Mutation of the p53 gene precedes aneuploid clonal divergence in colorectal carcinoma

PJ Carder¹, KJ Cripps², R Morris², S Collins², S White², CC Bird² and AH Wylie²

¹Academic Unit of Pathology, Alergon Firth Building, University of Leeds, Leeds LS2 9JT, UK; ²CRC Laboratories, Department of Pathology, University of Edinburgh, Tertiot Place, Edinburgh EH8 9AG, UK.

Summary To establish whether p53 mutation precedes or follows clonal divergence in human colorectal carcinomas, 17 tumours were analysed at multiple sites (2-5 each) for single-strand conformation polymorphisms (SSCP) within exons 5-8 of the p53 gene. A previous study had demonstrated subclones of differing DNA ploidy in these tumours, but all showed immunocytochemical evidence for p53 stabilisation, using the monoclonal antibody PAB 1801. Mutations within exons 5-8 of p53 were identified by the presence of an abnormally migrating band in 10 of the 17 carcinomas: five in exon 5, four in exon 7 and one in exon 8. In each of these positive cases, samples from different parts of the carcinoma showed identical gel migration patterns in SSCP analysis. Similarly, the remaining seven tumours were concordant for absence of band shift across all samples of each tumour. Six SSCP-positive cases contained multiple populations differing in DNA ploidy, while four were homogeneously diploid or aneuploid throughout. Very similar proportions were observed in the SSCP-negative cases. In four positive tumours the mutation was confirmed by sequencing or through alteration of nucleotide-specific restriction enzyme cleavage. Identical mutations appeared in every sample from the same tumour. The results provide unequivocal evidence that the same mutant allele of p53 is present throughout each tumour bearing a mutation, regardless of the clonal variation identified by analysis of DNA ploidy. We conclude that in colorectal tumorigenesis mutation of p53 occurs as a single event which precedes and may facilitate the aneuploid clonal divergence of carcinomas.

Keywords: colorectal carcinoma; p53; SSCP; clonal evolution

Inactivation of the tumour-suppressor gene p53 is an important step in the development of the majority of human cancers (Hollstein et al., 1991). Functional inactivation occurs most commonly as a result of missense mutation (Baker et al., 1989). We also report from interaction with oncogenic viral or cellular proteins (Mietz et al., 1992; Mornand et al., 1992). The p53 protein is a sequence-specific DNA-binding protein that is active as a transcription factor (Bargenetti et al., 1991) and can interact directly with the replication apparatus (Dutta et al., 1993). There is good evidence that its normal function is to establish G1 checkpoint control in response to DNA damage, so allowing time for DNA repair (Kastan et al., 1991) or the initiation of apoptosis (Clarke et al., 1993). Lack of functional p53 promotes genomic instability (Bischoff et al., 1990; Harvey et al., 1993), which is probably a key factor in acquisition of the multiple 'hits' required for carcinogenesis (Nowell, 1976).

We have recently shown that human colorectal carcinomas containing immunohistochemically detectable p53 are more likely to contain multiple aneuploid DNA stem lines than those which are p53 negative (Carder et al., 1993). This result is in keeping with a relationship between p53 and genomic instability. We also reported that in carcinomas containing stabilised p53 the abnormality was almost always present throughout the tumour even though DNA analysis revealed the presence of divergent stem lines. From these findings we concluded that p53 stabilisation is a critical early event in cancer evolution favouring the development of tetraploid and other aneuploid sublines. Since it is now clear that p53 stabilisation can be the result of the tumour cell environment, independent of mutation (Vojtsek and Lane, 1993), we felt it was important to re-examine these cases for more definitive evidence, firstly, that mutation of p53 had occurred and, secondly, that it preceded clonal divergence in tumour progression. To this end we used the technique of single-stranded conformation polymorphism (SSCP) analysis (Orita et al., 1989; Glavac and Dean, 1993) to identify mutant alleles of p53 and confirm that the same mutant allele was present in all of the samples taken from any one carcinoma, despite independent evidence of clonal divergence between these samples.

Correspondence: AH Wylie
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Materials and methods

Samples
Sixty-four samples from 17 colorectal carcinomas from our previously published series (Carder et al., 1993) were studied. An average of four samples per tumour (range 2-5) were analysed and compared with normal colonic mucosa from the same individual. All samples of all tumours contained stabalised p53 as determined by immunohistochemistry using PAB 1801 (Oncogene Science). Flow cytometry was performed on frozen tissue and immunohistochemistry was performed on tissue fixed in periodate lysine paraformaldehyde (PLFD), both as described previously. To minimise possible confusion between tumour and non-neoplastic stroma, samples were assessed as diploid only if tumour cells occupied in excess of 50% of tissue sections from which the corresponding flow cytometric analysis showed a diploid main peak. Samples were considered aneuploid if a separate peak, distinct from the diploid peak, was identified, and aneuploid samples from different populations of the same tumour were considered identical unless they differed in DNA index by more than 0.1. Samples were considered tetraploid if a separate peak with DNA index between 1.9 and 2.1 comprised more than 10% of the nuclei.

Single-strand conformation polymorphism (PCR-SSCP) analysis
The polymerase chain reaction (PCR) was performed on 0.1-1 μg of genomic DNA samples, in an 100 μl reaction containing 200 μM of each deoxynucleotide, 50 pmol of each primer and 2 units of a thermostable Tag polymerase in the relevant buffer. PCR was performed in a DNA thermocycler (Hybid) with the following temperature profile: one cycle of 94°C for 5 min, 30 cycles of 94°C for 1 min, 58°C for 1 min, 72°C for 1 min, and one cycle of 72°C for 10 min. The primers were as follows (5' to 3'): exon 5 up TTCCTCTTGCTCAGTACGTC and 5 reverse CGATGTTGAGCGC- TTGGG; exon 6 up CCTACTGATTGCTCTTAAAG and 6 reverse CTGGAGCTGTGGTGAAC; exon 7 up TGTGTTAATCCTCAAGGTTTG and 7 reverse GTCAGGACCCACTTGCC; exon 8 up TTCATCTCGATGGTTGCT and 8 reverse CGAGTTAGCGAAGCGAG. The
samples were extracted once with 24:1 chloroform/isoamylalcohol to remove any mineral oil and 5–10 μl was then denatured in 80 μM sodium hydroxide, 10 μM EDTA, at 48°C for 5 min. Two microlitres sequencing stop solution was added, and the whole sample loaded onto a 5% glycerol, 0.5× MDE Hydroblot gel (Hoefer Scientific). The gel was run in 1× TBE on the SE400 PAGE apparatus (Hoefer Scientific) at 25°C, 20 W, for 2–3 h.

Normal and tumour samples from the same individual were run together for comparison. Tumour samples were classified as SSCP positive if a discrete additional band was observed. Bands were visualised by a silver stain (BioRad) as per the manufacturer’s instructions with additional washes. The gel was dried on to 3MM paper and laminated.

**Sequence analysis**

A total of nine samples from two cases with band shift on SSCP throughout were sequenced by the dideoxy chain-termination technique as described previously (Cripps et al., 1994) using a commercial kit (Sequenase, Amersham).

**Restriction enzyme digestion using MspI**

A total of eight samples from two cases with band shift on SSCP throughout and known to contain a CGG→CAG mutation in codon 248 were analysed by restriction enzyme digestion using MspI. As described previously (Cripps et al., 1994), disruption of the CCGG recognition site by mutation results in loss of the 135 bp and 168 bp bands and creates an additional 303 bp band.

**Results**

Seventeen carcinomas which had been sampled at multiple sites and demonstrated immunohistochemical positivity for PAb 1801, and for which DNA ploidy values were available, were studied. In 14 cases the majority of nuclei stained intensely. Three cases showed the ‘mosaic’ pattern, in which positive nuclei were scattered sparsely throughout the tumour. Twelve cases contained multiple clonally distinct subpopulations as determined by assessment of DNA ploidy by flow cytometry, with eight containing two and four containing three variant subpopulations.

From each tumour sample exons 5–8 of the p53 gene were amplified individually and analysed by SSCP. Band shifts indicating the presence of a mutant p53 allele with altered conformation were detected by SSCP in 10 of the 17 carcinomas (59%). An average of four samples per tumour were studied, and the same band shift was present in all samples of each SSCP-positive case, suggesting a single clonal mutation event (Figure 1). The presence of a mutation in the amplified p53 fragment revealing the band shift was confirmed in nine samples of two cases by direct nucleotide sequence analysis. Six samples from one case contained a CGC→CAC mutation in codon 175 and three samples from the other case contained a CAT→TAT mutation in codon 179 (Figure 2). In two further cases a mutation in codon 248 was confirmed in a total of eight samples using the MspI restriction enzyme digestion technique (Figure 3). In all, five mutations occurred in exon 5, four in exon 7 and one in exon 8. Six cases with SSCP-confirmed mutations contained multiple divergent populations as detected by assessment of DNA ploidy, indicating clonal evolution subsequent to mutation (Figure 4). All samples of the SSCP-negative cases were concordant for absence of band shift, confirming that mutation had not occurred (at any rate in this commonly affected part of the p53 gene) during carcinoma progression in these cases. All cases with mutation demonstrated strong positive staining for PAb 1801 in a majority of tumour cell nuclei, but four cases with similarly strong staining were SSCP negative in exons 5–8. The relationship between immunocytochemistry and mutation is described more fully elsewhere (Cripps et al., 1994), in a larger number of cases including the present 17.

![Figure 1](image1.png)  
**Figure 1** Identical band shifts are revealed by SSCP analysis in all divergent subpopulations of a colorectal carcinoma. N, normal tissue; T1–3, multiple samples from the same carcinoma (DNA index: T1, 1.5/1.6; T2, 1.5; T3, 1.6).

![Figure 2](image2.png)  
**Figure 2** Dideoxynucleotide sequencing conforms a CAT→TAT mutation in codon 179 (exon 5) in all three samples from this tumour.

![Figure 3](image3.png)  
**Figure 3** MspI restriction digestion of exons 7–9 confirms a mutation in codon 248 in all tumour samples. Loss of the CCGG recognition site creates an additional 303 bp band and results in loss of the smaller 135 bp and 168 bp bands. The smaller bands are due to contaminating normal tissue.


| No. | D | T | A₁ | A₂ |
|-----|---|---|----|----|
| 1   |   |   | O  |    |
| 2   |   |   | O  | O  |
| 3   |   |   | O  | O  |
| 4   |   |   | O  | O  |
| 5   |   |   | O  | O  |
| 6   |   |   |    |    |
| 7   |   |   |    |    |
| 8   |   |   |    |    |
| 9   |   |   |    |    |
| 10  |   |   |    |    |
| 11  |   |   |    |    |
| 12  |   |   |    |    |
| 13  |   |   |    |    |
| 14  |   |   |    |    |
| 15  |   |   |    |    |
| 16  |   |   |    |    |
| 17  |   |   |    |    |

**Figure 4** Relationship between p53 mutation and clonal evolution. Over all sampled stem lines from any one tumour, band shifts on SSCP analysis of exons 5–8 were either uniformly present (closed symbols) or uniformly absent (open symbols): o, diploid; O, tetraploid; other symbols aneuploid with different aneuploid stem lines within the same tumour being designated by different stem lines. D, diploid; T, tetraploid: A₁ and A₂, aneuploid.

**Discussion**

We have used the technique of single-stranded conformation polymorphism (SSCP) analysis to detect clonality of p53 mutation in colorectal carcinomas. SSCP provided a sensitive method of detecting mutations, as mutant alleles generate characteristic band shifts on electrophoresis under non-denaturing conditions (Orita et al., 1989). The technique is sufficiently powerful to allow distinction between sequences differing in a single base pair (Cripps et al., 1994). While in theory one might expect a mutant allele to create at least two additional conformers (one from each strand), often only one is observed, and it appears that most new conformers derive from the purine-rich strand (Glavac and Dean, 1993).

Using SSCP we have demonstrated identical band shifts in the same exon in subclones of carcinomas that diverge in DNA ploidy, and also in carcinomas which are homogeneously diploid or aneuploid. We interpret this to indicate that p53 mutation occurs prior to divergence of clones differing in DNA ploidy, and hence any one tumour possesses a single p53 mutation throughout. In a large series of carcinomas reported separately (Cripps et al., 1994) we have confirmed by sequencing that band shifts detected by SSCP invariably denote mutation. And that the technique used here identifies more than 70% of all naturally occurring mutations in the region of p53 studied (exons 5–8). It may be argued that identical band shifts might still represent different point mutations within the same amplified fragment. We feel that this interpretation is unlikely for three reasons. Firstly, even minor differences in nucleotide substitution (e.g. in adjacent positions of the same codon) can lead to profound differences in band shift by SSCP analysis using the methods employed here (Cripps et al., 1994). Secondly, in the now considerable literature on p53 mutations in cancer, the occurrence of two distinct mutations in the same exon in any one tumour is uncommon. Finally, we have used direct sequencing to confirm identity of mutation throughout in two cases, and in a further two we have used a rapid restriction enzyme digestion technique to confirm identity of mutation site.

The observation that an identical mutation in p53 occurs throughout affected carcinomas, including those with and without clonal divergence in DNA ploidy, provides strong evidence for a single mutational event early in the development of these cancers. Around 20–30% of all colorectal carcinomas containing immunohistochemically stable p53 appear not to contain p53 mutations (Wynford-Thomas, 1993; Baas et al., 1994; Cripps et al., 1994) but examples of these in this series also showed divergent subclones concordant in p53 immunocytochemistry. Hence, even the non-mutational abnormalities that affect p53 stability may also be an early event. The data presented here emphasise the central role of wild-type p53 in preventing one type of genomic instability. This instability appears to result from continued DNA replication in the presence of DNA damage (Lane, 1992; Livingstone et al., 1992; Yin et al., 1993) and tends to produce near-tetraploid subclones (Carder et al., 1993). It is interesting that the selective growth advantage afforded by this instability appears to be a feature favouring carcinoma rather than adenoma growth, since p53 mutations are unusual in adenomas. Other lesions involved in colorectal tumorigenesis but unrelated to p53, such as mutation in the DNA repair genes implicated in hereditary non-polyposis colorectal cancer (Peltonäki et al., 1993), also initiate genomic instability, albeit of a different type, and associate preferentially with carcinoma rather than adenoma.

In conclusion we establish here by DNA analysis the impression gained from our previous immunohistochemical study: in colorectal tumorigenesis p53 mutation is a critical early lesion occurring as a single clonal mutational event which precedes and probably facilitates the emergence of divergent aneuploid tumour subpopulations.

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