Reviews

**The Neuropsychiatry of Hyperkinetic Movement Disorders: Insights from Neuroimaging into the Neural Circuit Bases of Dysfunction**

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**Abstract**

**Background:** Movement disorders, particularly those associated with basal ganglia disease, have a high rate of comorbid neuropsychiatric illness.

**Methods:** We consider the pathophysiological basis of the comorbidity between movement disorders and neuropsychiatric illness by 1) reviewing the epidemiology of neuropsychiatric illness in a range of hyperkinetic movement disorders, and 2) correlating findings to evidence from studies that have utilized modern neuroimaging techniques to investigate these disorders. In addition to diseases classically associated with basal ganglia pathology, such as Huntington disease, Wilson disease, the neuroacanthocytoses, and diseases of brain iron accumulation, we include diseases associated with pathology of subcortical white matter tracts, brain stem nuclei, and the cerebellum, such as metachromatic leukodystrophy, dentatorubropallidoluysian atrophy, and the spinocerebellar ataxias.

**Conclusions:** Neuropsychiatric symptoms are integral to a thorough phenomenological account of hyperkinetic movement disorders. Drawing on modern theories of cortico-subcortical circuits, we argue that these disorders can be conceptualized as disorders of complex subcortical networks with distinct functional architectures. Damage to any component of these complex information-processing networks can have variable and often profound consequences for the function of more remote neural structures, creating a diverse but nonetheless rational pattern of clinical symptomatology.

**Keywords:** Movement disorders, neuroimaging, psychopathology, frontostriatal

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**Introduction**

Movement disorders, particularly those associated with basal ganglia disease, have a high rate of comorbid neuropsychiatric illness.1 An early recognition of this fact can be found in Huntington’s classic description of hereditary chorea, in which he noted “a tendency to insanity and suicide” to be one of the “marked peculiarities” of the disease.2 Following his lead, McHugh described a triadic syndrome of “dyskinesia, dementia, and depression” in basal ganglia disorders, drawing on examples from Huntington disease (HD) and Parkinson disease in particular.3 Citing the seminal work of Alexander et al., McHugh went on to propose that damage to anatomically parallel but functionally distinct neural loops within the basal ganglia was responsible for the diverse manifestations of his syndrome. While aspects of McHugh’s remain pertinent today, our understanding of the biology of neuropsychiatric illness in movement disorders has advanced considerably in recent decades.5-12 This has been facilitated in part by developments in structural and functional neuroimaging techniques, especially in the field of magnetic resonance imaging (MRI) and positron emission tomography (PET).

This paper draws on neuroimaging research specifically in relation to hyperkinetic movement disorders, which, in their diversity, offer the
opportunity to refine and extend McHugh’s ideas. We argue that what unifies these disorders in terms of their common relationship to a range of cognitive, emotional, and behavioral symptoms lies in altered structure and function in subcortical loops and circuits with which basal ganglia structures are often, but not always, associated. The core neuropsychiatric syndromes of some of these disorders, such as HD and the neuroacanthocytoses for instance, are very likely to involve dysfunctional cortico-striatal circuits (Figure 1)\textsuperscript{11,13} other disorders, such as dentatorubropallidoluysian atrophy (DRPLA) and the spinocerebellar ataxias (SCAs), may involve additional dysfunction within cortico-spinal and cortico-cerebellar circuits (Figure 2).\textsuperscript{10,12,14} Our revised model synthesizes McHugh’s observations with modern pathophysiologic theories of cortico-subcortical circuits\textsuperscript{7–12} and finds support in a number of studies correlating structural and functional neuroimaging with the variable neuropsychiatric comorbidity of the hyperkinetic disorders. We argue that our approach provides novel explanations as to why certain psychiatric syndromes are more common in certain diseases, and may in fact offer a framework within which to hypothesize about the frequent comorbidity of psychiatric symptoms in other neurological disorders.

**Huntington disease**

HD (Online Mendelian Inheritance in Man 613004) is an inherited autosomal dominant condition defined by a pathological trinucleotide repeat affecting the huntingtin (HTT) gene on the short arm of chromosome 4. Its prevalence is estimated to fall between 4.1 and 7.5 per 100,000.\textsuperscript{15} The clinical onset of the disease is generally accepted as the manifestation of motor symptoms. This commonly occurs in the fourth or fifth decade, with an inverse correlation between age of onset and number of pathological trinucleotide repeats on the largest expanded allele.\textsuperscript{16}

Neurosychiatric symptoms have long been associated with HD.\textsuperscript{2} Depression appears to be the most common psychiatric comorbidity, with estimates of prevalence ranging from 33% to 69% of patients.\textsuperscript{17} This stands in contrast to the estimated lifetime prevalence of depression in the general population of about 16.5%.\textsuperscript{18} While one study found the rate of suicide to be up to eight times higher than that of the general population in HD patients over the age of 50,\textsuperscript{19} a more recent finding is that depressive symptoms decline with illness stage,\textsuperscript{20} a discrepancy that may be due to supervening of depressive symptoms by cognitive impairment in the later stages of the illness. Mania and psychosis are less common in HD, with estimates of prevalence ranging from 3–12% and 5–10%, respectively,\textsuperscript{21} which still stands in contrast to the estimated lifetime prevalence of up to 1% and 3%, respectively, in the general population.\textsuperscript{22,23} Patients with an earlier age of HD onset are at greater risk of psychosis.\textsuperscript{24} Formal cases of obsessive compulsive disorder (OCD) have rarely been reported, against a background prevalence of around 1.6% in the general population.\textsuperscript{18} Nevertheless, in one large study nearly one-quarter of patients presenting to a specialist clinic for the first time demonstrated obsessive or compulsive symptoms.\textsuperscript{25} Obsessive and compulsive symptoms have also been associated with increasing stage of illness\textsuperscript{26} and greater executive dysfunction.\textsuperscript{27}

The classic neuroimaging finding in HD is atrophy of the caudate and putamen with concomitant alterations in the ratios of frontal horn to intercaudate distance and intercaudate to inner table distance.\textsuperscript{28,29} This gives the frontal horns of the lateral ventricles a characteristically rounded or box-like appearance. T2-weighted MRI signal may also be
increased throughout the caudate, putamen, and globus pallidus due to gliosis and iron deposition.  

Several studies have used diffusion tensor imaging (DTI) to investigate changes in white matter connectivity in both patients and presymptomatic carriers of HD. Most studies have demonstrated disruption of multiple cortical and subcortical white matter tracts in patients and carriers alike, with robust correlations with stage of illness and selective cognitive and motor impairments.  

The origin of the changes lies in neuronal destruction in specific regions of the caudate and putamen.  

HD patients with depression have also been shown to exhibit hypometabolism in orbitofrontal and inferior prefrontal cortex on fluorodeoxyglucose (FDG)-PET compared with HD patients without depression.  

These changes are associated with subcortical but not cortical gray matter volume loss, suggesting that cortical functional changes may arise secondary to subcortical structural changes.  

We conclude that disruption to cortico-striatal circuits plays an important role, not just in the motor manifestations of HD but also in the cognitive and behavioral manifestations of the illness.  

**Neuroacanthocytoses**  
The neuroacanthocytoses are a group of disorders characterized by neuropsychiatric symptoms associated with spiculated red blood cells in the peripheral circulation. While genetically diverse, the neuroacanthocytoses include such disorders as chorea acanthocytosis (ChAc), McLeod syndrome (MLS), and Huntington disease-like 2 (HDL2), each of which will be considered below.  

**Chorea acanthocytosis**  
ChAc (OMIM 200150) is a rare autosomal recessive disorder caused by mutations in the VPS13A gene on chromosome 9q coding for the membrane protein chorein.  

As in HD, ChAc causes neuronal pathology that is most evident in the caudate and putamen, but can also be seen in the ventrolateral substantia nigra and globus pallidus.  

The clinical onset of the disease is usually between the ages of 25 and 45 years.  

The majority of ChAc patients experience neuropsychiatric symptoms, and these may precede overt neurological illness by more than a decade.  

The most common psychiatric problem in ChAc is OCD, which affects more than 50% of patients.  

A similar mechanism may account for the prevalence of OCD in ChAc given the established link between obsessive-compulsive and dysexecutive symptoms and impairments in the lateral orbitofrontal loop primarily centered on the caudate.  

**McLeod syndrome**  
MLS (OMIM 300842) is an X-linked genetic disorder characterized by weakly expressed or absent Kell red blood cell antigens, acanthocytosis, and elevated creatine kinase levels. It is caused by mutations in the KX gene and mainly affects males, although female carriers can show mild symptoms.  

Clinical onset is usually between the ages of 25 and 60 years, and up to 80% of patients demonstrate neuropsychiatric comorbidity at some stage of their illness.  

Like ChAc, the neuronal loss associated with MLS predominantly affects the caudate and putamen, and neuropsychiatric symptoms not uncommonly predate the neurological manifestations of the illness.  

Executive dysfunction is common in MLS and has also been described in female carriers.  

A high prevalence of OCD-like syndromes has been reported by a number of authors.  

Psychotic disorders may be just as frequent, with cases of schizophrenia-like illnesses being reported in which typical psychotic symptoms have preceded the onset of chorea.  

Marked caudate atrophy and increased T2 signal in the lateral putamen are common MRI findings in MLS.  

Reduced striatal D2 binding has been reported in PET studies along with hypometabolism of the basal ganglia and frontal lobes and altered N-acetylaspartate to creatine/choline ratios in the frontal and medial temporal cortex and thalamus.  

The absence of cortical neuronal loss on limited neuropathological studies is reminiscent of HD, and again suggests that functional cortical abnormalities are secondary to striatal neuron loss, presumably via the loss of mediating cortico-striatal pathways.  

**Huntington disease-like 2**  
HDL2 (OMIM 606438) is an autosomal dominant disease caused by a CTG expansion in the junctophilin-3 gene (JPH3) on chromosome 16q24.3.  

It is only found in patients of black African ancestry.  

Only a small number of HDL2 cases with comorbid neuropsychiatric illness have been reported in the literature, but they are notable for...
the heterogeneity of associated symptoms, including depression, anxiety, and psychosis, in addition to a frontal-dysexecutive syndrome leading to dementia. While neuropsychiatric comorbidity appears to be the rule rather than the exception in HDL2, it may differ from the other neuroacanthocytoses in that neuropsychiatric symptoms seem to be reported only after the onset of neurological illness.

In common with the other neuroacanthocytoses, neuroimaging reveals significant atrophy of both the caudate and the putamen, and to a lesser extent the substantia nigra and pallidum. The association of deep structural brain changes with such a broad range of neuropsychiatric symptomatology again illustrates the critical role of cortico-subcortical circuits in the behavioral manifestations of the illness.

**Wilson disease**

Wilson disease (WD—OMIM 277900) is an autosomal recessive disorder characterized by a mutation in the \(ATP7B\) gene coding for a copper transport protein, leading to copper deposition in multiple organ systems. In the central nervous system (CNS), it particularly affects the putamen and pallidum. While the initial presentation of WD may be hepatic, neurological, or psychiatric, about half of all patients have neuropsychiatric symptoms at a given time.

Affective disorders appear to be the most common neuropsychiatric illnesses in WD and often meet the criteria for bipolar disorder. Personality changes characterized by irritability and aggression are also very common. While neurologically asymptomatic patients rarely show cognitive deficits, those with neurological disease often display a range of difficulties, including impairments of executive function, memory, and visuospatial processing. Psychosis is relatively uncommon although cases of delusional disorder have been reported. OCD has only been reported once in association with WD, which is surprising given the high rate of OCD in other basal ganglia disorders, but may be hypothesized to reflect some sparing of the caudate relative to illness such as HD and ChAc.

About half of patients with WD demonstrate basal ganglia hypodensity on computed tomography (CT) and virtually all show abnormalities on MRI. Characteristic features may include T2 hyperintensities within the lenticular nuclei, ventrolateral thalamus, and hypothalamus, probably due to the paramagnetic effect of copper and iron deposition within these structures. T2 hyperintensities in the tegmentum often spare the red nuclei and medial margins of the parts reticula and superior colliculi, giving rise to the so-called “face of the giant panda” sign.

FDG-PET typically shows hypometabolism in the lenticular nuclei and in one study 19 of 25 patients examined by \(^{99m}\)Tc-ECD SPECT exhibited diffuse or focal hypoperfusion affecting the superior frontal, prefrontal, and occipital cortices, the temporal gyri, caudate, and putamen. The fact that the primary pathology associated with this disease is to be found in the basal ganglia once again suggests that proximal damage to components of cortico-subcortical circuits may have profound effects on distal brain function, such as cognition and emotion.

**Neurodegeneration with brain iron accumulation**

Neurodegeneration with brain iron accumulation (NBIA) encompasses a group of genetically diverse disorders characterized by neuronal death secondary to brain iron deposition. In the adult population, NBIA includes pantothentenate kinase-associated neurodegeneration, aceruloplasminemia, and neuroferritinopathy.

**Pantothenate kinase-associated neurodegeneration**

Pantothentenate kinase-associated neurodegeneration (PKAN—OMIM 234200) is a rare autosomal recessive disorder caused by mutations in the \(PAVK2\) gene. There are two main phenotypes. In its classic form, PKAN is characterized by a severe and rapidly progressive hyperkinetic movement disorder with an onset in the first decade leading to complete loss of ambulation. A second, atypical form exists that is characterized by a less severe and less rapidly progressive movement disorder with an onset in the second to third decade. Both phenotypes are associated with additional neurological symptoms including dysarthria, dystonia, and cortico-spinal abnormalities.

Cognitive decline is common in PKAN and tends to affect earlier-onset patients more severely. It is not uncommon for cognitive symptoms to predate motor signs and a pattern of impaired attention and executive function is characteristic. Up to half of all patients experience psychiatric problems, which include behavioral disturbances, OCD, tic disorders, psychosis, and depression.

The neuroradiological hallmark of PKAN is the so-called “eye of the tiger sign”, which is characterized by bilateral areas of hyperintensity within a region of hypointensity in the medial globus pallidus on T2-weighted MRI. The low signal intensity is said to result from excessive iron accumulation with the central high signal being attributed to gliosis and cavitation resulting in increased water content. It is important to note that the sign is not specific to PKAN: it can also been seen in other neurodegenerative disorders such as cortico-basal ganglionic degeneration, progressive supranuclear palsy, and neuroferritinopathy.

In terms of functional imaging, a \(^{99m}\)Tc-ECD SPECT study has described regional hypoperfusion of the bilateral frontoparietal lobes, lenticular nuclei, and ventriculus quartus in two siblings with adult-onset disease and an identical \(PAVK2\) mutation. The cortical metabolic findings are of particular interest given that PKAN is classically defined by its subcortical pathology, which again suggests that diaschisis via cortico-subcortical circuits plays an important role in the genesis of clinical neuropsychiatric symptoms.

**Aceruloplasminemia**

Aceruloplasminemia (AC—OMIM 604290) is a rare autosomal recessive disorder characterized by mutation in the ceruloplasmin (\(CP\) gene leading to iron deposition in multiple tissues due to impaired ceruloplasmin ferroxidase activity. The major sites of CNS iron deposition include the basal ganglia, cerebellar dentate nuclei, red nucleus, thalamus, and hippocampus. AC occurs in just 1:2 million non-consanguineous births and usually presents in the fifth or sixth decade.
About half of all patients with AC present with cognitive impairment \(116\) and most progress to develop a subcortical dementia characterized by executive dysfunction and cognitive slowing associated with frontal hypometabolism. \(114,117,118\) Only one case of major psychiatric illness associated with AC has been reported, in the form of a typical schizophreniform psychosis. \(118\)

MRI typically demonstrates marked T2 hypointensity corresponding to regions of pathological iron deposition including those noted above \(119–121\) and can be used to quantify iron accumulation in vivo. \(122\) Subtle hypointensities of posterior white matter tracts have also been reported, and subtle superficial cerebral and cerebellar cortical hypointensities may be detectable when sequences sensitive to the magnetic susceptibility effects of iron are used. \(121\)

Given the characteristic distribution of iron deposition in AC, the primarily dysexecutive cognitive syndrome associated with the illness may be the result of impairment to both cortico-striatal and cortico-cerebellar circuits at the subcortical level. \(117\)

**Neuroferritinopathy**

Neuroferritinopathy (NF—OMIM 606159) is a rare autosomal dominant disorder characterized by mutations in the ferritin light chain (FTL) gene. Patients commonly present in the fourth to sixth decade with features of chorea (50%), dystonia (42.5%), or Parkkinsonism (7.5%). \(123\) As with the other NBIAs, the globus pallidus is a common focus of iron deposition. \(124\)

In the small number of cases described, dementia and psychosis have been the most common neuropsychiatric comorbidities. While patients are generally said to have intact cognition until late in the disorder, \(125\) disinhibition and emotional lability may be the early symptoms of a dysexecutive cognitive syndrome typically leading to a frontosubcortical dementia. \(126,127\) In one pedigree, akinetic mutism was an additional late manifestation. \(126\) Psychosis has also been associated with NF: ataxia, rigidity-bradykinesia and neuroleptic-responsive psychosis was reported in an adolescent patient with a family history of schizophrenia, \(129\) while another patient who presented with severe generalized dystonia at the age of 22 years later developed delusional jealousy. \(130\)

MRI typically demonstrates iron deposition and cavitation in the basal ganglia with a characteristic loss of T2 signal. The red nucleus and substantia nigra appear to be the first structures affected, with subsequent involvement of the dentate nucleus, putamen, globus pallidus, thalamus, caudate nucleus, and prefrontal cortex. \(126\) Increasing R2* (the inverse of T2*) signal appears to correlate with increasing severity of dystonia, and may be a clinically useful method of tracking disease progression. \(131\) Although functional imaging studies are yet to be conducted, the anatomical distribution of injuries on structural MRI would be expected to result in abnormalities in cortico-striatal and cortico-cerebellar circuits, potentially correlating with the predisposition of NF patients to psychosis and dementia.

**Dentatorubropallidoluysian atrophy**

Dentatorubropallidoluysian atrophy (DRPLA—OMIM 125370) is a rare genetic disorder caused by a triplet repeat expansion of the atrophin gene \((ATN-1)\) on chromosome 12p13.31. \(132\) It is most prevalent in Asian pedigrees, and a review of cases within the Japanese literature found psychosis to be the most common psychiatric comorbidity, affecting about 10% of patients. \(133\)

Serial neuroimaging findings were reported in a Caucasian man who was first scanned at the age of 38 years with a history of seizures, tremor, ataxia, and dysarthria. \(134\) Interval imaging showed progressive atrophy of cerebellum and brainstem, with GRE and SWI sequences demonstrating marked susceptibility effect throughout the cerebellar hemispheres, vermis, and dentate nuclei. T2-weighted signal abnormalities have also been documented in both the subcortical white matter and brainstem white matter tracts in DRPLA. \(135,136\) A neuroradiological-pathological study of a father and son demonstrated signal intensity changes in white matter were due to a marked loss of myelinated fibers, while gray matter changes were largely due to a loss of neuropil. \(137\) Given the strong evidence that alterations to myelinated fibers underpin psychotic symptoms in schizophrenia, \(138–140\) it is not surprising that psychosis is a common psychiatric comorbidity in this group. It is difficult to explain why the degenerative changes of the cerebellum and brainstem have not been associated with cognitive symptoms, except to hypothesize that the potential link may not yet have been investigated.

**Spinocerebellar ataxias**

The SCAs are a large family of autosomal dominant neurodegenerative disorders associated with subcortical dementia syndromes, \(141–144\) depression, and personality change. \(145\)

Neuroimaging studies have mainly focused on SCAs 1 (OMIM 164400), 2 (OMIM 183090), 3 (OMIM 109150), 6 (OMIM 183086), and 17 (OMIM 607136), and typically reveal one of three patterns of atrophy: spinal atrophy (SA), olivopontocerebellar atrophy (OPCA), and cortico-cerebellar atrophy (CCA). \(146\) OPCA is typical of SCA 1, 2, and 3 and is characterized by diffuse T2 and proton density signal changes in the pons, middle cerebellar peduncle, and cerebellum in association with atrophy of the cerebellum, brainstem, and cervical spinal cord. \(147\) CCA is associated with SCA 6 and 7, and MRI typically shows atrophy of the cerebellar folia without signal change, while the brainstem and spinal cord volume is preserved. \(148\) A finding that is believed to be specific to SCA3 is linear T2-hyperintensity along the medial margin of the globus pallidus interna, and may be due to degeneration of the lenticular fasciculus. \(149\)

In MRI studies using diffusion-weighted and diffusion tensor imaging, both diffusivity and fractional anisotropy (FA) in the brainstem and cerebellum was found to be decreased relative to controls in SCA 1 and 2. \(150,151\) Decreased FA may be more robustly correlated with clinical severity in SCA 1 than measures of atrophy using conventional MRI. \(152\)

One voxel-based morphometry study of SCA2 patients correlated patterns of executive dysfunction with atrophy of the posterior cerebellum, and coordinative dysfunction with atrophy of the anterior cerebellum. \(153\) Another study demonstrated a similar dissociation of cognitive and motor functions in SCA17, as well as an inverse...
Metachromatic leukodystrophy

Metachromatic leukodystrophy (MLD—OMIM 250100) is a severe neurodegenerative metabolic disorder, also classified as a lysosomal storage disorder. It is caused by deficient activity of arylsulfatase A (ARSA) leading to accumulation of glycosphingolipid sulfatide and progressive demyelination in the central and peripheral nervous system. MLD is both genetically and phenotypically heterogeneous, with a variable age of onset. Some MLD mutations are associated with predominant motor presentations, others with cognitive and psychiatric features. Adult homozgyotes for p.P426L tended to present with gait disturbances followed by choreathetic movements, dysphagia, dysarthria, tremor, and nystagmus, whereas carriers of the less common p.I179S mutation present primarily with psychosis. The cognitive changes of MLD may resemble those of a generalized Alzheimer’s dementia, with features that include amnesia, visuospatial dysfunction, attentional deficits, and slowed processing speed.

MLD is typically associated with the distinctive imaging findings of diffuse periventricular and subcortical white matter hyperintensities sparing subcortical U-fibers on T2-weighted MRI. The pattern of white matter involvement is also characteristic with linear or punctate high signal radiating in the demyelinated white matter, sparing the perivascular white matter, resembling “tigroid” or “leopard-skin”. An increase in white matter myo-inositol on magnetic resonance spectroscopy (MRS) supports demyelination in the pathophysiology of MLD.

A large case review has previously hypothesized that the heavy burden of white matter disease in MLD, which particularly affects the subfrontal white matter, may account for the unusually high prevalence of psychosis in this disorder via disruptions to both frontotemporal cortico-cortical and cortico-striatal circuitry.=

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Niernmann–Pick disease type C

Niernmann–Pick disease type C (NPC—OMIM C1 257220, C2 607625) is a progressive neurodegenerative disorder caused by disrupted intracellular sterol trafficking. Most cases are associated with disruption of the NPC1 gene on 18q11, but around 5% are associated with disruption of the NPC2 gene on 14q24.2. The clinical manifestations of the disease are highly heterogeneous and may appear at any time between birth and late adulthood, although adult patients typically present with ataxia, dystonia, chorea, vertical gaze palsy, impaired cognition, and psychiatric illness.

Psychosis is the most common psychiatric presenting complaint among adult NPC patients. Presentations of depression, bipolar affective syndromes and OCD have also been reported. While up to 38% of NPC patients will first present with a psychiatric syndrome, psychiatric sequelae—other than those associated with dementia—in patients who first present with neurological symptoms is regarded as rare.

In one study of six adult NPC patients, VBM demonstrated gray matter reductions particularly affecting the hippocampus, thalamus, superior cerebellum, and striatum, regions where ganglioside accumulation is greatest. On DTI, widespread reductions in fractional anisotropy in major white matter tracts were also observed, with subanalyses suggesting pathological contributions from both impaired myelination and altered axonal structure. Other volumetric studies of NPC patients have demonstrated reduced callosal area and thickness and increased pontine-to-midbrain ratio in correlation with duration of illness, symptom score, and aspects of saccadic dysfunction; and reduced left hippocampal volume in correlation with symptom score and cognitive dysfunction. An FDG-PET study of two monozygotic children over a 2-year period demonstrated diffuse cortical hypometabolism initially in the medial frontal cortex that progressed to severe bilateral hypometabolism of frontal, parietal, and temporal cortices.

These structural and functional imaging studies in NPC provide a compelling model of disrupted cortico-cortical connectivity due to diffuse and extensive white matter changes, and cortico-subcortical and cortico-cerebellar disconnection secondary to direct gray matter disruption in these key relay zones.

Discussion

While hyperkinetic movement disorders are heterogeneous in terms of both their neuropathology and neuropsychiatric comorbidity, observations from neuroimaging studies suggest they share some common neural circuit bases in association with complex forms of cognitive, emotional, and behavioral dysfunction. By synthesizing a modern model comprising cortico-striatal-pallido-thalamic and cortico-cerebellar circuits with that of McHugh, we argue that damage to certain strategic components of these circuits provides unifying explanation for the apparently disparate functional manifestations of these disorders. The argument proceeds as follows:

Firstly, several neuropsychiatric syndromes are especially prevalent across hyperkinetic movement disorders: executive dysfunction and...
subcortical dementia, affective disorders including OCD, and psychotic disorders including schizophrenia-like syndromes. Many of these neuropsychiatric disorders have been independently associated with particular patterns of cortical dysfunction and dysfunction ascribed to particular cortico-subcortical circuits. For instance, it appears that early damage to the striatum strongly predisposes to OCD.54 The caudate in particular is recognized as a point of origin for the lateral orbitofrontal loop, and OCD may arise as a consequence of alterations to the normal neurodevelopment of this circuit.181 Corticostriatal and striato-limbic circuits are similarly implicated in the pathophysiology of psychosis,178 and cortico-cerebellar circuits in the cognitive abnormalities associated with schizophrenia.156,179

Secondly, almost all of the hyperkinetic movement disorders are associated with neuron loss in the basal ganglia, and some are also associated with neuron loss in subthalamic and cerebellar nuclei. These nuclei can be conceptualized as connective hubs or crossroads4 for both higher and lower brain functions, and within particular cortico-subcortical circuits.

Thirdly, the limited functional imaging studies that exist in this area points towards a dissociation between changes in cortical volume and changes in cortical function,36 suggesting that the cortical abnormalities associated with these disorders may, in some circumstances, arise secondarily to the loss of populations of subcortical neurons. Alterations to cortico-subcortical (Figure 1) and cortico-cerebellar (Figure 2) circuitry constitute a mechanism for diaschisis that is consistent with the manifestations of neuropsychiatric dysfunction observed.

While certain psychiatric syndromes may be more or less common in the hyperkinetic disorders, their heterogeneity also serves to provide some evidence of the involvement of cortico-subcortical circuitry: although the main circuits may be functionally distinct and subserve different motor, cognitive, and emotional behaviors,4 they are not anatomically separate; rather, they are parallel loops connected by white matter tracts converging at certain strategic hubs, such as the striatum, pallidum, thalamus, and cerebellum.7–12 Consequently, these circuits interact in the striatum and other hubs to modulate each other’s output,100 leading to symptoms across a broad range of domains. The strategic vulnerability of cortico-subcortical circuitry at multiple sites in the network may be a key factor in the pathophysiology of these disorders.181

In the case of the hyperkinetic disorders with pathology more strongly rooted in the basal ganglia (such as HD and the neuroacanthocytes), there is persuasive clinical and neuroimaging evidence of remote functional pathology detectable at the level of both white matter tracts and cortex. On the other hand, disorders associated with primary white matter pathology, such as MLD, clearly exert a strong functional influence on their proximal gray matter connections. This paper also extends the model of cortico-subcortical dysfunction to include nuclei in the cerebellum and brainstem, with similar observations to be made for subthalamic diseases such as DRPLA and the SCAs. Finally, those diseases in which cortico-subcortical loops are affected very selectively (such as HD or MLD) appear to demonstrate more robust associations to severe neuropsychiatric illness than those that are affected more diffusely (such as WD and NBIA).

McHugh proposed a triadic model of basal ganglia disorders characterized by symptoms of dyskinesia, dementia, and depression. His original hypothesis was that these disparate syndromes could be accommodated by virtue of shared anatomical pathology affecting functionally distinct subcortical loops. Our argument is that McHugh’s acknowledgement of depression must be extended to include a broader range of psychiatric syndromes associated with the cognitive-emotional domains subserved by such circuits, and that his understanding of dyskinesia must be extended to include a broader range of motoric dysfunction

We would also like to suggest that a neurodevelopmental model may help scaffold an understanding of the relative preponderance of these conditions at different stages of illness. For instance, it is generally the case that disorders known to be associated with departures from a normal neurodevelopmental trajectory tend to present with early neuropsychiatric illness such as OCD and schizophreniform psychosis. The disruption to crucial late neurodevelopmental processes by young-onset neurodegenerative disorders may thus result in neuropsychiatric syndromes that would otherwise tend to present in adolescence or early adulthood.182 In contrast, disorders that are associated with degeneration of the mature brain, and which result in cognitive decline or frank dementia, tend to be those which present later. One possible exception to this principle is the early appearance of executive dysfunction, which may be detectable in carriers and presymptomatic individuals long before the onset of clinical disease. Executive function may be a particularly cognitive vulnerable faculty given its phylogenetically more recent origin; this may also, however, reflect the rich interconnectedness between frontal cortex and subcortical gray matter, with disorders that either diffusely affect connecting structures or specifically impact basal ganglia structures showing a predilection for impairments in executive function.

In conclusion, we suggest that an expanded version of McHugh’s model incorporating modern models of cortico-subcortical circuitry7–12 can be extended beyond diseases of the basal ganglia to encompass a wide range of neurodegenerative disorders affecting cortico-subcortical loops. Damage to any part of these complex information-processing systems—including cortical grey matter, subcortical white matter, or subcortical grey matter nuclei—can have variable and often profound consequences for the function of more remote neural structures, creating a diverse but nonetheless rational pattern of clinical symptomatology.

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