shapes and sizes. The mean average Caucasian optic disc area, excluding high myopes, is between 2.1mm² and 2.8mm² with a range between 0.80mm² to 6.00mm².¹ The average vertical disc diameter is 1.8mm with a horizontal disc diameter of 1.7mm. Discs have been shown to be smaller on average in Caucasians (Figure 1), and progressively larger in Mexicans, Asians and African North Americans (Figures 2 & 3).² Disc size is also larger in myopes beyond –8 D and smaller in hyperopes greater than +4 D.² Oval discs (Figure 4) are often found with corneal astigmatism and the direction of the longest optic disc diameter can indicate the axis of astigmatism.³

Optic cupping
The central proportion of the nerve head usually contains a depression called the ‘cup’. This is often associated with an area of pallor due to the lamina cribrosa reflecting through in the absence of axons and their associated capillaries. However, in some cases the cup can extend beyond the area of pallor, so that this should not be used as an indicator of cup size during 2-D evaluations such as provided by direct ophthalmoscopy. Rather the kinking of blood vessels as they pass over the edge of the cup should be used as an estimate of the cup position.

As discussed above, discs can vary considerably in size, yet approximately the same number of axons (about 1 million) leave the eye via the optic nerve head. Therefore large optic discs typically have larger cupping because of the absence of axons in the middle of the disc as the neurons leave the retina in the larger rim tissue of larger discs. The physiological cup-to-disc ratio (CDR) is normally less than 0.60, but is relative to the size of the disc; so that smaller cupping should be seen in a small-sized disc and larger cupping is expected in large discs (Figs. 2 and 3). For this reason, a 0.30 CDR in a small disc may be more indicative of glaucoma than a 0.70 CDR if it is in a large-sized disc (e.g., Garway-Heath et al.4), highlighting the importance of assessing disc size. The CDR is typically measured in the vertical meridian as the cup tends to enlarge in the vertical meridian in glaucoma. A large physiological cup

Figure 1
A small, flat optic nerve head of a young Caucasian patient
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The optic nerve heads and cups of the two eyes are typically mirror images of each other and differential diagnosis of many optic nerve head anomalies is provided by an inter-eye comparison. The neural rim tissue should be thickest in the inferior region of the disc, then the superior and nasal, being thinnest in the temporal region, so that normal neural rim tissue follows the ISN'T rule. Lamina cribrosa

Seen in about 30% of eyes as grey dots at the bottom of the optic cup (Figure 5). It is a sieve-like structure of largely connective and glial tissue that is continuous, although embryologically distinct, with the scleral coat. Perhaps not surprisingly, it is more visible in larger discs and larger cups and this appears to be the only reason that visible lamina cribrosa has a higher prevalence in patients with open-angle glaucoma.

The vertical CDRs indicated for the photographs presented here are based on the 2-D diagrams. The optic nerve heads and cups of the two eyes are typically mirror images of each other and differential diagnosis of many optic nerve head anomalies is provided by an inter-eye comparison. The neural rim tissue should be thickest in the inferior region of the disc, then the superior and nasal, being thinnest in the temporal region, so that normal neural rim tissue follows the ISN'T rule.

Lamina cribrosa

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Drusen of the disc
A familial, typically bilateral condition, found in about 0.3% of patients, which becomes more obvious with age. In children, they may be buried in the nerve head and not seen and the disc appears swollen, so that the condition is sometimes called pseudopapilloedema. They are golden, autofluorescent, glowing, calcific globular deposits that sit in front of the lamina cribrosa (Figure 6). They are typically found in small discs with little or no cupping and this appearance can mask signs of early glaucoma. Although typically benign they can shear blood vessels and/or nerve fibres, leading to haemorrhages (2-10%) and visual field loss (~75%), some of which can be progressive. Visual field monitoring is essential. Areas of peripapillary pigment atrophy can be seen after resolution of any haemorrhages.

Myelinated nerve fibres
Found in about 1% of patients and represents myelin sheathing of the optic nerve fibres that extends beyond the lamina cribrosa and presents a superficial, white, feathery opacification which hides any underlying retinal blood vessels. They are usually continuous with the optic nerve head (Figures 7 & 8), although small discrete patches of myelinated nerve fibres can appear and may mimic a cotton wool patch. They are typically benign, although may cause visual field loss at threshold. A small number are characterized by ipsilateral extensive myelinated nerve fibres, anisometropic myopia, ambylophia, and strabismus. In most cases, myelinated nerve fibres remain unchanged over time. However, loss of myelinated nerve fibres may occur due to a central demyelinating process or the result of direct axonal destruction or nerve fibre layer ischaemia and has been reported in central retinal artery occlusion, uncontrolled glaucoma, anterior ischemic optic neuropathy, optic neuritis and Behçet disease and after plaque radiotherapy for choroidal melanoma and pars plana vitrectomy.

Nerve fibre layer striations
These are brightest at the superior and inferior poles, where the nerve fibre layer is thickest and are best seen in young patients, particularly those with heavily pigmented fundi (Figure 9). The striations are caused by the tubes of astrocytes that surround the retinal ganglion cell axon. Fundus photography, particularly digital, may provide a better assessment of the nerve fibre layer than fundus biomicroscopy. Nerve fibre layer striations are best seen with the green (red-free) filter as the lower wavelengths do not penetrate the nerve fibre layer and are more readily reflected back. Focal wedge-shaped defects in the nerve fibre layer striations can occasionally be detected in patients with glaucoma and although diffuse nerve fibre layer is more common in glaucoma, it is more difficult to detect.
Peripapillary atrophy (PPA)

PPA can be categorised into zone alpha and beta. Zone beta PPA is found adjacent (bordering) to the disc and is present in about 15% of normal eyes (Figure 10). It is more common in glaucoma, but more importantly it can be a sign of progressive disease if it gets larger. The RPE and choriocapillaris are lost and all that is visible are the large choroidal vessels and sclera. Zone alpha is present in nearly all normal eyes and is characterised by irregular hyper and hypopigmented areas in the RPE, either on their own or surrounding zone beta PPA. PPA is most commonly found at the temporal edge of the disc. It should be differentially diagnosed from high myopic atrophy and malinserted optic discs.

Tilted discs & optic disc malinsertion

The tilt can be seen with the 3-D view of fundus biomicroscopy. With direct ophthalmoscopy it is seen as an oval disc whose edges may not be exactly focussed simultaneously. In the tilted disc syndrome, the disc or discs are commonly tilted inferior nasally with a nasal staphyloma (bulging of the sclera) and situs inversus, where the temporal blood vessels first course towards the nasal retina before sharply changing course (tilted disc in Figure 11 and a normal disc in the fellow highly myopic eye in Figure 12). Tilted discs are thought to be caused by an incomplete closure of the embryonic foetal fissure, similar to the aetiology of a coloboma. The condition is benign, although the area of nasal staphyloma can produce a temporal visual field defect. Tilted disc syndrome can be in one or both eyes and the binocular condition can produce bi-temporal field defects. These can usually be differentiated from chiasmal lesions as the bi-temporal field defects cross the vertical midline and the field defect can be improved if retested with a higher myopic correction. Tilted discs are associated with corneal astigmatism and myopia and the direction of the longest optic disc diameter can indicate the axis of corneal astigmatism.

Much more common is optic disc malinsertion, which is a simple insertion of the optic nerve at an acute angle and without the appearance of rotation of the optic nerve, nasal staphyloma or situs inversus. The malinsertion is almost always bilateral and the malinserted discs are mirror images of each other, typically elevated nasally, tilting downwards temporally and with a temporal scleral and/or choroidal crescent. Photographs from the right and left eyes of a patient with malinserted discs are shown in Figures 13 and 14.

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Figure 11
- Tilted disc syndrome with situs inversus. This Caucasian eye is also highly myopic and astigmatic, with a thin retina and visible underlying choroid.

Figure 12
- The image from the right eye of the same patient shown in fig. 11. This eye is also highly myopic and has a thin retina and visible underlying choroid.

Figure 13
- The image from the right eye of an Asian patient with optic nerve malinsertion. There is some light scatter from the edge of the pupil.

Figure 14
- The image from the left eye of the same patient shown in fig. 13. Nerve fibre striations and macula pigment are also clearly visible.
Module questions
Please note, there is only one correct answer. Enter online by March 5 2008.

1. The average cup-to-disc ratio viewed by direct ophthalmoscopy is:
   A. The same size compared to when viewed by indirect ophthalmoscopy.
   B. Larger compared to when viewed by indirect ophthalmoscopy.
   C. 0.30 when using either direct or indirect ophthalmoscopy.
   D. Smaller compared to when viewed by indirect ophthalmoscopy.

2. Drusen of the disc can cause:
   A. Papilloedema.
   B. Relatively little as it is essentially a benign condition.
   C. Secondary glaucoma.
   D. Haemorrhage and progressive visual field loss.

3. Of the following, which is the most significant risk factor for primary open-angle glaucoma?
   A. 0.60 cup-to-disc ratio.
   B. Tilted disc.
   C. 0.50 cup-to-disc ratio in a small disc.
   D. Oval cup-to-disc ratio.

4. Of the following, which is the most significant risk factor for primary open-angle glaucoma?
   A. Visible large pores of the lamina cribrosa.
   B. Zone alpha peripapillary atrophy.
   C. Visible nerve fibre striations.
   D. Zone beta peripapillary atrophy.

5. Myelinated nerve fibres:
   A. Are benign and cause no visual loss.
   B. Can cause visual field loss, regress with certain conditions and should be monitored.
   C. Can cause visual field loss and should be given a non-urgent referral on first diagnosis.
   D. Can progress and regress with certain ocular conditions and should always be given a non-urgent referral on first diagnosis.

6. Nerve fibre layer striations are best seen:
   A. In undiagnosed glaucoma at the inferior and superior poles.
   B. In the peri-macular bundle.
   C. At the superior and inferior poles.
   D. In treated glaucoma at the inferior and superior poles.

Please complete on-line by midnight on March 5 2008 - You will be unable to submit exams after this date – answers to the module will be published in our March 7 issue.