Eye signs for the neurologist in the Intensive Care Unit

Sreenivas Umaiorubahan Meenakshisundaram1, Prabash P.R.2, Meenakshisundaram Umaiorubahan2

1Institute of Neurology, Madras Medical College, Chennai, India
2Apollo Hospitals, Chennai, India

ABSTRACT

Introduction. The eyes are a window to the brain’ is a maxim that holds true especially in the intensive care setting. Recognising specific eye signs aids rapid decision-making regarding diagnosis or prognosis. Eye signs play a pivotal role in intensive care for the neurologist.

State of the art. Eye signs have long been considered the best clinical clue for assessment of a comatose patient. In critically ill patients, the recognition of brainstem involvement hinges primarily on eye signs. The ability to recognise and interpret these signs goes a long way towards ensuring proper care of neurological illness in intensive care units.

Clinical implications. In this article we enumerate the various signs to be assessed in the ocular and periocular structures. We look at the various types of nystagmus and abnormal eye movements which help to localise lesions in the brainstem. This will aid better diagnosis and prognostication. We categorise eye signs as Category 1 or 2 according to whether they are periorbital and ocular signs or oculomotor abnormalities. Category 2 signs are further sub-classified into Category 2a – common and Category 2b – uncommon.

Future directions. Clinical anatomical correlation of specific signs such as ocular dipping has yet to be elucidated. Research that looks into specific eye signs may help with better anatomic correlation and localisation of lesions.

Key words: eye signs, nystagmus, coma, intensive care, pupil, oculomotor abnormalities

Introduction

Just as the ‘face is the index of the mind’, to a neurologist ‘the eyes are the window to the brain’. This is especially true in the setting of acutely ill patients encountered in the intensive care unit. Eye signs can be of diagnostic, prognostic and therapeutic value.

The key is to spend some time observing the eyes because some signs can be made out only on examination lasting for at least a few minutes (up to five minutes for some) as this is likely to be rewarding and exciting.

The eye signs in the ICU of interest to a neurologist can be divided into two broad categories (Tab. 1, 2):

1. signs involving the periorbital, ocular structures including the pupils, eyelids, conjunctivae, lens and cornea
2. signs of eye movement abnormalities – paresis and abnormal movements. This review, primarily emphasising the clinical value of eye signs, will focus on this category of signs because a detailed enumeration of the first category is beyond the scope of this article.

The second category can be further divided into:

2a. common signs including gaze preferences, deviations, dysconjugate eyes, and common types of nystagmus

Table 1. Localisation of pupillary abnormalities

<table>
<thead>
<tr>
<th>Pupil size and location</th>
<th>Site of lesion</th>
</tr>
</thead>
<tbody>
<tr>
<td>small and reactive</td>
<td>Diencephalon</td>
</tr>
<tr>
<td>Midposition and fixed</td>
<td>Midbrain</td>
</tr>
<tr>
<td>Dilated and fixed, down and out</td>
<td>3rd cranial nerve</td>
</tr>
<tr>
<td>Large and fixed</td>
<td>Tectal</td>
</tr>
<tr>
<td>Pinpoint</td>
<td>Pons</td>
</tr>
<tr>
<td>Small and reactive</td>
<td>Metabolic</td>
</tr>
</tbody>
</table>

Address for correspondence: Meenakshisundaram Umaiorubahan, “Vilvam”, 4, Yadavali Street, Virugambakkam, Chennai, India 600092, e-mail: drums.neuro@gmail.com
Table 2. Categorisation of eye signs

<table>
<thead>
<tr>
<th>Sign</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Category 1</strong></td>
<td></td>
</tr>
<tr>
<td>Exophthalmos</td>
<td>Retrobulbar lesion, thyrotoxicosis, sphenoid wing mass</td>
</tr>
<tr>
<td>Periorbital ecchymoses</td>
<td>Base of skull fracture</td>
</tr>
<tr>
<td>Conjunctival congestion</td>
<td>Carotico cavernous fistula, bleeding diatheses, endocarditis, trigeminal autonomc cephalgia</td>
</tr>
<tr>
<td>Band Keratopathy</td>
<td>Hypercalcemia</td>
</tr>
<tr>
<td>Kayser Fleischer ring</td>
<td>Wilson's disease</td>
</tr>
<tr>
<td>Ptosis</td>
<td>Myasthenia gravis, 3rd nerve palsy, Horner's syndrome</td>
</tr>
<tr>
<td>Pupillary changes</td>
<td>Brainstem</td>
</tr>
<tr>
<td><strong>Category 2a</strong></td>
<td></td>
</tr>
<tr>
<td>Down and out eye</td>
<td>3rd cranial nerve palsy</td>
</tr>
<tr>
<td>Down and out with dilated pupil</td>
<td>Posterior communicating artery aneurysm</td>
</tr>
<tr>
<td>Down and out with spared pupils</td>
<td>Ischaemic 3rd nerve palsy</td>
</tr>
<tr>
<td>Down and out with contralateral superior rectus weakness</td>
<td>3rd nerve nuclear lesion</td>
</tr>
<tr>
<td>Extorted and elevated eye</td>
<td>4th nerve palsy</td>
</tr>
<tr>
<td>Opposite head tilt</td>
<td>4th nerve palsy</td>
</tr>
<tr>
<td>Lateral head turn</td>
<td>6th nerve palsy</td>
</tr>
<tr>
<td>Complex ophthalmoplegia</td>
<td>Dysthyroid state, mitochondrial disorders, myasthenia</td>
</tr>
<tr>
<td>Internuclear ophthalmoplegia</td>
<td>Medial longitudinal fasciculus</td>
</tr>
<tr>
<td>One and a half syndrome</td>
<td>MLF and PPRF</td>
</tr>
<tr>
<td>Gaze preference</td>
<td>Supratentorial - I/L, Brainstem - C/L</td>
</tr>
<tr>
<td>Down and in eyes</td>
<td>Thalamus, upper midbrain</td>
</tr>
<tr>
<td>Wrong way eyes</td>
<td>Medial thalamic haemorrhage</td>
</tr>
<tr>
<td>Skew deviation</td>
<td>Lesion on side of hypotropic eye</td>
</tr>
<tr>
<td>Upbeat nystagmus that increases in upgaze</td>
<td>Cerebellar vermis</td>
</tr>
<tr>
<td>Upbeat nystagmus that decreases in upgaze</td>
<td>Medulla</td>
</tr>
<tr>
<td>Downbeat nystagmus</td>
<td>Cervicomedullary junction</td>
</tr>
<tr>
<td>Rebound nystagmus</td>
<td>Cerebellum</td>
</tr>
<tr>
<td>Multidirectional, gaze evoked nystagmus</td>
<td>Cerebellum</td>
</tr>
<tr>
<td>Unidirectional, mixed nystagmus</td>
<td>Peripheral vestibular apparatus</td>
</tr>
<tr>
<td>Direction changing nystagmus</td>
<td>Lateral medulla</td>
</tr>
<tr>
<td><strong>Category 2b</strong></td>
<td></td>
</tr>
<tr>
<td>Convergence retraction nystagmus</td>
<td>Periaqueductal grey</td>
</tr>
<tr>
<td>Ocular flutter</td>
<td>Dentate nucleus</td>
</tr>
<tr>
<td>Opsoclonus</td>
<td>Infectious / parainfectous / paraneoplastic</td>
</tr>
<tr>
<td>Ocular bobbing</td>
<td>Pons</td>
</tr>
<tr>
<td>Ocular bobbing</td>
<td>Brainstem</td>
</tr>
<tr>
<td>Seesaw nystagmus</td>
<td>Sellar / parasellar masses</td>
</tr>
<tr>
<td>Periodic alternating nystagmus</td>
<td>Nodulus</td>
</tr>
<tr>
<td>Oculogyric crisis</td>
<td>Postencephalitic parkinsonism / drug-induced</td>
</tr>
<tr>
<td>Roving eyes</td>
<td>Light coma</td>
</tr>
<tr>
<td>Ping-pong eyes</td>
<td>Bihemispheric dysfunction</td>
</tr>
</tbody>
</table>

2b. uncommon/ominous signs including ocular flutter, opsoclonus, ocular bobbing, ocular dipping and a few others. This article will then discuss ocular manifestations of stroke and of brain death. It is intended as a guide to the clinician when eye signs are encountered in the Intensive Care Unit.

Category 1 signs can be:

1. Exophthalmos (unilateral) – causes include thyroid eye disease, orbital mass lesion, orbital pseudotumour, cavernous sinus thrombosis, carotid cavernous fistula, meningocoele, sphenoid wing meningioma, orbital neoplasm,
mucormycosis, vascular malformation of the orbit, and neurofibromatosis. Horner’s syndrome causes apparent enophthalmos [1].

II. Periorbital ecchymosis (so-called ‘raccoon eyes’) indicates the presence of basal skull fractures [2].

III. Carotid cavernous fistula can cause tortuous (corkscrew) blood vessels in the conjunctiva. A subconjunctival haemorrhage may indicate bleeding disorders or endocarditis. Conjunctival injection may be a part of trigeminal autonomc cephalalgia, Horner’s syndrome [3].

IV. Copper deposition in the Descemet membrane results in he
cratically. Ptosis in Lambert-Eaton syndrome may temporarily
be seen in isolation or as part of dyskinesias and drug-
induced movement disorders. When orbicularis oculi is
contract, due to the left MLF lesion. As a result, gaze to the
right results in abduction of the right eye but no adduction
of the left eye. Typically the abducting eye shows nystagmus.
Slowness of the adducting saccades compared to the abducting
saccades may be an early sign of an INO due to demyelination,
and is a very useful sign for diagnosis. Patients with bilateral
INO have been said to have WEBINO syndrome (wall-eyed
bilateral INO) [12].

A medial pontine lesion (e.g. on the right) can affect both the right PPRF and the right MLF crossing from the contra-
lateral side. Because of the right PPRF lesion, the patient has a complete horizontal gaze palsy to the right side. Because of the right MLF lesion, the patient has an INO on the right side resulting in adduction failure of the right eye. The only eye move-
ment possible is abduction of the left eye. The constellation of
these findings is termed ‘one-and-a-half syndrome’. It should
be borne in mind that the 6th cranial nerve nucleus has both nuclear and internuclear components and a nuclear 6th cranial nerve can be associated with an INO (and 7th nerve involvement) [13].

● Gaze deviations/preferences

Unilateral frontal lobe seizure activity pushes the eyes to the opposite side. With destructive frontal lobe lesions, the patient is unable to move their eyes contralaterally. This is known as gaze paresis or preferential gaze to the opposite side. The normal PPRF pulls the eyes over, so destructive lesions of the PPRF impair the ability to gaze ipsilaterally, resulting in gaze deviation toward the intact side. Therefore patients with destructive frontal (supratentorial) lesions gaze away from the side of the hemiparesis, while patients with pontine (infratentorial) strokes gaze toward the hemiparesis. 'Downward and inward' eyes indicate a thalamic, upper midbrain lesion.

Category 2

2a Commonly encountered signs and their significance

- Lesions of 3rd, 4th, or 6th cranial nerves:
  - Complete 3rd cranial nerve palsy causes ptosis, with an outward and mildly downward eye (unopposed action of the 4th and 6th nerves) with a dilated pupil that has no light response, directly or consensually, or near response. Pupillary function is usually spared in ischaemic lesions. A nuclear lesion of the 3rd cranial nerve may also cause weakness of the contralateral superior rectus [10].
  - In 4th nerve palsy, there is extorsion and impairment of depression of the adducted eye. The characteristic head tilting to the opposite shoulder is known as Bielschowsky sign.
  - In 6th nerve palsy, there is impairment of abduction of the affected eye, and the eye deviates medially. The head is turned towards the side of the paretic muscle [11].

Causes of complex ophthalmoplegia are dysthyroid eye
disease, myasthenia gravis, and mitochondrial disorders.

- Internuclear ophthalmoplegia

Lesions of the medial longitudinal fasciculus (MLF) cause an internuclear ophthalmoplegia (INO). When, for example, the right PPRF and 6th nerve nucleus act to initiate lateral gaze towards the right, the left medial rectus receives no signal to contract, due to the left MLF lesion. As a result, gaze to the right results in abduction of the right eye but no adduction of the left eye. Typically the abducting eye shows nystagmus. Slowness of the adducting saccades compared to the abducting saccades may be an early sign of an INO due to demyelination, and is a very useful sign for diagnosis. Patients with bilateral INO have been said to have WEBINO syndrome (wall-eyed bilateral INO) [12].
Wrong way eyes is a term used to denote deviation of the eyes towards the side of weakness of limbs in a supratentorial lesion, and has been described with medial thalamic haemorrhages. This has been attributed to compression of the frontopontine fibres at the level of mesencephalon, irritation of intralaminar thalamic nuclei, and imbalance in the bihemispheric control of pursuit mechanisms [14].

- Skew deviation
  In skew deviation there is vertical deviation (malalignment) of one eye above the other that is caused by an imbalance of the vestibular inputs to the oculomotor system. It results from a pnenuclear lesion involving the brainstem or cerebellum, and the lesion is usually on the side of the hypotropic eye, except for lesions of the midbrain, where the ipsilateral eye is hypertropic. This is a component of the ocular tilt reaction and is also usually seen with an INO. Skew deviation may sometimes alternate with lateral gaze to either side. This indicates a lesion at the pretectal midbrain or the cervicomedullary junction. They can be differentiated based on associated findings [15].

- Nystagmus
  Nystagmus denotes an imbalance and an inability to maintain fixation, resulting in unwanted slow deviation of the eyes followed by a fast deviation in the opposite direction. The abnormality is the slow deviation, although the fast component is used to denote the direction of the nystagmus. Differentiating between central and peripheral causes of nystagmus is paramount in an ICU setting. The central ocular motor signs that are in favour of a central nystagmus are: skew deviation, central fixation nystagmus, saccadic smooth pursuit, gaze-evoked nystagmus contralateral to direction of spontaneous nystagmus, rebound nystagmus, acute vertigo and nystagmus with normal head impulse test, and impaired VOR suppression [16].

  In upbeat nystagmus, there is up-beating nystagmus on primary gaze. If the nystagmus increases in upgaze the localisation is in the cerebellar vermis, but if the nystagmus decreases in upgaze, the lesion is in the medulla.

  Downbeating nystagmus in primary gaze is downbeat nystagmus, maximal in eccentric and downgaze. The location of the pathology may be the cervicomedullary junction.

  Rebound nystagmus is horizontal gaze-evoked nystagmus that changes direction with refixation to the primary position – here the pathology is in the cerebellum or cerebellar connections.

  Nystagmus in posterior circulation stroke:
  - Medbrain – convergence retraction, dissociated (INO), rotary (torsional),
  - Pons – INO, ocular bobbing (although not exactly a nystagmus),
  - Medulla oblongata – may imitate peripheral type,
  - Cerebellum – peripheral type, multidirectional gaze-evoked.

  Lateral medullary syndrome is a treasure trove of eye signs. The characteristic nystagmus is the direction-changing type: horizontal-rotary to one side and horizontal to the other side, i.e. with features of peripheral vestibular dysfunction to one side and a central fixation type on looking to the other side [17].

2b Uncommon eye signs

In Parinaud’s syndrome, there is impaired upgaze. When patients attempt to look up, the eyes may spasmódically converge and retract backward into the orbits. This phenomenon is known as convergence-retraction nystagmus. This is in fact a misnomer, because there is no slow deviation to start with. Other features include eyelid retraction (Collier’s sign) with a poor, rarely absent, light response and much better near response (tectal pupils). This is due to a lesion involving the periaqueductal region [18].

Ocular flutter - intermittent, rapid, back-to-back horizontal saccades causing a quivering movement. Ocular flutter and opsoclonus are a continuum, and the location of the pathology is in the cerebellum or brainstem cerebellar connections, dentate nucleus. They are clinically differentiated by the fact that ocular flutter occurs only in the horizontal plane, while opsoclonus is multiplanar. Opsoclonus consists of involuntary, arhythmic, chaotic, multidirectional saccades, without intersaccadic intervals. Saccades may be horizontal, vertical, or torsional. It is present during fixation, smooth pursuit, convergence, and persists during sleep or eyelid closure. Opsoclonus differs from nystagmus in that the phase that takes the eye off the target is always a saccade, not a slow eye movement. Opsoclonus may be infective / parainfectious or paraneoplastic, and is likely to be a part of the opsoclonus-myoclonus-ataxia syndrome [19].

Ocular bobbing is a swift downward jerk with slow drift back to the primary position. The patient is often unconscious. These abnormal movements in the vertical plane indicate a lesion involving the horizontal gaze mechanisms, and hence localised to the pons. This needs to be differentiated from a downbeat nystagmus, which starts with a slow deviation upwards. Absent horizontal eye movements in ocular bobbing also helps to differentiate them [20].

Ocular dipping, reverse ocular bobbing, and reverse ocular dipping are other rare signs that occur with brainstem lesions and diffuse cerebral hemispheric dysfunction [20].

In seesaw nystagmus, there is pendular torsional-vertical oscillation; one eye rises and intorts, the other falls and extorts; this is sometimes associated with bitemporal hemianopia caused by sellar or parasellar masses [21].

Periodic alternating nystagmus is a remarkable horizontal jerking that periodically (roughly every 90 seconds) changes direction, interposed with a brief neutral period during which the eyes show no nystagmus, or a jerk downward. Periodic alternating nystagmus (PAN) is seen with lesions in the nodulus of the cerebellum. PAN can be abolished by the administration of baclofen [22].

Oculogyric crisis refers to attacks of involuntary conjugate upward deviation of the eyes, which may be transient or last for hours, and it is classically associated with postencephalitic parkinsonism. It can also be an acute dystonic reaction caused by dopamine receptor blocking agents [23].
Roving conjugate eye movements are characteristic of a light coma [9].

Ping-pong gaze, where slow horizontal ocular deviations shift every few seconds from side to side, is a form of roving eye movement that occurs with bihemispheric infarctions. In a similar phenomenon called ‘windscreen-wiper eyes’, there is slower, side-to-side pendular oscillation of the eyes. This is also associated with bilateral hemispheric lesions that have presumably released a brainstem pacemaker [9].

The absence of eye movements after VOR and caloric testing is an important step in documenting brain death. Lack of consistent visual following of objects is a sign of vegetative state.

The eye signs in brainstem dysfunction are: loss of spontaneous eye movements; midposition of the eyes; lack of response to oculocephalic and caloric (oculovestibular) testing; and the presence of dilated or midposition fixed pupils (no smaller than 3 mm) [9].

Conclusion

Examination of the eye is one of the most important steps in the assessment of a patient in the intensive care unit. The eye offers important clues in terms of localisation and prognosis. Future research may provide information about the clinical significance and localisation of signs such as ocular dipping and opsoclonus, which as yet do not have a specific anatomical substrate to explain the findings. Even in this age of advanced imaging and electrophysiological examination, clinical examination of the eyes remains a valuable tool in the diagnostic armamentarium of the neurologist.

Ethical permissions: No ethical approval was necessary for the preparation of this article

Conflict of interest: None

References


www.journals.viamedica.pl/neurologia_neurochirurgia_polska

Neurologia i Neurochirurgia Polska 2019, vol. 53, no. 6

www.journals.viamedica.pl/neurologia_neurochirurgia_polska

Neurologia i Neurochirurgia Polska 2019, vol. 53, no. 6

www.journals.viamedica.pl/neurologia_neurochirurgia_polska


