Frequency of myotonic dystrophy gene carriers in cataract patients

A M Cobo, J J Poza, A Blanco, A López de Munain, A Saénz, M Azpitarte, J Marchessi, J F Martí Massó

Abstract
DNA samples from 231 unselected patients with cataracts were studied to determine the frequency of the DM mutation in cataract patients. A previous epidemiological study established a high prevalence of DM in the population of Guipúzcoa (Basque Country, Spain), 26.5 cases/100,000. We have found two carriers (0.9%) of the DM mutation in patients who are not related to any previously known DM family. The screening of the DM mutation in cataract patients should be restricted to young patients or people with multicoloured and iridescent opacities, in which the risk of carrying the DM premutation could be higher. Our results suggest that subjects with 38 to 80 repeats could constitute the genetic reservoir of the DM mutation. (J Med Genet 1996;33:221-223)

Key words: myotonic dystrophy gene; cataracts.

In the absence of any evident clinical features, identification of typical subcapsular opacities in subjects at risk of myotonic dystrophy (DM) was used as an indicator of a minimally affected gene carrier before the characterisation of the DM mutation. The identification of a specific mutation involving an unstable CTG repeat sequence in DM patients has provided an accurate molecular test for the disorder, and has allowed the detection of minimally affected subjects with a premutation (50 to 100 CTG repeats). DM premutation carriers with cataracts and no family history of DM could constitute the genetic reservoir of the DM mutation in the population. In a recent and extensive epidemiological study, in which we have tested nearly all the members of the families with the disease, we have found a prevalence of 26.5 cases per 100,000 in Guipúzcoa, a province of the Spanish Basque Country. This is the second highest prevalence reported after the one found in the Saguény-Lac-St-Jean region (189/100,000) in Québec, Canada. The assessment of the DM premutation in an unselected sample of cataract patients from a regional Ophthalmology Service could help to determine the genetic reservoir of the DM mutation in this high prevalence area.

Materials and methods

PATIENTS
DNA was obtained from blood samples taken from 231 patients recruited prospectively, ranging in age from 21 to 91 years (mean age 72 years), and presenting at the eye clinic of the Ntra Sra de Arantzazu Hospital. Patients with evident signs of DM or related to previously known DM families were excluded. The presence of other risk factors for developing cataracts, such as arterial hypertension or diabetes, was assessed in all patients. The age at the time of cataract surgery and sex distribution are shown in the table. The patients were informed about the possible genetic cause of their cataract and the purposes of this study, which was approved by the ethics committee of our hospital. The patients with a positive result were informed by a neurologist (ALM) and genetic counselling was offered.

PCR ANALYSIS
PCR analysis of the CTG repeats in the DM gene was performed as described previously with the following minor modifications. Reactions were performed in a 30 μl volume containing 1 μmol/l of each primer, 1 × PCR buffer (50 mmol/l KCl, 10 mmol/l Tris HCl, pH 8.3, 0.01% gelatine, 1.5 mmol/l MgCl2), 200 μmol/l each of dATP, dCTP, dGTP, and dTTP, 700 ng DNA, and 1 U Taq polymerase. Cycling parameters were as follows: 94°C for three minutes, 30 cycles of 94°C for one minute 30 seconds, 67°C for one minute, 72°C for two minutes. Final extension was at 72°C for seven minutes. PCR products and the marker pBR322 (New England Biolabs) were run for two hours on 3% Metaphor agarose (FMC) gels at 200 mA and visualised with ethidium bromide. This protocol can detect from five (128 bp) to up to 130 to 140 (560 bp) CTG repeats (fig 1).

STATISTICAL ANALYSIS
Statistics were performed using SYSTAT package (PC version 5.0). ANOVA test and Pearson correlation matrix were used when appropriate.

| Age and sex distribution in cataract patients studied |
|---|---|---|---|---|
| Age | Total |
| --- | --- | --- | --- | --- |
| <45 | 13 | 45-60 | 61-75 | >75 |
| Men | 2 | 32 | 212 |
| Women | 4 | 61 | 63 | 132 |
| Total | 6 | 108 | 95 | 231 |

Results
As seen in fig 1A, the modified PCR protocol is very efficient for detecting clearly up to 140
CTG repeat expansions. The patients with minimal signs of DM (cataracts or myotonic discharges on EMG or both) have expansions in this range.\textsuperscript{7-9} Patients with larger expansions, except children and young people, present other clinical signs suggestive of DM, which constitute exclusion criteria for our study. Fig 1B shows an example of normal repeats in cataract patients. The marker used as a repeat number standard and the resolution obtained allow alleles of 31 and 32 repeats to be distinguished.

In our series, we have observed the same trimodal allele distribution in the normal size range (5 to 37 CTG repeats) as the one previously reported for white populations (fig 2).\textsuperscript{10} No relationship was found between the number of CTG repeats and sex (ANOVA p = 0·64), age (Pearson correlation matrix - 0·08), presence of diabetes (ANOVA p = 0·71), or arterial hypertension (ANOVA p = 0·94).

Two DM premutation carriers were found. One of them was a 74 year old man with diabetes and arterial hypertension, who had an expansion of 80 CTG repeats, and the other was a 47 year old woman with an expansion of 55 CTG repeats. Both of them came from the Bajo-Deba area of Guipuzcoa where the prevalence of the disease is the highest (30.9/100 000).\textsuperscript{5}

Discussion

Until now, only two studies have analysed the risk to cataract patients of being DM carriers.\textsuperscript{11,12} Harley et al\textsuperscript{11} found no DM carriers among 101 cataract patients.\textsuperscript{11} Kidd et al\textsuperscript{12} found six patients with the DM mutation out of 113 patients studied (17 were analysed retrospectively). However, their inclusion criteria were not as strict as ours, since three had signs, symptoms, and even an evident familial history of DM, whereas in our study these patients were excluded. Considering only the other three patients, the prevalence of DM among cataract patients found by Kidd et al\textsuperscript{12} is higher than ours. This could be because of the age difference in both series; the patients younger than 60 (inclusion criteria in the study of Kidd et al\textsuperscript{12} constitute only 11% of our sample. We have observed that 0-9% of our cataract patients carry the DM premutation. The percentage obtained is very low to recommend genetic screening for the DM mutation in all the cataract patients. However, we must consider the importance of the detection of DM premutation carriers, owing to the anticipation phenomenon. These subjects are at risk of having children with typical DM and grandchildren with congenital DM, especially when the DM premutation is transmitted by a male.\textsuperscript{13} The detection of DM premutation carriers could help to prevent the occurrence of congenital cases in families without a clinical history of typical DM. Screening for the DM mutation in cataract patients should be restricted to young patients or subjects with multicoloured and iridescent opacities, in whom the risk of carrying the DM premutation could be higher. It is interesting to note that in our study, a frequency of the DM mutation of

\begin{figure}
\centering
\includegraphics[width=0.5\textwidth]{figure1}
\caption{Frequency distribution of the CTG repeat in cataract patients.}
\end{figure}
1/26 in the age group less than 60 years was found, whereas in those over 60 years, the proportion was only 1/203. It has been postulated that "trinucleotide diseases" like myotonic dystrophy could be time bombs within the human genome.\(^{14}\) The strong tendency towards expansion of repeats, shown by Rubinstein et al.\(^{15}\) in Huntington disease,\(^{15}\) would lead to a steady increase over time in the frequency of alleles in the premutation range coming from alleles at the highest point of the normal range. However, in our cataract patients, we have not found subjects in the 33 to 50 repeat range, which could be ready to pass the threshold in few generations. Moreover, we have not found intergenerational variation in the transmission of normal CTG repeats (5 to 37) in 346 parent-child pairs, of which 62 were transmissions of alleles in the 19 to 33 repeat range.\(^{16}\) Unfortunately, we were not able to study the parents (dead) who transmitted the DM mutation to our two cataract patients. The 47-year-old woman with 55 repeats had no first-degree relatives and the offspring of the other patient declined to participate in the study. Even if the patients had not had a direct relationship with any known DM family, we suspect they could belong to asymptomatic unknown collateral branches of symptomatic DM families which were not included in our epidemiological study.\(^{17}\) In the first case, the patient was from a small village with a high prevalence of DM and the other patient shared the same rare family name as a known DM family from a neighbouring province.

The genetic reservoir of the disease could be the generally mildly affected patients (with cataracts as the only sign of the disease) in the 38 to 80 (CTG)\(_3\) range, derived from collateral branches of DM pedigrees. Some premutated alleles between 38 and 80 repeats can expand from 120 and up to 1000 repeats in some cases especially in male transmissions.\(^{17-19}\) While in other cases, the premutated alleles are transmitted silently over generations.\(^{19}\) The alleles which expand would replace the losses owing to the reduction in fitness and marriage produced by the disease.\(^{20}\) For a better understanding of the dynamics of the DM mutation in this range, the next step should be the study of young cataract patients with the DM premutation, their parents, and, if possible, their grandparents, in order to look for the ancestral origin of the premutated allele and try to determine the moment where the jump to instability occurred.

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