Movement Disorder Society Task Force Viewpoint: Huntington’s Disease Diagnostic Categories

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Huntington’s disease (HD) is traditionally considered as a triad of movement, cognitive, and emotional disorders.1–4 According to current clinical practice, “manifest HD” is diagnosed primarily according to motor criteria, that is, when a clinician has 99% confidence of “an otherwise unexplained extrapyramidal movement disorder” in someone with a family history of HD. It is desirable to incorporate other features of the triad into a new classification system for clinical care and research. Large observational studies, including TRACK-HD, PREDICT-HD, COHORT-HD, and PHAROS, have greatly expanded our understanding of HD natural history. Subtle motor, cognitive, and emotional changes begin years before motor-manifest HD, and brain changes likely begin even earlier, motivating the consideration of consistent definitions across a wide range including prior to the appearance of manifest HD.5–14

In a previous publication, 3 of us (C.R., B.L., and R.R.) suggested modifying current diagnostic criteria to more broadly incorporate clinical features of HD.15 That article provided background about the natural history of HD, the current diagnostic criteria, and our proposed new diagnostic categories. The Movement Disorder Society commissioned a task force to consider the issues in research and clinical definitions of HD and to develop a lexicon. Christopher Ross was selected as chair with cochairs Francisco Cardoso and Ralf Reilmann. The change was to “select and convene a committee of HD experts, with involvement of patient and family representatives, to discuss diagnostic categories for Huntington’s disease” based on the recent studies of natural history and biomarkers for HD and to “produce a set of recommendations for diagnostic classifications of HD.” The task force met in person on February 4 and 5, 2017, followed by several teleconferences. The task force proposed 3 categories, presymptomatic HD, prodromal HD, and manifest HD, with presymptomatic and prodromal together comprising the premanifest HD period (Fig. 1, Table 1). The criteria for these classifications were developed using an informal consensus approach and include both cognitive and motor components.

Manifest HD

We propose adding nonmotor signs, particularly cognitive signs, to current motor diagnostic criteria13 (see Table 1). HD is a clinical diagnosis made on the basis of family history, personal history, neurological and psychiatric examinations, and genetic and any other appropriate testing. Extrapyramidal movement disorder is diagnosed based on the neurological examination, and the severity can be quantified using the Unified HD Rating Scale (UHDRS) motor examination,16 which yields a “total motor score” ranging from 0 to 124. Motor abnormalities in individuals

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A range from about 8 to 35. For individuals not followed longitudinally but seen for the first time, a higher motor score may be appropriate to make a diagnosis of definite HD.

In contrast, cognitive disorder is more difficult to diagnose because cognitive abilities vary considerably in the general population. It is optimal to confirm cognitive changes with longitudinal detailed neuropsychological testing, although this is not always feasible or affordable and should not be required. Information from family members or coworkers can provide crucial data about whether there appears to be cognitive impairment with change over time.

The task force proposes that cognitive disorder in manifest HD should be diagnosed according to the criteria of the Diagnostic and Statistical Manual, Fifth Edition (DSM) for either major neurocognitive disorder or minor neurocognitive disorder. A summary of how major neurocognitive disorder is defined in the DSM is as follows: first, evidence is provided of significant cognitive decline from a previous level of performance in 1 or more cognitive domains (complex attention, executive function, learning and memory, language, perceptual-motor, or social cognition). This is based on concern of the individual, a knowledgeable informant, or the clinician that there has been a significant decline in cognitive function and a substantial impairment in cognitive function, preferably documented by standardized neuropsychological testing or, in its absence, another quantified clinical assessment. Second, the cognitive deficits interfere with the independence in everyday activities (ie, at a minimum, requiring assistance with complex instrumental activities of daily living such as paying bills or managing medications). An important aspect of the DSM criteria is that the cognitive difficulties represent a decline from a previous level of functioning.

The tests listed in the National Institute of Neurological Disorders and Stroke Common Data Elements for Huntington’s disease, cognitive domain (https://www.commondataelements.ninds.nih.gov/HD.aspx#tab=Data_Standards) provide a useful resource, with the Montreal Neuro Cognitive Assessment (MoCA) perhaps the simplest and most widely used screening test. The Symbol Digit Modalities Test, which is part of the UHDRS cognitive battery, is another well-validated test for HD. Several executive function tests are listed, with the Trail-Making Test perhaps the simplest to use for screening (an abbreviated trail-making test is part of the MoCA).

A summary of how mild neurocognitive disorder is defined is as follows: first, evidence is provided of modest cognitive decline from a previous level of performance in 1 or more cognitive domains (complex attention, executive function, learning and memory, language, perceptual-motor, or social cognition).

In typical clinical practice, there is often substantial overlap between the stages of cognitive impairment described above. At the presymptomatic stage, individuals are rated as 0, normal. At the prodromal stage, patients are rated as 1, possible HD ("50% probability" of onset). At the symptomatic stage, patients are rated as 2, probable onset ("90% confidence") or 3, definite ("99% probability") HD. There is a long history of use by HD clinicians and researchers. HD is a clinical diagnosis, and so a Total Motor Score (TMS) threshold for diagnosis is not suggested or implied. In the PREDICT-HD study, which followed 225 premotor cases through motor diagnosis, the mean TMS score at diagnosis was around 15, with a range from about 8 to 35. For individuals not followed longitudinally but seen for the first time, a higher motor score may be appropriate to make a diagnosis of definite HD.

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memory, language, perceptual–motor, or social cognition), based on concern of the individual, a knowledgeable informant, or the clinician. Second, there is a modest impairment in cognitive performance. In this case, the cognitive impairments do not interfere with independence in everyday activities, but greater effort, compensatory strategies, or accommodation may be required.

It is important to exclude cognitive disorder secondary to depression, which can appear clinically similar to the cognitive disorder of HD, both sometimes described as having "subcortical" features. Depression and its associated cognitive disorder are not infrequent in all individuals and can appear many years before the onset of other features of HD and subsequently resolve, and depression and other mood disorders are treatable, so the task force agreed that depression by itself should not be used to diagnose HD.

Other emotional disorders (irritability, apathy, and personality change) are more variable in their presentation, can occur at any time during the course of HD, and are common in individuals who do not have a CAG repeat expansion, thus making emotional changes challenging for establishing a diagnosis of HD.

One important complication for the diagnosis of HD is that individuals with HD often have reduced or absent awareness of their impairments, or anosognosia. It is vital that clinicians engage with family members or other informants who often can provide critical longitudinal information, beginning with the initial presentation and continuing throughout the course of HD.

Manifest HD causes functional disability. This might be relatively subtle in the earlier phases, for example, greater expenditure of time or effort on adequate work performance, but there is then progressive decline in social and occupational function, and finally difficulty with basic activities of daily living. The division of HD into 5 clinical "stages" based only on functional capacity scores, and not on any biological criteria, appeared arbitrary to the committee, and 3 broad clinical periods such as "early," "moderate," and "severe" were favored.

We propose that the combination of a motor diagnostic confidence of 3 (notionally "90%"), plus minor neurocognitive disorder, be sufficient for a diagnosis of manifest HD. Some individuals may already have major neurocognitive disorder when seen for diagnosis. Manifest HD could still be diagnosed by motor criteria alone, which would still require a diagnostic confidence of 4.

**Presymptomatic HD**

We believe that it is important to establish a category for individuals who have the CAG expansion but as yet have no signs or symptoms related to HD, that is, prior to prodromal HD. These individuals would have no significant motor signs on exam (diagnostic confidence 0 or 1, nonspecific findings) and no cognitive changes. Individuals in this group may be candidates for future disease-modifying treatment to delay or prevent the onset.

**Premanifest HD**

The term premanifest is defined as the period prior to manifest HD, that is, inclusive of both the presymptomatic period and the prodromal period.

**Issues for Discussion**

**Approach to Diagnosis in Individuals Who Have Not Had Genetic Testing**

There was an agreement that similar categories would be applicable in the absence of genetically confirmed HD. The following terms are suggested: at risk for HD, but nonmanifest (ie, no signs or symptoms); clinically prodromal HD, and clinically manifest HD.

The diagnostician would likely need more definitive clinical evidence, especially longitudinal data, in those at risk but without genetic testing when compared with individuals with confirmed CAG repeat expansion. There are numerous HD-like syndromes that need to be considered in the differential diagnosis, especially in those without a clear family history.

Thus, the criteria described in Table 1 are to be reserved for individuals with a positive genetic test for the HD CAG repeat expansion.
Role of Genetic Testing and Biomarkers for Diagnosis

Task force members generally agreed that the diagnostic categories should currently refer to the clinical status of the patient rather than to genetic or diagnostic testing or biomarker determination. HD is a clinical diagnosis. A combination of CAG repeat length and age, usefully summarized as the CAG age product or CAP score, roughly predicts onset in groups of patients and serves as a useful longitudinal index of exposure to the effects of the CAG repeat expansion. However, CAG length by itself only explains about 50% in the variance of onset age. Genetic modifiers provide additional information. Quantitative motor examination (eg, Q-Motor instrumentation) is useful in research, but it has not been evaluated within the diagnostic setting.

Many years of study have consistently shown that progressive changes in structural magnetic resonance imaging (and likely functional magnetic resonance imaging and magnetic resonance spectroscopy) begin well before manifest HD and perhaps even before the period of prodromal HD. Blood and cerebrospinal fluid biomarkers, especially mutant Huntingtin (Htt) and neurofilament light-chain levels, are becoming increasingly relevant. Imaging and other tests should be used to rule out other conditions (especially in older individuals) and to determine whether other conditions may be contributing to the clinical presentation. However, biomarkers, although very useful for studying groups of patients and likely for clinical trials, are currently insufficiently precise and well validated for the diagnosis of individual patients.

“Cognitive Onset” of HD?

It is increasingly clear that cognitive dysfunction is important in causing functional disability and that prominent cognitive impairment can occur with relatively less noticeable motor changes. It is less clear whether substantial cognitive impairment occurs frequently in the absence of any detectable change on motor exam. The use of a multidimensional diagnosis including the cognitive, motor, behavioral, and functional aspects of the UHDRS (question 80) resulted in a slightly earlier diagnosis of HD than when based on motor exam alone (question 17), although the difference may have reflected functional as well as cognitive changes. Furthermore, the subgroup described as “predominantly cognitively impaired” in that study actually had higher motor scores than the other subgroups. Longitudinal neuropsychological testing (although time consuming and expensive) can be useful for cognitive assessment, contributing to the diagnosis of both manifest HD and prodromal HD. Individuals with early cognitive and behavioral changes may have greater anosognosia, and thus may not present in a timely fashion for motor examination. Even in the same family and with the same repeat expansion size, HD does not have an identical-appearing onset and progression. Thus we retain motor criteria for the diagnosis of HD, but highlight the importance of cognition.

Emotional Aspects of HD

There was general agreement about the importance of emotional alterations in HD. Personality changes, especially apathy and irritability, are increasingly appreciated as important contributors to functional disability. The TRACK-HD study has shown that apathy can begin quite early in what we would now term the prodromal phase, consistent with clinical experience. However, great caution should be taken before considering emotional changes exclusively for diagnosis of HD: emotional changes are quite common in individuals without the CAG expansion and may be even more common in individuals at risk for HD because of the disruptive effects of illness on family life. Furthermore, depression may be mistaken for apathy, and a readily treatable diagnosis should be prioritized. Nonetheless, emotional changes in someone at risk or with a known CAG expansion should trigger close follow-up of both motor and cognitive features and additional testing as appropriate.

Possible Subtypes of HD

The committee discussed whether the diagnostic categories would apply equally well to all individuals with HD. Juvenile-onset patients often have more bradykinesia, dystonia, and rigidity, and less (sometimes essential no) chorea, than adult-onset cases, whereas late-onset patients often have chorea-predominant HD, but both variants are well described by the current criteria. Cognitive onset has been proposed to be the most relevant for HD subtypes such as juvenile-onset HD.

Application to Research and Clinical Practice

We believe that ultimately the same categories should be used in both clinical and research settings to facilitate the transition from clinical trials to clinical practice. The set of classification criteria proposed are the result of an informal consensus process. The criteria are the result of holistic considerations of a selective group of publications and expert opinion. For these criteria to be refined both in clinical practice and research, it is desirable to validate this classification for accuracy, representativity, and usability in future studies. The currently proposed criteria are primarily designed for research because they use the research-based examination, the UHDRS; however, as clinical trials of disease-modifying therapy are advancing rapidly and early intervention may be most beneficial, we believe that clinical application should come soon, and that it is especially timely to include the prodromal HD diagnosis. We urge diagnostic and clinical practices to continue to be based on research and evidence-based criteria.

The composition of the task force was designed to be international, and we hope that our ideas will have wide international application, but we also are aware of the importance of regional and cultural issues. In some countries, such as the United States, the diagnosis and treatment of HD is often restricted by insurance rules to neurologists. The task force recommends that any physician with relevant training and experience should be
considered qualified to diagnose and treat HD, including psychiatrists, especially given the increasing awareness of the importance of the cognitive and emotional aspects of the disease. We also highlight that sensitive clinical judgment is of course paramount in discussing diagnostic and prognostic issues with patients and families.

Topics for the Future

Basic and clinical research in HD is moving rapidly, with clinical trials in progress (e.g., clinicaltrials.gov, NCT02519036) or in planning stages for several strategies for huntingtin lowering. These disease-modifying strategies have the potential to be applied in the prodromal or presymptomatic periods, making the availability of these diagnostic categories especially relevant for clinical practice. In addition, biomarker research is advancing rapidly. The proposed categories are based on clinical examination and have the limitation of the use of the diagnostic confidence scale with its disadvantage of implying a “pseudo-precision” via the “probability” thresholds. For many reasons, therefore, we propose a frequent reexamination of these diagnostic categories and a reassessment in 2 to 3 years, especially in relation to biomarkers.

We also urge further research on the topics discussed previously. Some topics for further study include the nature of early cognitive and emotional changes and their correlation with imaging and other biomarkers and the question of which signs and symptoms are most important in causing functional disability. The combination of datasets from the large observational studies will facilitate these studies, for instance, the identification of early cognitive changes. An important question for the timing of treatment is the point at which neuronal cell death or irreversible neuronal changes begin, as treatment, if safe, should ideally begin before that point. Given the limited access to different brain regions of some of the large molecules currently under study, understanding of the regional differences (and possible spread within the brain) of pathology at different points in the natural history will be especially important. All of these questions could use the CAP score more systematically to help order and sequence clinical and neurobiological changes and help in designing clinical trials. It would be especially valuable to design measures of the earliest changes in functional abilities. In keeping with the international nature of the task force, it would also be useful to know the extent of regional differences in the impact of definitions and clinical diagnoses. Perhaps most important will be research to demonstrate the efficacy of Htt-lowering or other disease-modifying therapies given during the premanifest period to delay or even conceivably prevent the onset of manifest HD.

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Disclosures

Ethical Compliance Statement: The authors confirm that, since there were no subjects or patients involved, patient consent or the approval of an institutional review board were not relevant nor required for this work. We confirm that we have read the Journal’s position on issues involved in ethical publication and affirm that this work is consistent with those guidelines.

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Comments and Suggestions

We welcome comments and suggestions. They should be sent to the chair (caross@jhu.edu).
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