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Chapter

Eye Movement Abnormalities in Neurodegenerative Diseases

Roberto Rodríguez-Labrada, Yaiméé Vázquez-Mojena and Luis Velázquez-Pérez

Abstract

Neurodegenerative disorders consist in heterogeneous group of neurological conditions characterized by a wide spectrum of clinical features resulting from a progressive involvement of distinct neuron populations. Oculomotor abnormalities take a key place in the clinical picture of these disorders because the neurodegenerative processes involve the brain circuits of eye movements. The most common abnormalities include the saccadic dysfunction, fixation instability, and abnormal smooth pursuit. The clinical assessment of oculomotor function can help to differentiate diagnosis, while electrophysiological measures provide useful biomarkers for the understanding of disease physiopathology and progression. In this chapter, we review the state of the art of the eye movement’s deficits in some neurodegenerative diseases, such as Parkinson’s disease, Alzheimer’s disease, amyotrophic lateral sclerosis, Huntington’s disease, and the hereditary ataxias.

Keywords: eye movements, oculomotor abnormalities, neurodegenerative disorders, biomarkers, Parkinson’s disease, Alzheimer’s disease, dementia, hereditary ataxias

1. Introduction

Neurodegenerative disorders encompass a highly heterogeneous group of complex neurological disorder characterized by progressive dysfunction and loss of neuron populations leading a wide spectrum of clinical features that cause notable motor and/or intellectual disabilities regularly incompatible with the life [1]. Consequently, some of these conditions represent important public health concern and has been identified as a research priority. Although physiopathological mechanisms generally differ among neurodegenerative diseases, a great number of them are characterized by abnormal accumulation of misfolded protein resulting in the loss of their physiological function and/or the gain of toxic functions [2, 3].

Classification of neurodegenerative disorders can be established by both the cardinal clinical features and the disease proteins (Figure 1). The former characterization distinguishes those conditions characterized by dementia syndromes and the movement disorders. Among dementias, the most commonly recognized disorder is the Alzheimer’s disease. Other dementia syndromes include the frontotemporal dementia, the posterior cortical atrophy, the corticobasal syndrome, and others. Movement disorders comprise hypokinetic (such as Parkinson’s disease) and hyperkinetic (such as Huntington’s disease) conditions, as well
Eye Motility

Figure 1. Classification of neurodegenerative diseases according to cardinal syndrome (A) and disease proteins (B).

as cerebellar ataxias and motor neuron diseases (such as amyotrophic lateral sclerosis) [1, 4].

The protein-based classification includes the tauopathies, the α-synucleinopathies, the TDP-43 and FUS proteinopathies, the polyglutamine diseases, and the prion disease. Tauopathies are caused by abnormal accumulation of tau protein and B-amyloids and are represented by the Alzheimer dementia, whereas among the α-synucleinopathies are recognized as the Parkinson’s disease, dementia with Lewy bodies, and multisystem atrophy. Abnormal accumulation of TDP-43 and FUS proteins defines the physiopathology of the amyotrophic lateral sclerosis and frontotemporal lobar degeneration, whereas the polyglutamine diseases result from the accumulation of proteins with abnormally expanded polyglutamine domains and include the Huntington’s disease; the spinocerebellar ataxias 1, 2, 3, 6, 7, and 17; the dentatorubral-pallidoluysian atrophy; and the spinal and bulbar muscular atrophy. Finally, the Creutzfeldt-Jakob disease is classified as a prion disease [1, 4].

Although the phenotypical features of neurodegenerative disorders generally differ between distinct disorders due to the differential involvement of specific functional systems, most of these conditions are characterized by altered oculomotor function as a result of the high vulnerability of the oculomotor system to the toxic protein deposition and other physiopathological mechanisms causing neurodegenerative diseases [5, 6]. Accordingly, the assessment of oculomotor function has become a helpful approach to diagnose some of the neurodegenerative diseases. Besides, eye movements are usually used for monitoring of disease progression [6, 7].

This chapter is focused to review the state of the art of the eye movement’s deficits in some neurodegenerative diseases, such as Parkinson’s disease, Alzheimer’s disease, amyotrophic lateral sclerosis, Huntington’s disease, and the hereditary ataxias.
2. Brief overview of eye movements

Eye movements facilitate the clear vision stabilizing images on the retina, particularly against head and body movements, capturing and keeping specific stimuli on the fovea and aligning the retinal images in the two eyes to ensure the single vision and stereopsis. Ocular motility is guaranteed by five basic types of eye movements: the vestibulo-ocular reflex, the optokinetic reflex, the saccadic movements, the smooth pursuit movements, and the vergence [8]. Although they differ in various aspects, such as their velocity, reaction time, reflexivity/volitional degree, and their neurobiological substrates [9], all have generic kinematic properties and share a common final path represented by three cranial nerve nuclei and the three pairs of eye muscles that they control [8, 10]. Cranial nerve III (oculomotor) innervates the superior, inferior, and medial rectus muscles as well as the inferior oblique muscle, whereas trochlear (IV) and abducens (VI) nerves innervate the superior oblique and lateral rectus, respectively [10].

The vestibulo-ocular reflex (VOR) is elicited by the vestibular system in response to body/head rotations and consists on compensatory eye movements in opposite direction to body/head movements to guarantee the image stabilization on the retina [11]. When head/body rotations are very large and continued, the VOR is depressed, and thus, it is complemented by the optokinetic reflex (OKR), in which the speed and direction of a full-field image motion are computed to develop eye movements with two phases: a slow phase that alternates with resetting a quick phase [12].

Saccades are ballistic and conjugate eye movements that redirect the fovea from one object of interest to another, allowing to explore accurately the visual scenes. For that, saccadic system processes information about the distance and direction of a target image from the current position of gaze. Saccades are the fastest eye movements, reaching up to 800/s. Behaviorally, saccades may be classified as reflex-guided saccades and intentional or volitional saccades. The first ones are evoked by suddenly appearing targets, whereas the second ones, called also as higher-order saccades, are made purposefully. Therefore, intentional saccades involve high-cognitive processing and include voluntary, memory-guided and delayed saccades, as well as antisaccades [13, 14].

Smooth pursuit eye movements enable us to maintain the image of a moving object relatively stable on or near the fovea by matching eye velocity to target velocity [10]. Smooth pursuit performance is optimal for target speeds ranging between 150/s and 300/s, but pursuit velocity can reach up to 100/s [8, 15].

Vergence eye movements are disjunctive movements that provide the binocular alignment in response to changing fixation of target distances, requiring that both eyes point in contrary directions. These movements are elicited by retinal disparity (when a fixation target is not on both foveae) and retinal blur (when a target is not in focus). Therefore, these movements are closely related to the lens accommodation and pupillary reflexes [16].

3. Oculomotor disturbances in neurodegenerative diseases

3.1 Parkinson’s disease and other parkinsonian disorders

3.1.1 Parkinson’s disease

Parkinson’s disease is a progressive disorder pathologically defined by the degeneration of the dopaminergic neurons in the substantia nigra and formation
of α-synuclein-containing Lewy bodies in the residual dopaminergic neurons. Consequently, the clinical picture is characterized by progressive motor symptoms that include bradykinesia, muscular rigidity, rest tremor, as well as postural and gait impairment. The disease is also associated with many non-motor symptoms, some of which precede the motor dysfunction by more than a decade [17]. Global prevalence of PD ranges between 100 and 200 cases per 100,000 inhabitants, with an annual incidence around 15 cases per 100,000 [18]. Although the etiology of PD is commonly unknown, monogenic causes can be considered in 5–10% of the cases [19].

Findings about oculomotor function in PD are certainly inconsistent due to the reduced number of patients included in the majority of the studies and the heterogeneity of the disease phenotype [7]. Nevertheless, saccadic hypometria is recognized as the most striking oculomotor feature in PD patients, which can be documented both at bedside and by electrophysiological approaches even early in the disease course. As a result of the saccade hypometria, PD patients frequently require multistep sequences to reach the target [20]. This behavior is more pronounced during memory-guided saccades, and it is considered as a disease biomarker [21, 22]. The marked saccade hypometria in PD can be explained by the neurodegenerative changes in the basal ganglia causing the decrease of pre-oculomotor drive through the substantia nigra to the superior colliculus [21]. Alongside the saccade hypometria, PD patients also show abnormally prolonged latency of voluntary saccades such as the memory-guided saccades and the antisaccades; nevertheless, the latency of externally triggered saccades to visual targets is normal [23]. Distinct to the saccade hypometria, the deficits in the saccade initiation are detectable later in the disease course and are closely related with the cognitive impairments and the involvement of non-dopaminergic pathways such as the frontal and parietal eye fields, the premotor cortex, and the lateral prefrontal cortex [24].

The delayed prosaccade and the antisaccade tasks reveal an impaired inhibition of saccades as evidence of deficit of automatic response inhibition. PD patients show increased timing error rates in the delayed prosaccade paradigm, which are closely associated with abnormal neuropsychological performance, whereas antisaccade paradigm reveals higher directional error rates [25]. Antisaccade errors can be detected early in the disease course [26]. Beyond saccadic impairments, PD patients show slight alterations in other eye movements, such as reduced gain of the smooth pursuit movements [27] and slow and hypometric divergence movements, but normal convergence movements [28].

3.1.2 Other parkinsonian disorders

Oculomotor findings of patients suffering from other parkinsonian disorders are varied and usually distinctive to the PD. In cases with multisystem atrophy with predominant Parkinsonism (MSA-P), the clinical assessments of oculomotor function usually reveal increased square wave jerks, saccade hypometria, as well as abnormal smooth pursuit and vestibulo-ocular reflex [29, 30]. Less common oculomotor features in MSA-P include downbeat nystagmus, head-shaking nystagmus, and mild vertical supranuclear gaze palsy [29, 31].

In the progressive supranuclear palsy with Parkinsonism (PSP-P), the most important oculomotor feature is the slowing of vertical saccades, which progresses to supranuclear gaze palsy in the 70% of the cases but appear lately in the disease course than in the classic PSP [32]. In addition, these patients show reduced gains of the smooth pursuit movements and saccadic eye movements at similar extent that in classic PSP [27].
3.2 Alzheimer's disease and other dementias

3.2.1 Alzheimer's disease

Alzheimer's disease (AD) is the most common neurodegenerative disorder worldwide with a global prevalence above 20 million of affected people, which is estimated to grow notably in the next decades. The histopathological hallmark of the disease is the deposition of insoluble protein aggregates such as amyloid-β (Aβ) plaques and neurofibrillary tangles of tau in the brain, causing a significant brain atrophy and subsequent cognitive features such as memory disturbances, executive dysfunction, difficulties with language, and other cognitive skills that affect a person's ability to perform every day [33]. Similar to PD, the etiology of Alzheimer's disease (AD) is not fully understood, but several environmental and genetic factors are assumed to contribute to the disease etiopathogenesis [34].

Oculomotor testing in Alzheimer's disease reveals a varied group of eye movement abnormalities, but no specific oculomotor feature is distinguished. Among oculomotor features of AD patients, the saccadic intrusions are one of the most common [35, 36]. These unwanted microsaccades are mainly oblique and can be detected even in subjects with mild cognitive impairment which identify this oculomotor feature as a potential biomarker of Alzheimer's disease at early stages [37]. These microsaccades are more frequent in those patients with higher dementia scores [38], which support the notion that gaze-fixation instability in AD results from the involvement of cognitive processes such as the attention and working memory. Nevertheless, the impairment of the saccade pathways could also explain the high occurrence of saccadic intrusions, mainly at later disease stages [39].

 Reflexive and voluntary saccades of AD patients are usually characterized by prolonged latencies, reduced velocity, and hypometria. Antisaccadic paradigm reveals increased directional error rate alongside with the reduction of the error correction, which are closely associated with the severity of dementia [40]. Both prosaccadic and antisaccadic alterations in AD are proposed to result from impaired inhibitory control and attentional failures, as well as from the later involvement of saccadic circuitry at brainstem [39]. In addition, AD patients show increased latency to initiate smooth pursuit movements, with decreased gain velocity and increased catch-up (compensatory) saccades. Similar to the saccadic intrusions and antisaccadic deficits, the rate of compensatory saccades during the smooth pursuit is narrowly related with the dementia severity [40–42].

3.2.2 Other dementias

In the posterior cortical atrophy (PCA), an atypical variant of AD, the most frequent oculomotor abnormalities include increased saccade latency and decreased saccade amplitude, but the saccade velocity is normal. Also, the PCA patients show increased time to saccadic target fixation, even higher than subjects with typical AD. Moreover, these patients show large saccadic intrusions whose frequency is correlated with generalized reductions in cortical thickness. Smooth pursuit gain is slightly reduced in these patients [43, 44]. Moreover, individuals with frontotemporal dementia (FTD) show increased reflexive saccade latency and higher rates of antisaccadic errors, but the error correction abilities are preserved. In addition, the smooth pursuit movements are characterized by the reduction of gains and accelerations [40, 45, 46].
3.3 Huntington’s disease

Huntington’s disease is a neurodegenerative disorder caused by the abnormal expansion of cytosine-adenine-guanine (CAG) trinucleotide repeats in the huntingtin gene on chromosome 4, encoding the huntingtin protein. The mutation results in an excessively long polyglutamine stretch near the N-terminus of this protein, which identify this disorder as a polyglutamine disease. Mutant HTT affects some cellular processes, including protein-protein interaction, protein clearance, mitochondrial function, axonal trafficking, gene transcription, posttranslational modification, and others that ultimately cause the loss of striatal neurons [47].

Clinically, the disease is characterized by a progressive motor, cognitive, and psychiatric disturbance. The motor phenotype includes chorea as cardinal feature, as well as dystonia and Parkinsonism, whereas the cognitive dysfunction comprises dysexecutive signs, as well as memory and attentional dysfunction. Psychiatric features are usually depression, anxiety, apathy, obsessive-compulsive behaviors, and others. Similar to other polyglutamine disorders, the age at onset of HD is highly influenced by the CAG repeat length, but other genetic and environmental modifying factors are proposed to also control the age at onset variability [47, 48].

Oculomotor abnormalities of patients with HD include saccade slowing and deficits in the initiation and suppression of these movements. The reduction of saccade velocity appears in around 60% of patients and is commonly observed in the vertical plane, but in those cases in advanced disease stages, the saccade slowing reaches also the horizontal movements [49, 50]. Saccade latencies are significantly prolonged and show a marked variability, which is more pronounced in patients showing higher disease severity. Studies using the antisaccadic paradigm have revealed and increased rate of directional errors, which are also closely correlated with the severity of the disease. Moreover, increases of latency variability and timing errors are observed in the memory-guided saccade task. The deficits of the suppression and initiation of the saccades can be explained by the neurodegenerative changes in the frontal cortex and in the basal ganglia [51, 52]. So, a recent imaging research revealed a close association between the voluntary saccade inhibition deficits and the white-matter corticobasal atrophy in patients [53].

Several authors have evaluated saccadic eye movements in asymptomatic carriers of the HD mutation. These studies have found a significant delay in the initiation of voluntary eye movements, increase in the variability of saccadic latency, and increase in the rate of antisaccadic errors [54–56]. A longitudinal follow-up of these alterations demonstrated their usefulness as preclinical markers due to the high replicability and consistency of these measures [22]. Imaging studies in asymptomatic carriers of HD have shown a significant correspondence between alterations in saccadic latency and the decrease in the number of fronto-striatal fibers that project into the caudate nucleus and the atrophy of gray matter in cortical structures, which deepens in the pathophysiology of saccadic alterations in this disease [57, 58]. A recent paper demonstrated that the horizontal ocular pursuit item of the Unified Huntington’s Disease Rating Scale is useful for detecting differences between premanifest individuals and controls [59].

3.4 Amyotrophic lateral sclerosis

Amyotrophic lateral sclerosis (ALS) is the most common and devastating age-related motor neuron disease, characterized by a progressive loss of upper and lower motoneurons, causing paralysis and death in approximately 3 years. The pathological hallmark of ALS is the presence of abundant cytoplasmic inclusions containing ubiquitin and TDP-43, a RNA-binding protein. The clinical picture
comprises progressive muscle weakness alongside hyperreflexia and spasticity associated with fibrillations and fasciculations [60]. The disease has a global prevalence around five cases per 100,000 inhabitants. Most of ALS cases are sporadic, and only the 5% of patients are familial, with at least 12 genes implicated, such as the superoxide dismutase 1 (SOD1), trans-activate response DNA-binding protein (TARDBP), C9ORF72, FUS, and the ataxin-2 genes [61, 62].

Some evidences have demonstrated the involvement of the oculomotor system in ALS, leading a broad range of eye movement deficits affecting the saccades and the smooth pursuit movements [63–66]. The most prominent and early oculomotor alterations of ALS patients are related with abnormal executive oculomotor control as evidence of frontal lobe involvement. They primarily includes the increase of error rates in anti-saccades and delayed saccade paradigms as well as reduced voluntary gaze shift and increased number of saccadic intrusions. In general, these oculomotor alterations are correlated with the severity of the disease and the neurocognitive measures. In a following stage of oculomotor abnormalities, some ALS cases can show slow saccades, saccade hypometria, and interrupted smooth pursuit, as evidences of the involvement of the brainstem and pre-cerebellar/pontine circuits [67].

3.5 Hereditary ataxias

Hereditary ataxias consist in a heterogeneous group of genetic disorders phenotypically characterized by gait ataxia, limb incoordination, dysmetria, dysarthria, oculomotor disturbances, and other motor and non-motor features. These disorders are associated with atrophy of the cerebellum, which can be accompanied with the degeneration of other regions in the central and peripheral nervous system in various genetic subtypes [68].

Hereditary ataxias are classified into four main groups regarding their inheritance patterns: autosomal dominant (also referred as spinocerebellar ataxias), autosomal recessive, X-linked, and mitochondrial ataxias [68, 69]. Till now, 46 subtypes of spinocerebellar ataxias have been identified, which imply at least 37 distinct genes [70]. The most common subtypes are caused by polyglutamine (polyQ)-coding CAG repeat expansions (SCA1,2,3,6,7, DRPLA) [71]. Regarding the recessive ataxias, nearly 100 genes have been identified, with the highest prevalence for the Friedreich's ataxia (FRDA), caused by GAA repeat expansions or point mutations in the frataxin (FXN) [68, 72]. Global prevalence of hereditary ataxias is estimated around three cases per 100,000 inhabitants, but there are large regional variations of prevalence due to founder effects of some genes [73].

Oculomotor disturbances of SCA patients are varied and result from the cerebellar and/or brainstem involvement. The former abnormalities are the most common and include the presence of pathological nystagmus, abnormal smooth pursuit, and saccadic dysmetria, whereas the impaired VOR, saccadic slowing, and ophthalmoplegia are related with pontine degeneration. Nevertheless, the notable overlapping of oculomotor features between SCA subtypes implies the requirement of other clinical criteria or the genetic testing for sensitively discriminating among these diseases [74–78] (Figure 2).

In the case of SCA2, an early and severe saccadic slowing is observed even more than a decade before the ataxia onset [79], which identifies it as important preclinical biomarker of the disease. Interestingly, the SCA2 saccade slowing is tightly influenced by the expanded CAG repeats in the ATXN2 gene [80] and shows a significant familiar aggregation which leads to the suitability of this disease feature as endophenotype marker [81], with potential usefulness for the search of modifier genes and neurobiological underpinnings of the disease and as outcome measure.
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Figure 2. Cerebellar and/or brainstem origin of oculomotor features in SCAs.

In future neuroprotective clinical trials. Moreover, the saccade slowing in SCA2 progresses significantly along time providing novel insight into the cumulative polyglutamine neurotoxicity and supporting the usefulness of saccade peak velocity as a sensitive biomarker during the natural history of the disease [82]. Saccade pathology in SCA2 is also characterized by abnormal prolongation of reflexive and voluntary latencies and increases of the antisaccade error rate. The later feature is also detected in prodromal stage and is significantly correlated with the mutation size [83–85].

The main eye movement abnormalities of SCA1 patients include saccadic dysmetria, gaze-evoked nystagmus, and depressed smooth pursuit [86]. Saccadic hypermetria is observed in majority of cases, appears at an early stage of the disease, and progresses quickly [75, 76, 87]. SCA3 is characterized by a higher frequency of gaze-evoked and rebound nystagmus [88], in addition to decreased smooth pursuit gain and saccadic dysmetria. These patients also show decreased VOR gain, which correlated with the CAG repeats, suggesting the pathologic involvement of the vestibular nuclei in the lateral brainstem [74–76]. Divergence insufficiency and strabismus are also common oculomotor features of these patients [89, 90].

In SCA6, a higher frequency of spontaneous downbeat nystagmus and square wave jerks is detected [76, 91, 92]. The square wave jerks together with subtle abnormalities of saccades and smooth pursuit movements can be detected even before the disease onset [93]. The major saccadic alteration in SCA7 is the slowing of saccades, together with saccadic dysmetria [94, 95]. These alterations may precede cerebellar and retinal manifestations by some years [96]. Patients with SCA17 show hypometric saccades which are increased with disease duration but neither with ataxia score nor CAG repeats number [97].

Eye movement disturbances are frequent in FRDA. The most prominent abnormalities consist in fixation instability such as multiple square wave jerks and ocular flutter, which are also complemented by abnormal smooth pursuit, saccadic dysmetria, prolongation of saccade latency, gaze-evoked nystagmus, and impaired VOR. Interestingly, the prolongation of saccade latency and the square wave jerks are significantly correlated with the disease severity and age at disease onset, respectively [98, 99]. Moreover, antisaccades and memory-guided saccades are also abnormal in these patients as evidence of the disruption of the higher-order processes controlling the saccade movements [100].
4. Concluding remarks

Eye movement abnormalities are among the most common phenotypic manifestations of patients with neurodegenerative diseases. The prominent features include the saccadic abnormalities, fixation instability, and abnormal smooth pursuit. Thus, the examination of eye movements is a very useful, but not determinant, approach for the differential diagnosis of these disorders. For example, the increased square wave jerks and the slowing of vertical saccades may be useful features for the clinicians in order to distinguish between the MSA-P and the PSP-P from the idiopathic Parkinson's disease, respectively. In addition, the early and severe saccadic slowing with rare pathological nystagmus distinguishes SCA2 from other autosomal dominant ataxias, whereas the marked abnormalities of smooth pursuit, VOR and OKR, in association with pathological nystagmus and rare saccadic slowing may help to define a SCA6 phenotype. Nonetheless, the notable overlapping of oculomotor features between neurodegenerative disorders suggests the necessity of other diagnostic criteria for sensitively discriminating among diseases with similar symptomatology.

Besides, the assessment of oculomotor function in neurodegenerative disorders leads to the identification of disease biomarkers, which acquire key values in the clinical and research practice of neurodegenerations. Many eye movement markers of neurodegenerative disorders allow to assess the disease stage and disease progression, because their changes over time are significantly linked with clinical outcome of syndrome severity, and interestingly some oculomotor disturbances precede the clinical diagnosis of the disease, which identify them as useful preclinical markers to detect the early stages of the neurodegenerative process, to evaluate the genetic susceptibility of the asymptomatic relatives, and to identify individuals for enrolment in early intervention trials.

Moreover, the study of eye movements in neurodegenerative diseases offers valuable advantages to assess the cognitive functioning in these conditions, mainly those measures that reflect the high-order processes underlying the oculomotor functions such as the antisaccade and memory-guided saccade task outcomes, the saccade latency, and others.

In conclusion, although by decades the oculomotor system has been widely studied in neurodegenerative diseases, further efforts are warranted to study their involvement in other—less common—disorders, to understand the physiopathological mechanisms underlying oculomotor disturbances and to certify the role of oculomotor features as sensitive outcome measures in further neuroprotective trials.

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Conflict of interest

Authors declared no conflict of interest.

Appendices and nomenclature

AD Alzheimer’s disease
ALS amyotrophic lateral sclerosis
CAG  cytosine-adenine-guanine trinucleotide
DRPLA  dentatorubral-pallidoluysian atrophy
FRDA  Friedreich’s ataxia
FTD  frontotemporal dementia
FXN  frataxin
HD  Huntington’s disease
MSA-P  multisystem atrophy with predominant Parkinsonism
OKR  optokinetic reflex
PCA  posterior cortical atrophy
PD  Parkinson’s disease
PSP-P  progressive supranuclear palsy with Parkinsonism
SCA  spinocerebellar ataxia
VOR  vestibulo-ocular reflex

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