A single origin for the most frequent mutation causing late infantile metachromatic leucodystrophy

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Abstract

Metachromatic leucodystrophy is an autosomal recessive degenerative disease of the nervous system caused by the deficiency of the lysosomal enzyme arylsulphatase A (ARSA). We report here on the high incidence of late infantile MLD among Muslim Arabs originating from Jerusalem, most probably because of a founder effect. All the patients were found to be homozygous for 459 + 1 G→A, a mutation which destroys the splice donor site of exon 2 of the ARSA gene. This mutation has been reported to be the most common mutation causing MLD.

We studied the ARSA haplotype defined by three intragenic polymorphic sites in DNA samples from Muslim Arab patients from Jerusalem, a Christian Arab patient originating from the region, and eight other white patients, all homozygous for the 459 + 1 G→A mutation. All the alleles carried the same haplotype which is in complete linkage disequilibrium with the mutation. This finding indicates a common origin for the 459 + 1 G→A mutation which may have been introduced into Jerusalem at the time of the Crusades.

Material and methods

SCREENING FOR THE 459 + 1 A→G MUTATION

During the last 15 years 27 patients have been diagnosed as being affected with late infantile MLD in the Department of Human Genetics at Hadassah Medical Centre. The diagnosis was based on clinical investigations and deficiency of ARSA in leucocytes and fibroblasts. These 27 patients originated from 17 unrelated families, four Jewish and 13 Arabs. Three of the Muslim Arab patients originated from Jerusalem. This represents a very high incidence, since in this period fewer than 50 000 Muslim Arab children were born in Jerusalem.

DNA samples from these patients were tested for the 459 + 1 G→A mutation using an MvaI polymorphism which is abolished by the mutation. The primers used for the amplification were 5' AGC CCG TGC CAG TGG AGG AG 3' and 5' CAA CAG TGG GAT GGG GAC 3'. The cycling conditions were 94°C for 30 seconds, 56°C for 30 seconds, 72°C for 1 minute for 30 cycles with an extension in the last cycle at 72°C for 7 minutes. The amplified fragment of 350 bp was digested with MvaI according to the manufacturer's recommendations, and the products were analysed on a 2% agarose gel (figure).

DETERMINATION OF INTRAGENIC ARSA HAPLOTYPES USING THREE POLYMORPHISMS

Three intragenic polymorphic sites have been described within the ARSA gene: a BglI site in exon 3, a BsrI site in exon 7, and a BamHI site in intron 7. The polymorphisms were determined as previously described after the amplification of the ARSA gene in two fragments (Zlotogora et al, submitted). In a previous study we showed that the PD allele is in complete linkage disequilibrium with the haplotype (BglI (2), BsrI (2), BamHI (1)) and that the mean frequency of each of the polymorphisms in
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Results
Among the 27 patients with late infantile MLD originating from 17 unrelated families, four unrelated Arab patients (three Muslims and one Christian) were found to be homozygous for 459+1 G→A. This was determined by DNA analysis in three patients, while since no DNA was available from another unrelated Muslim Arab patient, both parents were examined and found to be heterozygous for the mutation.

All the three Muslim Arab patients originated from the same small village, which relatively recently has been included in the municipality of Jerusalem. Each of the patients was born to first cousin parents, and no relationship was known between the different families. The fourth patient was a Christian Arab from a small town situated a few kilometres from Jerusalem.

Statistical Analysis
In the analysis of the results the $\chi^2$ test was used.

Discussion
Late infantile MLD is a relatively rare disease and therefore, even though 459+1 G→A is the most frequent mutation, causing the disorder, it is still rare among white populations. This is the first report of a community with a high incidence of this mutation.

The three Muslim Arab patients homozygous for the mutation 459+1 A→G originated from a small village which was relatively recently included within the municipality of Jerusalem. In this part of the Middle East, most of the villages were founded by only a few persons and marriages are by tradition within the family; first cousin marriages are, by preference, the most frequent. Therefore, even though the families of the patients do not know of any relationship between them, it may be assumed that they have common ancestors and that the mutation originates from a common founder. On the other hand, the fourth patient who was from a close geographical region was born into a Christian family. Since inter-religious marriages are very rare, a common founder is improbable. In addition, since there are only a few thousand original inhabitants of the village, the incidence of the mutation among them is probably very high. However, there have been many changes in the distribution of the population since the village was included within the municipality of Jerusalem and the mutation probably began to spread among Muslims. In order to determine the frequency of the mutation, we plan a screening programme of
this population at risk. Such screening will also allow appropriate genetic counselling.

When first reported, the 459 + 1 G → A allele was completely sequenced in one compound heterozygote patient and the haplotype was found to be (BglI (1), BamHI (1), BsrI (1)). Therefore in 21 alleles (the allele sequenced and 20 alleles from 10 homozygotes) with the 459 + 1 G → A mutation, there was complete linkage disequilibrium between the allele carrying the mutation and a haplotype defined by three different intragenic polymorphic sites. This haplotype (BglI (1), BamHI (1), BsrI (1)) is rare in the general population (3-9% of alleles) and the linkage disequilibrium is highly significant. The finding of a single haplotype in all the 459 + 1 G → A alleles suggests a common origin for all the carriers. Since both Muslim and Christian patients have been diagnosed in the region of Jerusalem, this may suggest that the mutation was introduced in Jerusalem at a period when both migration and religious conversion were frequent. For instance, the mutation may have been introduced from Europe at the time of the Crusades in the 11th and 12th centuries. Further knowledge of the distribution of the mutation is needed in order to understand its origin better.

Three other mutations which cause late infantile MLD have been reported to be frequent, each in a particular population. The G99D mutation was found in eight out of 12 alleles from Japanese patients (66% of mutant alleles) but up to now has not been found in white populations. In Australia a mutation, T274M, was found in the homozygous state in six patients of Lebanese origin. The same mutation was also found in two Israeli Christian Arabs from the Galilee and most probably has a common origin in Lebanon (in preparation). The third mutation, L377P, which occurred on the PD allele background, was found to be very frequent because of a founder effect in an isolate, the Habbanite Jews (17% carrier frequency), and relatively frequent among the Yemenite Jews who were living in the same geographical region (Zlotogora et al, submitted). The other mutations causing late infantile MLD which have been reported so far are rare or unique.

From the analysis of the different mutations causing late infantile MLD it seems that the difference between the mutation 459 + 1 G → A and the other relatively frequent mutations is that 459 + 1 A → G first occurred in Europe, a continent in which and from which migration has been frequent. The other mutations occurred in populations which have been, up to recently, relatively isolated and therefore have not yet spread among other populations.

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