A genotypic mutation system measuring mutations in restriction recognition sequences

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Received March 7, 1991; Revised and Accepted May 15, 1991

ABSTRACT

The RFLP/PCR approach (restriction fragment length polymorphism/polymerase chain reaction) to genotypic mutation analysis described here measures mutations in restriction recognition sequences. Wild-type DNA is restricted before the resistant, mutated sequences are amplified by PCR and cloned. We tested the capacity of this experimental design to isolate a few copies of a mutated sequence of the human c-Ha-ras1 gene from a large excess of wild-type DNA. For this purpose we constructed a 272 bp fragment with 2 mutations in the Pvull recognition sequence 1727–1732 and studied the rescue by RFLP/PCR of a few copies of this ‘Pvull mutant standard’. Following amplification with Taq-polymerase and cloning into λgt10, plaques containing wild-type sequence, Pvull mutant standard or Taq-polymerase induced bp changes were quantitated by hybridization with specific oligonucleotide probes. Our results indicate that 10 Pvull mutant standard copies can be rescued from 10^8 to 10^9 wild-type sequences. Taq polymerase errors originating from unmodified, residual wild-type DNA were sequence dependent and consisted mostly of transversions originating at G.C bp. In contrast to a doubled ‘standard’ the capacity to rescue single bp mutations by RFLP/PCR is limited by Taq-polymerase errors. Therefore, we assessed the capacity of our protocol to isolate a G to T transversion mutation at base pair 1698 of the MspI-site 1695–1698 of the c-Ha-ras1 gene from excess wild-type ras1 DNA. We found that 100 copies of the mutated ras1 fragment could be readily rescued from 10^5 copies of wild-type DNA.

INTRODUCTION

Molecular characterization and quantitation of mutations is of fundamental importance to the understanding of evolution, differentiation and the etiology of hereditary disease. The frequent occurrence of single base pair changes in the activation of the ras protooncogenes (1) and in the inactivation of the tumor suppressor gene p53 (2–4) documents the involvement of mutations in human carcinogenesis. Ideally, methods are required which allow it to identify specific DNA sequence changes in relevant genes in tissue biopsies. For the elucidation of early pathogenetic processes it is necessary to detect these changes in a minute minority of cells without the clonal expansion of mutated cells in vivo or in vitro and without the need for the expression of a selectable mutant phenotype (5). To achieve this goal it is necessary to isolate specific mutated DNA sequences from a large excess of homologous wild type DNA by biochemical means. In contrast, all classical ‘phenotypic’ mutation systems rely on the isolation of a few mutated cells from a large, usually dividing cell population. This limits mutation analysis to a few genes encoding proteins which produce a selectable cellular phenotype.

A number of experimental approaches to ‘genotypic’ mutation systems have been developed (6). Several are based on Southern- or Northern hybridization often with sequence amplification by polymerase chain reaction (PCR) and in combination with the specific cleavage of hetero-duplexes at mismatched base pairs. Other protocols take advantage of differences in electrophoretic mobilities of hetero-duplexes or of mutated single stranded nucleic acids. The sensitivity of all these approaches is limited by backgrounds which arise from the presence of excess wild-type sequences. The RFLP/PCR approach to genotypic mutation analysis described here greatly reduces this problem. Base-pair (bp) mutations are detected which are located in a restriction recognition sequence and render this site resistant to cleavage by the corresponding endonuclease. The resistant DNA sequence containing the mutated site is amplified by PCR only after wild type DNA has been essentially eliminated by restriction digestion. Amplified DNA is cloned into λgt10 and mutants are quantitated by oligonucleotide plaque hybridization relative to an internal standard. The maximal sensitivity of the RFLP/PCR method is determined by the completeness of the removal of wild-type sequences together with the inherent error-rate at a particular base pair of the polymerase used in the PCR. The RFLP/PCR protocol for genotypic mutation analysis is applicable to any gene of known sequence. It is highly sensitive, but limited to the identification of point mutations (bp-changes, small deletions and insertions)

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which result in the elimination of restriction sites. We report here results involving bp-changes in the PvuII site 1727–1732 and the Mspl site 1695–1698 of exon 1 of the human c-Ha-rasl gene. The Mspl-site 1695–1698 covers 2 bp of codon 12 of c-ras which represents a mutation hot spot in human cancer (1).

METHODS

RFLP/PCR analysis of PvuII-site 1727–1732 of c-Ha-rasl

Preparation of authentic mutants at PvuII-site 1727-1732 of c-Ha-rasl. All 12 possible single bp mutations in the internal tetranucleotide AGCT of the PvuII recognition sequence (residues 1727–1732) of exon 1 of the human c-Ha-rasl gene were prepared using synthetic oligonucleotides and a PCR protocol (see Figure 1 for sequence information). Twelve different 20-mers corresponding to residues 1719–1738 which contained single base changes in the AGCT sequence were used as left-side amplimers. A common right-side amplimer corresponding to nucleotides 1765–1784 of c-Ha-rasl plus a 12 nucleotide tail containing an EcoRI recognition-sequence was used in 12 PCR reactions with pSVneo-rasl as template (note: pSVneo-rasl contains the BamHI fragment, residues 1–653, of human c-Ha-rasl (7) inserted into the BamHI sites of the SVneo cloning vector). The amplification conditions were as described below. The 12 resulting 78 bp fragments were purified on Quiaigen-5 tips as described by the supplier (DIAGEN, KONTRON, Switzerland) and then used as right-side amplimers in a second round of amplifications with a common left-side amplimer corresponding to residues 1646–1665 of c-Ha-rasl plus a 12 nucleotide tail containing an EcoRI recognition sequence. The resulting 163 bp fragments were purified on Quiaigen-5 tips, digested with EcoRI and cloned into λgt10.

All authentic mutant λgt10 constructs were plaque purified on E. coli C600 HfI. For this purpose plaques were lifted onto colony/plaque screens and probed with mutant specific 32P-end-labeled oligonucleotides. Positive plaques were picked, the phage eluted from the agar and plated onto E. coli C600 HfI. When confluent lysis had been achieved the phage was collected in 5ml buffer (100mM NaCl, 20 mM Tris.HCl pH 7.4, 10mM MgSO4) and after the addition of 100 μl CHCl3 the suspension was centrifuged at 10’000 rpm for 10 min at 4°C. Mutant phage contained in the supernatant was titered and then stored at 4°C in the presence of 0.3% CHCl3.

A mutant standard with two bp changes in the PvuII recognition sequence and an additional upstream mutation was prepared according to the same experimental design using the 20-mer 5’T-GAGCATCCGGTGTACG (residues 1719–1738 of c-Ha-rasl1; positions of mutations underlined) as left-side amplimer and a 20-mer corresponding to the sequence from residue 1888 to 1907 of c-Ha-rasl as right-side amplimer. The right-side amplimer contains a XbaI recognition site. The amplification conditions were as described below except that the DMSO concentration was only 6%. After 38 cycles the resulting 188 bp fragment was purified on a Quiaigen-5 tip and used as right-side amplimer in a second amplification reaction with a left-side amplimer corresponding to residues 1636 to 1655 of c-Ha-rasl1 which contains a SmaI site. After purification the final 272 bp fragment was digested with XbaI and SmaI and cloned into pSP64. This plasmid, referred to as ‘mutant standard’ (pSP64-rasl PvuII), was used as internal standard in mixtures with pSVneo-rasl.

PvuII - restriction and amplification. A stock of pSVneo-rasl was digested exhaustively with 4U PvuII/μg plasmid DNA in Tris buffer pH 7.5, 10mM MgCl2, 50mM NaCl, 1mM DTE for 3 h at 37°C. To 108 copies of restricted pSVneo-rasl was added 10 or 100 copies of PvuII mutant standard (see above) which had been predigested with XbaI and SmaI in order to release a 272 bp fragment containing the mutated sequence. The samples were prepared in the amplification buffer which was composed of 66.7mM Tris.HCl pH 8.8 (25°C), 6.7mM MgCl2, 16.6mM (NH4)2SO4, 1mM each of the 4 dNTPs, 12% DMSO, 1.25 μM of the left-side and right-side amplimers in a final volume of 25 μl. The amplimers used in the first 30 PCR cycles were 20-mers corresponding to the sequences 1646–1665 and 1765–1784 of c-Ha-rasl1 (see Figure 1). After addition of 0.5 U of Taq polymerase (from bacter. Thermus aquaticus, Cetus, Calif.) the samples were heated initially to 91°C for 4 min followed by 95 sec at 59°C. Consecutive cycles consisted of 85 sec at 91°C and 95 sec at 59°C and were carried out in an Ampliphen Apparatus, Moretronic, Switzerland. After 2,4,6 and 8 cycles 1 U PvuII was added and the samples incubated for 10 min at 52°C in order to eliminate any amplified wt-sequence. Amplification was then continued with 0.5 U fresh Taq-polymerase. After the first 8 cycles the amplification was continued without interruption up to 30 cycles when a 1 μl aliquot was withdrawn from each sample and added to 50 μl fresh amplification buffer. The mixtures contained 1.6 μM of clonable amplimers which in addition to the 20-mer sequences described above contained a 12-nucleotide tail with an EcoRI-recognition sequence (see Figure 1). Before the continuance of the amplification for 10 cycles the samples were treated with PvuII for 20 min at 37°C. In independent experiments 108 copies of PvuII digested pSVneo-rasl1 were mixed with 1 or 1000 copies of mutant standard and processed according to the protocol described above except that PvuII digestions during amplification were omitted.

Purification of amplified DNA and cloning into λgt10. The amplification mixtures were purified on Quiaigen-5 tips and the DNA precipitated with isopropanol after the addition of 1 μg E. coli tRNA as carrier. The samples were then treated with EcoRI in order to free the ends of the amplified fragment. After renewed purification on Quiaigen-5 tips, precipitation with isopropanol and washing of the precipitates with 75% ethanol 5ng/DNA aliquots were ligated to 500 ng λgt10 arms as outlined by the supplier (Promega Biotec, Madison, WI.) The phage DNA was packaged (Packagene', Promega Biotec, Madison, WI.) and E. coli C600 HfI infected. Plaques were lifted onto colony/plaque screens (NEF-978, New England Nuclear). At least 2 independent packagings were carried out and 1–3×109 plaques on 10 to 15 Petri-dishes were analyzed for each experimental condition.

Oligonucleotide plaque hybridization. Specific probes were prepared according to standard conditions by end-labeling with gamma-32P-ATP with T4 polynucleotide kinase of oligonucleotide 20-mers corresponding to the wild-type sequence 1719–1738 of c-Ha-rasl1, mutant standard containing 3 base changes and oligomers containing all 12 possible single base mutations in the internal tetranucleotide AGCT of PvuII recognition site 1727–1732. The colony/plaque screens to be analyzed were prehybridized for 1 h with 5×SSPE (1× 10mM Na-phosphate pH 7.4, 1mM EDTA, 0.15M NaCl), 0.3% SDS, 1% milk protein. After rinsing with water the discs were hybridized overnight with the radioactive probes at 54°C in
5×SSPE, 0.3% SDS. After a first wash at room-temperature with 5×SSPE selective washing temperatures between 60 and 63°C were employed in order to exclude non-specific hybridization. The selective washing temperatures were determined with known mixtures of λgt10 stocks containing authentic mutant inserts (for their preparation see above).

**RFLP/PCR analysis of MspI-site 1695–1698 of c-Ha-rasl**

*Preparation of authentic rasl-T1698 with a single base pair substitution in the MspI-site 1695–1698.* The c-Ha-ras1 gene with a G→T transversion at residue 1698 of exon 1 had been originally cloned from the EJ human bladder carcinoma line (8). A Smal fragment (1644–2246) of this gene was cloned into the vector puc18. From this the Xbal-EcoRI ras fragment 1644–1905 was removed and cloned into the pSP64 vector (9). This vector, referred to below as pSP64-rasl-T1698 was used in the mutant rescue experiments described below.

**MspI/HpaII restriction and amplification.** Stocks of pSVneo-ras1 were digested exhaustively with 10U MspI/μg and 2U HpaII/μg plasmid overnight at 37°C under the conditions described above for PvuII. pSP64-rasl-T1698 was digested with Xbal-EcoRI in order to release a 261 bp fragment containing the mutated MspI-site 1695–1698 CCGT. Three samples were prepared in the amplification buffer described above except that the DMSO concentration was only 6%: a) 10³ copies restricted pSVneo-ras1 b) 100 copies restricted pSP64-rasl-T1698 plus 10³ copies restricted pSVneo-ras1 c) 1000 copies restricted pSP64-rasl-T1698 plus 10⁶ copies restricted pSVneo-ras1. The amplimers used in the first 35 PCR cycles were 20-mers corresponding to the sequences 1646–1665 and 1765–1784 of c-Ha-ras1 (see Figure 1). The amplification cycles were as described above. After 2, 4, 6, 8, 10, 12 and 14 cycles 2U MspI were added and the samples incubated for 10 min at 55°C in order to eliminate any residual wild-type sequences. Amplification was then continued after the addition of 0.5U fresh Taq polymerase. After 14 cycles the amplification was continued to 35 cycles without interruption when a 1μl aliquot was withdrawn from each sample and added to 50μl fresh amplification buffer. The amplification was continued with 1.6μM nested clonal left-side amplimer 1673–1692 and clonal right-side amplimer 1765–1784. These amplimers possessed 12-nucleotide 5'-tails containing EcoRI-recognition sequences (see Figure 1). All subsequent steps consisting of purification of the amplified DNA, cloning into λgt10 and oligonucleotide plaque hybridization were as described above for the analysis of mutants in the PvuII-site 1727–1732. The results shown on Figure 5 are derived from 2 independent packagings and 600–1100 plaques on 4 to 6 Petri dishes.

**Radiolabeled amplimer incorporation**

In order to compare the efficiency of amplification of mutant standard and wild-type c-Ha-ras1 DNA the 20-mer corresponding to residues 1646–1665 was end-labeled with γ^32P-ATP and T4 polynucleotide kinase under standard conditions. This
radiolabeled oligonucleotide was used as left-side amplitiser in a PCR-reaction with a non-radioactive right-side amplitiser and 5×10^10 copies of XbaI and SmaI digested mutant standard pSP64-ras PvuIISt or of XbaI and SmaI digested pSVneo-rasl as templates, respectively. In a second experiment only 10^8 copies of the template DNAs were used. Amplification conditions were as described above with the exception that no PvuII digestions were performed. Aliquots of the reaction mixtures were withdrawn after increasing numbers of PCR cycles and applied to 5% polyacrylamide gels. After development, the dried gels were exposed to x-ray film for autoradiography. The intensities of the bands at 138 bp corresponding to the expected amplification product were determined by densitometry (Zeineth Scanning densitometer, Biomed Instr., CA.).

RESULTS

Efficiency of amplification of PvuII mutant standard and wild-type sequence 1646–1784 of c-Ha-rasl

In order to calibrate single bp changes by RFLP/PCR an internal ‘PvuII mutant standard’ is used with 2 bp-mutations in the chosen recognition sequence. We prepared a mutant standard pSP64-ras PvuIISt with 2 bp changes in the PvuII recognition sequence (1727–1732) and an additional mutation upstream (see Methods and Fig. 1). For the validity of the calibration the amplification efficiencies for mutant standard DNA and mutated sequences (or undigested wt-sequences) from genomic DNA have to be the same. This is demonstrated in Figure 2. The upper portion shows that radiolabeled left-side amplitiser (see Figure 1 for sequences) was incorporated into the desired 138 bp fragment with equal efficiency whether mutant standard or wt-sequence served as template. Either 5×10^10 or 10^8 initial copies of the templates were amplified for 10 or 14 PCR cycles, respectively. The lower portion of Figure 2 compares the amounts of 138 bp fragment synthesized from 10^5, 10^6 and 10 copies of the two templates after 29 and 34 PCR cycles directly by ethidium bromide staining on a 2% agarose gel. The results indicate comparable amplification efficiencies for PvuII mutant standard- and wt-template also at low numbers of initial copies and high cycle numbers.

Rescue of PvuII mutant standard from excess wild-type sequence 1646–1784 of c-Ha-rasl

The capability of the RFLP/PCR method to rescue a small number of copies of PvuII mutant standard from a large excess of wt sequence was investigated. Mixtures were prepared of 10 and 100 copies of XbaI/SmaI digested mutant standard pSP64-ras PvuIISt with 10^8 copies of exhaustively PvuII digested pSVneo-rasl and of 1000 copies of mutant standard with 10^9 copies PvuII digested pSVneo-rasl. The mutant standard was restricted with XbaI/SmaI in order to release a 272 bp fragment similar in size to the 256 bp-rs-fragment which is liberated by PvuII from pSVneo-rasl containing a mutation in the PvuII site 1727–1732. The mixtures were amplified with repeated restrictions with PvuII in 30 cycles with ‘non-clonable’ amplifiers and 10 final cycles with amplifiers containing tails with EcoRI recognition sequences (see ‘Methods’ and Figure 1). The 163bp amplified fragments were cloned into pGEM10 and plaques were analyzed by oligonucleotide hybridization for their content in wt-rasl sequence 1719–1738 and mutant standard sequence.

The results summarized in Table 1 show that as few as 10 copies of PvuII mutant standard were rescued from 10^8 copies of wt-sequence giving rise to 7.3% of total plaques. Increasing initial copy numbers of PvuII mutant standard gave rise to larger fractions of plaques containing the mutant standard sequence. The fractions of wt-plaques varied from 14 to 23%.

**ANALYSIS OF AMPLIFICATION MIXTURE BY λ-PLAQUE HYBRIDIZATION**

<table>
<thead>
<tr>
<th>Mutant standard</th>
<th>Wild type</th>
<th>Mutant standard % plaques</th>
<th>Wild type % plaques</th>
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<tr>
<td>10</td>
<td>10^8</td>
<td>7.3 ± 0.8</td>
<td>23.3 ± 1.2</td>
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<tr>
<td>100</td>
<td>10^8</td>
<td>13.9 ± 1.3</td>
<td>43.4 ± 1.4</td>
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<tr>
<td>1000</td>
<td>10^9</td>
<td>18.0 ± 1.7</td>
<td>20.4 ± 1.7</td>
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*Plaques containing mutant standard or wild-type sequence at the PvuII recognition sequences (1727–1732).* Plaques from at least 2 packagings were used and plaques on 10-15 petri dishes were analysed.

Figure 3. Analysis of λ-plaques for base-pair mutations in PvuII-recognition sequence 1727–1732 of c-Ha-rasl. Mixtures of a few copies of mutant standard pSP64-ras PvuIISt with 10^8 copies of wild-type pSVneo-rasl were processed by the RFLP/PCR protocol described in 'Methods'. The amplification product was cloned into pGEM10, plaques transferred to plate screens and analyzed by hybridization with 32-P-end-labeled mutant oligonucleotide probes for Taq-polymerase induced single bp mutations in the PvuII recognition sequence 1727–1732. The selectivity of the washing conditions was ascertained for each mutation with authentic mutant constructs. Upper half, left-side: plaques containing authentic mutant CAGCTG; right-side: plaques on 4 plates with same mutation originating from the amplification mixture. Lower half, left-side: plaques containing authentic mutant CAGGTT; right-side: plaques on 4 plates with the same mutation originating from the amplification mixture.

Taq-polymerase errors in PvuII-site 1727–1732 of c-Ha-rasl

Despite exhaustive restriction with PvuII before and during amplification a small amount of intact wt-sequences undoubtedly remains, will be amplified and can give rise to Taq-polymerase errors. Amplified sequences containing Taq-polymerase errors in the PvuII recognition sequence 1727–1732 are resistant to digestion and will be cloned. The ultimate sensitivity of the RFLP/PCR method for a particular bp in the chosen recognition site depends on the error-rate of Taq-polymerase for the replication of this bp. Therefore, we probed λ-plaques resulting from the mixtures of mutant standard with excess wt-sequence for their content of Taq-polymerase induced single bp changes in the internal tetranucleotide AGCT of the PvuII recognition sequence. The plaques were probed with specific oligonucleotides for all 12 possible mutations. Figure 3 shows an example in which discs had been probed for a G→C transversion giving rise to
The polymerase induced which of the amplification cycles of _be_ (Figure rasl to represents and CACCTG and constructs (lower portion). In all cases the specificity of the probing conditions was ascertained with mixtures of authentic mutant constructs and by assuring that each plaque only hybridized with a single oligonucleotide probe.

Figure 4 compares the frequencies of the 12 possible Taq-polymerase induced bp changes at the internal tetranucleotide AGCT of the PvuII site in 3 independent amplification experiments in which 10^9 (Figure 4A) and 10^8 initial copies of wt pSVneo-rasl1 (Figure 4B & C) were used, respectively. While there are small quantitative differences the following consensus can be reached for the relative abundance of Taq-induced errors: C→A(AGAT) ≡ C→G(AGGT)>G→T(ACTG). Since the amount of wt-copies which remain undigested during the amplification cycles is not known absolute Taq-error frequencies cannot be derived from our data. In contrast, from a comparison to the data with the mutant standard it is evident that less than 10 copies containing a single bp mutation in the presence of 10^9 copies of wt-sequence can readily be detected for mutations with low Taq-error frequencies, e.g. for A→C(GGCT); T→G(A-GCC); T→A(AGCA); A→T(TGCT); G→A(AACT).

**Rescue of a single base pair mutation in MspI-site 1695 – 1698 from excess wild-type c-Ha-rasl DNA**

The rescue of mutant standard such as pSP64-rasl PvuIISt with multiple bp changes in a chosen restriction site from excess wt sequences is not affected by Taq-polymerase errors. In contrast, the sensitivity of the detection of single bp mutations is limited by polymerase errors originating from the amplification of residual undigested wt sequences. We assessed the capacity of the RFLP/PCR protocol to rescue a few copies of the c-Ha-rasl1 exon 1 fragment 1646 – 1784 containing a G→T transversion at residue 1698 which is part of the MspI-site 1695 – 1698 from a large excess of wt c-Ha-rasl1 DNA. The chosen mutation affects the middle bp of codon 12 and was detected in the c-Ha-rasl1 gene of the EJ human bladder carcinoma line (8) (see Figure 1 for sequence information). Three samples were analyzed by RFLP/PCR (1) 10^9 copies of exhaustively MspI/HpaII digested wt pSVneo-rasl1 (2) 100 copies mutant pSP64-rasl1-T1698 (XbaI/EcoRI restricted) plus 10^5 copies MspI/HpaII digested pSVneo-rasl (3) 100 copies mutant pSP64-rasl1-T1698 (XbaI/EcoRI restricted) plus 10^4 copies MspI/HpaII digested pSVneo-rasl1. The samples were amplified with repeated MspI restrictions in 35 cycles with ‘non-clonable’ amplimers and 10 final cycles with amplimers containing tails with EcoRI recognition sequences (see Figure 1 and Methods). The amplified 136 bp fragments were cloned into _E.g_. and plaques analyzed by hybridization with 20-mer oligonucleotides corresponding to the wt c-Ha-rasl1 sequence 1688 – 1707 and with the mutant sequence containing a T-residue at position 1698. From the results shown in Figure 5 it is evident that Taq-polymerase amplification of residual wt ras1 DNA which escaped restriction at the MspI-site 1695 – 1698 generated 12% T-plaques with G→T transversions at residue 1698 for the sample containing only 10^9 copies of pSVneo-rasl1 at the outset. The percentage of plaques containing G→T transversions at residue 1698 was 23% for the initial mixture of 100 copies mutant pSP64-rasl1-T1698 with 10^8 copies of wt pSVneo-rasl1 corresponding to a 2 fold increase over.
the fraction produced by Taq-polymerase infidelity. At 1000 initial copies of mutant pSP64-ras1-T1698 plus 10^8 copies wt pSVneo-ras1 the percentage of mutant plaques had increased to 34%. The Figure also gives the percentage of plaques which contained wt-sequence 1688–1707.

**DISCUSSION**

The aim of the present work was to test the feasibility of using the RFLP/PCR approach for the development of a genotypic mutation system. Such a system should be capable of measuring low frequency mutations in any gene of known sequence without the need for the phenotypic selection of mutated cells (6). In the RFLP/PCR approach wt-sequences are selectively removed by exhaustive restriction and resistant sequences are amplified by PCR. We have chosen the tetranucleotide AGCT of PvUII recognition site for our studies because it contains all 4 nucleotides. While the composition of the PCR product reflects the initial mixture of mutations it does not allow the estimation of absolute mutation frequencies. Therefore, we are using an internal ‘mutant standard’ for calibration. We have constructed a mutant standard with 2bp changes in PvUII recognition site and a third mutation upstream. This allows the distinction between λ-plaques containing mutant standard from plaques containing single bp mutations in the final analysis with specific probes. In the present experiments we have used the mutant standard pSP64-ras PvuIISt to evaluate the capacity of the RFLP/PCR protocol to rescue a few copies of mutated sequences in the PvuII site 1727–1732 from a large excess of wt c-Ha-ras1 without interference by Taq-polymerase errors. We have ascertained that mutant standard and wt-sequences are amplified with equal efficiencies (see Figure 2). Our results indicate (Table 1) that 10 copies of mutant standard were readily rescued from 10^8 to 10^9 copies of wt-DNA. While the fraction of mutant standard plaques increased with increasing copy number there is no simple relationship between the two parameters.

In contrast to the rescue of mutant standard the sensitivity of the detection of single bp mutations depends on the frequency of polymerase errors during the replication of a particular bp. Only polymerase errors originating from wt-sequences will interfere since errors in already mutated sequences will give rise to double mutations which are not detected with oligonucleotide probes containing a single altered base. Therefore, the final polymerase error frequencies will depend on the completeness of the removal of wt-sequences by restriction and our protocol employs several steps of restriction before and during amplification. We used conditions for the amplification reaction which were expected to yield relatively low Taq-polymerase errors (10) but retained high amplification efficiency, i.e. MgCl2 concentration of 6.7 mM, a total concentrations of dNTPs of 4 mM but a relatively high pH of 8.2 (70°C). The data in Figure 4 attests to the large differences in Taq-polymerase errors depending on the type of bp and the sequence. The replication of G.C bp produced more errors then the replication of A.T bp and transversions were considerably more frequent than transitions. Sequence dependence is indicated by the fact that the C→G transversion (AGGT) was more frequent than the G→C transversion (ACCT) and the C→A transversion (AGAT) more frequent than the G→T transversion (ATCT). Comparison of the three panels in Figure 4 indicates that the fraction of plaques with Taq-polymerase errors did not change significantly when the initial number of wt-copies was increased from 10^6 to 10^9. Under the latter conditions a single copy of mutant standard resulted in 0.3% mutant standard plaques. We estimate that at best RFLP/PCR analysis at PvuII-site 1727–1732 is capable of rescuing 1–5 copies of a single bp mutation from 10^6 copies of c-Ha-ras1 for mutations with low Taq-polymerase error frequencies, e.g. for A→C (CCGG), T→G (AGGG), A→T (TGCT) and T→A (AGCA). It is conceivable that polymerase errors can be further reduced by modifying the amplification conditions (10,11) and consequently that the sensitivity of our RFLP/PCR protocol can be increased.

We have further assessed the sensitivity of the RFLP/PCR protocol for the rescue of a single bp mutation in the PvuII-site 1695–1698 of exon 1 of c-Ha-ras1. Unlike the rescue of a doubly mutated standard such as pSP64-ras PvuIISt the capacity to rescue a single bp mutation is limited by backgrounds caused by polymerase errors. The experimental conditions were analogous to the PvuII-site 1727–1732 with the exception that nested ‘clonable’ left-side amplimer was used for the final 10 PCR cycles. The chosen mutant results from a G→T transversion at bp 1698 which represents the middle bp of codon 12. Mutations in this ras-codon are frequently detected in the DNA from human tumor tissues (1) and the particular G→T transversion studied in the present work had been first detected in the EJ human bladder carcinoma line (8). Our results show that 100 copies of a ras1 fragment containing a mutated PvuII-site 1695–1698 with a T-residue in position 1698 could be readily rescued from 10^6 copies of wt ras1 DNA. As expected from the analysis of the PvuII-site 1727–1732 under similar experimental conditions the G→T transversion at bp 1698 represents a relatively frequent Taq-polymerase error. The fraction of plaques containing the T1698 mutation was 12% for the sample composed initially of only wt DNA. This value is comparable to 11% for the frequency of Taq-polymerase induced G→T transversions at bp 1729 in the PvuII-site 1727–1732 for the sample containing initially only 1 copy of PvuII mutant standard but 10^8 copies of wt ras1 DNA.

The predominance of transversions involving G.C bp in our experiments disagrees with published work in which A.T→G.C
transitions were reported to be the most frequent Taq-polymerase errors (10,11,12,13). Differences in the amplification conditions may be responsible for these discrepancies; e.g. high ratios of concentrations of Mg\(^{2+}\)/total dNTPs in several published reports relative to a ratio of 1.7 in the present work; the use of relatively high concentrations of DMSO in our work in contrast to previous authors etc. It should be stressed that our data does not allow it to calculate absolute error-frequencies for Taq-polymerase because the amount of residual, unrestricted wt-sequence in the amplification mixture cannot be assessed by the RFLP/PCR protocol.

In conclusion, the RFLP/PCR protocol described here allows the analysis of point mutations (bp-changes, small deletions and insertions) in any gene of known sequence without the need to select mutated cells on the basis of an altered phenotype. The method is highly sensitive and relatively simple but limited to mutations which result in the elimination of a restriction site. The experimental design of the RFLP/PCR approach to genotypic mutation analysis has been discussed previously (14,15).

ACKNOWLEDGEMENTS

This work was supported by the Swiss National Science Foundation and the Swiss Association of Cigarette Manufacturers and the Association for International Cancer Research.

REFERENCES