Frequency of the Tay-Sachs disease splice and insertion mutations in the UK Ashkenazi Jewish population

Eleanor C Landels, Ian H Ellis, Anthony H Fensom, Peter M Green, Martin Bobrow

Abstract

Tay–Sachs disease is a lethal neurodegenerative disorder caused by deficiency of the lysosomal enzyme β-hexosaminidase A and inherited in an autosomal recessive fashion; carriers of the disease are 10 times more frequent in the Ashkenazi Jewish community than in the general population. Over 90% of North American Ashkenazi carriers tested have been shown to have either a splice site mutation at the boundary of exon 12 and intron 12 in the β-hexosaminidase α subunit gene, or a 4 base pair insertion in exon 11. We describe simple assays involving amplification of DNA across these two mutation sites by polymerase chain reaction and the results of screening 75 subjects are given.

The frequencies of the splice and insert mutations in 41 UK Ashkenazi carriers (20% to 80%, respectively) were similar to those found in the North American community. Twelve Ashkenazi subjects classified as non-carriers on enzyme assay were found to be negative for both mutations tested. All Ashkenazi carriers tested (both obligate carriers and those picked up by population screening) had either the splice or insert mutations; in contrast to this, only 21% of the non-Ashkenazi carriers had one or other of these mutations.

It is concluded that within the carrier screening programmes for the Ashkenazi community, assays for the splice and insert mutations, together with an assay recently described for a mutation causing the rarer adult onset form of the disease, will prove useful as confirmatory tests for subjects who give positive or borderline results when screened on enzyme assay.

Tay–Sachs disease (TSD), one of the most severe forms of the GM2–gangliosidoses, is an incurable neurodegenerative disorder inherited in an autosomal recessive fashion. Damage to the nervous system is caused by accumulation of ganglioside GM2, the metabolism of which is blocked because of deficient activity of the lysosomal enzyme β-hexosaminidase A (Hex A). The cause of TSD is mutation in the gene, located on chromosome 15, for the α subunit of Hex A. Carriers of the TSD gene, who have reduced Hex A activity, are estimated to have a frequency of 1/167 in the general population. However, in the Ashkenazi Jewish population TSD carriers are approximately 10 times more frequent and so carrier screening programmes, based on measuring Hex A activity in serum or leucocyte samples, have been made available to Ashkenazi Jewish communities, with the aim of reducing the incidence of TSD births.

The hexosaminidase α subunit gene has been cloned and several mutations causing various GM2–gangliosidoses have been characterised. In the Ashkenazi community there appear to be two mutations accounting for over 90% of TSD carriers. A splice junction mutation of a G to C transition in the first nucleotide of intron 12, fortuitously creating a DdeI site, was present in 11 of 40 obligate TSD carriers. The more common mutation, found in 14 of 20 obligate TSD carriers, is a 4 bp insertion in exon 11. Assays have been described for these two mutations, based on the polymerase chain reaction (PCR), which involve radiolabelling during DNA amplification or the use of radiolabelled oligonucleotides of mutant or normal sequences to probe patient DNA. More recently, Triggs–Raine et al have used amplification of DNA by PCR followed by restriction enzyme digestion to assess the incidence of the splice and insert mutations in the North American Ashkenazim. They also screened this population for the Gly to Ser mutation in exon 7 which causes adult onset GM2–gangliosidosis.

We have independently developed similar assays for carriers of the splice and insert mutations, avoiding radioisotope labelling, based on amplification of DNA by PCR followed by digestion with DdeI for detection of the splice mutation (restriction enzyme digestion is not necessary to detect the insert mutation.) DNA fragments were then separated by electrophoresis and viewed directly under UV light. Here we present data on the frequency of these two mutations in the UK Ashkenazi community. We have

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Received for publication 27 July 1990.
Revised version accepted for publication 20 September 1990.
tested 75 subjects, including both Jewish and non-Jewish obligate carriers for TSD, and subjects picked up by population screening programmes whose carrier status has been determined by enzyme assay.

Materials and methods
Population screening was based on measuring percentage Hex A in serum and leucocyte samples. The assay was essentially as described by Kaback but adapted to be carried out on a centrifugal analyser.13

DNA AMPLIFICATION
Approximately 100 ng of genomic DNA, isolated by phenol-chloroform extraction using a 340A Nucleic Acid Extractor (Applied Biosystems) from blood samples (whole EDTA blood or white blood cell pellets, stored at −20°C), were amplified by the PCR method.14 The oligonucleotide primer sequences used to amplify the region of the splice mutation were as published previously.6 For the region of the insertion mutation we designed primer sequences based on the published cDNA sequence5: TS1–5′: 5′–TTCCGGG-CCTTCTCTTGCC–3′ and TS1–3′: 5′–TTCAAA-TGCCAGGGTTCCAC–3′. The reaction mix contained 2-5 units of Taq polymerase (Amplitaq, Cetus), 0-5 mmol/l dNTPs, 300 ng of each primer, 67 mmol/l Tris-HCl, pH 8-8, 16-6 mmol/l (NH₄)₂SO₄, 6-7 mmol/l MgCl₂, and 170 µg/ml bovine serum albumin.

The reaction was carried out in a Perkin–Elmer Cetus thermal cycler for 35 cycles of one minute at 93°C, three minutes at 70°C. For detection of the splice mutation, products of the PCR were then digested with DdeI at 37°C for one hour. The products of reactions were electrophoresed in 12% polyacrylamide gel. There is also a further band which is probably a heteroduplex of normal and mutated DNA.15 The two types of DNA will pair exactly except for the 4 bp insert which probably loops out, causing the heteroduplex to migrate anomalously. This 'hybrid' band is reproducible and straightforward. Fig 2, track 1 shows the 96 bp DNA amplified in a non-carrier; heterozygotes for the insert will have both a 96 bp band and one of 100 bp (fig 2, track 2). The 4 bp difference is resolvable in a 12% polyacrylamide gel. There is also a further band which is probably a heteroduplex of normal and mutated DNA.15 The two types of DNA will pair exactly except for the 4 bp insert which probably loops out, causing the heteroduplex to migrate anomalously. This 'hybrid' band is reproducible and straightforward.

Results
The results of the splice mutation assay are shown in fig 1. Track 1 shows the 135 bp DNA band amplified by PCR. After digestion with DdeI, non-carrier DNA is reduced in size to a 120 bp fragment and two smaller fragments which are not seen as they run off the gel, shown in track 3. Carriers of the splice mutation have an extra DdeI site and so after digestion have four DNA fragments, 85 bp, 35p, and the two smaller fragments, of which only the 85 bp fragment is seen (track 2). As well as the 85 bp fragment, the unmutated DNA of the normal allele has also been amplified and on digestion with DdeI gives the 120 bp fragment (also visible in track 2).

Detection of the 4 bp insert mutation is also

![Figure 1](http://jmg.bmj.com/)

**Figure 1** Detection of the splice site mutation: normal and heterozygote DNA patterns after amplification by PCR and DdeI digestion. (1) Amplified DNA (before DdeI digestion), (2) heterozygote DNA, (3) normal DNA.

![Figure 2](http://jmg.bmj.com/)

**Figure 2** Detection of the 4 bp insertion mutation: normal and heterozygote DNA patterns after amplification by PCR. (1) Normal DNA, (2) heterozygote DNA.
Results of screening 75 subjects for the splice and insertion mutations.

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<tr>
<td>Non-Jewish</td>
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makes detection of heterozygotes for the insert mutation very easy. The results of screening various subjects using these two assays are shown in the table.

Discussion

The frequencies of the splice and insertion mutations among UK Ashkenazi TSD carriers tested so far (20% and 80% respectively) are similar to those found in the North American community. In the most recent study, TSD carriers picked up by population screening were also tested for the Gly\(^{69}\) to Ser mutation in exon 7 of the Hex A α subunit gene, which causes adult GM\(_2\)-gangliosidosis; five out of 216 carriers had this mutation. We studied 26 Ashkenazi carriers picked up through population screening. If the frequency of the exon 7 mutation in the UK Ashkenazi population is similar to that in the North American Ashkenazim, we would only expect to find 0.6 carriers to be negative for both the splice and insert mutations; thus it is not significant that none was found.

Twelve Ashkenazi Jewish subjects classified previously by enzyme screening as non-carriers were negative for both the splice and the insert mutations. Three subjects, two of whom were Ashkenazi, who repeatedly gave borderline results on enzyme assay, were also negative for both mutations. Work is in progress to test these subjects for the exon 7 mutation.

In the study of North American Ashkenazim, 39 of 216 carriers picked up by population screening were found not to carry the splice, insert, or exon 7 mutations, an apparent false positive rate of 18%. Assuming a similar false positive rate, we would expect to have found four or five of the 26 carriers picked up by population screening to be negative for both the splice and insert mutations; we found none. We feel this almost certainly reflects differences in the enzyme assays of different testing laboratories.

All 41 Ashkenazi carriers (as classified by enzyme assay of serum and leucocyte samples) tested were positive either for the splice or insert mutations. In contrast to this, only 21% of the 19 non-Jewish carriers tested proved positive for one or other of these mutations; 5-3% had the splice mutation and 15-8% had the insert. The remaining 79% presumably carry unknown mutations.

Although in this study 100% of Ashkenazi carriers tested have had either the splice or insert mutation, other workers have found an Ashkenazi TSD patient who was heterozygous for the insert and an unknown mutation. It is thought that Hex A α subunit mutations other than the splice and insert occur at the same low frequency in Ashkenazim as in the general population. Thus, while it can be envisaged that these assays, together with the assay for the exon 7 mutation, will be useful in addition to the enzyme assay, if used alone for carrier screening within the Ashkenazi community approximately 2% of carriers will be missed.

At present the need to extract DNA before PCR amplification adds considerably to the time and cost of these assays. We are further developing the assays to allow amplification by PCR directly on DNA released from various tissues, avoiding the DNA extraction step, as has been done for blood and cells collected using clinical swabs. Another way of reducing the cost of the assays may be to combine them, so that the regions of both the insert and splice mutations are amplified in the same reaction mixture; this should be possible since the same temperature cycles are used for both sets of primers. The resulting products, or an aliquot of them, would then be digested with DdeI to show the splice mutation. A disadvantage is that the band patterns would be more complex but the convenience of testing for both mutations at the same time should outweigh this.

In conclusion, an optimum strategy might be to screen on the basis of serum Hex A levels, using PCR assays to confirm all borderline and positive results, with leucocyte enzyme assays held in reserve for problematic cases.

We are grateful to Roland Roberts for helpful discussions and primer sequence design skills, Simon Dear and Suzanne Clark for help with DNA extraction, Adrienne Knight for help in preparing the manuscript, and to the British Tay–Sachs Foundation for generous financial support.


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doi: 10.1136/jmg.28.3.177

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