Many ocular and neurological diseases and conditions are known to exhibit distinct visual field loss patterns, and thus, visual field testing may assist in the differential diagnosis process. The ability to map the depth, extent and change of visual field defects should be considered in clinical management decisions. Results must be interpreted critically (reliability and repeatability) and in conjunction with other clinical signs, symptoms and examination findings. At present, standard automated perimetry of visual field defects should be considered in clinical management decisions. Some common types of visual field defects and their more common differentials are outlined below. Results must be interpreted critically (reliability and repeatability) and in conjunction with other clinical signs, symptoms and examination findings. At present, standard automated perimetry of visual field defects should be considered in clinical management decisions.

### VERTICAL FIELD LOSS PATTERN

Vertically oriented field defects should always raise the suspicion of pathologies on the visual pathway beyond the retina, particularly if it respects the vertical midline.

**Differentials:**
- Retinal disease
- Pre-chiasmal or anterior chiasmal lesion (e.g. compressive lesions)

**Bilateral (homonymous):**
- Post-chiasmal lesion (e.g. compressive lesions, stroke, injuries)

**Bilateral (bitemporal/binasal):**
- Chiasmal lesions (pituitary adenoma, meningioma, parasellar carotid artery aneurysm, meningioma, craniopharyngioma, glioma)
- Tilted disc syndrome

Vertical field loss can be classified into the following patterns:

1. **Vertical Step:**
   - Generally respects the vertical midline with at least 2 points outside 15° of fixation.

2. **Quadrantopia:**
   - Visual field loss that respects both the vertical and horizontal midline. Suggestive of neurological involvement if bilateral. All points within the quadrant must be P<5%.
   - **Differentials:**
     - Branch retinal artery/ vein occlusion
     - Advanced glaucoma
     - Cortical disease (if bilateral, rare)

3. **Hemianopia:**
   - Loss of the vertical hemifield respecting the vertical midline either partially or completely.
   - **Note:** Monocular temporal hemianopia may occur if the lesion is more anterior and only affecting the nasal crossing fibres from the ipsilateral eye.

4. **Three Quadrants:**
   - Three quadrants with all points at least P<5%. Partial three quadrant losses do not have all points P<5% but is greater than a complete hemianopia.
   - **Note:** Multiple lesions or pathologies may need to be considered.

### CENTROCAECAL

**Description:** Field loss extending from blindspot to fixation. Must include fixation and does not obey horizontal midline. Usually due to damage of the papillomacular bundle.

**Differentials:**
- Optic neuritis
- NAION/AION
- Macular disease
- Retinal disease

### NASAL STEP

**Description:** Field loss respecting the nasal horizontal midline with at least 1 abnormal point outside 15°. No more than 1 point may be on the temporal side.

**Differentials:**
- Glaucoma
- Chronic papilloedema
- Optic nerve drusen
- Retinal disease

### ALTIMUDUAL

**Description:** Field loss that respects the horizontal midline.

**Differentials:**
- Branch retinal artery/ vein occlusion
- NAION/AION
- Retinal disease

### ARCULATE

**Description:** Field loss extending from the blind spot to the nasal field with at least one point outside 15° nasally and at least one abnormal point temporally.

**Differentials:**
- Glaucoma
- Chronic papilloedema
- Optic nerve drusen
- Optic neuritis

### TEMPORAL WEDGE

**Description:** Small visual field defect temporal to blind spot.

**Differentials:**
- Optic neuritis
- Glaucoma (rare)
- Retinal disease

### PARACENTRAL

**Description:** A small visual field abnormality not contiguous to the blind spot and within 15° of fixation obeying the horizontal midline.

**Differentials:**
- Glaucoma
- Chronic papilloedema
- Optic nerve drusen

### ENLARGED BLIND SPOT

**Description:** Visual field loss involving at least two points contiguous to the blind spot.

**Differentials:**
- Early papilloedema
- Glaucoma (rare)
- Large peripapillary atrophy
- Optic disc drusen
- Optic nerve coloboma
- Staphyloma
- Megalopapillae
- Titted disc syndrome

### CLOVER LEAF

**Description:** Diagonal paracentral points show normal or near-normal sensitivity but all other points reduced. This is often due to patient responding normally at the start of the test only as the visual field instrument generally test these points first. Often accompanied by high fixation loss and false negatives.

**Differentials:**
- Inattentive patient
- Poor supervision
- Malingering
- Retinal disease

This reference is based on the current literature and evidence at the time of writing. This reference is designed a guide to aid diagnosis and management decisions however individual cases must be assessed in the context of all available clinical data.
### CHAIR-SIDE REFERENCE: VISUAL FIELD

<table>
<thead>
<tr>
<th>STRUCTURE/FUNCTION</th>
<th>VISUAL PATHWAY</th>
<th>CLINICAL PEARLS</th>
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| The relationship between the retinal nerve fibre layer location and corresponding visual field is complex due to significant individual variations (Lamparter et al. 2013). The following shows a ‘typical’ structure/function relationship for the right eye adapted from Ferreras et al. 2008. Note that atypical anatomical configurations such as tilted discs and high refractive error can change the structure-function relationship. | There are many ocular and neurological conditions that can lead to field defects with the following diagram showing the possible location of a visual pathway defect based on the pattern of field loss. Note, however, that:  
• field loss often does not precisely follow the pattern as outlined below;  
• partial losses or losses that are not entirely symmetrical are common. | RELIABILITY  
• Unless there is a correlating structural finding, field defects need to be repeatable before they can be considered to be clinically significant due to large variability, especially in the periphery.  
• False positive errors (>15% should be concerning) have a greater effect on visual field reliability than the fixation loss or false negative errors.  
• Increased false negative errors are correlated with the severity of visual field loss, even with reliable visual field takers (Bengtsson and Heijl 2000) and thus should not be used for assessing reliability in isolation.  
• Blind spot based fixation monitoring is generally ineffective, and other forms of fixation monitoring such as gaze-monitoring and practitioner observation needs to be used instead. |
| Figure 1. A map showing the relationship between RNFL sectors and test points on a 24-2 field test adapted from Ferreras et al. IOVS 2008. | | VISUAL FIELD DEFECTS  
• A visual field area with “complete loss” (e.g. <0dB) is not necessarily completely blind. A target with a greater luminance or size may still be visible.  
• Remember that checking the raw sensitivity results should be performed in conjunction with the probability maps: the former gives depth information whilst the later presents statistically significant anomalies and patterns of loss. |
| Figure 2. A diagram showing the visual pathway and field loss that may result from different injuries. Grey denotes scotoma on the right hand diagrams. (Zangerl et al Clin Exp Optom 2017) | | GLAUCOMA  
• In glaucoma, either structural loss or functional loss can occur first depending on the sensitivity of the devices used to detect the loss (Keltner et al. 2006), i.e. do not rely solely on imaging for “pre-perimetric glaucoma”  
• Central field loss may be seen in as many as 50% of glaucoma cases (Schiefer et al. 2010) and thus, a 10-2 field or equivalent may be useful.  
• 24-2 is designed for glaucoma assessment; if a non-glaucomatous defect, especially in neurological assessments, is suspected, utilise a 30-2 instead. |

This reference is based on the current literature and evidence at the time of writing. This reference is designed a guide to aid diagnosis and management decisions however individual cases must be assessed in the context of all available clinical data.