Deletion mapping on chromosome 1p in well-differentiated gastric cancer

T Ezaki1,4, A Yanagisawa2, K Ohta1, S Aiso4, M Watanabe3, T Hibi1, Y Kato2, T Nakajima3, T Ariyama6, J Inazawa6, Y Nakamura1 and A Horii1

Departments of 1Biochemistry, 2Pathology and 3Surgery, Cancer Institute, 1-37-1 Kami-Ikebukuro, Toshima-ku, Tokyo 170; Departments of 4Anatomy, and 5Internal Medicine, Keio University School of Medicine, 35 Shinhonouchi, Shinjuku-ku, Tokyo 160; 6Department of Hygiene, Kyoto Prefectural University of Medicine, Kamigyō-ku, Kyoto 602, Japan.

Summary To define the region on the short arm of chromosome 1 that is thought to include one or more tumour-suppressor genes for gastric cancers, we carried out loss of heterozygosity (LOH) studies in 26 gastric adenocarcinomas, using three restriction fragment length polymorphism (RFLP) markers and nine microsatellite markers. All tumours were informative with at least one locus; three revealed replication errors (RERs) at multiple microsatellite loci, and interstitial or telomeric allelic deletions were observed in 12 cases. Deletion mapping of these tumours defined a commonly deleted region between two loci, D1S201 and D1S197, that are 13 cM apart. As two loci within the commonly deleted region, D1S57 (pYNZ2) and D1S62 (pTH154), were mapped respectively to 1p35 and 1p34.3 by fluorescence in situ hybridisation, we conclude that a locus likely to contain a tumour-suppressor gene for gastric cancer is located within a 13 cM region encompassing these two chromosomal bands.

Keywords: chromosome 1p; LOH; human gastric cancer

Although gastric cancer is the most common malignancy in the world (Parkin et al., 1988), the genetic pathway of gastric carcinogenesis is not well understood. To date only a few genetic alterations associated with gastric cancer have been reported; these include amplifications of the erbB-2 (Yokota et al., 1988) and K-sam (Hattori et al., 1990; Nakatan et al., 1990) genes and mutations of the APC (Horii et al., 1992; Nakatsu et al., 1992), ras (Deng et al., 1987; Kihana et al., 1991) and p53 (Tamura et al., 1991; Strickler et al., 1994) genes. Amplification of the erbB-2 gene and mutations of the APC and ras genes are found frequently in well-differentiated adenocarcinomas but not in poorly differentiated adenocarcinomas (Yokota et al., 1988; Kihana et al., 1991; Nakatsu et al., 1992). In contrast, amplification of the K-sam gene and replication errors (REs) at microsatellite loci have been detected preferentially in poorly differentiated adenocarcinomas (Hattori et al., 1990; Han et al., 1993). These observations imply that the genetic pathways involved in development of these two histopathologically distinguished forms of gastric adenocarcinoma are likely to be different.

Recent results of LOH studies have suggested that loci containing tumour-suppressor genes associated with gastric carcinogenesis exist on chromosome arms 1p, 1q, 5q, 7q, 12q, 17p and 18q (Sano et al., 1991; Uchino et al., 1992; Kuniyasu et al., 1995). As part of a strategy to identify these putative tumour-suppressor genes we began by attempting to define the region on chromosome 1p that is commonly deleted in gastric cancers. Here we report results of LOH studies in 26 well-differentiated adenocarcinomas of the stomach.

Materials and methods

Preparation of samples and DNA

A total of 26 paired samples of tumours and corresponding normal tissues removed from Japanese patients with well-differentiated adenocarcinomas of the stomach were obtained at the Cancer Institute Hospital, Tokyo. In 18 cases the tissue samples were fixed in formalin, embedded in paraffin and attached individually to glass slides. Genomic DNA was extracted according to methods described elsewhere (Goeltz et al., 1985; Yanagisawa et al., 1991). The remaining eight samples were frozen in liquid nitrogen after surgical resection and stored at −80°C until isolation of DNA. Genomic DNA was extracted from the frozen tissues according to methods described elsewhere (Sato et al., 1990).

RFLP markers for LOH analysis

The three RFLP markers used in this study, D1S77 (pMCT58), D1S57 (pYNZ2) and D1S62 (pTH154), were described previously (Nakamura et al., 1988a,b; Holm et al., 1988). A 5μg aliquot of each DNA sample was digested with appropriate restriction enzymes, electrophoresed in either a 1.0 or a 0.7% agarose gel and transferred to a nylon membrane. Membranes were hybridised with probes radio-labelled with [3P]dCTP by the random hexamer priming method (Feinberg and Vogelstein, 1984). After hybridisation, membranes were washed under stringent conditions and exposed to X-ray film for 1–3 days at −80°C.

Microsatellite markers for LOH analysis

The nine polymorphic microsatellite markers used in this study, D1S24 (AFM147v8), D1S232 (AFM19xb4), D1S199 (AFM078yg5), D1S201 (AFM094tb7), D1S255 (AFM260 zg5), D1S197 (AFM073xe9), D1S246 (AFM225g27), D1S219 (AFM161xb2) and D1S207 (AFM116xb2), were described previously (Weissenbach et al., 1992). The polymerase chain reaction (PCR) to amplify these loci from genomic DNA was carried out using 20 ng of DNA, 67 mM Tris, 16.6 mM ammonium sulphate, 6.7 μM EDTA, 10 mM β-mercaptoethanol, 1.5 mM of each deoxynucleotide, 5 mM magnesium chloride, 800 nM each of unlabelled primer and primer labelled with [3P]dATP (> 5000 Ci mmol−1) and 0.5 units of Taq DNA polymerase in a total volume of 25 μl. The PCR was carried out for 35 cycles of 30 s at 95°C, 30 s at 55°C and 30 s at 72°C. An aliquot of 2.5 μl of each amplified DNA was mixed with an equal volume of 95% formamide containing 0.3% xylene cyanol and 0.3% bromophenol blue, and was then denatured and electrophoresed in a 6% polyacrylamide gel containing 8 M urea and 32% formamide. After electrophoresis gels were fixed for 30 min in a solution containing 5% acetic acid and 5% methanol, dried and exposed to X-ray film for 12–24 h.
Localisation of DNA markers by FISH

FISH was performed as described elsewhere (Inazawa et al., 1991). Metaphase chromosomes were prepared by the thymidine synchronisation–BrdU release technique (Takahashi et al., 1990) for delineation of replicated G-bands. The DNA probes were prepared as described elsewhere (Inazawa et al., 1993). Hybridisation was carried out at 37°C for 16 h, and post-hybridisation washing was performed as described elsewhere (Inazawa et al., 1992). Hybridisation signals for the cosmid probes were detected cytochemically with FITC-avidin. The chromosomes were counterstained with 1 μg ml⁻¹ propidium iodide (PI) and anti-fade solution containing 1% DABCO [1,4-diazabicyclo-(2,2,2)-octane](Sigma). Microscopy was performed with a Nikon Y2F-EDF2 fluorescent microscope. PI-stained chromosomes and FITC signals were

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Figure 1 Allelic loss in cases 214 and 17 demonstrating typical patterns of LOH. Case 214 showed LOH in the tumour at D1S255. Case 17 showed LOH at D1S201 and D1S57. N and T denote DNA from normal and cancer tissue respectively. (M) and (R) denote microsatellite and RFLP markers respectively.

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Figure 2 LOH analysis at 12 loci on chromosome 1p in 26 gastric adenocarcinomas. (M) and (R) denote microsatellite and RFLP markers respectively. Top of each lane, tumour number. DNA samples 201–218 were obtained from tissue fixed in formalin and embedded in paraffin; the rest were extracted from frozen tissues. ●, LOH; ○, heterozygosity retained; –, uninformative; RER, replication error; blank, not examined.
Results

All 26 tumours were informative at one or more of the tested loci; 13 of them showed LOH for at least one locus, of which 11 instances were considered to represent partial or interstitial deletions. Our criterion of allelic loss was that more than 50% reduction in the intensity of a band of the tumour was observed when compared with corresponding band of the normal tissue. At least two sets of experiments were performed to confirm our results in ambiguous cases. Some examples of these results are shown in Figure 1, and the losses or retentions of alleles in all the tumours are summarised in Figure 2. The order of the marker loci was based on a previous study (Dracopoli et al., 1994). The deletion mapping indicated that the deletion in tumour 17 encompassed a region no greater than the distance between D1S199 and D1S246. Similarly, maximum regions of deletion were defined between D1S228 and D1S219 in case 203, between D1S199 and D1S246 in case 206 and between D1S201 and D1S197 in case 214. The combined deletion data allowed us to define a commonly deleted region between D1S201 and D1S197. These two loci lie approximately 13 cM apart.

We also performed FISH of cosmids representing two loci in the commonly deleted region, D1S57 (pYNZ2) and D1S62 (pTHI54). This procedure mapped D1S57 to lp35 and D1S62 to lp34.3 (Figure 3). These results suggest that the 13 cM commonly deleted region that includes a putative tumour-suppressor gene associated with gastric carcinogenesis is physically located in the chromosomal region lp35 to lp34.3.

Replication error (RER) at microsatellite loci was detected in three cases (12%) as shown in Figure 4. Tumour 207 gained a number of CA repeats at both D1S219 and D1S207; tumour 208 gained repeats at the D1S219 locus and lost repeats at the D1S207 locus. Interestingly, all three tumours revealed RERs at more than one of the microsatellite loci examined; in particular, in tumours 207 and 208 (see Figure 2), the RER was detected at more than half of the loci tested.

Discussion

In the present study, we performed LOH analyses for polymorphic loci on chromosome 1p in 26 gastric adenocarcinomas of the well-differentiated type. Our results demonstrated that a region commonly deleted in gastric adenocarcinomas is located between D1S201 and D1S197. Recent studies based on allelic deletions in various types of
tumours have implied the presence of a putative tumour-suppressor gene(s) on chromosome 1; these results have been reported in colorectal cancers (Leister et al., 1990; Bardi et al., 1993), hepatocellular carcinomas (Simon et al., 1991; Yeh et al., 1994), neuroblastomas (Fong et al., 1989; Weith et al., 1989; Schleiermacher et al., 1994; Takeda et al., 1994), phaeochromocytomas and medullary thyroid carcinomas (Mathew et al., 1987; Mole et al., 1992) and breast cancers (Bieche et al., 1993; Dracopoli et al., 1994; Loupart et al., 1995). As the commonly deleted regions we have observed in gastric cancers overlaps with the commonly deleted regions reported in various other tissues (Mathew et al., 1987; Fong et al., 1989; Leister et al., 1990; Mole et al., 1992), inactivation of the same unidentified tumour-suppressor gene(s) on the short arm of chromosome 1 may be associated with more than one type of cancer.

Sano et al. (1991) indicated a difference in the frequency of LOH between two types of gastric cancer; they reported that LOH on chromosome 1p was frequent in poorly differentiated adenocarcinomas of the stomach but not frequent in well-differentiated adenocarcinomas, whereas LOH on chromosome 1q was frequent in well-differentiated adenocarcinomas. However, we detected LOH on the short arm in a high proportion of well-differentiated adenocarcinomas of the stomach. This discrepancy may be due to the number and locations of the markers used in the two studies.

We detected RERs at microsatellite loci, a phenomenon that is considered to reflect some defect in the DNA mismatch repair system, in three (12%) of the 26 gastric tumours studied here; two of them revealed RERs at five of the nine microsatellite loci tested. We reported previously that tumours from patients with multiple primary cancers showed frequent RERs at microsatellite loci (Horii et al., 1994). We speculate that those gastric cancer patients in which genetic instability was detected may carry a defect in one or more of the genes involved in the DNA-mismatch repair system, and therefore may bear a higher risk than the other patients of developing a second primary cancer.

**Abbreviations**

LOH, loss of heterozygosity; RFLP, restriction fragment length polymorphism; RER, replication error; PCR, polymerase chain reaction; FISH, fluorescence in situ hybridisation; FITC, fluorescein isothiocyanate.

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