A general method for the detection of large CAG repeat expansions by fluorescent PCR

Jon P Warner, Lilias H Barron, David Goudie, Kevin Kelly, David Dow, David R Fitzpatrick, David J H Brock

Abstract
The expansion of a tandemly repeated trinucleotide sequence, CAG, is the mutational mechanism for several human genetic diseases. We present a generally applicable PCR amplification method using a fluorescently labelled locus specific primer flanking the CAG repeat together with paired primers amplifying from multiple priming sites within the CAG repeat. Triplet repeat primed PCR (TP PCR) gives a characteristic ladder on the fluorescence trace enabling the rapid identification of large pathogenetic CAG repeats that cannot be amplified using flanking primers. We used our method to test a cohort of 183 people from myotonic dystrophy families including unaffected subjects and spouses. Eighty five clinically affected subjects with expanded alleles on Southern blot analysis were all correctly identified by TP PCR. This method is applicable for any human diseases involving CAG repeat expansions.

Key words: fluorescent PCR; CAG repeat; myotonic dystrophy; TP PCR.

The expansion of the trinucleotide sequence CAG is the mutational mechanism for several human genetic diseases including myotonic dystrophy, Huntington's disease, spinocerebellar ataxia I and III, and dentatorubral-palludoluysian atrophy.1,2 Currently, for these diseases, PCR using flanking primers allows amplification up to approximately 100 CAG repeats but is unreliable above this size. The diagnostic threshold for this series of diseases is at between 40 to 50 repeats with alleles above this size being in the affected range. The myotonic dystrophy CAG repeat can expand to give alleles of greater than 5 kb.1,3 PCR amplification of DNA from an affected person carrying such an expansion gives a single normal allele and fails to pick up a larger allele. For people apparently homozygous for small common repeat size, the presence of a large unamplifiable allele can be excluded by performing a Southern blot and probing with a locus specific fragment flanking or containing the CAG repeat. We have developed a simple fluorescent PCR system that can rapidly identify but not size the largest alleles for any disorder with a CAG repeat expansion. This will reduce the number of Southern blot analyses required by allowing the identification of people homozygous for smaller alleles. Samples with large CAG expansions identified by TP PCR will, however, still require Southern blotting if an accurate estimation of size is needed.

Materials and methods
PCR DESIGN
Traditionally, PCR amplification of these unstable DNA sequences involves the use of a pair of locus specific primers (P1, P2) that flank the variable trinucleotide sequence. Our PCR assay uses a specific flanking fluorescently labelled primer (P1) in concert with a pair of primers (P3, P4) which have a common 5' sequence (tail) (fig 1). The common 5' sequence or tail primer (P3) was selected from an artificially produced random DNA sequence. The selection criteria for the sequence of the tail primer were (1) that it should contain little or no self complementarity, (2) no complementarity with either (GCA), or (TGC), and (3) no homology to known human sequences. The paired primer (P4) has the sequence (GCA), or (TGC), at its 3' terminus depending on the strand of the CAG repeat to be amplified.

In the early amplification cycles the repeat specific 3' terminus of P4 primes at multiple sites within CAG repeat alleles giving rise to a mixture of products. Specificity is dictated by the fluorescent locus specific primer (P1). A 1:1 molar ratio of P3 to P4 ensures that primer P4 is exhausted in the early amplification cycles. This reduces priming at (CAG)n sites internal to the PCR products produced in earlier rounds, which would result in a gradual shortening of the average PCR product size (fig 1). The phenomenon of maintaining the allele size distribution by using a combination of tailed primer pairs was first described by Neil and Jeffreys4 in the context of satellite DNA repeat amplification. The primer P3 preferentially binds to the end of products from previous amplification rounds owing to the stabilising effect of the 5' tail sequence. A long extension time is used to allow complete extension of the larger sized products within the PCR product mixture.

POLYMERASE CHAIN REACTION
DNA used for PCR amplification was extracted from venous blood leucocytes using standard DNA extraction methods. Primer sequences are shown in table 1. The sequences of HD1 and DRPLA 2 were selected from the HD1 and DRPLA1 gene sequences respectively. P1...
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and P2 were selected from the 3' non-coding region of the myotonic protein kinase gene DNA sequence.1 The primers flank the myotonic CAG repeat, have matched annealing temperatures, and are also appropriate for amplification in conjunction with the primers P3R, P4CTG, and P4CAG. PCR reactions were performed in 25 μl reaction volumes. The following buffer components were found to give optimal amplification: 1.5 mmol/l MgCl, 10 mmol/l Tris HCl, pH 8.3, 50 mmol/l KCl, 10% DMSO, 200 μmol/l dATP, 200 μmol/l dCTP, 200 μmol/l dGTP, and 200 μmol/l dTTP. Genomic DNA (200 ng-1 μg) and approximately 2 U of Taq polymerase (Cetus) were used for each reaction. The following primer combinations were used. Mix A: P1 (5' fluoresceinylated 1 μmol/l) and P2 (1 μmol/l) for conventional amplification of the myotonic dystrophy CAG repeat, mix B: P1 (5' fluoresceinylated 1 μmol/l) with P3R (1 μmol/l) and P4CTG (0.1 μmol/l), and mix C: P2 (5' fluoresceinylated 1 μmol/l) with P3 (1 μmol/l) and P4CAG (0.1 μmol/l) for amplifications priming from the CAG repeat allele. Samples were overlaid with mineral oil (Sigma) and incubated for four minutes at 94°C before the addition of the Taq polymerase. The reactions were subjected to 30 cycles of 94°C for one minute, 60°C for one minute, and 72°C for two minutes followed by a 10 minute extension at 72°C in a Perkin Elmer DNA thermal cycler.

Five microlitres of each PCR product were added to 5 μl of distilled water and 10 μl of loading buffer (10 mg/ml blue dextran in deionised formamide) and incubated at 95°C for three minutes. Tubes were cooled on ice before loading on a denaturing 6% polyacrylamide 7 mol/l urea gel. PCR products were resolved by electrophoresis in 0.6 x TBE and visualised using an automatic laser fluorescent sequencer (ALF Pharmacia). Traces were analysed using the Pharmacia fragment manager software.

Table 1 Primer sequences

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence (5'-3')</th>
</tr>
</thead>
<tbody>
<tr>
<td>P1</td>
<td>AGA AAG AAA TGG TTC TGT GAT CCC</td>
</tr>
<tr>
<td>P2</td>
<td>GAA CGG GGC TCG AAG GGT CCT TGT AGC CG</td>
</tr>
<tr>
<td>P3R</td>
<td>TAC GCA TCC CAG TTT GAG ACG</td>
</tr>
<tr>
<td>P4CTG</td>
<td>TAC GCA TCC GAG TTT GAG ACG TCG TGC TGC TGC TGC T</td>
</tr>
<tr>
<td>P4CAG</td>
<td>TAC GCA TCC GAG TTT GAG ACG CAG CAG CAG CAG CAG CAG</td>
</tr>
<tr>
<td>HD1</td>
<td>ATG AAG GCC TTC GAG TCC CTC AAG TCC TTC</td>
</tr>
<tr>
<td>DRPLA2</td>
<td>TCA CCA TCA CCA CCA GCA ACA GCA A</td>
</tr>
</tbody>
</table>

**Results and discussion**

Typical traces obtained using mix A are shown in fig 2. The PCR amplification across the myotonic CAG repeat gives reliable amplification up to alleles of approximately 100 CAG repeats but fails to amplify above this size. Analysis of severely affected myotonic dystrophy patients by genomic Southern blotting typically shows expansions between 1.5 and 4 kb. PCR amplification of DNA from an affected subject carrying such an expansion gives a single normal allele and fails to pick up a larger allele, and is thus indistinguishable from patients genuinely homozygous for a small allele.

Typical traces obtained using mix B are shown in figs 3 and 4. Mix B gives similar traces to mix C (not shown). Amplification products show a characteristic ladder with a three base pair periodicity when products are analysed on a laser fluorescence DNA sequencer. The ladder peaks at the size of the largest allele present or, for very large repeat alleles, the peak height diminishes gradually.
with mix A gave the characteristic ladder which extended beyond 50 CAG repeats with the peak height diminishing with increasing product size (figs 3 and 4, bottom trace). These 76 people were scored as affected with myotonic dystrophy with CAG repeat alleles of greater than 50 CAG repeats. The remaining 107 DNA samples did not give the long characteristic ladder seen for large CAG expansions with mix B. Amplification with mix A showed that 76 people were heterozygous for CAG repeat alleles in the normal range with 31 patients homozygous for alleles in the normal range.

The clinical status of the patients and the results of the PCR and genomic Southern analysis reported by the Aberdeen laboratory were consulted. A total of 74 of the 76 people with CAG repeat alleles in the affected range were symptomatic and classed with an affected disease status. Two patients had alleles just within the affected range and were as yet asymptomatic. Both had either children or close relatives affected with myotonic dystrophy in whom we had also detected expanded CAG repeat alleles. The results of the conventional DNA analysis were that 75 patients had expanded CAG alleles in the affected range, from 50 CAG repeats to expansions of >5 kb. All 75 were in the group of 76 patients for whom our PCR assay had given evidence for the presence of a large CAG repeat allele. The one patient we identified who had not been ascertained by conventional testing was said to be clinically affected. The result of the Southern analysis for this DNA had been ambiguous because of poor DNA quality and a fresh sample will be collected for retesting.

On the basis of this series of tests we are confident that our TP PCR test for myotonic dystrophy is robust and reliable. For this large cohort of patients we obtained no false negative results and the results we obtained showed absolute concordance with the results obtained by conventional means. The only difference we saw suggested that our method was able to identify expanded alleles in poor quality DNA samples where Southern analysis failed to give a reliable result. This method does not measure the size of large expanded CAG repeat alleles but reliably identifies their presence. DNA samples can be screened using this method and only samples with large expansions would need to be analysed further using genomic Southern blotting in order to establish the exact size of large expanded alleles. The recently described XL PCR based method of Cheng et al could be used for samples where insufficient DNA for Southern analysis is available. This method is, however, technically demanding and time intensive as PCR products are resolved by agarose gel electrophoresis, the gel being then blotted and probed with a locus specific probe. Our TP PCR method works if the fluorescent primer P1 is replaced by 32P end labelled P1. These radiolabelled products are, however, more difficult to size accurately in the 30 to 60 CAG repeat size range (data not shown). Obviously silver staining is inappropriate for TP PCR as many non-specific inter(CAG), products are generated.

![Figure 2: PCR amplification at the myotonin locus using flanking primers. The axis shows migration time in minutes. CAG allele sizes shown with the arrows. (A) Trace obtained from a heterozygous normal subject. (B) Trace obtained from a heterozygous subject with a small expansion. (C) Trace obtained from a patient with myotonic dystrophy and an expanded allele size of >4 kb as determined by Southern blot analysis. The larger allele fails to amplify.](http://jmg.bmj.com/Downloaded from http://jmg.bmj.com/)
We have carried out some pilot TP PCR experiments using specific flanking primers for other diseases where expanded CAG repeats are the mutational mechanism. For Huntington's disease the same PCR conditions as described above for mix B replacing the primer P1 with HDI give reliable results (not shown). It has been suggested that for people with juvenile onset of Huntington's disease, where an expanded allele appears to be absent and the patient is homozygous for a smaller allele, a Southern analysis should be performed to exclude the presence of an unamplifiable large allele (Clinical Molecular Genetics Society of Great Britain, personal communication, 1995). Despite never having failed to amplify both alleles for juvenile Huntington's disease referrals up to 96 CAG repeats using flanking primers in Edinburgh, we would recommend the use of TP PCR to show the presence of these large, more difficult to amplify alleles. Similarly, we have shown that reliable results are obtained for large alleles for DRPLA when the primer DRPLA replaces P2 in mix C.

In conclusion we have developed a novel PCR assay that can be used for screening for expanded CAG repeat alleles including full mutation alleles of >5 kb in myotonic dystrophy. TP PCR is robust and reliable and can in theory also be used in conjunction with locus specific flanking primers for any disease with a similar mutational mechanism.

We would like to thank all the clinicians who have sent us the samples over the years and Aileen Crosbie for help with clinical records.

(cont. overleaf)

Figure 4 Close up of traces shown in fig 3. The axis shows migration time in minutes. CAG allele sizes shown with the arrows. (A) Detail from trace shown in fig 3B. (B) Detail from trace shown in fig 3C.
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