Attitudes of neurologists, psychiatrists, and psychotherapists towards predictive testing for Huntington’s disease in Germany

U Thies, B Bockel, V Bochdalofsky

Abstract
Predictive testing for Huntington’s disease (HD) in Germany is performed by genetic counsellors, neurologists, psychiatrists, and psychotherapists. In order to evaluate the attitudes of neurologists, psychiatrists, and psychotherapists in Germany towards predictive testing for HD, a postal questionnaire was sent to this group. Two German Bundesländer were chosen, Baden-Württemberg (BW) and Niedersachsen (NS). Of 469 persons interviewed the response rate was 32.6%. The questionnaire consisted of 17 items assessing sociodemographic data, acquaintance with HD patients, lay organisations, attitudes towards genetic counselling, presymptomatic and prenatal DNA testing, and reproduction of persons at risk for HD.

More than 70% of the subjects were well informed about predictive DNA testing but knowledge about the details of the test procedure, especially the World Federation of Neurology (WFN) and International Huntington Association (IHA) recommendations, was quite low (11.8%). Nevertheless, the majority would recommend predictive testing for HD although they anticipated problems for the probands. The majority of our respondents favoured psychological test and post-test counselling for those tested. Concerning reproduction, most subjects favoured prenatal testing or that persons at risk should refrain from having children. We found that the opinions of practitioners and at risk persons differed with respect to the predictive DNA test and, particularly, to prenatal testing. Therefore the testing procedure could be improved if practitioners were better informed about the DNA test in general and about the attitudes and wishes of their patients.

Huntington’s disease (HD) is an autosomal dominant, neurodegenerative disease with late onset that results in a progressive movement disorder accompanied by psychiatric alterations and cognitive impairment. It affects 1 in 10 000 persons in most European populations. The onset of HD occurs in or around the fourth decade of life and leads to death within 15 to 20 years. Although the gene has now been identified, the biochemical basis of the disease is not yet understood and there is no effective treatment in delaying or preventing the onset or progression of the disease.

In 1983 Gusella et al. localised the genetic defect to the tip of the short arm of chromosome 4. After finding close genetic linkage between the G8 probe and the HD gene, predictive molecular testing became available for at risk persons from HD families. Until the identification of the HD gene, which was published in March 1993, only an indirect approach was available to predict whether or not persons at risk were gene carriers. The error rate of the indirect approach with closely linked polymorphic markers was around 2%.

In Germany there are over 8000 persons affected with HD. Presymptomatic and prenatal testing for at risk persons was started in 1989 at the Institute of Human Genetics, Göttingen. Predictive testing in HD raises ethical and moral questions. Therefore predictive testing was performed within a structured protocol, as follows. The recommendations for predictive testing published by the International Huntington Association (IHA) and the World Federation of Neurology (WFN) are part of our test procedure. Furthermore, the cooperation of the proband, genetic counsellor, neurologist, psychiatrist, and psychotherapist is required. For the indirect approach it is essential to have the diagnosis of HD confirmed by at least one neurologist/psychiatrist. Clinical examination of the proband is necessary to exclude early signs of the illness, because the goal of predictive testing is to ascertain the level of risk of developing HD in currently healthy subjects. In probands with serious risk of suicide or other psychiatric symptoms the test is postponed. The test result is only given to the proband if professional support in the test and post-test phase is ensured. The test result is disclosed in a session with the genetic counsellor, the psychotherapist, and the proband when all three parties agree. Up to February 1993 in our Institute 52 indirect predictive and 10 prenatal tests had been performed.

The direct test approach now available will revolutionise predictive testing in subjects at risk for HD and make it possible in families in which the family structure was unsuitable for indirect testing or where the diagnosis of HD remains uncertain. Nevertheless, the above mentioned problems associated with preclinical testing for a late onset disorder will remain and therefore the current guidelines for testing are also valid for the future.

Whereas a large number of surveys concerning the attitudes of persons at risk towards
predictive testing have been performed, there have only been a few studies on the attitudes of professionals to presymptomatic testing in HD.

In the present study we tried to assess the knowledge and attitudes of neurologists, psychiatrists, and psychotherapists in Germany towards predictive DNA testing. The results of the study suggest that knowledge of this group in specific areas needs to be enlarged. Along with genetic counsellors, this group is mainly involved in the test procedure and the decision making process of the probands. Therefore we tried to find out if this group would support at risk persons in the test and post-test phases.

Sample composition and methods

Subjects

Questionnaires were sent to 469 neurologists, psychiatrists, and psychotherapists from Baden Württemberg (BW) (256) and Niedersachsen (NS) (213) listed in the Ärztebuch (medical directory) 1990.

Questionnaire

The questionnaire consisted of 17 questions assessing sociodemographic data, acquaintance with HD patients, lay organisations, knowledge about recent developments in HD research, attitudes towards genetic counselling, presymptomatic and prenatal testing, and reproduction of persons at risk for HD. The questions were closed or open ended (appendix). The anonymous questionnaire was accompanied by a personal letter giving a minimum of information about prenatal and presymptomatic DNA testing which allowed those who were not well informed about the disorder to answer the questions.

Statistical analysis

Using the $\chi^2$ test or Fisher’s exact test, statistical analyses were carried out with the SAS statistical package. The Bonferroni multiple testing procedure was used in all cases. In the following sections the values are only mentioned if the results are significant.

Results

Respondents

Replies were received from December 1991 to March 1992; 153 questionnaires were returned, 56 from Niedersachsen (return rate 26.3%) and 97 from Baden Württemberg (return rate 37.9%). The returns were identified by the postmark. No apparent reason exists for the lower return rate from NS. The respondents’ qualifications in the different areas of neurology, psychology, and psychotherapy are shown in table 1. There are no differences in their answers and 127 (83%) stated that they had either counselled or treated at least one patient with HD (median 2, 25% quantile 1 and 75% quantile 4).

<table>
<thead>
<tr>
<th>Qualification</th>
<th>No</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neurologist</td>
<td>16</td>
<td>10.5</td>
</tr>
<tr>
<td>Psychiatrist</td>
<td>1</td>
<td>0.7</td>
</tr>
<tr>
<td>Psychotherapist</td>
<td>10</td>
<td>6.5</td>
</tr>
<tr>
<td>Neurologist/psychiatrist</td>
<td>65</td>
<td>42.5</td>
</tr>
<tr>
<td>Neurologist/psychologist</td>
<td>4</td>
<td>2.6</td>
</tr>
<tr>
<td>Psychiatrist/psychologist</td>
<td>4</td>
<td>2.6</td>
</tr>
<tr>
<td>Neurologist/psychiatrist/psychologist</td>
<td>51</td>
<td>33.3</td>
</tr>
<tr>
<td>No information</td>
<td>2</td>
<td>1.3</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>153</td>
<td>100</td>
</tr>
</tbody>
</table>

Table 2: Circumstances under which the predictive test is recommended (answered by 44% of the respondents)

<table>
<thead>
<tr>
<th>Circumstances</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Family planning</td>
<td>94%</td>
</tr>
<tr>
<td>Sake of partnership</td>
<td>64%</td>
</tr>
<tr>
<td>Psychological problems</td>
<td>57%</td>
</tr>
<tr>
<td>Personal lifestyle planning</td>
<td>14%</td>
</tr>
<tr>
<td>Other reasons (including answers: only after identification of the gene, only in case of a cure)</td>
<td>18%</td>
</tr>
</tbody>
</table>

Table 3: Anticipated problems of predictive testing (answered by 86% of the respondents)

<table>
<thead>
<tr>
<th>Problems</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Psychological problems</td>
<td>93%</td>
</tr>
<tr>
<td>No protection for the privacy of personal data</td>
<td>92%</td>
</tr>
<tr>
<td>Problems with social life (friends, etc)</td>
<td>89%</td>
</tr>
<tr>
<td>Problems with insurance companies</td>
<td>89%</td>
</tr>
<tr>
<td>Family problems</td>
<td>83%</td>
</tr>
<tr>
<td>Risk of suicide</td>
<td>81%</td>
</tr>
<tr>
<td>Other problems</td>
<td>14%</td>
</tr>
</tbody>
</table>

Lay organisations

Addresses of lay organisations (Deutsche Huntington Hilfe e.V and Deutsche Huntington Gesellschaft e.V) were only known by 41 respondents (26.8%), but 82 (53.6%) would be willing to arrange contact between at risk persons and lay organisations. The willingness to arrange contact was different between Baden Württemberg (46 respondents, 47.4%) and Niedersachsen (36 respondents, 64.3%).

Knowledge about predictive testing

A total of 112 respondents (73.2%) knew that presymptomatic DNA diagnosis is possible, 89 (58.2%) knew that for the indirect approach the whole family must be analysed, and 18 (11.8%) were familiar with the WFN and IHA recommendations for predictive testing for HD. Eight (5.2%) respondents stated that they would counsel at risk persons themselves, 106 (69.3%) would forward these persons to a genetic counselling unit, and 37 (24.2%) would cooperate with a genetic counsellor (not answered 3, 1.3%). It turned out that knowledge of the details of the test procedure of those who counsel by themselves was insufficient. Sixty-eight (44.4%) would recommend predictive DNA testing for at risk persons under certain circumstances (table 2), whereas 26 (16.9%) would recommend testing in all cases. One respondent would not recommend testing at all. The answer rate to this question was low because the subjects who chose to refer probands directly to genetic counselling units were not asked to answer this question (58, 37.9%). Concerning predictive testing and the related problems, which are listed in table 3, 132 (86.3%) physicians anticipated
Table 4 Opinions about reproduction and prenatal testing in HD.

<table>
<thead>
<tr>
<th></th>
<th>Yes (%)</th>
<th>No (%)</th>
<th>No answer (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;Probands who are given a raised risk of developing HD should not have children&quot;</td>
<td>80-4</td>
<td>12-4</td>
<td>7-2</td>
</tr>
<tr>
<td>&quot;Would you approve of prenatal testing in the case of a pregnancy of an at risk person?&quot;</td>
<td>88-9</td>
<td>7-2</td>
<td>3-9</td>
</tr>
<tr>
<td>BW (56)</td>
<td>49 (87-5)</td>
<td>5 (8-9)</td>
<td>2 (3-6)</td>
</tr>
<tr>
<td>NS (56)</td>
<td>45 (87-6)</td>
<td>6 (8-2)</td>
<td>6 (6-2)</td>
</tr>
</tbody>
</table>

problems whereas 11 (7-2%) did not (not answered 10, 6-5%).

TEST PROCEDURE
Psychotherapeutic support for persons at risk who want predictive testing was favoured by the majority of the respondents and about 94-8% (145) of the subjects favoured post-test counselling. However, 82 (53-9%) were of the opinion that it is only necessary in cases of a raised risk, whereas 60 (39-2%) thought that post-test counselling should take place irrespective of the test result. Two (1-3%) respondents considered post-test counselling unnecessary (not answered 9, 5-9%).

REPRODUCTION AND PREGNATAL TESTING IN PERSONS AT RISK
Concerning reproduction and prenatal testing (table 4), 123 (80-4%) respondents thought that persons at risk should refrain from having children of their own (not answered 11, 7-2%). The majority (136, 88-9%) was in favour of prenatal DNA testing if one parent is an HD gene carrier, whereas 11 (7-2%) rejected prenatal molecular testing (not answered 6, 3-9%). A total of 35-9% (55) would tolerate an abortion only in those at 98% risk, another 51% (78) would justify it at 50% risk, and 9-8% (15) would refuse an abortion at all (no answer 5, 3-3%).

Discussion
Predictive testing in HD raises ethical, legal, social, and psychological questions, which are well known by neurologists, psychiatrists, and psychotherapists. Our survey showed that they are aware that genetic testing for this late onset, incurable disease needs special support and counselling for at risk persons.

In general it turned out that the respondents are well informed about predictive testing and DNA analysis in HD. Most of the physicians know about the possibility of DNA testing in HD (73-2%). This rate is higher than in previous studies; for example, only 24% of Scottish practitioners were familiar with the test. However, the Scottish study was performed two years ago and practitioners are now, in general, better informed about DNA testing. A Dutch study performed in 1992 showed that 59% of general practitioners (GPs) who had persons at risk as patients were familiar with the test. However, details of the test procedure were not well known by our respondents; 58-2% of the physicians knew that predictive testing at the time was only possible by family study (indirect approach). Only a few (11-8%) were aware of the international guidelines of the IHA and WFN. This might explain the general willingness of the respondents to cooperate with a genetic counselling unit instead of counselling HD persons by themselves. However, the knowledge of the eight physicians who would not cooperate with genetic counselling units was less than the other respondents. We think that genetic counselling and especially DNA testing counselling should remain the responsibility of genetic counselling units.

The main reasons given by the respondents for recommending predictive testing to at risk persons were family planning, the sake of partnership, and psychological reasons. At risk persons would take the test mainly because they want more certainty about their carrier status, family planning, for example, has a lower priority in their opinion. Furthermore, the number of at risk persons opting to take part in the test procedure is lower than the recommendations of professionals to take part in the test procedure. This shows that the decisions of at risk persons are not always influenced by their physicians. The different ranking of the reasons for taking part in the test might result from the more general view of practitioners and from the fact that they are not personally involved. The difference in the attitudes of physicians and at risk persons is even more pronounced regarding the problems which may arise during or as a consequence of the test procedure. Most respondents (more than 86%) believe that many problems might arise, such as risk of suicide, psychological and social problems, problems with insurance companies, and lack of confidentiality. Most at risk persons do not rank psychological problems or the risk of suicide highly, whereas the other problems do rank highly in their opinion. The physicians seem to be well aware of the fact that persons who received raised risks were more shocked than they themselves had expected.

Willingness of the respondents to give support after disclosing the test result was not as great as expected (40%). The majority (54%) thought that it was only necessary in the case of an increased risk. In fact, even persons with a decreased risk have great difficulty in coping with this information and need professional support, as the surveys of Tibben et al and Huggins et al indicated.

There is an opinion contained in German textbooks for neurologists that persons at risk for HD should refrain from having children and about 80% of our respondents would agree with this. Only a few of them believe that the decision whether to have children should be left completely to the persons at risk and their partners. These answers suggest that the physicians want to eradicate the disease.

About 90% of our respondents, the same rate as in the Dutch study, approved of prenatal testing for HD and termination of a high risk pregnancy. In opposition to that finding, only 29-4% of at risk persons would use prenatal testing, as shown in the Vancouver study. In the Dutch study the number of at
Another area for improvement will be to keep the manuscript for advice and helpful discussion, questionnaires. We also thank Professor W Engel for helpful discussion concerning the questionnaire. A novel gene containing a trinucleotide repeat that is expanded and unstable on Huntington’s disease chromosomal. Nature 1983;306:234-8.


Appendix  Questionnaire.

(1) Are you a practising 
Neurologist  Psychiatrist  Psychotherapist 
Private practice 
Doctor in a university hospital 
Doctor in a district hospital 
Doctor in a psychiatric hospital 
(2) Have you counselled patients with HD? 
Yes  (number)  No 
(3a) Are you familiar with HD lay organisations? 
Yes  No 
(3b) Do you know addresses of lay organisations and would you make contact with them? 
Yes  No 
(4) Were you aware of the possibility of DNA diagnosis for HD? 
Yes  No 
(5) Are you aware that DNA diagnosis is only possible within a family investigation? 
Yes  No 
(6) If an at risk person wishes to have presymptomatic DNA testing carried out, would you: 
Counsel them yourself? 
Send them to a genetic counselling unit? 
(7) If you carry out the counselling yourself: 
Would you recommend presymptomatic DNA diagnosis to persons at risk for HD? 
No 
I would recommend DNA diagnosis only in particular circumstances, which are: 
If the person wishes to have children 
If risk modification is important for the partnership 
If risk modification seems to be important for physical reasons 
For professional reasons 
For the following reasons: 
Only when a cure becomes available 
Only when, once the gene has been identified, investigation of at risk patients is possible without reference to other family members 
(8) Do you believe that, as a result of participation in presymptomatic DNA testing, problems could arise which would require special attention? 
Yes  No 
If yes, which problems do you believe would arise? 
Negative influence on the family 
Negative psychological effect on the patient 
Increased risk of suicide 
Negative social reaction (loss of job, stigma of being 'sick', etc) 
Problems with health and life insurance 
No protection of the privacy of personal data 
Other problems 
(9) Are you acquainted with the WFN (World Federation of Neurology) and IHA (International Huntington Association) recommendations concerning DNA diagnosis for HD (see J Neurol Sci 1989;94:327–32) 
Yes  No 
(10) DNA diagnosis should only be carried out in laboratories which work in cooperation with a genetic counselling unit and with the neurologist/psychiatrist/psychotherapist 
Yes  No 
DNA diagnosis could also be carried out by a commercial laboratory 
Yes  No 
(11) Should at risk persons who are taking part in a DNA diagnosis programme receive counselling from a neurologist/psychiatrist/psychotherapist with regard to the test results 
Yes  No 
(12) Should the appointment for the test result be set by the supervising doctor? 
Yes  No 
(13) The test result should only be given to the patient in the presence of the supervising doctor (neurologist/psychiatrist/psychotherapist) and the genetic counsellor 
Yes  No 
(14) After receiving the test result, should the proband be looked after by the supervising doctor? 
Only if the proband receives a high risk 
In every case 
Yes  No 
(15) Do you believe that probands who receive a high risk should abstain from having children? 
Yes  No 
(16) One parent carries the gene for HD. Every child has a risk of 50%. In case of a pregnancy the risk can be more accurately determined by molecular diagnosis (2% or 98%). Do you think prenatal diagnosis is justified in this case? 
Yes  No 
(17) Do you think an abortion is justifiable? 
In the case of a 50% risk 
In the case of a 98% risk 
No