

Development of a Facile Fluorescent Assay for the Detection of 80 Mutations Within the p53 Gene

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Accepted March 3, 2002

Abstract

Background: Alterations in the p53 tumor suppressor gene constitute one of the most frequent genetic events associated with the development of human cancers. Determination of an individual's p53 status may be of value in early diagnosis, prediction of response to treatment, and for the detection of minimal residual cancer. Recent studies have also revealed that specific mutations affecting the p53 gene are associated with a poor outcome. The majority of tumor biopsies that are sent for study in the laboratory contain neoplastic cells intermingled with stroma, such that the detection of alterations in the p53 gene requires a tumor enrichment technique and/or highly sensitive mutation detection technologies. Thus, it is desirable that a clinically useful assay for detecting point mutations in the p53 gene function in the presence of significant

quantities of wild-type sequence and identify the critical sequence aberrations.

Materials and Methods: We utilized molecular beacons in a real-time allele-specific PCR format to obtain reference data on samples of quantitatively known p53 mutation status. These data have been statistically analyzed and the results used to detect p53 mutations, indicating the presence of occult tumor.

Results: We describe validation of a simple, rapid, sensitive, and quantitative ARMS assay for identifying the levels of 80 point mutations within the p53 gene that, when mutated, constitute at least 1% of the total p53 sequences.

Conclusions: The assay successfully identifies rare p53 gene mutations in clinical samples and overcomes many of the limitations of current technologies.

Introduction

Mutation of the p53 tumor suppressor gene is one of the most frequently reported events in neoplastic cells, occurring in approximately 50% of human cancers (1). There have been many reports concerning the utility of p53 gene alterations for tumor diagnosis (2,3) and management of cancer patients (4–8). Such studies have come to the fore as our understanding of the pathways involved in apoptotic cell death and the response to DNA damage has increased. Tumors harboring a subset of mutations in the conserved core region, encoded by exons 5–8, have been associated with reduced survival or poor response to therapy, when compared to mutations that lie outside of this region, or wild-type sequence (7–16).

However, results from studies evaluating specific tumor types, for example squamous tumors in the head and neck region, often conflict with reports indicating that p53 mutations are associated with sensitivity to chemotherapy or radiotherapy (16–22). It is important to recognize that these differences reflect the fact that it is impossible to separate sensitivity to a

given treatment from the overall biological behavior of a tumor (23). Thus, unraveling the relationship between genetic aberrations and response to treatment will require in-depth study of the genetic background and clinic pathologic features before substantive conclusions can be reached.

Mutations affecting the p53 gene have been shown to have diagnostic value for patients with squamous tumors of the head and neck. Recent studies, using a molecular diagnostic tool based on finding the same p53 gene mutation in a primary tumor and the surrounding normal tissue, pronounced tumor-free after conventional light microscopy, detected occult tumor in approximately one-third of all patients tested (2,3,24). In our latest report, molecular evidence of residual cancer was found in both mucosal and deep muscle surgical margins (3). At present, the significance of finding a p53 gene mutation in mucosal margins is unclear, but the presence of this aberration in muscle biopsies clearly identifies residual tumor, which is frequently responsible for the patient's demise. Based on these findings, a multinational study has begun to establish whether the molecular diagnostic identifies more cases with residual disease than conventional light microscopy, since detection of sub-microscopic residual disease at an early stage would enable clinicians to ensure that these patients are targeted to receive adjuvant

treatment at an early stage, when it would be most cost-effective and reduce morbidity and mortality.

At present, the results obtained using the molecular test are not taken into account when planning adjuvant treatment, typically postoperative radiotherapy, and a proportion of cases with occult p53 mutation-positive residual tumor develop a recurrence at the operative site, despite receiving full-dose adjuvant treatment. Whether this treatment fails because these patients have tumors which are clinically aggressive, or because they lack functional p53, or harbor other genetic abnormalities which modulate radiosensitivity, is not clear at present. However, this observation highlights the need for prospective studies to investigate the molecular determinants of response to adjuvant treatments, and also to develop novel approaches to deal with the problem of radio-resistant minimal residual cancer (25).

Current approaches to screen for p53 mutations include immunohistochemistry (IHC) (26), a functional assay of separated alleles in yeast (FASAY) (27) and genetic analyses such as enzymatic or hybridization sequencing, and single-strand conformation polymorphism analysis (SSCP) (28,29,30). The current experiment, to establish the prognostic significance of molecular detection of tumor-positive surgical margins, utilizes FASAY and a p53 phage-plaque assay. In essence, the techniques involve screening p53 cDNA for the presence of a p53 mutation by cloning into an indicator yeast strain, sequencing cDNA to identify the nature of the p53 aberration, and then designing a mutant-specific oligonucleotide probe to detect the presence of the same mutation in phage libraries prepared from surgical margins which are pronounced tumor-free by the histopathologist. Although highly sensitive, this methodology is extremely time consuming and the result is not available for 4–6 weeks. The assay often needs to be repeated to ensure sensitivity and specificity in cases where only low levels of mutant are detected. Critical decisions about the need for adjuvant treatment need to be made at an early stage; it is well-recognized that early repopulation of the tumor at the operative site contributes to local relapse (31). Any assay must be sufficiently sensitive to directly identify mutations in the presence of significant quantities of normal tissue, preferably incorporate closed tube technology to avoid cross-contamination of patient samples, be automatable to increase sample throughput, and be sufficiently rapid to provide quantitative data to aid post-operative patient management.

We developed a panel of sensitive assays for specific p53 mutations using real-time allele-specific amplification (ARMS). ARMS is a powerful technique for detecting mutations and polymorphisms in DNA (32). Primers are designed to match the mutant sequence at their 3' termini. Facile amplification results from a mutant template and significantly inhibited amplification from a 3' mismatched, wild-type sequence. In a real-time PCR assay, the

production of PCR product results in a fluorescent signal that is monitored throughout PCR. The threshold cycle (Ct)—the point at which signal becomes distinguishable from background noise—indicates the input copy number of target sequence (33). In a real-time ARMS reaction, the Ct is also affected by the proportion of the sample matched to the 3' end of the primer. Therefore, by performing both a PCR and an ARMS reaction on a sample, the proportion of mutant sequence present can be quantified. For any given control Ct (indicating total copy number) the proportion of mutant sequences can be determined by comparing the ARMS Ct for the sample with reference data from samples of known composition.

In the present study, we employed a molecular beacons format (34) to generate sequence-specific signals during thermal cycling. Molecular beacons are single-strand nucleic acids composed of self-complementary sequences flanking a central region. The central region is designed to be complementary to sequences anticipated to be produced on amplification. In the absence of specific amplification product, the flanking regions remain self-hybridized and hold a terminal fluorophore and quencher molecule in close proximity. In the presence of the amplification, product a more stable interaction between it and the central region of the beacon predominates and physically separates fluorophore and quencher. The reduction in fluorescent resonance energy transfer results in an increase in fluorescent signal. We used this methodology to produce an assay for the 80 mutations occurring in the p53 gene. The assay covers 42% of reported mutations, insertions, and deletions in head and neck cancer (see Table 1 and [35]). We describe the production of the assay and the acquisition and analysis of reference data that enable the rapid, quantitative, and sensitive detection of specific p53 mutations in clinical samples. We demonstrate the use of the assay for detecting minimal residual cancer in biopsies of normal tissue following surgical ablation of squamous tumors of the head and neck. The assay may be of widespread utility as a sensitive marker of cancer cells and for patient stratification based on the functional consequences of the p53 mutation detected.

Materials and Methods

Instrumentation

PCR reactions without real-time monitoring were carried out on a PE 480 thermal cycler (Applied Biosystems, Foster City, CA, USA). PCR reactions with real-time fluorescent monitoring were carried out in optical thermal cycling plates on the ABI prism 7700 thermal cycler (Applied Biosystems).

Reagents and Solutions

DNA size marker (50 bp) and dNTPs were from Pharmacia Biotechnology (Piscataway, NJ, USA), DNA polymerase (AmpliTaQ Gold) from P.E. Applied

Table 1. Sequences of ARMS primers used to detect 80 mutations in the p53 gene

Well	Codon	Change	Frequency	ARMS Primer (5' to 3')	Reverse PCR Primer/ Molecular Beacon
3	132	AAG to AGG	0.357	ggt ctt gac cag ttg gca aaa cat tc	861-98/5aU
4	132	AAG to CAG	0.071	tcc tac agt act ccc ctg ccc tca tcc	862-98/6aL
5	135	TGC to TAC	0.071	gta ctc ccc tgc cct cac caa gat gtt gta	862-98/6aL
6	141	TGC to TAC	0.143	aac aag atg ttt tgc caa ctg gcc aag aca ta	862-98/6aL
7	151	CCC to TCC	0.214	tgg cgt gga tgc ggg tgc cgg gcg gag a	861-98/5aU
8	151	CCC to ACC	0	cct gtg cag ctg tgc gtt gat tcc ata a	862-98/6aL
9	152	CCG to CTG	0.143	gca gct gtg ggt tga ttc cac acc tct	862-98/6aL
10	156	CGC to CCC	0.071	cca cac ccc cgc ccg gca ctc c	862-98/6aL
11	157	GTC to TTC	0.929	aca ccc ccg ccc ggc acc ctc t	862-98/6aL
12	158	CGC to CTC	0.286	cac acc ccc tcc cgg cac ccg cgt gct	862-98/6aL
15	158	CGC to CAC	0.571	gat tcc aca ccc ccg ccc ggc acc ctc gta ca	862-98/6aL
16	161	GCC to ACC	0.143	gcc agg cac ccg cgt ccg cgc cag ga	862-98/6aL
17	163	TAC to TGC	0.429	tcc gtc atg tgc tgt gac tgc tgg c	861-98/5aU
18	173	GTG to ATG	0.143	cta caa gca gtc aca gca cat gac gga gga ta	862-98/6aL
19	173	GTG to TTG	0.357	aca agc agt cac agc aca tgc cgg agg att	862-98/6aL
20	175	CGC to CAC	5.071	cac agc aca tga cgg agg ttg tga ggg a	862-98/6aL
21	175	CGC to AGC	0	cac agc aca tga cgg agg ttg tga cga	862-98/6aL
22	175	CGC to TGC	0.071	cac agc aca tga cgg agg ttg tga agt	862-98/6aL
23	175	CGC to CCC	0	aca gca cat gac gga ggt tgt gag ccc	862-98/6aL
24	175	CGC to CTC	0	aca gca cat gac gga ggt tgt gag tct	862-98/6aL
27	176	TGC to TAC	0.286	cat gac gga ggt tgt gag gcg gta	862-98/6aL
28	176	TGC to TTC	4.214	gca cat gac gga ggt tgt gag gcg gtt	862-98/6aL
29	177	CCC to CGC	0	cat gac gga ggt tgt gag gcg ctg ctg	862-98/6aL
30	179	CAT to CGT	0.429	agg ttg tga ggc gct gcc ccc act g	862-98/6aL
31	179	CAT to TAT	0.500	tga cgg agg ttg tga ggc gct gcc ccc att	899-97/6aL
32	192	CAG to TAG	0.214	gat tgc tct tag gtc tgg ccc ctc ttt	694-98/6bL
33	193	CAT to CGT	0.286	gct ctt agg tct ggc ccc tcc tca ccg	694-98/6bL
34	195	ATC to ACC	0.500	ctt agg tct ggc ccc tcc tca gca tct gac	694-98/6bL
35	196	CGA to TGA	1.000	ata ctc cac acg caa att tcc ttc cac aca	861-98/5aU
36	205	TAT to TGT	1.000	cga aaa gtg ttt ctg tca tcc aat c	863-98/6aL
39	213	CGA to TGA	0.786	gga gta ttt gga tga cag aaa cac ttc tt	694-98/6bL
40	220	TAT to TGT	1.571	aca tag tgt ggt ggt gcc gtg	694-98/6bL
41	228	GAC to AAC	0	gcc tgt gtt atc tcc tag gtt ggc tgt a	644-98/7aL
42	234	TAC to TGC	0.714	ccc atg cag gaa ctg tta cac atg tag atc c	643-98/7bU
43	237	ATG to ATA	0.571	gac tgt acc acc atc cac tac aac tac aaa	644-98/7aL
44	238	TGT to TAT	0.214	ctg tac cac cat cca cta caa cta cat cta	644-98/7aL
45	241	TCC to TTC	0.214	tcc agt tca tgc cgc cca tgc aca	643-98/7bU
46	242	TGC to TTC	0.500	acc atc cac tac aac tac atg tgt aac agt tct tt	644-98/7aL
47	244	GGC to TGC	0.214	cta caa cta cat gtg taa cag ttc ctg cat ct	644-98/7aL
48	245	GGC to AGC	1.714	tac gtg tgt aac agt tcc tgc atg gtc a	644-98/7aL
51	245	GGC to GAC	0.786	cat gtg taa cag ttc ctg cat ggg cta	644-98/7aL
52	245	GGC to TGC	0.429	act aca tgt gta aca gtt cct gca tgg tct	644-98/7aL
53	245	GGC to GTC	0.214	cat gtg taa cag ttc ctg cat ggg agt	644-98/7aL
54	245	GGC to CGC	0	aca tgt gta aca gtt cct gca tgg ccc	644-98/7aL

(Continued)

Table 1. (Continued)

Well	Codon	Change	Frequency	ARMS Primer (5' to 3')	Reverse PCR Primer/ Molecular Beacon
55	245	GGC to GCC	0	cat gtg taa cag ttc ctg cat ggg agc	644-98/7aL
56	248	CGG to CAG	2.571	aca gtt cct gca tgg gcg gca tga aac a	644-98/7aL
57	248	CGG to TGG	2.071	aac agt tcc tgc atg ggc ggc atg aag t	644-98/7aL
58	248	CGG to CTG	0.357	taa cag ttc ctg cat ggg cgg tat gaa tct	644-98/7aL
59	248	CGG to GGG	0.143	gta aca gtt cct gca tgg gcg gcg tga agg	644-98/7aL
60	248	CGG to CCG	0.071	tgt aac agt tcc tgc atg ggc ggc atg aac ac	644-98/7aL
63	249	AGG to AGT	0.357	ttc ctg cat ggg cag cat gaa ccg gtg t	644-98/7aL
64	249	AGG to ATG	0	ttc ctg cat ggg cga cat gaa ccg tat	644-98/7aL
65	249	AGG to AGC	0	gca tgg gcg gca tga acc gga ac	644-98/7aL
66	249	AGG to GGG	0.286	ttc gag tgt gat gat ggt gag gat ggg cgc	643-98/7bU
67	249	AGG to TGG	0.214	ttg cag tgt gat gat ggt gag gat ggg aca	643-98/7bU
68	249	AGG to ACG	0.071	aac agt tcc tgc atg ggc gac atg aac cgg tc	644-98/7aL
69	249	AGG to AAG	0	atg ggc agc atg aac cga aa	644-98/7aL
70	258	GAA to AAA	0	ggc cca tcc tca cca tca tca cac tta	644-98/7aL
71	258	GAA to GGA	0.071	cca tcc tca cca tca tca cac tgc g	644-98/7aL
72	266	GGA to GTA	0	tcc tat cct gag tag tgg taa tct act tgt	629-98/8aL
75	266	GGA to AGA	0.286	ttc cta tcc tga gta gtg gta atc tac gga	629-98/8aL
76	266	GGA to GAA	0.214	tcc tat cct gag tag tgg taa tct act gaa	629-98/8aL
77	272	GTG to ATG	0	atc tac tgg gac gga aca gct ttg tga	629-98/8aL
78	273	CGT to CAT	2.429	ggg acg gaa cag ctt tga ggt gaa	629-98/8aL
79	273	CGT to TGT	1.143	act ggg acg gaa cag ctt tga ggt ct	629-98/8aL
80	273	CGT to CTT	0.714	tac tgg gac gga aca gct ttg agg tac t	629-98/8aL
81	273	CGT to AGT	0	tac tgg gac gga aca gct ttg agg aga	629-98/8aL
82	273	CGT to GGT	0.143	tac tgg gac gga aca gct ttg agg agg	629-98/8aL
83	273	CGT to CCT	0.071	act ggg acg gaa cag ctt tga ggt gac	629-98/8aL
84	275	TGT to TAT	0.357	acg gaa cag ctt tga ggt gcg tgt tga	629-98/8aL
87	278	CCT to CTT	0.214	aac agc ttt gag gtg cgt gtt tgt gcc tgt gt	629-98/8aL
88	278	CCT to TCT	0.786	gaa cag ctt tga ggt gcg tgt ttg tgc cta tt	629-98/8aL
89	280	AGA to ACA	0.071	ggt gcg tgt ttg tgc ctg tcc tgg aac	629-98/8aL
90	282	CGG to TGG	1.643	tgc gtg ttt gtg cct gtc ctg gga gag agt	629-98/8aL
91	282	CGG to CCG	0	gtg cgt gtt tgt gcc tgt cct ggg aga gac gc	629-98/8aL
92	282	CGG to GGG	0.071	ggt gcg tgt ttg tgc ctg tcc tgg gag agt cg	629-98/8aL
93	282	CGG to CAG	0	gtg cgt gtt tgt gcc tgt cct ggg aga gac ga	629-98/8aL
94	285	GAG to AAG	0.571	gcc tgt cct ggg aga gac cgg cgc aga a	629-98/8aL
95	286	GAA to AAA	0.214	tcc tgg gag aga ccg gcg cac agc ga	629-98/8aL
96	298	GAG to TAG	1.000	ccg caa gaa agg gga gcc tca cca gt	629-98/8aL

Key to legend:

Well = position on assay plate,

codon = p53 codon

change = mutation in p53 gene sequence

Frequency = frequency of mutation occurrence in p53 database [34] as a percentage of all mutations reported for head and neck cancer (excluding data from cell lines)

ARMS primer = primer sequence used to detect mutation

Reverse PCR Primer/Molecular beacon = reverse PCR primer used with ARMS primer and molecular beacon used to detect PCR amplification (see table 3 for sequences).

Biosystems, NuSieve Agarose from FMC Bioproducts (Rockland, MD, USA), and X174 RF DNA HaeIII digest molecular size marker from Life Technologies. Spin columns for PCR product purification were from Qiagen (Hilden, Germany). All other chemicals were from Aldrich Chemical Company Co. (Milwaukee WI). Molecular beacons were prepared by Oswel Research Products (Southampton UK). The remaining oligonucleotides were synthesized on a P.E. Applied Biosystems 394 DNA/RNA synthesizer with P.E. Applied Biosystems reagents according to the manufacturers protocols and were purified by size exclusion filtration through NAP-10 columns (Pharmacia). ARMS buffer consists of 1.2 mM MgCl₂, 10 mM Tris-HCl, 50 mM KCl, pH 8.3, and cassette dilution buffer consisting of 10 mM Tris-HCl, 50 mM KCl, and 1g/L BSA, pH 8.3.

Samples

Ethical Committee Approval for the minimal residual cancer study was granted at King's College Hospital. Biopsies of primary oral SCCs were harvested at the time of primary surgery and stored at -70°C until required. Following excision of the primary tumor, four mucosal and two deep surgical margins, each approximately 1 cm × 8 mm × 8 mm were harvested. Each margin was divided into two; the inner portion was processed for conventional light microscopy, the outer half frozen for molecular analysis. Cases reported as having tumor-positive margins by the histopathologist were not forwarded

for molecular analysis. All samples were coded and analyzed blind.

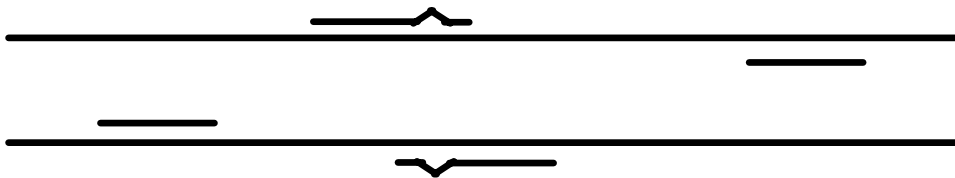
DNA Preparation

DNA was prepared from tumor frozen portion of the surgical margins using DNA STAT 50 (Biogenesis, Poole, UK) according to the manufacturer's instructions.

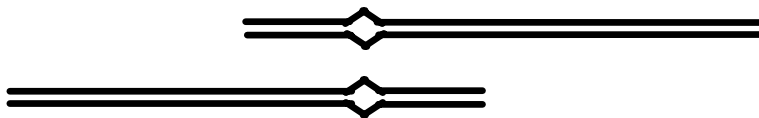
Template Production for Real-Time Analyses

A multiplex PCR was performed on neat and 1 in 1000 diluted DNA samples derived from peripheral blood or tumors to amplify 477-, 236-, and 415-bp fragments incorporating p53 exons (5,6), 7 and 8,9, respectively. The following primers were used in combination at the final concentrations shown; 5' tga ctt tca act ctg tct cct tc (419 nM), 5' acc cgg agg gcc act gac aac (419 nM), 5' cac agg tct ccc caa ggc gca c (93 nM), 5' ggg ccc agg ggt cag cgg caa (93 nM), 5' ttc ctt act gcc tct tgc tc tct (488 nM), and 5' aag aag aaa acg gca ttt tga gtg (488 nM). The PCR mix also contained dNTPs (100 mM), Taq DNA polymerase (1 unit), and the appropriate DNA template (2.5 μl) in ARMS buffer (total volume 25 μl). Reactions were cycled 1× (94°C for 20 min), 40× (94°C for 1 min, 60°C for 1 min), 1× (72°C for 10 min). Mutant p53 fragments or cassettes were prepared by the method of Higuchi (37) (Fig.1). The cassettes were examined by agarose gel electrophoresis to a) check that a single fragment of the correct size had been prepared and b) permit approximate quantitation of the cassette

1st Round. Synthesis of mutant half cassettes in two tubes



2nd Round. Megaprimering of half cassettes to produce full length mutated template



2nd Round. Nested primers produce final cassette



Final mutated cassette



Fig. 1. Preparation of synthetic mutated templates. The method is based on the method of Higuchi et al. (37). In the first round, two fragments are created in separate reactions, each harboring the primer-induced mutation. In the second round, overlapping strands from the two fragments hybridize and produce a cassette with the mutation in the center. Nested primers, also in the second round, then amplify the desired product.

Table 2. Sequences of control primers used in the p53 assay

Wells	Exon	Primer (5' to 3')	Reverse PCR Primer/ Molecular Beacon
1,2,13,14	6	ccc agg gtc ccc agg cct ctg att	862-98/6aL
25,26,37,38	6	ccc agg gtc ccc agg cct ctg att	694-98/6bL
49,50,61,62	7	cac agg tct ccc caa ggc gca c	644-98/7aL
73,74,85,86	8	ttc ctt act gcc tct tgc ttc tct	629-98/8aL

Key to legend:

Well = position on assay plate,

exon = p53 exon

Primer = Forward primer sequence used to perform control reaction

Reverse PCR Primer/Molecular beacon = reverse PCR primer used with forward primer and molecular beacon used to detect PCR amplification (see table 3 for sequences).

by comparing the band intensities with those of the ?X174 RF DNA HaeIII digest, which contains known amounts of DNA. Cassettes were diluted to a working concentration in cassette dilution buffer and admixtures of mutant cassettes in a wild-type p53 multiplex product background were created by mixing samples with equivalent Cts in control real-time PCRs.

Design of ARMS Primer and Molecular Beacons

ARMS and reverse primers (Tables 1 and 3) were designed to detect the 80 prevalent mutations (excluding cell lines) of human p53 in the database maintained by T. Soussi (35). Each ARMS reaction could be correlated with a further mutation independent control PCR (Tables 2 and 3), which utilized the same multiplex product as template. The ARMS primers incorporate additional mismatches at position -2 or -3. Molecular beacons (34) were designed to anneal to sequences between amplification primer sites. The open and closed forms of the beacons were modeled for stability on the website of Michael Zuker (38). Beacons generally comprised 5-mer self-complementary hairpins and 24–27 base central probe regions.

Real-Time Amplification Conditions

Each of the 96 wells on the optical plates contained a reverse and a control or ARMS primer at 500 nM and a molecular beacon at 400 nM (800 nM for beacon 6bL) (see Tables 1 and 2 for plate configuration). In addition, each well contained 100 nM dNTPs, 60 nM Rox passive reference (Perkin Elmer, Warrington, UK), 1× ARMS buffer, 1 U of AmpliTaq Gold, and DNA template in a total volume of 25 µl. For use on unknown samples, plates containing a 2× concentrate of well-specific reagents were stored frozen and a mixture of template and common reagents added by multipipette or immediately prior to use. Plates were cycled as follows: 1× (94°C for 20 min), 50× (94°C

for 45 sec, 60°C for 45 sec). Rox on and quencher off were selected and FAM signals were collected during the 60°C-annealing phase.

Preparation of 80 Reference Data Sets

Data points were collected, upon independent runs on two separate 7700 machines, for each of the 80 ARMS and 4 control reactions on dilutions of samples of known composition. Wild-type multiplex templates from 35 individuals were used. Mutant data, a minimum of nine independent data points, were generally

Table 3. Sequences of reverse primers and molecular beacons

ID	Sequence (5' to 3')
629-98	gaa tct gag gca taa ctg cac cct t
643-98	cac agg tct ccc caa ggc gca c
644-98	agc ggc aag cag agg ctg ggg c
861-98	tga ctt tca act ctg tct cct tc
862-98	tct gtc atc caa ata ctc cac acg caa
694-98	acc cgg agg gcc act gac aac
5aU	ggc cgc tct tcc tac agt act ccc ctg ccc ggc c
6aL	cgc ccg gat aag atg ctg agg agg ggc cag ggc g
6bL	cgc cca gtt gca aac cag acc tca ggc ggg ggc g
7aL	cgc ccc aag tgg ctc ctg acc tgg agt ctg ggc g
7bU	ccg gcg tgt tat ctc cta ggt tgg ctc tga ctg ccg g
8aL	cgc cgc acc gct tct tgt cct gct tgc ttc ggc g

Key to legend:

ID = primer identifier

Sequence = reverse primer and molecular beacon sequences. All molecular beacons contain 5' FAM (6-carboxyfluorescein) fluorophore and 3' methyl red quencher modifications. Self-complementary stem sequences are highlighted in bold.

obtained from dilutions originating from one mutant cassette stock. Data were analyzed on the 7700 using the ROX-off setting, the ROX signal being used only to confirm template and reagent delivery. The cycle threshold was obtained for each well and the Ct of ARMS reactions correlated to the Ct of the corresponding control reaction to give a single well-specific data point. Data sets arising from the dilutions of wild-type and mutant templates were analyzed by linear regression to calculate the lines of best fit and 95% confidence intervals for each of the 80 tests on 1% mutant and wild-type templates. The resultant data were used to construct a diagnostic MS Excel macro that extracts the Ct of control reactions from an unknown sample and generates confidence limits around the expected ARMS reaction Ct for known wild-type and mutant harboring samples using the 80 reference data sets. For proof of principle with this prototype assay, the calculations assumed that the data obtained were representative of a continuous population whose y variance is independent of x . The macro then simultaneously plots the observed and confidence limits for Ct from each of the 80 tests.

Assay of Primary Tumor and Surgical Margins

Real-time analyses were only performed on multiplex products from samples for which >1000 genomic copies had been input to the multiplex (multiplex functional on 1/1000 dilution of sample). Multiplex products from samples were diluted to add an estimated 10^5 copies per well and subjected to real-time amplification as described. The 80 test-specific data points arising from each sample were analyzed with the diagnostic macro. Mutations were called where the ARMS Ct was earlier than both the lower confidence limit for a wild-type sample and the upper confidence limit for a sample harboring 1% mutant sequence. The results of the real-time assay were compared with those obtained using the FASAY/phage plaque approach (3).

Results

p53 Multiplex

Three product bands of the expected sizes were clearly observed on agarose gel electrophoresis subsequent to multiplex amplification from all neat and 1/1000 dilutions of genomic samples. The ability to produce PCR products from 1/1000 dilutions of sample demonstrates that sufficient copies of DNA were present in the neat sample to ensure (a) no false negatives are called due to insufficient DNA and (b) no false positives arise at the 1% level due to polymerase error in the initial round of amplification.

Creation of Reference Data Sets

Each of the 80 ARMS tests gave test (y) Ct data that were plotted against mean control (x) Ct data from which lines of best fit and confidence intervals were calculated. Validation data from one of the 80 tests are illustrated in Figure 2. Each of the four control reactions were present in quadruplicate on every 96-well plate. To eliminate variability due to any preferential amplification of individual multiplex products, ARMS reaction Ct were always correlated with control PCR, which employed the same multiplex fragment as substrate. Data from cassettes harboring the same mutation, but of different lengths, independently created from different DNA stocks and primers, were indistinguishable (results not shown). The reference data sets permit the control Ct obtained from an unknown sample to be used to predict the ranges of expected Ct from each of the 80 ARMS tests if the sample were to be wild-type or contain 1% mutant sequence.

Use of the Assay on Tumor and Surgical Margin Samples Derived from Patients with Head and Neck Cancer

Real-time data obtained in the four control reactions (four replicates of each) on a clinical sample were used to generate 95% confidence intervals for

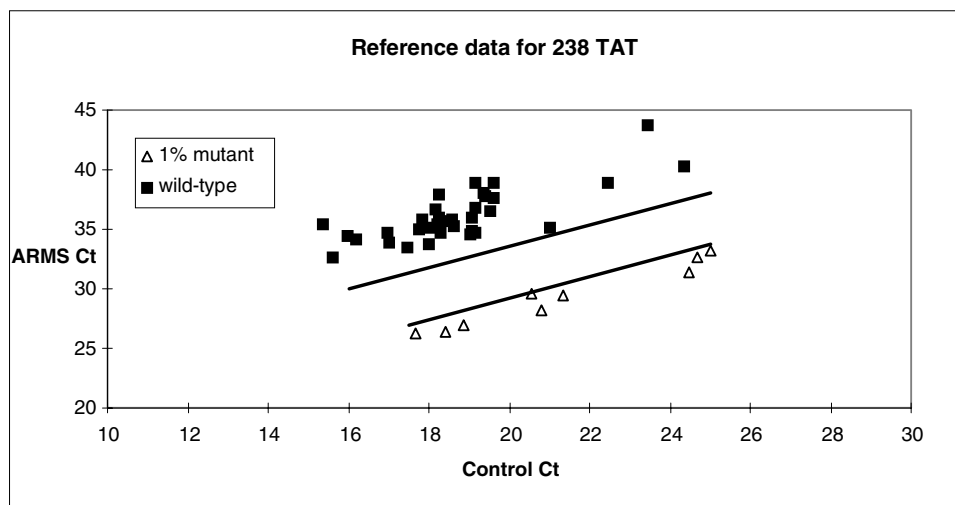


Fig. 2. Graph of example test data with confidence limits. Data were produced from wild-type DNA and samples harboring 1% mutant p53 sequences (238 tat). Each data point was acquired from a separate amplification reaction plate. The solid lines represent the lower and upper 95% confidence limits for the lines of best-fit for the wild-type and mutant data, respectively. For simplicity the data were assumed to be representative of a continuous population whose y variance is independent of x .

expected wild-type and 1% mutant Ct values for the 80 ARMS reactions run on the same 96-well plate. Sequence sites were diagnosed as wild-type unless the ARMS Ct was both excluded from the 95% wild-type confidence interval and included in the 1% mutant 95% confidence interval. In certain tests, where ARMS specificity was at its lowest, the 95% confidence intervals for wild-type and 1% mutant samples overlapped slightly. In these instances, a small proportion of samples harboring p53 mutations as 1% of total p53 sequences go undetected. Where ARMS Cts are out of the wild-type range but not early enough to be included within the 1% confidence interval, the multiplex may harbor a mutation at less than 1% of total p53 sequences (see Discussion).

The graphical output from the use of the assay on DNA derived from a representative head and neck tumor and the matched surgical margins is illustrated in Figure 3. The tumor contains a high proportion of p53 sequences harboring the 175 cac mutation (well 20). This mutation is also present in the surgical margin at a much lower level. No other test Ct from the tumor or margin gives a value which is both out of the 97.5% confidence intervals for normal sequence and within the confidence interval for mutant levels of 1% or greater. The results of the FASAY/phage plaque assay were disclosed subsequently and concurred with diagnosis by the real-time assay. The margin was estimated to contain ~2% mutant sequence via the phage plaque assay.

Discussion

The major cause of morbidity and mortality following treatment for head and neck cancer is locoregional recurrence due to persistent or recurrent disease. Microscopic minimal residual cancer at the surgical margin has been shown to correlate with a high locoregional recurrence rate and decreased survival (39,40). However, an ultrasensitive diagnostic test, which depends on finding the same p53 gene mutation in a primary tumor and the surrounding normal tissues considered to be tumor-free after conventional light microscopy, may allow detection of more cases with occult tumor at risk of local recurrence, such that these patients can be targeted to ensure that they receive early adjuvant treatment to eliminate any remaining malignant cells and improve their chance of cure (2,3). However, this approach is a complex procedure with many sources for error. Self-ligation of the vector, due to partially degraded cDNA, is a significant problem. In addition, the protocol is very time consuming, such that incorporation of this technique, together with the p53 phage-plaque assay, is not practical in terms of developing a test that is quick, reliable, and can be automated to increase sample throughput and be used to influence decisions about patient treatment.

Many studies have used IHC detection of p53 as a proxy for detecting mutations in the p53 gene. The significance of such studies is open to debate; interpretation is subjective, about 25% of p53 mutations

do not result in protein overexpression, p53 protein levels may be elevated for other reasons, and the nature of the mutation present is not determined (15,41). These difficulties may account for conflicting data concerning the relationship between immunoreactive p53 and prognosis or response to therapy. Direct DNA sequencing and hybridization techniques will only detect mutant p53 sequences present at levels greater than 25% of the total, unless they are applied to tumor-enriched samples. All tumor enrichment procedures involve a significant time/cost element and include visual selection (15), laser capture microdissection (42), and antibody binding (43). SSCP is compatible with scanning PCR products for mutations in a relatively routine, yet sensitive, manner and is currently the most widely used technique for genetic screening for p53 mutations (29,30). However, SSCP is not quantitative and does not directly identify the mutation responsible for an observed band shift. What is required for routine application is a test that is quick, simple, and reliable, and that is not customized for each patient but is suitable for automation and amenable to quality control.

In the present study we describe a simple, quantitative, nucleic acid-based mutation detection system, capable of identifying specific p53 gene mutations against a background of tissue harboring wild-type alleles, likely to be very valuable in terms of prognostication and treatment planning for these patients. The 80-test panel described detects over 40% of p53 mutations reported in head and neck cancer (Table 1). The method relies entirely on the use of reference data sets to establish statistically validated "fingerprints" of the p53 status of tissues. The production of the reference datasets is relatively cost and labor intensive but, once generated, they provide a powerful diagnostic and quality control for data generated from clinical samples. Confidence in an accurate diagnosis of the composition of the multiplex products is provided by the presence of a Rox passive reference peak in each well, indicating the delivery of enzyme, template, and so on, as well as the conforming of the vast majority of ARMS test Cts to their expected wild-type values.

The assay produces highly reproducible data on any given multiplex product (data not shown) such that the performance of the multiplex is the critical step in the described procedure. It is imperative that the real-time assay data be interpreted in the knowledge of the minimum initial gene copy number input to the multiplex. If no bands are produced in the multiplex from a 1/1000 dilution of tumor or margin DNA, then false negatives arise from the fact that the 1% detection limit is meaningless in the absence of >100 copies, and false positives may arise from the detection of a replication error from early in the amplification process. The multiplex is therefore necessary if samples are to be interrogated at large numbers of sites: for example, a test for 1000 mutations would require a minimum of 1000 ×

1000 = 1,000,000 copies of the gene to add 1000 copies of the gene per test well. This corresponds to a minimum of 3 μg of genomic material, which is unlikely to be forthcoming in many clinical situations. When bands are produced from the multiplex on dilute samples, similar false positives could arise if a simple non-wild-type analysis were performed, as certain ARMS tests can easily detect one mutant allele in the presence of 100,000 wild-type. It is therefore essential that, if data from a single, isolated run are to be interpreted, inclusion in the 1% data set is the criterion defining positivity, ensuring that even replication errors arising in the first round of amplification would not be accorded positive status.

The real-time ARMS assay successfully identified p53 mutations in clinical samples in blind trials. In the example shown, analysis of surgical margins from patients with head and neck cancer for the presence of tumor cells missed by conventional light

microscopy detects the same mutation (175 CAC) in paired tumor and margin samples as independently detected in the phage-plaque assay. A further margin, in which an even lower level of mutant was detected by the yeast assay, was diagnosed as wild-type by the real-time ARMS assay. However, it is pertinent to note that the ARMS assay is highly quantitative, and thresholds may be set subsequent to data collection, such that clinical cut-offs can be optimized at a later date. The question of sufficient sensitivity for clinical utility can only be answered in an experiment in which the quantitative and objective data produced by such assays can be correlated to clinical outcome. The real-time assay described is amenable to use in such experiments because it is extremely easy to use, takes a day from sample to results, and requires less than 15 min of user intervention time to run; assuming stocks of multiplex primers and assay plates. We have also determined

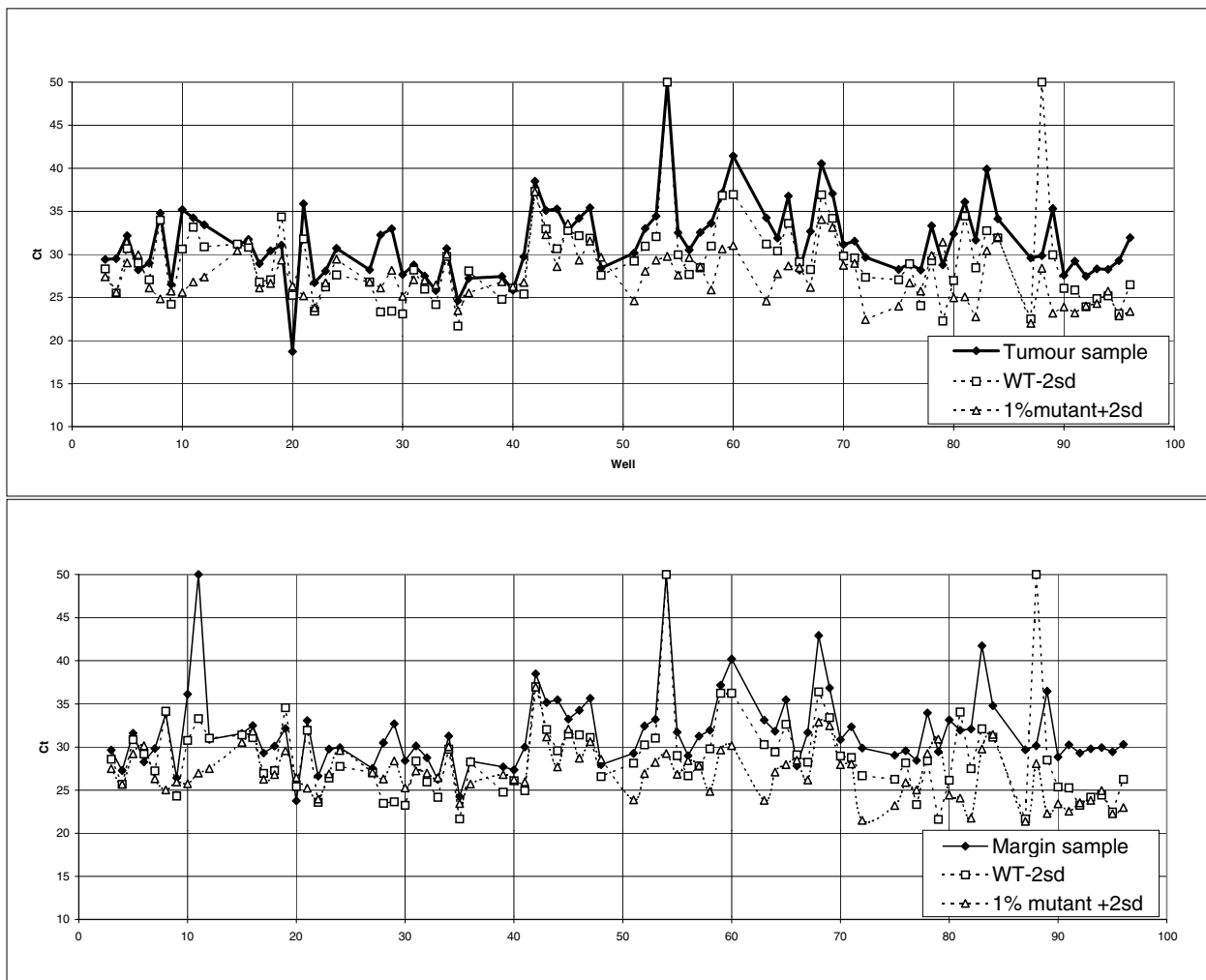


Fig. 3. Use of the assay on tumor and associated margin tissue. The Cts from the tumor and margin fall within the expected 97.5% confidence limits for all of the 80 tests except that in well 20, 175 cac. The mutation is clearly present as a larger proportion of the p53 sequences isolated from the tumor, but is also present within the tumor margin, where its presence implies a high probability of tumor tissue remaining within the patient after surgery.

that real-time assay ready reagent plates can be stored frozen or dried without affecting the data produced (data not shown). In comparison with the yeast assay, the approach reduces the time taken to test a patient sample for residual tumor from 12 to 1 working day, and means that the tumor and each surgical margin can be screened for a p53 gene mutation to produce data sufficiently rapidly to be used to influence postoperative therapy. The closed-tube technology obviates the need for a gel, minimizes the risk of sample contamination, and ensures quality control. The cost of the raw materials used in the assay is around US \$60.00 per sample, but could be reduced considerably by miniaturization onto technology platforms currently reaching the market. The ARMS assay is sensitive, quick, simple, reliable, and robust enough to allow automation. In addition, as the assay uses closed-tube technology, the risk of cross-contamination of patient samples is minimized, a consideration paramount when translating molecular diagnostics from a research to a routine diagnostic setting.

The test panel described detects approximately 42% of p53 mutations reported in head and neck cancer and contains tests for the 80 most prevalent p53 mutations. Increasing the number of tests gives greater coverage of mutations. Table 4 shows that by increasing the number of tests to include the 240 most prevalent mutations, 75% coverage could be obtained. This approach is feasible using a suitable real-time PCR platform capable of handling 384-well plates such as the Applied Biosystems 7900.

Table 4. Coverage of all p53 mutations obtained by increasing the number of tests included in the panel in order of their reported prevalence. Data obtained from the database maintained by T. Soussi, Laboratoire de Genotoxicologie des Tumeurs, Paris [35]

Number of p53 Mutations Tested	Percentage Coverage of all p53 Mutations Observed in Tumours
80	42
90	48
100	51
120	56
160	64
240	75

Key to legend:

Number of p53 mutations tested—Number of mutations included in a test panel.

Percentage coverage of all p53 mutations observed in tumours—Maximum percentage coverage of all p53 mutations that would be obtained by test panel.

In the present example, a mutation affecting the p53 gene is used as a “tumor fingerprint” to detect occult tumor. Such molecular analysis will play an increasingly important role in establishing whether margins are tumor-free or not. However, at this stage further studies are needed to clarify whether any existing and emerging treatments will be more efficacious if patients are stratified on the basis of p53 status. Based on the key position of the p53 protein in cell cycle regulation and current reports (44–48) it seems highly likely that incorporation of assays such as described here into routine clinical practice will enable information about the genetic aberrations associated with tumors to be used to plan treatment as well as providing an aid to diagnosis.

Acknowledgments

We thank Dr. Jayne Fox and Dr. Simon Clayton for helpful discussions. This work was supported by the AstraZeneca Strategic Development Fund.

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