Clinical approach to vision loss: a review for general physicians

Authors: Antony Raharja\textsuperscript{A} and Laurence Whitefield\textsuperscript{B}

Visual loss describes temporary or permanent reduction in visual acuity and/or field. Its aetiology is diverse due to the contributions of the different neuro-ophthalmic structures (eye, optic nerve, and brain) to image formation and perception, but may be categorised into ocular causes (corneal, lenticular, vitreoretinal and macular) or optic neuropathies. Clinical evaluation of visual loss relies on thorough history and examination to guide further tests. In this article, we provide a practical overview of visual loss assessment for general physicians.

Introduction

Visual loss is a serious clinical problem, commonly presenting first to non-ophthalmic healthcare professionals. It typically describes a reduction in visual acuity (sharpness/clarity) or field. This may be acute- or gradual-onset, monocular or binocular and transient or permanent, and may affect central and/or peripheral vision. The work-up for vision loss begins by characterising the nature of visual disturbance, eliciting any associated ocular, neurologic or systemic symptoms, and is followed by clinical examination to guide further tests and investigations. This article provides a brief overview of a practical approach to evaluating visual loss.

Background

The importance of vision to health and quality of life cannot be overstated. Of the different human senses, vision is frequently the most valued.\textsuperscript{1,2} In fact, participants within a UK-based study would trade off 10 years of life without sight for 4.6 years in perfect health.\textsuperscript{1} In another US-based nationwide poll, sight loss was deemed worse than life-changing diagnoses such as Alzheimer’s disease, cancer and AIDS/HIV.\textsuperscript{2} Beyond its influence on quality of life, vision loss has indirect repercussions, including productivity losses and costs of informal care.\textsuperscript{3}

Worldwide, the leading causes of blindness in adults over 50 years old are cataract, glaucoma, uncorrected refractive error, age-related macular degeneration (AMD) and diabetic retinopathy.\textsuperscript{4} These remain the top five causes in the UK, with an estimated 2 million people being blind or partially sighted.\textsuperscript{5} Among working-age adults in England and Wales, diabetic retinopathy/maculopathy was previously the leading cause of vision loss, but this has been overtaken by inherited retinal disorders.\textsuperscript{5}

Requirements for clear vision

For a clear image, light must pass through clear media (tear film, cornea, aqueous, lens and vitreous) and be refracted by the cornea and lens onto the retina. Photoreceptors (rods and cones) then sense and transmit light as nerve impulses along the optic nerve. Fibres from nasal hemiretinae (temporal visual field (VF)) decussate at the optic chiasm and join the contralateral optic tracts, whereas fibres from temporal hemiretinae (nasal VF) join ipsilateral optic tracts. As such, optic tracts carry fibres from contralateral visual fields.

Optic tracts synapse at ipsilateral lateral geniculate ganglia within the thalamus before continuing as optic radiations. Parts of the inferior

Key points

Assessment of visual function (acuity, colour vision and field) is the initial step to examining patients with vision loss.

Improvement in visual acuity with pinhole indicates undercorrected refractive error.

Ocular pathologies (corneal, lenticular, vitreoretinal and macular) and prechiasmal optic neuropathies typically cause monocular vision loss.

Urgent ophthalmology discussion is recommended for any patient with reduced vision and recent eye surgeries.

Amaurosis fugax represents ischaemia-related transient monocular vision loss, and must be recognised as it may herald an impending stroke.

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optic radiations (contralateral superior quadrant VF) loop around the temporal horn of lateral ventricle (Meyer loop), whereas superior optic radiations pass through parietal lobes to arrive at primary visual cortices along calcarine fissures within the occipital lobes.

Due to the contributions of the different neuro-ophthalmic structures (eye, optic nerve and brain) to image formation and perception, the aetiology of vision loss is diverse but can be divided into ocular causes (eg corneal, lenticular, vitreoretinal and macular) and optic neuropathies (Table 1).

### Key points in history taking

#### Monocular versus binocular visual loss

Monocular visual loss is typically caused by ocular pathologies or prechiasmal optic neuropathy unless the underlying disease affects both eyes; for instance, systemic conditions (eg diabetic macular oedema, malignant hypertension and nutritional optic atrophy) and inherited diseases (eg retinitis pigmentosa).

Binocular visual disturbance should raise a suspicion of chiasmal, retrochiasmal or other intracranial pathologies causing raised intracranial pressure (ICP) and papilloedema. Due to the retinotopic organisation within the visual pathway, the nature of VF loss may localise the lesion. In particular, bitemporal hemianopia is classically caused by chiasmal lesions, whereas homonymous field loss (congruent VF loss in both eyes) indicates retrochiasmal lesions; for example, temporal lobe lesions compressing Meyer loop may cause contralateral superior homonymous quadrantanopia (‘pie in the sky’ VF defect).

#### Central versus peripheral vision loss

Poor central vision sparing the peripheries indicates macular involvement. Macular diseases (eg AMD and central serous

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**Table 1. Differential diagnoses for vision loss**

| Mechanism of reduced vision | Clinical features | Examples of causes |
|-----------------------------|------------------|--------------------|
| **Ocular causes**           |                  |                    |
| Refractive errors (light is not focused onto the retina) | > Blurry vision corrected with pinhole, glasses or contact lens | > Myopia (short-sighted) |
| Media opacity (light is obstructed from reaching the retina) | > Notable structural opacity on examination > Red reflex may be reduced or absent altogether | > Hyperopia (far-sighted) |
| Retina (failure of photoreceptors or its supporting structures) | > Visual acuity is reduced with macula involvement > Field loss corresponds to the area of retina affected > Medical comorbidities and family history are clues > Diagnosis is largely based on clinical appearance | > Astigmatism |
| **Optic nerve pathologies (dysfunction of optic nerve to transmit signals)** | > Dyschromatopsia > Relative afferent pupillary defect (if unilateral or asymmetric) > Visual field deficit > Optic disc appearance: pale, swollen or large cup-to-disc ratio | > Unilateral optic disc swelling: demyelinating (MS), giant cell arteritis, non-artenitic anterior ischaemic optic neuropathy or compressive |
| Optic neuropathy (anterior to the optic chiasm) | > | > Bilateral disc swelling: demyelinating (neuromyelitis optica) or raised ICP (IIH, CVST, intracranial lesion, hydrocephalus or medication-induced) |
| Optic chiasmal lesion | > Bitemporal hemianopia | > Optic disc atrophy: nutritional (B12 or folate deficiency), previous optic disc swelling or compressive lesion |
| Retrochiasmal lesion | > Contralateral homonymous hemianopia | > Glaucoma a |

*AMD = age-related macular degeneration; BRAO/CRAO = branch/central retinal artery occlusion; BRVO/CRVO = branch/central retinal vein occlusion; CSR = central serous choriotretinopathy; CMV = cytomegalovirus; CVST = central venous sinus thrombosis; DMO = diabetic macular oedema; DR = diabetic retinopathy; HSV = herpes simplex virus; ICP = intracranial pressure; IIH = idiopathic intracranial hypertension; MS = multiple sclerosis; VZV = varicella zoster virus. a Rarely causes dyschromatopsia and relative afferent pupillary defect.*
Contact lens use
Retinitis pigmentosa
Systemic symptoms: eg weight loss, Rheumatological diseases, eg Medications, eg tamoxifen,
Central or peripheral
Occupation/hobbies or driving needs
Diabetes and other cardiovascular risk
Migraine
Ocular symptoms: red eyes, pain,
Temporal arteritis symptoms: new-
Glaucoma

A thorough history may identify those with ocular pathologies if vision loss is secondary to ocular or optic nerve pathologies. A useful framework to narrow the differentials is to determine

Ocular pathology or optic neuropathy?
A useful framework to narrow the differentials is to determine if vision loss is secondary to ocular or optic nerve pathologies. A thorough history may identify those with ocular pathologies further.

Constricted peripheral vision is usually caused by optic neuropathy; the magnitude of field loss may be subtle and only detected during screening (eg chronic glaucoma), or large and cause difficulties with navigation (eg frequently bumping into things due to bitemporal hemianopia). Although uncommon, retinal causes of tunnel vision, such as retinitis pigmentosa or extensive bilateral panretinal photocoagulation, should also be considered.

Speed of onset
Speed and duration of onset may reflect the underlying pathophysiological processes. Vascular occlusions of retinal, optic discs or cerebral vessels usually cause rapid-onset vision loss. Inflammatory processes such as optic neuritis or scleritis generally cause subacute visual reduction over days. Degenerative processes (such as cataract) typically cause gradual-onset visual impairment, but the disturbance may not be realised in the early stages. In addition, serious insidious pathologies (eg compressive optic neuropathy from an intracranial mass) could cause slowly progressive visual loss.

Transient visual disturbance
Transient monocular visual loss (TMVL) is a worrying symptom, especially in older individuals with cardiovascular risk factors, as it could be ischaemic in origin (amaurosis fugax), heralding an impending stroke. Classically, TMVL in amaurosis develops in a downward curtain-like manner and lasts 2–30 minutes with an upward resolution. However, other patterns have been described.6 Amaurosis fugax typically results from temporary ischaemia caused by embolism originating from the carotid circulation, but may occur in chronic vascular insufficiency secondary to severely stenotic internal carotid artery (oculocerebral ischaemic syndrome). In the latter, amaurosis may be induced by light, exertion or change in posture.7 Other causes of TMVL are giant cell arteritis, impending central retinal vein occlusion and retinal migraine. An important differential for TMVL is migraine-associated visual aura. This presents as episodic zig-zag lines or scotoma, lasts 5–60 minutes and typically, but not always, precedes the headache; its slow and gradual development is a key feature, differentiating it from ischaemia-related causes eg transient ischaemic attack.8 For transient binocular disturbance, the precipitating and relieving factors are clues to the underlying causes. Transient visual obscurations in papilloedema are brief binocular visual blurriness or darkening lasting for a few seconds and precipitated by bending forward or lying down (or any actions that increase ICP). Transient blurry vision on standing up may be related to postural hypotension. Frequent episodic blurry vision that clears with blinking may be due to ocular surface diseases that disrupts the air-tear film interphase.

(Table 2; for instance, vision loss preceded by floaters and photopsia (brief sudden and flash-like) could be related to posterior vitreous detachment causing retinal detachment (Fig 1). Blurry vision in someone with severely painful red eye could be due to corneal ulcers, acute angle-closure glaucoma, scleritis or other ocular problems. However, most retinal causes of vision loss are painless and do not cause other symptoms; the diagnosis is based on the clinical appearance. The identification of optic neuropathy is also largely clinical: dyschromatopsia, relative afferent pupillary defect (RAPD) and characteristic VF defect. The underlying cause may be obvious, eg orbital cellulitis, thyroid-associated ophthalmopathy or trauma (suspected retrobulbar haemorrhage). If not, the optic disc appearance (swelling or pallor) will focus subsequent work-up (Table 1). Of note, giant cell arteritis, non-arteritic anterior ischaemic optic neuropathy and optic neuritis (eg multiple sclerosis) are frequent causes of optic disc swelling.

Assessment of visual function
Prior to examining the eye, visual function deficits should be established. Visual function includes distance and near acuity, colour vision and visual field. Measuring distance visual acuity (VA) using standardised charts (eg Snellen and logMAR) is the key initial step to quantifying visual deterioration. Each eye needs to be

Table 2. Salient points in the history taking when assessing vision loss

| Nature of visual complaint | Monocular or binocular |
|---------------------------|-----------------------|
|                           | Transient or permanent |
|                           | Central or peripheral  |
|                           | Speed of onset         |
| Other associated features | Ocular symptoms: red eyes, pain, purulent eye discharge, photophobia, floaters or photopsia |
|                           | Neurologic symptoms: headache, weakness or numbness |
|                           | Temporal arteritis symptoms: new-onset headache in individuals over 50 years old, pulseless and tender temporal artery, jaw claudication or scalp tenderness |
|                           | Systemic symptoms: eg weight loss, night sweat or rash |
| Social history            | Occupation/hobbies or driving needs |
| Past ophthalmic history   | Contact lens use       |
|                           | Recent ophthalmic surgery, eg cataract surgery |
| Past medical history      | Diabetes and other cardiovascular risk factors |
|                           | Rheumatological diseases, eg polymyalgia rheumatica |
|                           | Migraine               |
|                           | Medications, eg tamoxifen, hydroxychloroquine or ethambutol |
| Family history            | Glaucoma               |
|                           | Retinitis pigmentosa   |
tested separately, unaided, with glasses and with pinhole occluder.
VA improvement with pinhole suggests a degree of undercorrected refractive error. In infants and toddlers, the inability to fix and follow a target, or aversion to occlusion of one eye more than the other, may be indicative of poor vision.

Near vision can be crudely evaluated using small prints, but this is rarely useful in non-ophthalmic settings. Using a red object and comparing each eye in turn may assess for red desaturation (colour appearing pink or ‘washed out’). If available, colour vision can be quantified using Ishihara charts. The presence and location of VF deficits can be determined by the confrontation method. Formal perimetry testing may be subsequently required for monitoring progression and to assess driving eligibility.

**Essential eye examinations**

Eye examination starts by inspecting the eyelids (for cellulitis, lid retraction or ptosis), globe position (for proptosis or enophthalmos) and eye movements (for restricted or painful extraocular movements). The conjunctiva and sclera are then examined for any generalised, sectoral or circumcorneal injection. Ten per cent phenylephrine will blanch episcleritis but not scleritis.

The cornea is best visualised using a slit lamp. However, any focal opacities, foreign bodies or generalised haziness impairing VA are usually readily noticeable. Fluorescein staining can further highlight any epithelial defect under blue cobalt light, available on most handheld ophthalmoscopes. Macroscopic build-up of red cells (hyphaema) or white cells (hypopyon) is also easily noticeable without a slit lamp (Fig 2).

Next, pupils should be assessed for size asymmetry and response to light. Red reflex should be examined for leucokoria (eg retinoblastoma) in children and media opacity (eg dense cataract or vitreous haemorrhage) in adults.

Finally, the retina and optic discs can be examined using an ophthalmoscope; the view can be improved with pharmacological dilation, eg using cyclopentolate. The optic disc should be examined for its cup-to-disc ratio, colour and contour.

**Referring to ophthalmology**

Although the majority of eye presentations to the emergency department can be managed without ophthalmic input, vision loss warrants discussion with ophthalmology especially for those with recent eye surgeries. Further examination and tests (such as optical coherence tomography, ultrasound and electrodiagnostics) may be required. Neuroimaging is necessary in certain cases eg suspected compressive or demyelinating optic neuropathy and bilateral optic disc oedema. In the absence of organic disease, functional visual disturbance should be considered.

**Conclusion**

In summary, vision loss can be caused by ocular or optic nerve pathologies. Its initial evaluation can be done well without any sophisticated equipment. While non-ophthalmologists may not always reach a firm diagnosis, a thorough initial assessment is key to determining the urgency of referral.

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Address for correspondence: Dr Antony Raharja, Royal Free Hospital, Pond Street, London NW3 2QG, UK.
Email: antony.raharja@nhs.net