Nystagmus in pediatric patients: interventions and patient-focused perspectives

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Abstract: Nystagmus refers to involuntary, typically conjugate, often rhythmic oscillations of the eyes. The most common cause of nystagmus in children is infantile nystagmus syndrome (INS). INS presents within the first few months of life and is sometimes accompanied by an ocular condition associated with sensory impairment. Because this condition affects a person throughout life, it is important to understand the options available to manage it. This review focuses on the underlying nystagmus etiology, psychosocial and functional effects of nystagmus, as well as current principles of management, including optical, pharmacological, surgical, and rehabilitative options. Currently, the neural mechanisms underlying INS are not fully understood. Treatment options are designed to increase foveation duration or correct anomalous head postures; however, evidence is limited to mainly pre- and post-study designs with few objective comparisons of treatment strategies. Management of INS should be individualized. The decision on which treatment is best suited for a particular patient lies with the patient and his/her physician.

Keywords: nystagmus, infantile nystagmus syndrome, vision impairment, pediatric, quality of life

Introduction
Nystagmus refers to involuntary, typically conjugate, often rhythmic oscillations of the eyes. There are three types of nystagmus that are most likely to be encountered in children: infantile nystagmus syndrome (INS), fusion maldevelopment syndrome nystagmus (previously known as latent/manifest latent nystagmus), and spasmus nutans. This review focuses on INS. Some believe that fusion maldevelopment syndrome nystagmus represents a monocular form in the same spectrum as INS.1 The term congenital nystagmus is often used synonymously with INS; however, it is technically incorrect as nystagmus does not typically develop at birth but more likely at 2–3 months of age. INS may be associated with retinal or optic nerve maldevelopment (previously known as sensory nystagmus) or may occur in isolation (previously known as congenital motor nystagmus). Conditions commonly associated with nystagmus include albinism, aniridia, achromatopsia, cone dystrophy, optic nerve hypoplasia, foveal hypoplasia, congenital cataracts, corneal opacities, retinopathy of prematurity, Leber congenital amaurosis, and syndromic causes associated with early-onset retinal degenerations. The nystagmus associated with visual sensory deficit is identical to that which presents in isolation; however, acuity is typically worse in those with sensory deficit.2

Assessment of nystagmus
When assessing an infant or child with nystagmus, it is important to determine the age of onset as well as the child’s birth, developmental, medical, and family history. It is imperative to identify an underlying etiology if present, as the associated
ocular or associated systemic condition may require intervention. Newer high-resolution imaging modalities such as optical coherence tomography (OCT) are increasingly being used for determining the cause of INS in children. The ability to use imaging to augment the clinical evaluation is important as patients are often misdiagnosed as having idiopathic INS, when in fact there is an underlying ophthalmic diagnosis and foveal maldevelopment can be found on OCT.^3^ Neuro-imaging should be considered when nystagmus onset is after 3 months and not associated with an underlying sensory deficit or when associated with optic nerve hypoplasia.~4~

Clinically, the evaluation of nystagmus includes measurement of best-corrected visual acuity and a description of characteristics of the waveform (eg, direction, type, amplitude, frequency) as well as documentation about any head turns or tilts and location of the null point, if present. To further characterize nystagmus, eye movement recordings are utilized. These recordings may be used to determine change after medical or surgical intervention. There are at least three methods used to quantify foveation characteristics in patients with nystagmus. The first, developed in Dell’Osso’s lab, is the Expanded Nystagmus Acuity Function (NAFX) (Figure 1).~5~ Two additional measures (the nystagmus optimal fixation function~6,7~) have been developed that are fully automated and correlate well with the NAFX.

Dell’Osso and Daroff~2~ identified at least 12 waveforms that range from pure pendular to pure jerk. While not clinically evident, the intermediate waveforms are often punctuated by embedded saccades that increase the time the eye position is on the fovea. Nystagmus with acceleration of movement during the slow phase is considered characteristic of infantile nystagmus.~8~ The earliest form of infantile nystagmus seen tends to be pendular and develops into a jerk form in the first 2 years of life.~9~ The presence of a null point or zone is also characteristic of infantile nystagmus. The null point is typically within 10° of fixation with lateral head turns the most common adaptation.~9~ How the null point develops is poorly understood and has received surprisingly little research attention. The absence of illusory movement of the visual world (oscillopsia) is a characteristic of infantile nystagmus, which separates it from acquired forms. Visual perception is normally suppressed during saccades to prevent smear of the visual image.~10~ It has been postulated that the eye movement generation systems produce a copy of the nystagmus eye movement to be executed and send it to higher cortical areas. This “efference copy” serves as the template for the expected movement at the cortical level.~11~ Since there is a match between the true eye movement and the expected eye movement, no perception of visual motion is generated. In acquired nystagmus, a mismatch between the expected and actual location occurs, or possibly, multiple signals are misinterpreted, resulting in the illusion of motion of the visual world.

Genetic analysis may also further define what appear to be idiopathic cases. INS may be inherited in a dominant, recessive, or X-linked fashion. Five different chromosomal abnormalities (NYS1–5) have been linked to INS. Mutations in the X chromosome at locus q26 have been associated with typical clinical INS as well as periodic alternating nystagmus. The FRMD7 gene associated with this defect is found in areas of the brain involved in eye movement control and may play a role in neurite development. The defect has a relatively high penetrance in females. Mutations in FRMD7 have been found in less than 10% of sporadic cases.~12~

### Mechanisms for development of nystagmus

The underlying mechanism of infantile nystagmus is not fully understood, although theories abound that attempt to explain its development.~13,14~ Most implicated defects are found in fixation, saccades, the optokinetic reflex, the neural integrator, or the pursuit eye movement systems. No single theory has as yet been able to explain the development of the condition in all its complexity in an oculomotor movement system that largely has intact saccades, pursuits, and vestibulo-ocular reflex (VOR) movements outside the nystagmus.~15,16~ Three systems play a role in maintaining a stable image on the retina: fixation, the VOR, and gaze control. During the first few weeks of life, fixation is poor as the fovea is maturing,
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and orientation to targets is largely mediated by subcortical pathways through the superior colliculus. This rapidly changes as top-down pathways from the cerebral cortex establish greater control of eye movements at approximately 2–3 months of age. Larger waveform saccadic oscillations and intrusions have been reported to precede the onset of the infantile nystagmus waveform.

The vestibulo-ocular system holds the eyes stable during head movement. The VOR is one of the earliest developing of the image stabilization control systems. In patients with INS, the VOR system appears to function normally with a superimposed nystagmus. Invoking the VOR, head movement, or head shaking does not suppress movement or improve visual acuity in nystagmus.

The gaze control system holds the eyes stable when they are in an eccentric position and is coordinated by the neural integrator. The neural integrator is proposed to convert the velocity of the head’s movement generated by the VOR to a neural signal that is to be sent to the eye muscles for target tracking. In gaze-evoked nystagmus, it is presumed that the neural integrator becomes “leaky” allowing the eyes to drift away from the object of regard with a correcting saccade back to the target. During early infant development, it is likely that the neural integrator is “calibrating” itself. Inappropriate input or feedback during this calibration period could lead to a lifelong error signal. A number of theories of the development of nystagmus have postulated that errors in the inputs, outputs, or feedback loops within the neural integrator are at fault in the development of infantile nystagmus.

The optokinetic system is fundamental for maintaining a steady image as image flow moves across the retina. It is functionally present shortly after birth for large slow-moving targets. Optokinetic nystagmus (OKN) is nystagmus in response to stimuli drifting across the subject’s visual field. OKN’s similarity to nystagmus waveforms has led to theories that propose it as a possible driver of INS. The OKN

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Figure 2 Depiction of the brain, as viewed from above, showing normal cortical and subcortical projections during early human development. 

Notes: (A) Early in infancy, horizontal optokinetic stimuli (shown as leftward or rightward motion) from each nasal retina are transmitted via a subcortical pathway to the contralateral nucleus of the optic tract-dorsal terminal nucleus (NOT-DTN) of the accessory optic system (solid red arrow), which is directionally sensitive to ipsiversive motion (ie, nasalward for the contralateral eye). During this early stage of development, the cortical pursuit pathways (shown as corticofugal projection from the middle temporal area–medial superior temporal area [MT–MST] to the ipsilateral NOT-DTN) have not yet become functional (interrupted green arrows). (B) Later in infancy, horizontal optokinetic responses become encephalized, binocular cortical pursuit pathways become fully operational (solid green arrows), and subcortical optokinetic pathways regress (interrupted red arrows). Reproduced from Brodsky MC, Dell’Osso LF. A unifying neurologic mechanism for infantile nystagmus. JAMA Ophthalmol. 2014;132(6):761–768.1

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Abbreviations: L, left eye monocular cells; LGN, lateral geniculate nucleus; R, right eye monocular cells; R+L, cortical binocular cells; SCC, splenium of the corpus callosum; V1, primary visual cortex.
response is mediated through the dorsal terminal nucleus, lateral terminal nucleus, and medial terminal nucleus in the pretectum which collectively comprise the accessory optic system (AOS) as well as the nucleus of the optic tract (NOT). Symmetry in the monocular OKN develops by months 3–5 as binocular-driven motion pathways from the middle temporal and medial superior temporal cortex provide greater control and effectively override any direct retinal input (Figure 2).24 Brodsky and Dell’Osso have recently proposed that delays in cortical pathway maturation could allow the retinal–AOS–NOT system to act unimpeded with the resulting development of infantile nystagmus.1 They have posited the unimpeded AOS–NOT system activity as the unifying process of infantile nystagmus. Tychsen et al has suggested that a failure of binocular cortical connections could also lead to nystagmus in and of itself.25 These two opposing theories are referred to as bottom-up (Brodsky) or top-down mechanisms (Tychsen). While connections may not be fully developed for several months, the cortex does appear to have some very early connectivity with brainstem structures. Infants born without cortical structures do not develop OKN, and OKN is severely impaired in early cortical damage.26,27

The role that the eye muscles might play in nystagmus has received little attention. Biopsy of ocular muscle from nystagmus surgery patients has shown that there are more centrally nucleated muscle fibers with greater variation between them than is seen in normal extraocular muscle tissue.28 Unique proprioceptors are found within the extraocular muscles and appear to be altered in INS compared to normal. It is possible that proprioceptor function changes after muscle surgery for nystagmus.29

Brain imaging studies are also relatively rare. Hüfner et al using a voxel-based morphology analysis, found that the areas associated with motion processing in middle temporal lobe and areas near the cerebellar vermis had greater volumes in those with higher amplitude infantile nystagmus than those with lower amplitudes.30 Schlindwein et al reported positron emission tomography scan results in a patient with infantile pendular nystagmus that showed increased activity in the cerebellar nodulus, an area thought to be associated with the torsional optokinetic response.31 Leguier et al report that activity in the cerebellar declive, a subregion of the vermis, shows increased activity when the gaze is focused away from the null zone compared to when the gaze is directed into the null zone.32 Harris has pointed out that care needs to be taken while interpreting these results and that changes in function in the cerebellum may be a result of, rather than a cause of, nystagmus.

**Functional implications and quality of life**

Relatively little is known about vision-specific, health-related quality of life in children with vision impairment. The few studies that do exist evaluated groups of children with varied etiologies for vision impairment, only some of which are associated with nystagmus. Most studies of the effects of nystagmus on quality of life and visual function have been conducted using adult participants. There is clearly a need for more research in this area.

DeCarlo et al investigated the effects of pediatric vision impairment on quality of life using parallel focus groups of children with vision impairment and their parents. Nystagmus was present in 76% of participating children.33 The focus groups utilized a semi-structured script that covered topics related to general vision, mobility, school functioning, social relations and activities, self-care, medical treatment, and family impact. In focus groups of children, the largest number of negative comments pertained to the psychosocial aspects of their vision impairment. One child was noted to say “It’s weird when people look at my eyes. They’re like what’s wrong with your eyes, stop doin’ that and I can’t…” Parents also reported difficulties with teasing because of nystagmus but expressed the most negative comments about school functioning. The negative effects of nystagmus are twofold: it is an easily observable manifestation of their ocular condition that makes them different from their peers, and it is associated with decreased vision which can also have negative effects if appropriate accommodations are not made. However, there were also a substantial number of positive comments, specifically with regard to things they could do, and the beneficial effects of adaptive technology and accommodations as well as positive peer support from friends.

In a study of adults with nystagmus using a structured interview, 18 out of 21 participants specifically mentioned difficulties with the cosmetic appearance of nystagmus as well as other’s avoidance responses. Participants also reported poor self-esteem and increased dependency. In contrast to the previously discussed study of children33 that reported both positive and negative experiences, this study revealed universally negative experiences about living with nystagmus.

Another study using the VF-14 and a social function questionnaire mailed to members of the Nystagmus Network...
UK found that the impact of nystagmus on visual function was low compared to other chronic eye conditions that result in low vision. They also found that children with nystagmus had poorer social function than adults with nystagmus and that poor visual function was associated with poor social function. However, the authors caution that membership in the Nystagmus Network UK may be biased toward those with poorer visual function.

Children with vision impairment on average read slower than their normally sighted peers. The majority of children with vision impairment have nystagmus; however, no studies have focused solely on those children with nystagmus. Barot et al’s study of reading in adults with nystagmus found that maximum reading speeds were 18.8% slower in participants with albinism and 14.7% slower in those with idiopathic infantile nystagmus when compared to normally sighted controls. Woo and Bedell also studied adults with nystagmus and found that they read more slowly than controls when using either continuous text or rapid serial visual presentation (RSVP) text. Continuous text was read more quickly, presumably due to a lack of contextual clues in RSVP text. The most significant finding of this work was that reading speed exceeded the nystagmus frequency, suggesting that reading is still occurring during non-foveating periods. Readers with INS sometimes read much better than expected; this may be due to the strategies they use to manipulate their nystagmus to best obtain information from text. Nystagmus in and of itself cannot be the root cause of reading difficulties in children with nystagmus.

Nystagmus waveforms have been reported to be favorably, negatively, and unaffected by increased visual demand. The differences in these findings have been suggested to be related to the importance of the visual task. This has implications for how the relationship between nystagmus characteristics and visual acuity should be measured post-intervention. Typically, the two are not measured simultaneously, where the changes of interest might be noted.

There are many anecdotal reports of increased nystagmus intensity under conditions of stress, but only few studies have quantified it. Cham et al induced stress by having participants perform mental arithmetic orally, while they were responding to a Landolt C visual acuity task via a keyboard. A reward/penalty system was utilized for correct/incorrect responses. Heart rate as well as amplitude, frequency, and intensity of nystagmus increased during the task, although the two did not correlate. To further investigate this relationship, Jones et al used a two-alternative forced choice staircase procedure for Landolt C visual acuity under conditions of acclimation and relaxation as well as when subjects received a small electrical shock for incorrect answers (task demand condition) or at random intervals (anticipatory anxiety). They found that although nystagmus amplitude and intensity significantly increased while foveation periods decreased, visual acuity was not reduced. Shock increased reaction times, and INS participants were slower to respond than controls, consistent with reports that nystagmus contributes to a “slow-to-see” phenomenon.

Interventions for nystagmus
Refraction management
Interventions for INS all strive to increase fixation stability (by lengthening foveation time and increasing accuracy during foveation). Improving cosmesis by decreasing the nystagmus intensity (amplitude × frequency) is a secondary goal but one that is often very important to patients. The first intervention to be considered is optimal refractive correction, as children with nystagmus commonly have refractive error, especially with-the-rule astigmatism. This is important even at young ages, as the prevalence and magnitude of with-the-rule astigmatism increase with age, and there is little evidence of emmetropization through age 8. Adding yoked prism to the spectacles of patients with anomalous head posture (apex pointed in the direction of the patient’s gaze) can be helpful in alleviating the anomalous head posture when the angle is small. We have found that prism greater than 10 prism diopters becomes heavy and cosmetically unacceptable to most patients even when using high-index lenses. Using Fresnel prism will degrade image quality, so this technique is typically reserved for patients with small head turns. Prism may also be useful for patients with fusion whose nystagmus dampens in convergence. In this case, base-out prism may be trialed binocularly to stimulate convergence (additional minus lens power must be incorporated into the spectacles to compensate for convergence-induced accommodation). Positive response to prism treatment is a good predictor of response to surgical management.

Contact lenses can also be considered. Many patients needlessly worry that due to their nystagmus, contact lenses will not be an option. Contact lenses have the advantage that they maintain the optical center of the correction on the patients’ visual axis, which may make them preferable to spectacle lenses, especially for those with high refractive errors. Contact lenses broaden the high foveation-quality field, presumably through afferent feedback from the
ophthalmic division of the trigeminal nerve. Most reports are case studies and have conflicting results. However, the only randomized controlled trial published to date found no differences in nystagmus intensity comparing soft contact lenses, rigid gas-permeable lenses, and spectacles using an unmasked cross-over study design. They did, however, find a statistically but not clinically significant decrease in acuity with soft contact lenses compared to spectacles or gas-permeable contact lenses. Pharmacological and/or surgical treatment of INS is typically initiated after optimal refractive correction has been in place, but visual acuity or function is still not satisfactory or there is an anomalous head posture present.

Pharmacological treatment

Several case reports have described the benefit of various drugs in treating INS. Diethylpropionate was reported to improve visual acuity, stereopsis, exotropia, and ocular motility recordings in a female adult with INS. Ocular motility recordings showed increased foveation time and broadened null zone. Another report showed improvement in visual acuity and nystagmus intensity after smoking 10 mg of cannabis. Botulinum toxin injections have also been shown in a case series to be beneficial.

While several studies have been done on pharmacological treatment of acquired nystagmus, little research has been conducted on the treatment of INS. One such study looked at the effect of baclofen on infantile periodic alternating nystagmus. The proposed mechanism of action is to restore inhibitory control of the gamma-aminobutyric acid (GABA)-mediated vestibular nuclei with baclofen, a GABA agonist. Baclofen was started at 15 mg and increased until the patient felt a significant improvement clinically. Four out of eight patients had improved binocular Snellen visual acuity, and four reported improvement in head posture; however, four patients had to withdraw due to side effects. The authors concluded that a trial of baclofen in periodic alternating nystagmus might be valuable before considering surgical intervention. A cross-over, double-masked trial of topical brinzolamide (Azopt) versus placebo in treating INS showed improvement in nystagmus waveform characteristics and NAFX in all five patients. Four out of the five patients also had improved visual acuity. The mechanism of action of carbonic anhydrase inhibitors on INS is not well understood.

Gabapentin and memantine have also been shown to be beneficial in acquired nystagmus. Gabapentin binds a subunit of voltage-dependent calcium channels. Memantine selectively blocks excess glutamatergic activity. However, the mechanism by which these drugs reduce nystagmus is not known. One small study reported improvements in visual acuity and nystagmus amplitude in both acquired nystagmus and INS. These early reports of improvement led to a randomized, placebo-controlled trial of gabapentin and memantine in 48 adults with INS. The starting dose of memantine was 5 mg and could be increased up to 40 mg, while gabapentin was started at 300 mg and increased up to 2,400 mg. There was a statistically significant improvement in visual acuity for both treatment groups over the placebo group. Interestingly, the idiopathic infantile nystagmus group had a significant visual acuity improvement, while those with secondary nystagmus (infantile nystagmus due to ocular pathology) did not. Eye movement recordings also showed a statistically significant improvement in foveation and decrease in nystagmus intensity in all positions of gaze for both the drugs. However, self-reported subjective improvements in vision were equal among the three groups.

Surgical treatment

Surgery for nystagmus has been performed since the 1950s when Kestenbaum and Anderson first proposed surgical alleviation of abnormal head posture (AHP). The aim of surgery for nystagmus is to correct AHP or to improve visual acuity by decreasing nystagmus amplitude or frequency. Patients with INS are candidates for surgery when they have AHP because of an eccentric gaze null point and strabismus, or when nystagmus and visual acuity improve with convergence. The most common surgeries performed involve extraocular muscle recession, resection, tenotomy, or a combination of these.

Another less commonly performed procedure called artificial divergence surgery is based on the observation of dampened nystagmus with convergence. The surgery aims to create a divergent resting position (exophoria) that must be overcome by fusional convergence, thus decreasing the nystagmus intensity. Preoperative fusional amplitudes must be measured to assure adequate potential.

Recession and resection procedures

Kestenbaum and Anderson’s recession and resection procedures were initially intended to correct AHP; however, visual acuity also improved in some cases. Anderson
recommended recession of the recti muscles whose action was in the direction of the face turn. Kestenbaum performed resections and recessions on all four horizontal recti. The goal was to move the eyes in the direction of the face turn, thus placing the eccentric null zone in primary position to decrease the AHP. Their procedures were modified by Parks and Calhoun and Harley to reduce the rate of recurrence due to undercorrection. These alterations have led to the most commonly performed procedure, known as the augmented modified Kestenbaum procedure. The appropriate goal of this procedure is to improve cosmesis and reduce AHP with a small potential for improvement in visual acuity.

Several studies have looked at the effects of multiple variations of the Kestenbaum procedure on AHP, visual acuity, and eye movements in infantile nystagmus. Kumar et al used an Anderson–Kestenbaum procedure or a modified Anderson (recession of yoke muscles) procedure with or without tenotomy in 28 patients (mean age 10.9). After excluding five patients with 20/20 vision, 12 patients had improved acuity, and the rest were unchanged. There was no difference in visual acuity or residual AHP between the surgical techniques. In another prospective study, 58 patients with infantile nystagmus were assigned to one of three groups based on preoperative measures of AHP, visual acuity, and heterotropia. Each group underwent a different variation of recession and resection procedure to correct the AHP. Overall, they saw significant improvement in distance monocular visual acuity in all groups. Interestingly, mean distance binocular visual acuity only improved when two horizontal rectus muscle recessions were performed and in INS subjects with sensory impairment. Eye movement recordings showed changes in nystagmus intensity in all groups; however, the change was only significant for the four recti muscle recession surgery.

Other variations of recession and resection procedures have also been reported for INS, including symmetric recessions of all four horizontal recti muscles with new insertion posterior to the equator causing less rotational effect on the globe. Initial reports showed significant improvement in visual acuity, amplitude, and AHP without significantly limiting ocular motility. Twelve patients underwent this procedure, and 63.6% had increased visual acuity, and 81.8% had decreased nystagmus amplitude and frequency with no limitations on ocular motility. The effects were more beneficial to the subjects with idiopathic INS than sensory INS. Recession procedures also have the advantage of being reversible. A retrospective study of this procedure showed an improvement in AHP in addition to visual acuity. Out of 20 patients, AHP was present in seven patients preoperatively, and only two surgeries were planned to address the AHP. Still, AHP improved in all patients. Because AHP was not treated in five of the subjects, these findings may be due to a decrease in primary gaze nystagmus.

Most studies look at the effect of horizontal rectus muscle recession and resection on horizontal AHP. There are fewer reports on the effects of vertical rectus or oblique muscle surgery on AHPs with INS. Hertle et al looked at the effect of surgery in 24 patients with chin up or down AHP of greater than 10°. Each patient had horizontal nystagmus with decreased intensity in vertical eccentric null gaze. Thirteen patients with chin-down position (up gaze null) received bilateral superior rectus recession, bilateral inferior oblique recession, resection, or tenotomy of one horizontal rectus in each eye. Eleven patients with chin-up position (down gaze null) had bilateral inferior rectus recession, bilateral superior oblique tenectomy, and the same treatment on horizontal recti. Postoperatively, there was a 12%–15% symmetric, bilateral limitation in vertical gaze in the direction of the recessed vertical rectus. There was a significant improvement in vertical head posture after surgery; however, none showed complete resolution. In addition, there was a statistically significant improvement in both visual acuity and foveation time.

There is a risk of anterior segment ischemia with recession and resection of all four horizontal recti. When resection is utilized, blood vessels supplying the anterior segment can be severed which can complicate further procedures for AHP. One retrospective study utilized a tucking procedure instead of resection as a variation of Kestenbaum procedure. On long-term follow-up (36/42 subjects), median head turn was 10°. Out of the 36 patients, 72% had residual head turn less than 10°, and 83% had head turn of 15° or less. Importantly, there were no reported long-term anterior segment alterations.

Tenotomy

Improved visual acuity and foveation with Kestenbaum procedures led to the hypothesis that tenotomy with reattachment at the original insertion would have a similar effect. The mechanism of action is believed to be an interruption of the afferent proprioceptive loop, possibly tied to recently discovered nerve endings at the tendino-scleral interface in the extraocular muscles. This interruption will produce a dampened peripheral oculomotor response to the nystagmus signal. Hertle et al looked at the effect of horizontal rectus muscle tenotomy in five children with INS. Postoperatively, there was improvement in NAFX immediately and up to 1 year after surgery in two patients. Four out of five patients
had improvement in visual acuity. Another study by Wang et al examined the effects of tenotomy on foveation and gaze angle in patients with INS.\(^8\) For practical purposes, testing NAFX improvement across all gaze angles is necessary because the eyes do not remain in primary position. They used the NAFX in nine subjects at different gaze angles to determine if tenotomy increased potential acuity across these gazes. Eight out of nine subjects had increased NAFX at primary position. Seven out of nine patients showed improved NAFX at the various gaze angles. Six patients had improved visual acuity; acuity was unchanged in three. They concluded that tenotomy can broaden the area of heightened visual function and those patients with more impaired visual function have greater postsurgical improvement.

**Timing of surgical intervention**

Because there are currently no cures for the visual sensory deficits leading to INS, surgical treatment should be aimed at reducing nystagmus intensity, increasing foveation at null position and various gaze angles, or decreasing velocity of slow phase during foveation.\(^2\) It is believed that these alterations to the nystagmus waveform will allow for improved visual acuity and visual function. INS usually manifests by 3 months, but surgery is typically deferred until 7 years or 8 years of age to be sure that the AHP has stabilized.\(^62\) Still, there is a chance of recurrence of an AHP. Felius and Muhanna\(^81\) found that nystagmus-induced partial binocular visual deprivation during the sensitive period of visual development contributes to decreased vision in idiopathic INS and suggest that timing of interventions should consider this.

**Low-vision rehabilitation**

Visual acuity ranges widely in patients with nystagmus, with some patients having 20/20 vision, while others are legally blind. For those with decreased acuity that impairs their ability to perform everyday tasks, vision rehabilitation, or vision habilitation in the case of people born with vision impairment, is an important treatment modality to maximize visual function. Vision habilitation may include optical or electronic magnification devices as well as training in their use and other adaptive strategies. One common strategy used by children with nystagmus is large print. The use of electronic devices such as tablet computers for reading is increasing. These devices can provide variable amounts of magnification, which make them ideal for children with nystagmus and vision impairment. There are no large-scale studies of the outcomes of vision habilitation in children. A recent systematic review of low-vision rehabilitation in children identified only 22 studies, mostly case series.\(^82\) All studies reviewed had favorable outcomes supporting the use of low-vision rehabilitation in children.

**Intervention summary**

Since nystagmus has negative psychosocial and functional consequences as discussed earlier, there is a great deal of interest in its treatment. Intervention should be individualized based on patient age, function, needs, and concerns. The first step in intervention is provision of optimal refractive correction, possibly with the inclusion of prism to stimulate convergence or correct an AHP. If this does not meet the patient’s needs, then pharmacological or surgical intervention should be considered. Unfortunately, there are no randomized controlled trial data comparing pharmacological treatment to surgical treatment or comparing surgical techniques. In fact, most reports contain pre–posttreatment data for small sample sizes. The literature does not provide sufficient evidence to establish a preferred practice pattern for the treatment of nystagmus, although it does offer treatment choices. The decision on which treatment is best suited for a particular patient lies with the patient and his or her physician. While treatments are focused on decreasing nystagmus intensity and improving acuity, vision habilitation should also be recommended for those with vision impairment that affects the ability to perform everyday tasks.

**Disclosure**

The authors have no conflicts of interest.

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