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Standardized Assessment of Hereditary Ataxia Patients in Clinical Studies

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Abstract: Background: Hereditary ataxias are a heterogeneous group of degenerative diseases of the cerebellum, brainstem, and spinal cord. They may present with isolated ataxia or with additional symptoms going beyond cerebellar deficits. There are an increasing number of clinical studies with the goal to define the natural history of these disorders, develop biomarkers, and investigate therapeutic interventions. Especially, early and preclinical disease stages are currently of particular interest.

Methods and Results: Evidence-based, we review standards for sampling and storage of biomaterials, clinical and neuropsychological assessment, as well as neurophysiology and neuroimaging and recommendations for standardized assessment of ataxia patients in multicenter studies.

Conclusions: DNA, RNA, serum, and, if possible, cerebrospinal fluid samples should be processed following established standards. Clinical assessment in ataxia studies must include use of a validated clinical ataxia scale. There are several validated clinical ataxia scales available. There are no instruments that were specifically designed for assessing neuropsychological and psychiatric symptoms in ataxia disorders. We provide a list of tests that may prove valuable. Quantitative performance tests have the potential to supplement clinical scales. They provide additional objective and quantitative information. Posturography and quantitative movement analysis—despite valid approaches—require standardization before implemented in multicenter studies. Standardization of neurophysiological tools, as required for multicenter interventional trials, is still lacking. Future multicenter neuroimaging studies in ataxias should implement quality assurance measures as defined by the ADNI or other consortia. MRI protocols should allow morphometric analyses.

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Hereditary ataxias are a clinically and genetically heterogeneous group of degenerative diseases of the cerebellum, brainstem, and spinal cord. Their prominent clinical features are progressive ataxia comprising impairment of gait, balance, and limb coordination, as well as disturbances of speech swallowing and oculomotor control. Hereditary ataxias may present as purer forms with more isolated ataxia or as complicated forms with additional neurological symptoms.

Hereditary ataxias are subdivided into the autosomal-recessive ataxias with Friedreich’s ataxia (FRDA) as the most prevalent entity—except East Asia—, the autosomal-dominant ataxias designated as spinocerebellar ataxias (SCAs), and X-linked ataxias with fragile X-associated tremor ataxia syndrome (FXTAS) as the most common form. Currently, more than 50 recessive ataxias (http://neuromuscular.wustl.edu/ataxia/recatax.html), and approximately 40 SCAs, have been genetically delineated (http://neuromuscular.wustl.edu/ataxia/domatax.html). Additionally, ataxias can be caused by mitochondrial DNA mutations as well. A subgroup of potentially hereditary ataxias are the sporadic adult-onset degenerative disorders with no identified etiology share many clinical features with the hereditary ataxias, but the causes of these ataxias are unknown. An overview of hereditary ataxias and causative gene mutations can be found online (http://neuromuscular.wustl.edu/).

With the genetic classification of hereditary ataxias, an increasing number of clinical studies aim to define the natural history of these disorders, develop biomarkers, and investigate therapeutic interventions. Early and preclinical disease stages are currently attracting particular interest. For such studies, generally accepted standards for sampling and storage of biomaterials, clinical and neuropsychological assessment, quantitative performance tests, as well as neuroimaging are highly needed. In this article, we review the available evidence and give recommendations for standardized assessment of ataxia patients in multicenter studies.

**Methods**

Searches were conducted in the MEDLINE, EFNS Guidelines, and EMQN databases. The review period was from January 1990 to 2014. Original articles, review articles, and guideline recommendations were reviewed. Search terms were hereditary ataxia, cerebellar ataxia, autosomal-recessive ataxia, Friedreich’s ataxia, spinocerebellar ataxia, autosomal-dominant cerebellar ataxia, and Machado-Joseph disease and were used in various combinations with terms like clinical assessment scales, functional score, MRI, PET, gait analysis, genotyping, biomarker, biomaterial, morpho*, and neuropsychos*. In the next step, the identified assessments tools, such as “Scale for the Assessment and rating of Ataxia” (“SARA”), “International Cooperative Ataxia Rating Scale” (“ICARS”), “Composite Cerebellar Functional Severity Score” (“CCFS”), were used as search terms in combination with disease terms. Reference lists of publications of interest were screened for other relevant studies. B.P. performed the literature search and wrote the first draft. T.K. reviewed all versions. A preliminary version of the manuscript was then circulated among all authors. This version was discussed and consented by conference calls, before the final version was drafted and circulated.

**Ataxia Research Consortia and Study Groups**

In recent years, a number of large national and international research consortia have received public funding to perform clinical research in hereditary ataxias. SPATAX is a consortium of European and North African investigators under French leadership that focuses on clinical and genetic studies in recessive ataxias and hereditary spastic paraplegias (HSPs) (http://spatax.wordpress.com/). The EUROSCA Clinical Network that was funded by the European Union (EU) between 2004 and 2008 developed clinical assessment tools for ataxia, established an electronic database, and initiated a natural history study of dominantly inherited spinocerebellar ataxias (SCAs) (http://www.euroasca.org). To continue clinical research activities, investigators of the EUROSCA Clinical Network in 2008 founded the Ataxia Study Group (ASG) (http://www.ataxia-study-group.net/html/about). Currently, there are two large EU-funded consortia involved in clinical ataxia and HSP research. Major initiatives to study FRDA are performed by EFACTS (http://www.e-facts.eu) in Europe and the Cooperative Clinical Research Network (CCHR-N) in the United States, Canada, Brazil, and Australia. NeurOmics is using -omics methods to find disease causing genes, improve diagnostics, and develop novel therapies for 10 rare neurodegenerative and neuromuscular disease groups, including ataxia (http://rd-neuromics.eu/). The Cerebellar Ataxia Group (CAG), which has evolved into the Clinical Research Consortium for Studies of Cerebellar Ataxias (CRC-SCA), comprises North American ataxia investigators. Clinical studies initiated by the CRC-SCA received funding from the National Institutes of Health (NIH). The Japanese Ataxia Study Group receives funding from the Japanese government. It is running a large patient registry.

**Common Data Elements**

Starting in 2006, the National Institute of Neurological Disorders and Stroke (NINDS) and other federal agencies and international organizations have the common mission of developing data standards for clinical research. On their webpage (http://www.commondataelements.ninds.nih.gov/ProjReview.aspx#tab =Introduction), common as well as disease-specific data elements are published as a resource to build study specific common data element (CDE) data collection sheets. General CDEs include medical history data; scores on neurological assessments; demographic information (e.g., date of birth/age, race, and ethnicity); and details about medications used by participants throughout a study. For FRDA, disease-specific recommendations of tools and CDEs are given by NINDS (http://www.commondataelements.ninds.nih.gov/Doc/)
**Biomaterial Sampling and Storage**

Availability of biomaterials is an essential prerequisite for progress in the understanding of neurodegenerative diseases. Genetic testing is performed for genetic classification of study participants. On http://www.scabase.eu, information about ataxia genetics regarding current best practices protocols for molecular genetic testing, lists of repeat sequences, primers, and normal and pathogenic repeat sizes on the main loci are maintained.

Orphanet and—in Europe—EuroGenTest are data sources for genetic tests and relevant laboratories.

DNA is needed for the proper classification of study subjects with known diseases and for identification of novel disease genes and genetic modifiers.

Fibroblast or peripheral blood mononuclear cell (PBMC) samples are valuable sources for various experiments, in particular, generation of induced pluripotent stem cells (iPSC). Because sampling of PBMCs is less invasive and easier than that of fibroblasts, PBMCs are the currently recommended source of iPSCs. Given the enormous potential of research with iPSCs, routine sampling of PBMCs in clinical studies, if feasible, is highly recommended.

There are several systematic efforts for brain collections under standardized protocol. Here as well, effort and money are limiting factors. Additionally, the trust of potential donors and their families plays an important role in the success of building up brain collections.

Biomaterials, such as blood samples, urine, and cerebrospinal fluid (CSF), are required for the development of fluid biomarkers, which—other than in common neurodegenerative diseases, such as Alzheimer’s disease (AD)—are currently not available in ataxia.

Usefulness of biomaterials critically depends on the clinical information on the subjects from whom the materials are derived. Biomaterial collections that are not linked to reliable clinical data are therefore of limited use. Another important aspect that is critical for the usefulness of biomaterials is the biological quality of the samples. All procedures, including acquisition, handling, and preprocessing before final storage, storage, and retrieval, strongly influence the composition and stability of the samples. These steps should be therefore performed following standard operation procedures, as are generally recommended for biomaterial banking. Furthermore, a documentation of the biomaterial quality is necessary and should be performed as detailed and internationally comprehensive as possible. An example of such documentation is the standard preanalytical (SPREC) code (http://www.isber.org/?page=SPREC). For further information see supplemental material.

Overall, DNA, RNA, serum, and—whenever possible—CSF and PBMC samples comprise a set of biomaterials that should be taken.

**Clinical Scales**

Clinical scales are the most important component of assessment in clinical studies. They serve as outcome measures in clinical interventional trials and are therefore key instruments that have a strong impact on future research and patient care. Clinical ataxia scales have been recently reviewed in detail. We will describe here the most widely used scales and comment on their properties with respect to use in multicenter and longitudinal studies. Their basic psychometric properties are summarized in Table 1.

**ICARS**

The ICARS was the first ataxia scale and is widely used in observational studies as well as in interventional trials. It consists of 19 items grouped into four subscales that contribute to a total score of 100 points. Subdivisions of different ataxia components are postural and gait disturbance, limb ataxia, dysarthria, and oculomotor disorders.

ICARS has been evaluated in six published studies in MSA, SCA, and FRDA patients. Standard psychometric variables, such as total Crohnhach’s α, inter-rater reliability, and test-retest reliability, were reported to be good to very good. However, several investigators observed interdependencies between several items. Furthermore, principal component analysis showed that the ICARS score is determined by four different factors that do not coincide with the subscales. The ICARS also lacked linearity and showed floor and ceiling effects. However, ICARS shows good sensitivity to change.

There are two modified versions of the ICARS, the Brief Ataxia Rating Scale (BARS) for use by movement disorder specialists and general neurologists, and the modified ICARS (MICARS), in which further items were added. Both scales lack further validation. Recently, the effect of age on ICARS, BARS, and SARA scores has been assessed in healthy children.

**SARA**

The SARA is a clinical scale based on a semiquantitative assessment of cerebellar ataxia. It has eight unequally weighted items evaluating to gait, stance, sitting, speech, finger-chase test, nose-finger test, fast alternating movements, and heelf-shin test. Oculomotor functions are not included in the SARA.

The SARA was initially validated in two studies of 167 and 119 SCA patients. Subsequent evaluation studies were done in mixed populations of ataxia patients and in FRDA. The SARA has been translated into Iberia-Brazilian, Japanese, and Chinese versions, and the translated versions were also validated. Recently, the SARA was also tested in healthy children of various age groups.

Overall, psychometric properties were shown to be as good as for the ICARS. In contrast to the ICARS, the SARA behaves linearly without ceiling or floor effects. In FRDA
patients, sensitivity to change of SARA was greater than that of the Friedreich Ataxia Rating Scale (FARS) and ICARS. In addition, clinically relevant changes were defined, and it was demonstrated that SARA is a major determinant of health-related quality of life in SCA patients. From the practical point of view, SARA has less repetitive items, is less time-consuming to administer, and easier to use than ICARS.

FARS

The FARS comprises neurological signs that specifically reflect FRDA symptoms assessed in three functional domains: (1) functional staging (overall mobility); (2) activity of daily living (ADL); (3) neurological assessments (bulbar, upper and lower limbs, peripheral nerve, upright stability, and gait functions). The FARS has been validated in two studies of 14 and 155 FRDA patients, respectively. Regner et al. discussed that for longitudinal measurement in unselected cohorts, a 2-year period may be the minimum period to gain reasonable power.

Psychometric properties were good and comparable to ICARS and SARA. However, ceiling effects were observed. FARS meets essential criteria for validity for the measurement of disease progression. As a measure of sensitivity, FARS has greater effect size in FARDA patients than ICARS.

Neurological Examination Score for Spinocerebellar Ataxia

The Neurological Examination Score for Spinocerebellar Ataxia (NESSCA) was developed specifically for SCA3 patients based on a standardized neurological examination. Thus, the 18 items selected comprise a global inventory of general neurological signs and of signs specific for SCA3. It has been validated in a cross-sectional study of 99 SCA3 patients. Inter-rater reliability (20 patients) was high. Scores correlated with SARA scores. Sensitivity to change was not investigated.

Scales for the Assessment and Rating of Nonataxia Features

Inventory of Non Ataxia Signs

The Inventory of Non Ataxia Signs (INAS) comprises a list of various nonataxia signs that often occur in ataxia patients and of cerebellar oculomotor signs that are not considered in the SARA. INAS has 30 items related to neurological signs, such as spasticity, and to reported abnormalities, such as dysphagia. The items are grouped into functional categories, reflecting 16 nonataxia and cerebellar oculomotor signs. As a simple quantitative measure, the INAS count reflects the presence or absence of...
nonataxia signs, when at least one item related to a nonataxia sign is present.

The INAS performed very well in terms of inter-rater and test-retest reliability. There were mild floor and no ceiling effects. Responsiveness and sensitivity, however, were less satisfactory. Another limitation is that the INAS lacks precise definitions and instructions for investigators, which may cause variability in judgments of symptom severity. However, the INAS is an inventory—not a rating—scale, so the score depends on the presence, not on the severity of symptoms and signs. The INAS is easy to use, only requiring tests or procedures that are routinely performed in the neurological examination. Overall, it has proved to be a useful supplement to the SARA, but it is not an appropriate primary outcome measure in interventional trials in ataxia.

Other Scales

In some types of ataxias, specific symptoms are frequently encountered. Examples are spasticity in autosomal recessive spastic ataxia of Charlevoix-Saguenay (ARSACS) and autonomic dysfunction in SCA3. In these instances, it is useful to apply other scales that assess specific symptoms more accurately than the INAS. In SCA3 patients, the Unified Multiple System Atrophy Rating Scale (UMSARS) has been validated. In ARSACS and other spastic ataxias, the Spastic Paraplegia Rating Scale (SPRS), which has been developed for hereditary spastic paraplegia patients, can be applied. However, further validation is desirable.

Patient-Reported Measures

Outcome measures that are patient reported may be valuable for complementing physician-based outcomes. Examples for generic health status measurement tools or disease-specific tools are Short Form Health Survey version 2 (SF-36 V2) or the Friedreich Ataxia Impact Scale (FAIS). The SF-36 is a general measurement of quality of life. Quality of life is an important secondary endpoint in clinical interventional trials. The SF-36 does not address many ataxia-related issues, though. The FAIS contains 126 items covering eight functional areas. It was developed and validated in over 400 FRDA patients grouped into different levels of disease severity. In a recent longitudinal study, the FAIS was found to provide valuable insight into the perspective of individuals with FRDA on their health status and morbidity. However, responsiveness to change appears limited and its use in intervention studies is questionable.

The ADL part of FARS is another quality-of-life measure, which is used frequently. As an outcome in the EFACSTS study, it showed effect sizes comparable to those of the SARA.

Assessment of Neuropsychological and Psychiatric Symptoms

Because the cerebellum has a prominent role in motor control, ataxia patients mainly present with motor symptoms. The cerebellum plays a role also in nonmotor functions. Neuropsychological and psychiatric symptoms of variable severity are encountered in a number of ataxias. The reasons for this are 3-fold: (1) Because of cerebellar connections to nonmotor cortical areas, cerebellar disease can be associated with neuropsychological symptoms, mainly disturbances of executive function; (2) in many ataxias, neurodegeneration extends beyond the sites that result in ataxia by involving forebrain structures. This results in intellectual impact and cognitive decline; (3) furthermore, as with many other chronic brain diseases, ataxia can be associated with depression.

There are no instruments that were specifically designed for assessment of neuropsychological and psychiatric symptoms in ataxia. Thus, in studies of ataxia patients, various instruments are used that have been previously developed and validated in other populations. A comprehensive overview of such studies is given by Almeida-Silva et al.

Although neuropsychological and psychiatric abnormalities have been investigated in numerous studies, there are only a few multicenter and longitudinal studies. A major problem in the use of timed neuropsychological tests in ataxia patients is their motor slowness, which is a confounding factor in assessing cognitive function. This should be taken into account when selecting specific tests. Neuropsychological tests recommended in order to minimize motor skills’ interference with special emphasis on covering executive, visuospatial, and verbal domains are given in Table 2.

Quantitative Performance Tests

For ataxia, a number of tests are in use that measure performance in specific coordinative tasks in a quantitative way. These tests yield metric, continuous data that facilitate analysis. In addition, inter-rater reliability is usually higher than in clinical scales. Assessments range from simple time measurements, for example, the time needed to complete a 25-foot (ft) walk, syllable repetition rate, and pegboard tests, to complex assessments that require specific equipment, for example, stride length variability measured with the GAITRite system, acoustic speech analysis systems, and systems to analyse the kinematics and forces of reaching and grasping movements. Each of the obtained measures reflects only performance in the respective task. But the results of various tests can be combined to composite measures that are thought to provide a more comprehensive account of disease severity.

Ataxia Functional Composite

The Ataxia Functional Composite (AFCS) was specifically designed for FRDA. It is derived from the Multiple Sclerosis Functional Composite (MSFC) and composed of a timed 25-ft walk, the 9-hole pegboard test (9HPT), and low-contrast visual acuity (LCVA). It has been validated in 20 hereditary ataxia patients, showing a strong correlation to ICARS. Test results are given as calculated Z scores.
The SCA Functional Index (SCAFI) is very similar to the AFCS, but the LCVA was replaced by a speech test (PATA repetition rate). The walking test is designated as an 8-m walk, but is essentially identical to the 25-ft walk of the AFCS. The SCAFI score is derived from the arithmetic mean of the Z scores of the three tests. The SCAFI reached the criterion for retest reliability and showed, in general, favorable measurement precision. However, in longitudinal tests, SCAFI deteriorates within the stable group of patients during the follow-up period. The clinical relevance of the longitudinal SCAFI changes seeks further investigation.

The Composite Cerebellar Functional Severity Score (CCFS) is a performance-based scale that combines two tests of upper limb function, the 9HPT and the click test. Measurements are adjusted for age and can be used for both dominant and recessive ataxias and are adapted for children. A version of the CCFS includes a handwriting test (CCFSw). However, patients with a SARA score of 30 or above may not be able to perform the CCFS(w).

Posturography is a technique that quantifies postural control in upright stance in either static or dynamic conditions. It is widely used for diagnosis of balance disorders. Posturography using gold-standard laboratory (Codamotion 3D motion analysis system with Visual 3D offline processing software) has been used to quantitatively investigate balance (static conditions) in adults with SCA6. Measures of overall sway speeds as well as pitch and roll directional components of sway have been used to comprehensively describe overall and directional components of instability. Quantitative reports of mean sway speeds correlate with same-day measures of disease severity (using the SARA), where an increase in SARA is associated with linear increases in sway speed. Posturography was additionally used to explore levels of axial instability. Laboratory investigations of sensory mechanisms of balance control (dynamic conditions) have incorporated the use of posturography in order to determine sensorimotor abnormalities in adults with SCA6. Posturography requires careful control of sensory conditions to ensure reliability of highly sensitive measures, but, when achieved, is feasible as a measure to either track disease progression or to evaluate the treatment effect of therapies in both clinical and home settings. In addition to standardization of stance width, foot splay angle, and underfoot surface, standardization of the visual environment is of considerable importance during testing given that hypermetric balance responses and increased overall measures of instability are notable when patients observe moving scenery in their visual environment. Validated portable measures of posturography (body sway speeds and directional velocities) include the XSens MTx monitor, but posturographic methods, incorporating other accelerometry-containing devices, are similarly possible.

### TABLE 2 Recommended neuropsychological tests for assessing cognitive functions in ataxia patients

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<th>Function</th>
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*Considering to minimizing the need for manual response and ataxia as a confounding factor.
Quantitative Movement Analysis

Computerized gait analyses systems offer an objective and quantitative measure of gait as well as directed movements. It has been used in ataxia as an outcome measure in several nonpharmacological interventional studies. A spatiotemporal gait analysis system based on infrared diode recording (Vicon) has been shown to pick-up changes in movement performance and scores correlate with clinical ataxia scales and with relevance for everyday life. GAITRite is a portable, instrumented electronics carpet system that allows recording of stride length and variability during short walking sequences. GAITRite was validated in study of 25 healthy adults and showed strong concurrent validity and high test-retest reliability. In ataxia, it was used in only a few single-center studies. One study showed that dynamic properties of locomotion in cerebellar ataxia patients were markedly impaired and that shifting walking speed away from preferred walking speed resulted in a loss of dynamic stability. The other study showed that discrete gait abnormalities precede manifest ataxia in SCA6 mutation carriers compared to manifest SCA6 patients. In FRDA, the sensitivity in measuring disease severity was investigated. There are no studies that validate GAITRite as an assessment instrument in multicenter studies of ataxia patients. However, the feasibility of quantitative gait analysis as an outcome measure for clinical trials in SCAs has been assessed in a small monocentric study on a total of 20 SCA1, 2, 3, and 6 patients at two time points, and data indicated that a high test-retest reliability was demonstrated. GAITRite has been shown to pick-up changes in movement performance and scores correlate with clinical ataxia scales and with relevance for everyday life. GAITRite is a portable, instrumented electronics carpet system that allows recording of stride length and variability during short walking sequences. GAITRite was validated in study of 25 healthy adults and showed strong concurrent validity and high test-retest reliability. In ataxia, it was used in only a few single-center studies. One study showed that dynamic properties of locomotion in cerebellar ataxia patients were markedly impaired and that shifting walking speed away from preferred walking speed resulted in a loss of dynamic stability. The other study showed that discrete gait abnormalities precede manifest ataxia in SCA6 mutation carriers compared to manifest SCA6 patients. In FRDA, the sensitivity in measuring disease severity was investigated. There are no studies that validate GAITRite as an assessment instrument in multicenter studies of ataxia patients. However, the feasibility of quantitative gait analysis as an outcome measure for clinical trials in SCAs has been assessed in a small monocentric study on a total of 20 SCA1, 2, 3, and 6 patients at two time points, and data indicated that a high test-retest reliability was demonstrated. Vicon would also allow quantifying the kinematics of reaching and grasping movements.

Easy-to-apply hand-held objects that allow measurement of grip forces, an objective parameter of manual performance, have been used in single-center trials of patients with mixed cerebellar pathology. They need to be validated in multicenter studies. The same is true for automatic speech analysis systems.

Electrophysiology (Nerve Conduction Studies, Evoked Potentials)

Peripheral nerves, dorsal columns, and pyramidal tracts are frequently affected in degenerative ataxias and can be quantitatively assessed by neurophysiological tools, such as nerve conduction studies, somatosensory evoked potentials, and motor evoked potentials. Neurophysiological tests show characteristic findings in different subtypes of SCA, being present even in preclinical stages of SCA2. In FRDA, nerve conduction and evoked potential studies demonstrate the sensory neuropathy, dorsal column degeneration, and pyramidal tract involvement that characterize the disease. Nerve conduction studies, by revealing the presence of a peripheral neuropathy and its type, provide an important support to the differential diagnosis of recessively inherited ataxias. However, multicenter approaches and longitudinal studies in ataxias are rare and face major challenges in standardization to assure reproducibility and intercenter comparability of quantitative neurophysiological measures, as required for multicenter interventional trials.

Eye Movement Tracking

Oculomotor disturbances are a common and often early sign in most hereditary ataxias. Specific oculomotor abnormalities, such as saccade slowing, are helpful in the diagnosis of hereditary ataxias. Presence or absence of oculomotor signs is recorded by the INAS together with nonataxia signs. Quantitative recording of eye movements has been performed in several monocentre studies. However, multicenter and longitudinal approaches face major challenges in standardization to assure reproducibility and intercenter comparability. However, given the potential of oculomotor recording for detection of early or even preclinical abnormalities, efforts to establish reliable multicenter recording techniques are strongly encouraged.

Neuroimaging

Neuroimaging allows assessment of the morphological and functional alterations of the brain and spinal cord in ataxia patients. Most neuroimaging studies in ataxia use MRI, but there are also molecular imaging studies with radiotracers. A recent consensus paper discussed application of these neuroimaging methods in ataxia.

Magnetic Resonance Imaging (MRI)

Numerous MRI studies have been performed in ataxia patients, most of which were cross-sectional and done in a single center. Multicenter longitudinal studies, on the other hand, face numerous methodological problems, which require rigorous measures of quality assurance and control. These issues have been carefully addressed by large consortia investigating patients with common neurodegenerative diseases, mainly AD, but the respective standards and recommendations have not yet been applied to ataxia studies. Structural T1-weighted MRI enables quantification of focal brain atrophy, in particular, of the cerebellum, but also of other parts of the brain. Images have been analyzed with region of interest–based volumetry and voxel-based morphometry. A multicenter study of SCA1, 3, and 6 patients showed brain tissue loss in the cerebellum and brainstem with a genotype–specific pattern. In a 2-year longitudinal study of this cohort, the rate of tissue loss in these structures and basal ganglia nuclei was significantly greater in patients compared to controls. Some of the volumetric measures had larger effect sizes than the SARA clinical scale. Similarly, there was progressive gray and white matter loss in a single-center longitudinal study of SCA2 patients.

Diffusion imaging is the method of choice to study white matter abnormalities, either with diffusion tensor imaging (DTI) or with more recent mathematical models, for example,
high-angular resolution diffusion imaging (HARDI), that describe the 3D displacement distribution of water diffusion. Although multicenter DTI studies are feasible and there is evidence for white matter involvement in ataxias, DTI has not yet been assessed in larger ataxia studies. Quantitative MRI techniques have also potential interest to study tissue properties, including myelin and iron content with relaxometry and susceptibility-weighted imaging. Similarly, functional MRI in ataxia has been limited to small, single-center, cross-sectional studies. The challenge for multicenter studies lies in the experimental paradigms used in most functional MRI studies. MR spectroscopy allows measurement of brain metabolites, its usefulness is limited by the large voxel size in which measurements are made and the difficulty to standardize acquisitions across sites. Again, multicenter studies are lacking. Such studies will also need to account and correct for different scanner types.

Molecular Imaging

In molecular imaging, radiotracers are used to assess brain metabolic activity or label neurotransmitter receptors. Binding and distribution of the radiotracers in the brain are detected with PET or single-photon emission computed tomography (SPECT). PET and SPECT have been used to study glucose metabolism, dopamine receptors, dopamine transporters, benzodiazepine receptors, and acetylcholine esterase activity in ataxia patients. Although these studies provided important insights into the neurobiology of ataxia disorders, standardization and validation of the molecular imaging methods as assessment tools in multicenter ataxia studies is lacking. However, for AD, a framework for multicenter fluorodeoxyglucose (FDG) PET studies has been defined and applied by the ADNI consortium.

Conclusions and Recommendations

Biomaterials

Given the importance of availability of biomaterials for the development of biomarkers, biomaterial sampling should take place in all ataxia patient studies. DNA, RNA, and serum or plasma and—if possible—PBMC samples should be taken for analysis of molecular processes and alterations on several molecular levels. Although there are not many publications on molecular properties of CSF samples from ataxia patients, we suggest CSF sampling for future investigations as well.

To be able to compare biomaterials from different studies, it is recommended that acquisition, handling, and storage be performed according to generally accepted standards of biobanking, as outlined above. This includes documentation of biomaterial quality. The value of biomaterial samples critically depends on the availability of standardized clinical data.

Clinical Assessment

Clinical assessment in ataxia studies must include application of a validated clinical ataxia scale (see Table 1). To facilitate comparability between studies, generic ataxia scales, such as SARA and ICARS, are preferable to scales that are applicable only to one specific ataxia disease, unless there are particular reasons to choose a disease-specific scale. In many ataxia disorders, ataxia is accompanied by additional nonataxia symptoms. Currently, INAS is the only available instrument to assess these symptoms comprehensively.

Patient-Reported Measures

Quality of life and the perspective of individuals on their health status are important outcomes in clinical trials. FAIS or the ADL part of FARS may be suitable tools.

Assessment of Neuropsychological and Psychiatric Symptoms

There are no instruments that were specifically designed for assessment of neuropsychological and psychiatric symptoms in ataxia. Thus, in studies of ataxia patients, various instruments are used that have been previously developed and validated in other populations. A list of tests that may be valuable in monitoring ataxia patients is given in Table 2. For ataxia, these instruments need validation in longitudinal and multicenter studies.

Quantitative Performance Tests

Quantitative performance tests have a great potential to serve as supplementary outcome measures in addition to clinical scales because they provide objective and quantitative information. It is therefore hoped that they are more sensitive than clinical scales. AFCS, SCAFI, and CCFS are each sets of simple timed tests that do not require expensive equipment or extensive training. Although they have been sufficiently validated so that they can be used in multicenter longitudinal studies, there is currently no evidence that they are more sensitive than clinical scales. Validated posturographic approaches using gold-standard laboratory methods or portable home-based devices are feasible options with the potential to be sensitive continuous outcome measures for use in multicentered studies. Associated with disease severity and balance impairment, sway speeds as measures of instability require careful standardization of sensory conditions to ensure short- and long-term test-retest reliability. Oculomotor readings have a great potential for detection of early abnormalities. Efforts to establish reliable multicenter recording techniques are encouraged.

Neuroimaging

Future multicenter neuroimaging studies in ataxias should implement quality assurance measures following standards, as
defined by the ADNI consortium or comparable consortia. Given that MRI morphometric studies in ataxia yielded robust results, MRI protocols should include T1-weighted magnetization-prepared rapid acquisition gradient echo (MP-RAGE) sequences that allow morphometric analyses. Because multicenter studies using other imaging modalities are lacking, no recommendations concerning specific sequences or protocols can be given. As an outlook, multimodal integration could be the way to generate significant makers for follow-up combing several imaging and clinical tools.

Electrophysiology

Effects on peripheral nerves, dorsal columns, and pyramidal tracts can be quantitatively assessed by neurophysiological tools, such as nerve conduction studies, somatosensory evoked potentials, and motor evoked potentials. Multicenter approaches to longitudinal studies in ataxias are rare and face major challenges in standardization to assure reproducibility and intercenter comparability.

A brief summary of all conclusions is given in Table 3.

Author Roles


Disclosures

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