Chapter 2

Psychopathology in verified Huntington’s disease mutation carriers

E. van Duijn, E.M. Kingma, R.C. van der Mast

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Abstract

Huntington’s disease is characterized by motor, cognitive, and neuropsychiatric symptoms. This study reviews original research on psychopathology in Huntington’s disease that uses standardized instruments in verified mutation carriers. Frequently reported neuropsychiatric symptoms are depressed mood, anxiety, irritability, and apathy, with prevalences of 33% to 76%. Obsessive-compulsive symptoms and psychosis occur less often with prevalences of 10% to 52% and 3% to 11%, respectively. Available research provides little insight into the true prevalences of psychopathology in Huntington’s disease due to small sample sizes, use of different methodologies, and lack of comparison groups. Future research requires larger cohorts stratified to disease stage, consistent methodologies, and adequate comparison groups.

Introduction

Huntington’s disease, a progressive autosomal-dominant neurodegenerative disorder, is traditionally characterized by choreiform motor disturbances. In addition to motor symptoms and cognitive deterioration, neuropsychiatric symptoms comprise part of the Huntington’s disease phenotype.\(^1\) \(^2\) The genetic defect underlying Huntington’s disease is an unstable and expanded CAG repeat in exon 1 of gene IT15 on the short arm of chromosome 4, which is expressed as a mutant polyglutamic tract in the protein huntingtin (Htt).\(^3\) \(^7\) The mechanisms by which the mutant Htt protein induces a cascade of cellular changes, leading to cell dysfunction and degeneration, have not yet been fully elucidated. Modulation of genetic functioning through the IT15 gene, neuronal death in relation to intranuclear inclusions of aggregated mutant Htt, and progressive cerebral degeneration starting in the caudate nucleus and the putamen may all be part of the pathophysiology of Huntington’s disease.\(^8\) \(^11\)

Estimated rates for lifetime prevalence of psychiatric disorders among Huntington’s disease patients vary widely between 33% and 76%.\(^8\) \(^10\) The investigated neuropsychiatric symptoms include depressed mood, anxiety, irritability, apathy, obsessions and compulsions, and psychosis. This variation in prevalences can be explained by the use of different assessment methods with varying definitions of neuropsychiatric phenomena. No follow-up studies covering a longer period have been performed.

For many patients and their relatives, these neuropsychiatric symptoms constitute the most distressing aspect of Huntington’s disease and often constitute reason for hospitalization.\(^11\) Whereas severity of motor and cognitive dysfunction is only moderately related to the severity of functional decline, behavioral symptoms and psychiatric disorders seem to have a more severe negative effect on daily functioning.\(^11\) Previous findings have suggested, although inconclusively, that psychopathology as well as cognitive dysfunction may precede the onset of motor symptoms in many patients.\(^14\) \(^16\)

To get more insight into the occurrence and prevalence of behavioral problems and psychiatric disorders in Huntington’s disease, we review the literature on psychopathology in verified Huntington’s disease mutation carriers, with particular reference to its relationship with disease onset and progression, as well as possible underlying neuropathological pathways. We conclude with several recommendations for future research.

Data sources

We searched two literature databases, Embase and PubMed, for prevalence of psychopathology in Huntington’s disease. We used a variety of search terms, all synonyms for Huntington’s disease and various neuropsychiatric phenomena. Where possible, these were mapped onto the following standard database terms (subject headings/MeSH terms): ‘Huntington’s disease’, ‘Huntington’s chorea’, ‘mood disorder(s)’, ‘affect’, ‘anxiety disorder(s)’, ‘obsessive behavior’, ‘compulsive behavior’, ‘schizophrenia and disorders with psychotic features’, ‘psychosis’, ‘thought disorder’, ‘dissociative disorder(s)’, ‘neurotic disorder(s)’, ‘neurosis’, ‘impulse control disorder(s)’, ‘impulsive behavior’, ‘irritable mood’, ‘apathy’, ‘behavioral symptoms’, ‘behavior disorder(s)’, and ‘personality disorder(s)’. Animal studies and studies on pathophysiology were
excluded and the language was limited to English. The references of the resulting articles were
hand searched for further relevant literature. This search resulted in 134 articles, including 59
articles describing original research, 29 review articles, 25 articles on psychopharmacological
treatment, 19 case reports/series, and two editorials.

In order to estimate the cumulative prevalences and 95% confidence intervals (CI) of psycho-
pathological phenomena, the 59 articles on original research were further selected for meeting
the following conclusive set of criteria:

1) The study was original and measured the prevalence of psychopathology in a motor
symptomatic Huntington’s disease population.
2) The study applied standardized instruments with defined validity and reliability.
3) The study used samples with verified CAG repeat expansions, which implied publication after
1993 when the Huntington’s disease mutation was identified.

Results
A total of seven original articles met the final set of strict criteria (Table 1). The other 52 articles
were excluded for the following reasons: 22 articles did not cover research on the prevalence
of psychopathology; one article only concerned alcohol abuse; one exclusively concerned
sexual abuse; and seven others concerned suicide/suicidal behavior. In three studies, solely
pre-motor symptomatic subjects were included, and in five studies subjects were offspring of
Huntington’s disease mutation carriers and had not been genetically verified. One article was
excluded because only patients in a nursing home were examined. Of the remaining articles,
10 did not mention standardized instruments with defined validity and reliability and, in two
articles, subjects were clinically suspected for Huntington’s disease, but CAG repeat numbers
were not verified. The remaining seven articles employed the following instruments for the
assessment of behavioral and psychiatric symptoms: the Neuropsychiatric Inventory (NPI),6,18
the Structured Clinical Interview for the DSM-III (SCID),12 the behavioral section of the Unified
Huntington Disease Rating Scale (UHDRS),2,21 the Yale-Brown Obsessive Compulsive Symptom
(Y-BOCS) scale,22 and the more recently developed Problem Behavior Assessment Scale for
Huntington’s Disease (PBA),23 which rates severity and frequency of behavioral problems in
Huntington’s disease.

A broad range of symptoms portraying a chronically progressive course and fluctuating clinical
picture are reported as neuropsychiatric features of Huntington’s disease. These neuropsychiatric
phenomena are de noted by an array of terms, for example: behavioral problems or symptoms,
personality changes, and psychiatric problems or disorders. The characteristic behavioral changes
in early stage Huntington’s disease, including depression, irritability, mental inflexibility, and
apathy, have in earlier days been described as ‘choreopathy’.24 We have limited our subsequent
analyses to the most frequently reported symptoms of depressed mood, anxiety, irritability,
apathy, obsessive and compulsive symptoms, and psychosis. Originally, we intended to estimate
the cumulative prevalences of the different neuropsychiatric phenomena. In spite of our strict
inclusion criteria, however, large inconsistencies in methodology remained. This would lead
to neither reliable nor valid results; studies used different assessment methods with strongly
varying definitions of neuropsychiatric phenomena. For example, the definition of depression
is much stricter according to the SCID than to the UHDRS.25,26 Also, out of two studies using the
same neuropsychiatric assessment measure in populations of comparable disease duration and
cognitive function, one study27 consistently reported lower prevalences than the other.28 This
suggests a strong bias. Furthermore, the criteria for the onset of Huntington’s disease are not
always given, the comparability of reported disease durations is highly questionable, and finally,
not all populations are well-described. We therefore assumed them to be outpatients unless
otherwise noted.

Table 1. Overview of included articles on psychopathology in Huntington’s disease

<table>
<thead>
<tr>
<th>Year</th>
<th>Author et al.</th>
<th>n</th>
<th>Disease duration*</th>
<th>Measure</th>
<th>Symptoms</th>
</tr>
</thead>
<tbody>
<tr>
<td>2001</td>
<td>Anderson et al.24</td>
<td>27</td>
<td>Unknown</td>
<td>Y-BOCS</td>
<td>Obsessive and/or compulsive symptoms</td>
</tr>
<tr>
<td>2001</td>
<td>Craufur et al.19</td>
<td>78</td>
<td>9 ± 5</td>
<td>PBA-HD</td>
<td>Apathy, irritability, depressed mood, anxiety, obsessive and/or compulsive symptoms, psychotic symptoms</td>
</tr>
<tr>
<td>2001</td>
<td>Kulisevsky et al.17</td>
<td>29</td>
<td>5.6 ± SEM 1.6</td>
<td>NPI</td>
<td>Apathy, irritability, depressed mood, anxiety, psychotic symptoms</td>
</tr>
<tr>
<td>2001</td>
<td>Murgod et al.20</td>
<td>26</td>
<td>5.5 ± 3.9</td>
<td>UHDRS</td>
<td>Apathy, irritability, depressed mood, anxiety, obsessive and/or compulsive symptoms, psychotic symptoms</td>
</tr>
<tr>
<td>2001</td>
<td>Paulsen et al.18</td>
<td>52</td>
<td>4.7 ± 4.4</td>
<td>NPI</td>
<td>Apathy, irritability, depressed mood, anxiety, psychotic symptoms</td>
</tr>
<tr>
<td>2002</td>
<td>Leroi et al.19</td>
<td>21</td>
<td>12.0 ± 6.6</td>
<td>SCID</td>
<td>Depressive disorder</td>
</tr>
<tr>
<td>2005</td>
<td>Paulsen et al.21</td>
<td>2835</td>
<td>7.6 ± 6.0</td>
<td>UHDRS</td>
<td>Depressed mood, anxiety</td>
</tr>
</tbody>
</table>

* Disease duration in years (mean ± standard deviation; SEM: Standard Error of the Mean).
Y-BOCS: Yale-Brown Obsessive Compulsive Scale; PBA-HD: Problem Behavior Assessment for Huntington’s disease; NPI: Neuropsychiatric In-
ventory; UHDRS: Unified Huntington Disease Rating Scale; SCID: Structured Clinical Interview for DSM

Depressed mood
Six original studies investigate the prevalence of depressed mood in motor symptomatic
patients.13,21,22 Though two studies used the UHDRS to assess the prevalence of ‘low mood’,20,21
and two others used the NPI,13,18 results vary strongly, from 33% to 69% (Figure 1). The only
study using formal DSM criteria reports a prevalence of 43% for mood disorders: 29% for major
depression and 14% for non-major depression.19
Anxiety

Five studies assess the prevalence of anxiety (Figure 2).\textsuperscript{17,18,20,21,23} The lowest prevalence (34\%) was reported with the NPI.\textsuperscript{17} The prevalence almost doubled (61\%) in a small study, using the UHDRS, in 26 Huntington's disease patients at their first hospital visit because of manifesting motor symptoms.\textsuperscript{20}

Irritability

Irritability, varying in description from ‘difficult to get along with’ to ‘aggression’, is characterized by a reduction in control over temper that may result in verbal or behavioral outbursts.\textsuperscript{27} In four original studies that assessed irritability as a separate behavioral phenomenon in Huntington's disease, prevalences varied between 38\% and 73\% (Figure 3).\textsuperscript{17,18,20,23}

Apathy

Apathy, characterized by reduced energy and activity, lack of drive, and impaired performance of everyday tasks, may be a separate clinical entity distinct from depression, especially in neuro-psychiatric disorders.\textsuperscript{19,20} In three original studies, prevalences of apathy in Huntington's disease patients varied from 34\% to 76\% (Figure 4).\textsuperscript{17,18,23} Using the PBA, a cluster of symptoms reflecting apathy syndrome was found, with ‘loss of energy’ (88\%), ‘impaired performance of everyday life’ (76\%), and ‘lack of initiative’ (76\%) as the most prevalent behavioral abnormalities.\textsuperscript{23}
Observe and compulsive symptoms
The three studies investigating obsessive or compulsive behavior (Figure 5) reported prevalences of 10% to 52%.[^20-23] Out of 27 Huntington's disease patients visiting an outpatient clinic, 52% scored either on compulsions or obsessions on the Y-BOCS.[^25] The prevalence of obsessive symptoms was twice that of compulsive symptoms, while all patients with compulsive symptoms also had obsessive symptoms. Only two out of the 27 patients fulfilled formal DSM criteria for obsessive compulsive disorder. The two remaining studies reported lower prevalences in larger study populations.^[20-23]

![Figure 5: Prevalences of obsessive and/or compulsive symptoms (with 95%CI)](image)

Psychotic symptoms
Prevalences of psychotic symptoms varied between 3% and 11% in four studies.^[17,18,20,23] Because of small sample sizes, three of the four studies report 95% confidence intervals that include a prevalence of 0% (Figure 6).^[17,20,23]

![Figure 6: Prevalences of psychotic symptoms (with 95%CI)](image)

Discussion
These results confirm that behavioral problems and psychiatric disorders are major constituents of the clinical spectrum of Huntington's disease. This is an important finding because these neuropsychiatric symptoms have a substantial impact on daily functioning, possibly even more so than motor and cognitive dysfunctions.^[24] Nevertheless, more and better designed studies are necessary.

The studies up to date use a variety of assessment methods in Huntington's disease populations of different disease stages. This ensures that their results are impossible to compare and that reliable prevalence estimates cannot be made. Definitions of neuropsychiatric phenomena are often unclear, and differences in definition can strongly influence the prevalences found. For example, in the UHDRS only one item refers to 'low mood', whereas the SCID uses the stringent DSM criteria for depression. These different methodologies limit the generalizability of the reported findings, which is further impaired by small sample sizes. Importantly, none of the seven studies found used a representative comparison group and, to our knowledge, no follow-up studies have been performed to relate the incidence of behavioral problems and psychiatric disorders to disease onset and disease progression.

Psychopathology
The prevalence of depressed mood in Huntington's disease, varying from 33% to 69%, may be comparable to that of anxiety, irritability, and apathy. The development of depressive symptoms in Huntington's disease could be a direct result of cerebral degeneration, for which several neuropathological mechanisms have been proposed proposed.^[20,21] Depression could, however, equally well be a psychological reaction to being at risk for Huntington's disease, having grown up in an insecure and harmful environment, and/or the awareness of disease onset.

Many studies have found that depressive symptoms precede the onset of motor symptoms,[^20-24] but no relation between the occurrence of depressive symptoms and disease duration has so far been reported.[^25] Depression may, however, negatively correlate to cognitive decline,[^26] which is possibly the result of concurrent decreasing illness awareness. Anxiety has not been a main research interest in patients with Huntington's disease. Nevertheless, 'worrying', which could be part of a generalized anxiety disorder (GAD), is often reported in Huntington's disease patients, although it is mostly limited to worries about Huntington's disease.[^27] Since no studies were found that systematically investigate the prevalences of different anxiety disorders, this should be an important focus for future research.

Irritability without a prior history of short temper occurs in most Huntington's disease patients, and seems to precede motor symptoms in mutation carriers.[^27-29] A tendency for irritability occurring more frequently in late stage patients whose neurological symptoms have been present for 6 to 11 years has also been described.[^29] This is confirmed by a cross-sectional observational study of 27 nursing home residents with Huntington's disease (disease duration 7 to 11 years), which reports aggression in one-third of all patients over a 3-day period.[^20]

We contend that increasing degeneration of the striatum and the orbito-frontal-subcortical...
circuit in Huntington’s disease contributes to the development of socially inappropriate behavior, which initially may be manifested as subtle irritability and, in late-stage Huntington’s disease, as aggressive behavior.41

Of all neuropsychiatric symptoms only apathy consistently appears to be positively related to disease progression.72,42,43 Apathy is also strongly related to the decline of everyday functioning and, once present, tends to persist or worsen.15 Damage to structures of the anterior cingulate-subcortical circuit has in particular been associated with motivational disorders, including apathy,42 which may also be the case in Huntington’s disease.44

The occurrence of obsessive and compulsive symptoms in Huntington’s disease is of particular interest because obsessive-compulsive disorder and Huntington’s disease possibly share a similar neuropathology of the basal ganglia and (orbito)frontostriatal circuits.45,46 Many Huntington’s disease mutation carriers show personality changes with mental inflexibility in early stages,45 possibly heralding future obsessive and compulsive symptoms. Though in certain families obsessive and compulsive symptoms have shown an early phenotype of Huntington’s disease,45 they are not often identified as a manifestation of Huntington’s disease.47

Psychotic symptoms usually occur when movement symptoms are already clearly manifest. This could explain why in earlier days, when Huntington’s disease was diagnosed at a later disease stage, psychosis was usually described as the main psychiatric feature of Huntington’s disease.48 Even so, Huntington’s disease patients were often misdiagnosed with dementia praecox or schizophrenia until the first half of the 20th century. Nowadays, rather low prevalences (3% to 11%) of psychotic symptoms are reported, which is most probably due to earlier and better diagnoses of Huntington’s disease and a shift in research from inpatient to outpatient populations.

**Recommendations for future research**

The causal pathways leading to psychopathology in Huntington’s disease are unclear and should receive priority on the research agenda. Since Huntington’s disease families with multiple cases of schizophrenia and schizotypal/schizophreniform symptoms have been described,49,50 as well as families with obsessive-compulsive disorders,49 and families with a high occurrence of affective syndromes in both mutation carriers and non-carriers,51 it is highly probable that other genes than the Huntington’s disease gene itself, as well as environmental factors, play a role in the development of psychopathology in Huntington’s disease.52 Previous findings indeed suggest that both neuropathology and environmental stress contribute to the occurrence of neuropsychiatric phenomena in Huntington’s disease: a case series among 37 Huntington’s disease patients and 167 relatives reported significantly more psychiatric admissions and diagnoses in patients than in their relatives.53 Thus at least some psychopathology will be due to the etiology of Huntington’s disease, though not solely the Huntington’s disease mutation. Since the same study showed that relatives of Huntington’s disease patients also had more psychiatric diagnoses and admissions than the general population, stressors such as growing up with an affected parent and with the uncertainty about one’s own disease status are also likely causes of psychopathology in Huntington’s disease.

To correct for environmental stress, prevalences of psychopathology in Huntington’s disease should be compared to healthy family members, particularly siblings of patients who do not carry the Huntington’s disease mutation, since they share the same psychosocial family background as mutation carriers. There is need for prospective research covering all different stages of Huntington’s disease with the use of such a comparison group, just as has been carried out before in pre-motor symptomatic mutation carriers.73,26 Though this comparison group cannot correct for all potential biases affecting research in this difficult population (e.g., self-selection for genetic testing and difficulties with the staging of disease progression), use of the proposed comparison group will significantly increase the validity and interpretability of the test results. Also, a comparison of psychopathology in Huntington’s disease with other neurodegenerative disorders, such as Parkinson’s disease, could increase our understanding of the pathological mechanisms that affect the brain and behaviors of these patients.73,16,24 Nevertheless, such a comparison should be explorative in nature as neither Huntington’s disease patients nor patients with Parkinson’s seem an adequate comparison group to the other.

As the Huntington’s disease mutation does not have a full penetrance for developing specific behavioral problems or psychiatric disorders, research should also focus on the contribution of other biological factors, as well as environmental factors, to the behavioral phenotype of Huntington’s disease. Identification of endophenotypes, which do not depend on what is obvious to the unaided eye, could help to resolve questions about etiological models.51 Such an endophenotype-based approach has the potential to assist in the genetic dissection of psychopathology. These endophenotypes should be researched on the level of neurobiology, neuropsychology, and neuroendocrinology. An example of a possible neurobiological endophenotype is disturbance of the hypothalamic pituitary adrenal (HPA) axis functioning with hypercortisolism; the stress hormone cortisol plays a major role in psychiatric disorders, particularly depressive disorders that have a high prevalence in Huntington’s disease. Disturbances in HPA-axis functioning have already been found in Huntington’s disease patients but have not yet been linked to behavioral or psychiatric morbidity in Huntington’s disease.54,57 Another possible endophenotype is dysfunction of the immune system,58 with altered secretion of cytokines. These have also been related to the presence of depression.59 Vulnerability to psychopathology may be determined by genetic polymorphisms of the HPA-axis and the immune system, which is another important area of research in Huntington’s disease.53

All future research should improve upon current methodologies. Some potential sources of current variation in test results, such as low incidence of Huntington’s disease with resulting small sample sizes and self-selection for testing and research, are difficult to avoid; researchers should be aware of this. Other sources of variety, mainly differences in methodology, can be eliminated if consensus on terminology and staging methods, as well as standardization of instruments are achieved. This is a necessary requirement for the comparability and generalizability of results. We propose that DSM criteria are used as the gold standard for psychiatric diagnosis, and standardized instruments for other neuropsychiatric symptoms, such as apathy and irritability.

For disease progression, we propose the motor section of the UHDRS. Although the motor score is not perfectly correlated with disease progression, a functional assessment with the Total
Functional Capacity scale of the UHDRS, which is related to disease progression, is not a good tool for this kind of research, as it is directly influenced by psychopathology. Because the onset of Huntington’s disease is so gradual, disease duration is also not an adequate measure, and motor assessment is therefore the most objective, reliable, and comparable method of disease staging for research.

Research should lead to an increased understanding and recognition of psychopathology in Huntington’s disease and its causes. This is essential for adequate treatment of those symptoms that could improve the overall functioning and quality of life of the Huntington’s disease patient and his or her direct environment.

References
Chapter 3

Cross-sectional study on prevalences of psychiatric disorders in mutation carriers of Huntington’s disease compared with mutation-negative first-degree relatives

E. van Duijn, E.M. Kingma, R. Timman, F.G. Zitman, A. Tibben, R.A. C. Roos, R.C. van der Mast

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