REVIEW

The differential diagnosis of Huntington’s disease-like syndromes: ‘red flags’ for the clinician

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ABSTRACT
A growing number of progressive heredodegenerative conditions mimic the presentation of Huntington’s disease (HD). Differentiating among these HD-like syndromes is necessary when a patient with a combination of movement disorders, cognitive decline, behavioural abnormalities and progressive disease course proves negative to the genetic testing for HD causative mutations, that is, IT15 gene trinucleotide-repeat expansion. The differential diagnosis of HD-like syndromes is complex and may lead to unnecessary and costly investigations. We propose here a guide to this differential diagnosis focusing on a limited number of clinical features (‘red flags’) that can be identified through accurate clinical examination, collection of historical data and a few routine ancillary investigations. These features include the ethnic background of the patient, the involvement of the facio-bucco-lingual and cervical district by the movement disorder, the occurrence of cerebellar features and seizures, the presence of peculiar gait patterns and eye movement abnormalities, and an atypical progression of illness. Additional help may derive from the cognitive—behavioural presentation of the patient, as well as by a restricted number of ancillary investigations, mainly MRI and routine blood tests. These red flags should be constantly updated as the phenotypic characterisation and identification of more reliable diagnostic markers for HD-like syndromes progress over the following years.

INTRODUCTION
Huntington’s disease (HD) is the most common hereditary neurodegenerative illness with onset before mid-life and a distinct phenotype characterised by movement disorders (mainly chorea, dystonia and Parkinsonism), incoordination, cognitive deterioration and behavioural changes. Progressive neural cell loss within cortico-striato-thalamocortical circuits, predominantly in the neostriatum (caudate and putamen), leads to this clinical presentation.1 HD is a trinucleotide-repeat disorder with autosomal dominant mode of inheritance. The causative mutation is a prolongation (above 35 repeats) of a trinucleotide (CAG) stretch within the IT15 gene, associated with full penetrance when this exceeds 40 and incomplete penetrance between 36 and 39. As in several other trinucleotide-repeat disorders, there is a strong association between repeat length and age at disease onset, with higher repeat length leading to earlier onset. Paternal transmission facilitates this anticipation phenomenon. Age at onset is strongly linked to the clinical expression of both motor and cognitive features of the disease.1 Juvenile HD (ie, with onset before age 20) is dominated by a hypokinetic-rigid syndrome (Westphal variant) and a more rapid course, whereas adult-onset HD classically presents with a choreatic syndrome and a less severe progression of motor and non-motor features.1

An increasing number of progressive heredodegenerative illnesses may mimic the presentation of HD. It has been estimated that between 1% and 7% of subjects with a HD-like syndrome do not have HD.2 Given the relationship between motor phenotype and age at onset in HD, these HD-like syndromes are, with some exceptions, either diseases manifesting with a predominantly choreatic syndrome and onset in adulthood (online supplementary table 1A) or diseases presenting with predominant dystonia and Parkinsonism and onset in the first 2 decades (online supplementary table 1B). The comprehensive understanding of the phenomenology and diagnosis of progressive HD-like syndromes is crucial for an efficient and cost-effective diagnostic work-up and guides further genetic testing once the diagnosis of HD has been genetically excluded. This work-up may include thorough clinical examination, neuropsychological and neuropsychiatric assessment, metabolic screening, imaging and electrophysiology.

To date, progressive hereditary HD-like syndromes were classified mainly according to their mode of inheritance. Although this criterion can be very helpful in the presence of a well-established family history, it may not be practically useful in the numerous cases in which family history is unavailable or inaccurate. In this review, we discuss the differential diagnosis of HD focusing on a restricted number of clinical features (‘red flags’) that might help clinicians to narrow the spectrum of working diagnoses and improve efficiency and cost-effectiveness of the diagnostic process.

ETHNICITY
HD and the majority of progressive HD-like syndromes can be encountered in most geographical areas and ethnic groups. There are, however, a few important exceptions to consider in the differential diagnosis.

Huntington’s disease-like 2 (HDL2) is an autosomal dominant disorder described exclusively in families of Southern African origins, with the exception of one Brazilian family of Spanish/Portuguese ancestry. This condition accounts for about 0.7% of all HD-like presentations. HDL2 is caused by a CTG stretch in the JPH3 gene coding...
for junctophilin-3, leading to ubiquitin-immunoreactive intranuclear inclusions in the cortex and neuronal loss in the striatum and cortex. Unlike in HD, repeat expansions are more unstable and the anticipation phenomenon is more likely when the disorder is maternally inherited. HDL2 starts at the same age as HD, but can present either in a Parkinsonian or in a choreatic form. The choreatic form is strikingly similar to typical adult-onset HD, although both can present with cognitive deterioration, psychiatric features, eye movement abnormalities and MRI changes highly similar to HD. Overall, HDL2 should be suspected in every patient of South African ancestry with a HD-like presentation, accounting for 24%–50% of these cases. A HD-like syndrome in patients of Japanese ethnicity should raise the suspicion of dentato-rubro-pallido-luysian atrophy (DRPLA), another autosomal dominant disorder (summarised in detail below). In Japan, the prevalence of DRPLA is estimated to be similar to the prevalence of HD. DRPLA has been reported in non-Japanese patients, and a single haplotype has been found in both Japanese and Caucasian patients, in line with a common founder effect shared across ethnic groups.

Other conditions entering the differential diagnosis with HD could be more represented in some geographical areas than in others. Neuroferritinopathy is an autosomal dominant disorder presenting with progressive adult-onset chorea or dystonia initially affecting the limbs and mild cognitive deterioration. Molecular genetic testing for the FTL gene, coding for the ferritin light chain, confirms this diagnosis. This disorder was initially described in Cumbria, a region in the northwest of England. The majority of reported cases were originally from this region. Additional families reported in France and an additional North-American patient with German ancestry do not exclude the possibility of a founder effect in northwestern Europe. Other examples of ethnic segregation in HD-like syndromes include spinocerebellar ataxia 8, especially common in Finland, and ataxia with oculomotor apraxia (AOA) type 2, mainly seen in French–Canadian and Anglo-Norman populations.

**BODY SITE DISTRIBUTION OF MOTOR SYMPTOMS**

Most progressive HD-like syndromes display a generalised movement disorder, involving in different combinations face, neck, torso and limbs. A disabling involvement of the facio-bucco-linguo-masticatory region should, however, lead the clinician towards a few specific conditions.

HD-like syndromes presenting in adulthood with chorea or other movement disorders, including dystonia and tics, with marked involvement of the facio-bucco-linguo-masticatory region comprise ‘neuroacanthocytosis syndromes’, mainly chorea-acanthocytosis and McLeod syndrome. Chorea-acanthocytosis is an autosomal recessive disorder caused by mutations in the VPS13A gene coding for chorein, a large protein implicated in intracellular protein sorting. Facio-bucco-linguo-masticatory chorea, dystonia and dyskinesias are typically precipitated by feeding, with tongue protrusion and self-mutilating tongue and lip biting. These patients may also present with violent neck spasms with sudden flexion/extension that can lead to characteristic head drop and head banging (see online supplementary video 1). Vocalisations and dysarthria are also present, whereas Parkinsonism is rare. The co-occurrence of areflexia, weakness and muscle wasting, suggestive of concomitant peripheral neuropathy and/or myopathy, may further help in the diagnosis. McLeod syndrome is named after a Harvard medical student, Hugh McLeod, who did not express the Kell antigen on erythrocytes, as all affected male subjects with this condition. It is an X linked recessive disorder, present in most geographical regions, caused by different mutations in the XK gene. McLeod syndrome is a multi-system disorder, affecting basal ganglia, muscles, peripheral nerves and myocardium. It presents with chorea that may involve the facio-buccal region and vocalisations, but almost never with tongue or lip biting, dysphagia, or Parkinsonism; feeding dystonia has also been reported. Importantly, 60% of patients develop a cardiomyopathy with atrial fibrillation and risk of cardiac sudden death, thus requiring periodical cardiac monitoring.

When the movement disorder is mainly characterised by early dystonia or a hypokinetic-rigid syndrome with relevant facio-bucco-lingual involvement, the main differential diagnosis to consider is pantothenate kinase-associated neurodegeneration (PKAN), an autosomal recessive disorder with a frequency of 1–3 per million, caused by mutations of the PANK2 gene coding for pantothenate kinase, an enzyme relevant to the synthesis of coenzyme A from vitamin B5 and associated with lipid metabolism. PKAN is also known as neurodegeneration with brain iron accumulation type 1 (NBIA1). In almost 90% of cases, this condition starts before the age of 6, with rapid progression towards negative outcome within 10 years. PKAN presents with generalised dystonia with buccofacial and lingual involvement, often in the form of severe jaw opening dystonia, although not specifically related to feeding as in chorea-acanthocytosis (online supplementary video 2). Parkinsonism may be a presenting symptom, and choreoathetosis, tics and pyramidal signs may also occur. About a third of these patients develop retinitis pigmentosa. Sometimes, PKAN has later onset and less severe presentation, with rigidity, focal arm dystonia or cognitive–behavioural onset. A similar orofacial action-specific dystonia, associated with speech and dysarthrophonia, is seen in other forms of NBIA, including Kufor–Rakeb syndrome (autosomal recessive illness caused by ATP13A2 mutations), aceruloplasminemia and neuroferritinopathy, although the last typically presents in adulthood. Wilson’s disease (WD) may also present with severe orooral dystonia associated with Parkinsonism and in the setting of a generalised dystonia, thus sometimes posing a diagnostic challenge with HD. Finally, Lesch–Nyhan syndrome can also manifest in the first decade with severe action dystonia and chorea, often involving the facio-lingual region.

**ATAXIA**

Under rare circumstances, HD patients may exhibit a mild degree of cerebellar atrophy throughout disease progression. Marked cerebellar involvement has been reported only very rarely in juvenile HD. When present, cerebellar signs are generally very mild, and the occurrence of clinically more severe cerebellar dysfunction suggests alternative diagnoses.

Probably the most common HD-like syndrome in Caucasian populations is spinocerebellar ataxia 17 (SCA17, also known as HDL4). This accounts for 0.5%–1.8% of all HD-like syndromes, and its age of onset is between 19 and 48 years, rarely starting during childhood. Like HD, SCA17 is an autosomal dominant trinucleotide-repeat disorder, in which the mutated TBP gene encodes for the TATA-box binding protein, an important general transcription initiation factor. Intergenerational instability, especially during paternal transmission, and anticipation have been recognised. Cerebellar ataxia is the most common feature, but SCA17 may be very heterogeneous. Dystonia and chorea are the two most frequent movement disorders. When Parkinsonism is present, SCA17 may resemble multiple system atrophy, although without clear autonomic
**Movement disorders**

dysfunction, and the dopamine transporter single-photon emission computer tomography (DaTScan, GE Healthcare) may be abnormal. Pyramidal signs occur in more than a third of cases. Overall, in most families, a true HD-like presentation is observed only in one or few affected members, although intrafamilial phenotypic homogeneity has been described.17

Another important HD-like syndrome with cerebellar involvement is DRPLA. This is an autosomal dominant disease caused by CAG-repeat expansion in the ATN1 gene coding for atrophin-1.18 Like HD and SCA17, DRPLA shows the phenomenon of anticipation, and its typical age at onset is the third decade. DRPLA presents with three main features, chorea, myoclonus and ataxia. The differential diagnosis with HD and neuroacanthocytosis syndromes can be very challenging, especially when chorea is a predominant clinical feature (online supplementary video 3). When myoclonus is the most disabling symptom, generally this is associated with an earlier onset (<20 years), and with the coexistence of severe progressive myoclonic epilepsy and cognitive decline. When onset occurs after the age of 40, cerebellar ataxia is usually the predominant characteristic.18

Unlike ataxic syndromes may present rarely with choreatic movements. Unlike in SCA17, chorea presenting in young adults is very uncommon in other autosomal dominant SCAs. Choreatic movements of the limbs are exceptional in SCA1 and SCA2, and slower than in HD.19 20 Patients with SCA7, another trinucleotide disorder involving the ATXN7 gene coding for ataxin 7, may rarely manifest small choreatic movements in the distal limbs or in the orofacial district. Rare chorea of the hands may rarely be present in different combination and lead to peculiar gait. The association of chorea and predominantly ataxic gait, with widened base and difficulties performing the tandem gait, is an aspecific feature of both autosomal dominant (SCAs, DRPLA) and autosomal recessive ataxias (FA, AT, AOAs). In chorea-acanthocytosis and McLeod syndrome, the co-occurrence of limb chorea, dystonia and peripheral neuropathy contributes to the peculiar gait and postural changes of these disorders.10 11 These are characterised by unsteadiness of stance and gait, leading to frequent falls, and caused by impaired postural reflexes, sudden knee buckling, and equinovarus foot deformities secondary to dystonia and atrophy of peroneal muscles. The occurrence of flinging movements of arms and legs, pelvic thrusts and violent trunk spasms further compromise the stability of gait in these patients. The coexistence of dystonia, rigidity, bradykinesia and spasticity typifies the gait pattern in juvenile HD-like syndromes, for example, PKAN, WD, PLA2G6-associated neurodegeneration (PLAN), Kufor–Rakeb syndrome or the Parkinsonian form of HDL2.3 13 15 Particularly, gait disturbance is a common presenting symptom in PKAN, originating from the combination of lower limb rigidity, dystonic foot posturing in plantar flexion and inversion, and spasticity, as well as restricted visual field in those children with retinopathy. Marked freezing of gait has also been described in most syndromes mimicking juvenile HD.

**EYE MOVEMENT ABNORMALITIES**

A detailed examination of ocular movements can be very informative. HD patients manifest early abnormalities in initiating saccades, which progressively become slow and hypometric, with superimposed dysfunctional gaze fixation. In several progressive HD-like syndromes, primarily chorea-acanthocytosis,11 PKAN14 and WD,28 we find similar changes. A supranuclear gaze palsy with marked impairment of upgaze saccades, associated with oculogyric dystonic spasms and L-dopa-responsive akinetic-rigid Parkinsonism, is typical of patients with Kufor–Rakeb syndrome.29 In chorea-acanthocytosis, fractionated saccades and square-wave jerks are also found.30 HD can present also with a mild form of oculomotor apraxia, which is, however, much more severe in AOA types 1 and 2 and AT.25–27 When asked to look to one side, subjects with oculomotor apraxia turn their heads first, with eye contraversion, after which their eyes follow to the same side in several slow saccades with head thrusts to the point that, when the head is immobilised, eye movements are virtually impossible. Finally, it is very important to point out that an early manifestation of ‘cerebellar’ eye movement abnormalities, for example, dysmetric saccades, square-wave jerks, ocular flutter, saccadic pursuit and gaze-evoked nystagmus, is typical of the most common spinocerebellar degenerative illnesses, and hence very useful to differentiate HD particularly from SCA-17 and FA.

**SEIZURES**

In HD, seizures are present only in the juvenile variant, occurring in 30%–50% of patients with onset before age 10. At difference, in a number of progressive HD-like syndromes seizures are relatively common, presenting in 22% of SCA-17 cases, 60% of cases with chorea-acanthocytosis, 40% of cases with McLeod syndrome31 and the majority of cases with Huntington’s disease-like 1 (HDLI) syndrome, a prion disease resembling adult-onset HD (detailed below). Seizures are the initial manifestation in almost half of chorea-acanthocytosis cases. Temporal lobe seizures, even simulating a familial
temporal lobe epilepsy, are seen in chorea-akanthocytosis, whereas they are mostly generalised in McLeod syndrome.\textsuperscript{33} Seizures are also a cardinal feature of DRPLA, present in virtually all cases with onset before the age of 20, less frequently when onset is between ages 20 and 40, and rarely with onset above the age of 40.\textsuperscript{18} Seizures in DRPLA can be tonic, clonic, tonic-clonic, absences, atomic, and in the form of myoclonic epilepsy, with or without a progressive myoclonic epilepsy phenotype. Epilepsy is rare in all the other HD-like syndromes.

### SPEED OF PROGRESSION

The rate of progression of HD can vary considerably across subjects, being, at least in part, related to age of onset and length of trinucleotide-repeat expansion. HD-like syndromes with adult onset may vary in their speed of progression. Importantly, some conditions may progress more rapidly. Among the latter, HDL1, a very rare autosomal dominant prion disease with onset in early adulthood (third–fourth decades), should be kept in account. HDL1 is caused by a 192-nucleotide insertion in a region of the prion protein (PRNP) gene coding for an octapeptide-repeat. HDL1 is recognised as the fourth genetic prion disease phenotype, although not associated typically with spongiform degeneration pathology, but with atrophy and prion deposition within basal ganglia, frontal and temporal lobes, and cerebellar cortex.\textsuperscript{32} Although this condition progresses more slowly than other prion diseases, case series reported that most subjects survive between 1 and 10 years from onset. Patients develop prominent personality changes, cognitive decline, chorea, rigidity, limb and truncal ataxia, dystarthishia and seizures. It is useful to remind that chorea may be a feature also of the most common prion disease, that is, Creutzfeldt-Jakob disease.\textsuperscript{33} Other forms characterised by higher speed of progression are HDL2, relentlessly leading to death within 10–20 years,\textsuperscript{3} and DRPLA, generally leading to death within 10–15 years.\textsuperscript{18}

Like juvenile HD, HD-like syndromes with earlier onset progress rapidly. An outstanding exception is benign hereditary chorea (BHC), which, unlike other HD-like syndromes with onset in the first 2 decades, is characterised by a non-progressive or very slowly progressive course dominated by chorea. BHC is now considered a genetically heterogeneous spectrum, first described as an early-onset form of chorea not associated with intellectual deterioration. It is suspected in cases of focal or generalised chorea with autosomal dominant inheritance, and is often described as socially embarrassing (online supplementary video 4). Disease onset occurs in early infancy. A relevant proportion of these cases bear mutations in the \textit{NKX2-1} gene encoding for thyroid transcription factor-1 (TTF1), which is a very important transcription factor involved in the organogenesis of the brain, thyroid and lungs.\textsuperscript{34} An allelic disorder to BHC presents with a more severe phenotype characterised by choreoathetosis, congenital hypothyroidism and neonatal respiratory distress, also referred to as ‘brain-thyroid-lung syndrome’, and therefore very different from HD. These cases have also been associated with loss of TTF1-mediated striatal interneurons.\textsuperscript{35} Given its more benign outcome, it is very important to distinguish BHC from HD, although not always easy, due to its variable expression. Less frequently than chorea, these patients can manifest intention tremor, dystarthishia, gait impairment and exceptionally even psychosis, psychomotor developmental problems and short stature.\textsuperscript{16} Albeit usually non-progressive, the phenotype of BHC might change with time, shifting from chorea to myoclonus and dystonia.\textsuperscript{37} Reduced striatal volumes on volumetric studies have also been reported.\textsuperscript{38} However, genetic heterogeneity has been observed, with two Japanese families expressing a similar phenotype but with onset between ages 40 and 66, and for this reason resembling adult-onset HD; these subjects, however, lacked cognitive impairment and exhibited an association with a different locus on chromosome 8q. This picture has been named BHC 2, and a postmortem report in one of these patients surprisingly showed pathological changes remarkably similar to those typical of progressive supranuclear palsy (neurofibrillary tangles and tufted astrocytes immunoreactive for four-repeat isoforms of \(\tau\)).\textsuperscript{39}

### COGNITIVE AND BEHAVIOURAL FEATURES

Abnormalities of cognitive functioning and behaviour are cardinal features of HD. Classical, adult-onset HD is characterised by a global, progressive cognitive decline. In early stages, cognitive dysfunction may be subtle and detectable mainly through formal neuropsychometry. It may present as loss of mental flexibility, executive dysfunction, memory difficulties (especially of retrieval), and attention or concentration defects. Throughout the course of illness, patients develop forgetfulness, bradyphrenia (ie, slowness of thought processes), impaired visuospatial abilities and ability to manipulate knowledge, and language disturbances leading to reduced syntactic complexity, cortical speech problems, reduced word finding and paraphasic errors. Psychopathology includes early personality changes, depression (which may lead to increased suicidal ideation and suicide risk), hostility, obsessive–compulsive symptoms, anxiety, heightened interpersonal sensitivity and phobias. As disease progresses, patients may exhibit affective or schizophrenic psychoses, particularly with paranoia delusions, as well as explosive and aggressive outbursts, apathy, alcohol abuse, sexual dysfunction and increased appetite.

A formal neuropsychometric and psychiatric assessment may assist in the differential diagnosis in a minority of cases. Some of the most common adult-onset progressive HD-like syndromes are indistinguishable from HD in terms of their cognitive–behavioural profile. SCA17 leads to a fronto-subcortical dementia in 75%–90% of cases, and to psychiatric symptoms very similar to those in HD in 25%–30%.\textsuperscript{17} The choreatic form of HDL2 presents with a cognitive–behavioural syndrome of milder severity and slower progression, compared with HD and to its Parkinsonian form, in which more severe fronto-subcortical dementia, depression, apathy and irritability are observed.\textsuperscript{3} HDL1, on the other hand, is characterised by a more rapid progression of a similar cognitive dysfunction.\textsuperscript{32}

A fronto-subcortical dementia is a cardinal feature of DRPLA, present in 90% of patients, irrespective of their age at onset. Although a psychiatric onset is rare, up to 80% of DRPLA patients present with behavioural problems. Also, when the onset of DRPLA occurs before the age of 20, we can detect psychotic symptoms, hyperactivity/impulsivity, restlessness, aggressive behaviours, autism-like features, and developmental delay or regression.\textsuperscript{18}

The cognitive–behavioural profile of chorea-akanthocytosis is very similar for course and phenomenology to that of classical adult-onset HD, being characterised by a frontal lobe syndrome, apathy, depression, bradyphrenia and obsessive–compulsive symptoms, which are often subtle at onset. Memory impairment, dysexecutive problems and a schizophreniaiform psychosis may also occur at onset. Some additional features are, however, typical of chorea-akanthocytosis, especially particular compulsive behaviours like trichotillomania, or self-mutilating actions such as chronic head excoriations. Similarly severe, although at an earlier age of onset, the persistent self-injurious behaviours

of Lesch–Nyhan syndrome consist of biting of fingers, hands, lips and cheeks, and head- or limb-banging against hard objects. Fronto-subcortical cognitive features are milder in McLeod syndrome, whereas psychiatric problems are very common, consisting in personality disorder, anxiety, depression, obsessive–compulsive disorder, and bipolar and schizoaffective disorder. These psychiatric features may precede the movement disorder by many years, whereas cognitive decline appears later in the course of illness. Juvenile HD differs from adult-onset HD in that it may exhibit a more rapid and severe cognitive deterioration, associated with speech and language delay. The intellectual impairment observed in early onset dystonia/Parkinsonism syndromes, such as PKAN, may be indistinguishable from that of juvenile HD. Atypical forms of PKAN, which begin after the first decade, may be recognised, however, by some cognitive features. Speech problems including palilalia, tachylalia/tachylogia and dysarthria are frequent onset symptoms, which may be associated with personality changes with impulsivity, violent outbursts, depression and emotional lability. Similar features can be observed in all the main metal accumulation disorders resembling juvenile HD, including PLAN, Kufor–Rakeb syndrome and WD. In all these disorders, cognitive–behavioural abnormalities become less severe as age of onset increases.

ANCILLARY INVESTIGATIONS

Basic ancillary investigations should be performed before any genetic test in the diagnostic work-up of HD-like syndromes. A number of routine blood tests are necessary to differentiate HD from non-progressive causes of chorea. Most of these causes, however, present with an acute or subacute onset, and therefore display a very different course from heredodegenerative HD-like syndromes. A detailed overview of these non-genetic choreas is beyond the scope of this work and has been the object of other review articles.

Some laboratory tests are, however, useful also when facing a clearly progressive hereditary HD-like syndrome. An increased serum activity of creatine kinase and liver enzymes is frequently observed in chorea-acanthocytosis and McLeod syndrome. Low serum activity of creatine kinase and liver enzymes is frequently observed in chorea-acanthocytosis and McLeod syndrome. Low serum ferritin levels can be found in a proportion of patients with neuroferritinopathy, whereas abnormal copper studies—particularly increased 24 h urinary copper excretion—support a diagnosis of WD, or aceruloplasminemia if ceruloplasmin is undetectable in the serum. Abnormal lipoprotein electrophoresis can be found in some patients with PKAN. Among autosomal recessive ataxias, hyponatraemia is found in AOA type 1, hypercholesterolaemia in both AOA types 1 and 2, whereas increased serum levels of α-1-fetoprotein are found in AOA type 2 and AT.

Neuroacanthocytosis syndromes (chorea-acanthocytosis, McLeod syndromes, HDL2, PKAN) are obviously characterised by the presence of increased percentages of acanthocytes in the peripheral blood. However, the determination of an increased acanthocyte percentage in peripheral blood may not be sensitive enough with routine traditional methods, and may require phase contrast microscopy. An additional test that can be used to identify patients with chorea-acanthocytosis is the demonstration of absent chorein expression in erythrocytes using western blotting, but this test is not commercialised and is performed only by very few laboratories in Europe and North America.11

Conventional brain MRI may be useful under certain circumstances. The typical adult-onset form of HD is characterised by early striatal atrophy, particularly of the caudate, whereas a certain degree of cortical and cerebellar atrophy may be seen later on in the course of illness. Several adult-onset progressive HD-like syndromes are indistinguishable from HD based on MRI findings. Examples are HDL2, chorea-acanthocytosis and McLeod syndrome. In the last two, the atrophy of the caudate is accompanied by dilatation of the anterior horns of the lateral ventricles, visible particularly on coronal sections, as often seen also in HD.11 In addition, T2-weighted high signal lesions in the striatum, mild generalised cortical atrophy (less in McLeod syndrome), and hippocampal sclerosis and atrophy (especially in chorea-acanthocytosis) can be detected.31

The presence of cerebellar atrophy at various degrees is very important to help differentiating HD from all forms of SCAs, in particular SCA-17. MRI can be very useful to differentiate DRPLA from HD, in that DRPLA patients exhibit cerebellar and brainstem atrophy, particularly involving the pontine tegmentum, which correlates with age and number of CAG-repeats;18 also, diffuse T2-weighted high signal lesions in the deep subcortical white matter can be observed.41 Autosomal recessive ataxias like FA, AOA types 1 and 2, and AT may not exhibit cerebellar atrophy early in the course of illness. However, FA may progress with atrophy of the cervical cord, cerebellum and superior cerebellar peduncles,42 increased diffusion in white matter tracts on diffusion-weighted imaging,43 and reduced N-acetyl-aspartate peak height on magnetic resonance spectroscopy.44 BHC, conversely, does not show pathological changes on conventional imaging.

MRI can be of great use in guiding the differential diagnosis between juvenile HD and metal accumulation disorders. PKAN is associated with the pathognomonic ‘eye-of-the-tiger’ sign, characterised by a central region of hyperintensity surrounded by a rim of hypointensity on coronal or axial T2*-weighted images in the globus pallidus.13 45 This abnormality is highly correlated with the presence of PANK2 gene mutations, even in the presymptomatic phase. Other brain metal accumulation disorders, such as PLAN, Kufor–Rakeb syndrome, aceruloplasminemia (iron) and WD (copper) show mainly hypointense pallidal signal without central hyperintensity and, less commonly, iron deposition in the substantia nigra, especially in PLAN and Kufor–Rakeb–Syndrome. In NBIA, typically, globus pallidus and substantia nigra appear involved on T2*- and fast spin echo MRI scans, with also dentate involvement seen on T2*-weighted images.45 Interestingly, nigral pathology in PLAN and Kufor–Rakeb syndrome seems related to the presence of a PD-like Parkinsonism which responds to L-dopa, and may even be complicated with L-dopa-induced dyskinesias.15 MRI, however, may also be completely normal in these disorders.

Finally, other ancillary investigations provide limited additional aid to the diagnostic work-up. The confirmation of a concurrent peripheral neuropathy or myopathy in chorea-acanthocytosis or McLeod syndrome may require a routine electromyographic or electroneurographic study. Evoked potentials may be of some use to demonstrate degeneration of posterior columns in spinocerebellar degenerative ataxias, mainly SCA17 and FA. Muscle imaging, particularly CT, may unveil fatty changes, associated with neurogenic atrophy on biopsy, in patients with chorea-acanthocytosis and McLeod syndrome.

CONCLUSIONS

The number of progressive heredodegenerative illnesses that need to be considered in patients presenting with HD-like syndrome who tested negative for pathological IT15 gene trinucleotide-repeat expansions has remarkably increased over the next decade. The identification of disease-causing variants in
monogenic disorders is becoming increasingly powerful. Methodologies such as exome sequencing, currently applied on a limited basis, can provide specific genetic diagnoses also in some HD-like syndromes, such as neuroacanthocytosis syndromes, improving the potential of genotype–phenotype correlations. Genetic studies of these diseases remain in general, however, very costly and not widely available in routine diagnostic laboratories. There is a substantial overlap in the clinical phenotype of HD and HD-like syndromes, making the differential diagnosis between these different conditions difficult in several cases. However, a few clinical features, summarised in flowchart in figure 1, can help clinicians to rationalise the differential diagnosis, and avoid unnecessary expensive investigations. These red flags represent at present a simple guide that should be constantly updated as the phenotypic characterisation and the identification of more reliable diagnostic markers for HD-like syndromes progress over the following years. The list of novel experimental treatment approaches for HD-like syndromes is also growing.47–49 Further understanding of the phenotype–genotype correlation of this heterogeneous group of disorders will help patient selection and outcome prediction with respect to new therapeutic strategies.

Contributors DM: conception and execution of the research project; writing of the first draft. MS and KB: conception and design; acquisition of data; interpretation of data; drafting the article. MS: acquisition of data; critical revision for important intellectual content. KB: conception and design; critical revision for important intellectual content.

Competing interests None.

Patient consent Obtained.

Provenance and peer review Commissioned; externally peer reviewed.

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Movement disorders


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*J Neurol Neurosurg Psychiatry* 2013 84: 650-656 originally published online September 19, 2012
doi: 10.1136/jnnp-2012-302532

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