Molecular analysis of single isolated glands in gastric cancers and their surrounding gastric intestinal metaplastic mucosa

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Abstract. The biological properties and underlying genetics of gastric cancer and gastric intestinal metaplasia evolve with neoplastic progression from the genetics of the original gland cell. PCR assay with crypt isolation was used in tumors from 20 patients to examine microsatellite alterations (allelic imbalance at 17p, 5q, 18q, 3p, 4p, and 9p, and microsatellite instability) in glands from each tumor and from intestinal metaplastic lesions. Tumor specimens were processed as either pooled-gland samples or single-gland samples. Pooled gland sample was composed of 10-20 tumor glands, intestinal metaplastic glands, or nonmetaplastic glands. Single gland sample was 10 tumor glands from tumor and single gland sample was 5 gastric intestinal metaplastic and 5 nonmetaplastic glands from its surrounding metaplastic mucosa. Multiple genetic alterations were found in individual tumor glands, with various subclonal expansions seen within the same tumor. Although microsatellite instability was found in 2 of 20 tumor single-gland samples, none was detected in metaplastic single-gland samples. Most cancers appear to have a heterogeneous composition. On the other hand, microsatellite alterations were also detected within the nonmetaplastic as well as intestinal metaplastic single-gland samples. In conclusion, the present data on tumor and corresponding intestinal metaplastic and nonmetaplastic glands suggest that genetic alterations already occur within the surrounding of the noncancerous mucosa.

Introduction

It has long been established that gastric cancers develop from a single gland cell that undergoes an accumulation of genetic changes (1), and which eventually results in a malignant tumor with monoclonal character (2,3). Recent molecular genetic evidence also supports this concept (1). Despite the monoclonal origin of such cancerous tumors, any single tumor consists of numerous tumor glands, which may have various genetic alterations relative to the cell of origin. According to established theory, these accumulated genetic alterations can be identified by loss of heterozygosity (LOH) at many chromosomal loci within the tumor (4). It is also true that differences between tumor glands are thought to cause genetic heterogeneity within the same tumor (1-3), and some studies stated that genetic heterogeneity is frequently found in individual gastric cancers (5-7). Such genetic heterogeneity within the same tumor complicates the development of a tumor-treatment strategy based on tumor pathogenesis (8,9). Analyzing the accumulation of multiple genetic alterations in a single tumor gland is thought to be useful for assessing genetic differences among the cells of a single tumor (2,3).

Gastric intestinal metaplasia (GIM) is frequently found in the mucosa surrounding a gastric cancer (10), and is commonly thought to be a precancerous condition for gastric cancer, especially differentiated-type gastric cancers (11,12). However, an alternative hypothesis, the ‘paracancerous’ condition, as distinct from the ‘precancerous’ condition, has been described, especially in Japan (13). The paracancerous hypothesis derives from the fact that, based on routine pathological examination of surgical specimens, no pathologist has yet been able to identify the feature proving that differentiated-type gastric carcinomas arise directly from intestinal metaplastic glands. Accordingly, it remains unclear whether GIM is truly a precancerous lesion. In addition, GIM is not a single entity but rather a heterogeneous group of metaplastic glands (14). Previous studies suggested that GIM by itself is associated with carcinogenesis of differentiated-type gastric cancer and that GIM has a heterogeneous composition with different
biological characteristics (14-17). Identification of 1 or more biomarkers to reliably differentiate GIM types associated with gastric cancer and the molecular alterations associated with various types of GIM may be very valuable clinically as a tool for identifying patients who may be at higher risk for gastric cancer. However, molecular alterations contributing to the development or neoplastic progression of GIM have not been clarified (18). In addition, a non-neoplastic gland which is seen in the gastric intestinal metaplastic mucosa is not necessarily an intestinal metaplastic gland. Nonmetaplastic glands are often observed in gastric metaplastic mucosa. It is likely that the nonmetaplastic glands are also associated with gastric carcinogenesis. However, the molecular alterations of nonmetaplastic glands within the gastric intestinal metaplastic mucosa are still not identified.

For genetic evaluations of an individual gland taken from tissue of a single gastric cancer and the surrounding GIM and nonmetaplastic glands (intestinal metaplastic and non-intestinal glands), it is necessary to isolate single glands from the larger tissue mass. The crypt isolation method can be used to obtain individual tumor and non-neoplastic glands from a given tumor or surrounding tissue (19,20). In the current study, we used crypt isolation methodology to examine molecular alterations of single tumor glands from a gastric cancer and single glands from surrounding GIM tissue. The aim of this study was to verify the role of intratumoral molecular differences in sporadic gastric cancer and to further our understanding of molecular alterations in GIM tissue that result in gastric tumorigenesis.

Materials and methods

Tissue samples. Tissues were obtained from 20 patients with sporadic primary gastric adenocarcinoma of differentiated type, who had undergone gastrectomy. Of the 20 patients, there were 17 men and 3 women (mean age: 64.2 years). Tumor histological type and stage were classified according to the Japanese Research Society criteria for cancer of the stomach (21). The location of the gastric cancer was determined and the tumors were subclassified into 2 groups: proximal or distal. Clinicopathological data for the 20 patients in our study are shown in Table I. In addition, in order to clarify genetic alterations in non-neoplastic glands not demonstrating intestinal metaplasia (nonmetaplastic gland), nonmetaplastic glands associated with gastric adenocarcinomas were analyzed in the 10 tumors in which they were available.

Fresh tumor specimens and adjacent tissue were obtained from resected gastric cancers. Gastric mucosa most distal to the tumor that was regarded as normal was removed from the submucosa with scissors and discarded. In contrast, intestinal metaplastic mucosa was obtained from antral mucosa. Tissue tumor samples were obtained primarily from the central area of the tumor.

Crypt isolation technique. Isolation of tumor and mucosal glands was performed as previously described (22). Briefly, fresh tumor and non-neoplastic mucosa were separated from the underlying tissue layer and cut with a razor into minute pieces, and then incubated at 37°C for 30 min in Hanks’ balanced salt solution (HBSS), which is calcium- and magnesium-free, containing 30 mmol/l ethylenediaminetetra-acetic acid (EDTA). Specimens were stirred in HBSS for 30-40 min to allow isolation of cancerous and normal glands from the lamina propria mucosa or fibrous stroma. Isolated specimens were immediately fixed in 70% ethanol and stored at 4°C until analysis.

Identification of isolated gastric intestinal metaplastic glands. In the present study, GIM was recognized by the presence of goblet cells stained by Alcian blue. Detailed histological identification of intestinal metaplasia was performed on sections of paraffin-embedded tissue located adjacent to the tissue used for crypt isolation. Most tissue sections showed incomplete intestinal metaplasia.

Identification of isolated gastric intestinal nonmetaplastic glands. Nonmetaplastic glands which were not stained by Alcian blue were isolated from antral mucosa stained with Alcian blue (pH 2.5).

Identification of isolated normal gastric glands. Normal mucosal tissue was defined as glands obtained from Alcian blue-negative fundic mucosa, and was confirmed by histological examination. Normal mucosa was used to examine microsatellite alterations in normal (negative) controls.

DNA extraction. Isolated tumor glands were handled as follows: Ten to 20 isolated glands (tumor, intestinal metaplastic, and nonmetaplastic glands) were obtained from each
tissue sample. DNA extraction was performed on these glands to create the pooled-gland samples (3). Separately, 10 tumor glands and 5 intestinal metaplastic glands or nonmetaplastic glands were obtained from each tumor examined and its surrounding antral mucosa. DNA was extracted from each individual tumor, intestinal metaplastic, and nonmetaplastic gland, respectively, to create the single-gland samples, using techniques described previously (2,3). The DNA concentration of single tumor glands was estimated by the TagMan real-time PCR method using the ribosomal protein P0 (3684) gene as a reference (3). In addition, the amount of DNA content that we examined varied from 98.6 to 1034.8 ng (mean: 410.1 ng). On the other hand, that of DNA content examined in single intestinal metaplastic glands varied from 112.6 to 253.4 ng (mean: 198.7 ng).

Microsatellite analysis. Microsatellite analysis included 15 microsatellite markers (2 mono- and 13 dinucleotide repeats). These markers were selected either because of their location at chromosomal sites in or near genes known to be involved in gastric carcinogenesis (3p, 4p, 5q, 9p, 17p, and 18q), or because they are very sensitive markers for determination of microsatellite instability (MSI) (BAT25 and 26). The markers used for analysis of LOH in this study were: 3p (D3S2402, D3S1234), 4p (D4S2639, D4S1601), 5q (D5S107, D5S346, D5S299, D5S82), 9q (D9S171, D9S1118), 17p (TP53), and 18q (D18S487, D18S34).

PCR reactions were performed as described previously (23,24). One of the primers used for amplification was fluorescently labeled. PCR products were separated and detected with an automated sequencing system as previously reported (19,23).

Scoring of allelic imbalance. Allelic imbalance (AI) was determined using a calculation method described previously (25). A tumor was considered to have AI if the allelic peak ratio was <0.6, representing an allelic signal reduction of at least 40%. We interpreted this allelic imbalance as LOH with the provision that, in some cases, the change in allele peak ratio may have resulted from allelic amplification. When AI was observed in at least 1 locus of the markers examined, imbalances of the examined loci were confirmed. Finally, tumors exhibiting MSI at a given locus were not evaluated for LOH.

The overall extent of LOH for each single-gland sample (tumor, intestinal metaplastic, or nonmetaplastic single-gland sample) was calculated as follows: the number of single glands showing LOH divided by the number of informative (excluding uninformative cases, samples showing MSI, and cases that were not examined) single glands for each tumor or each intestinal metaplastic gland sample.

Scoring of microsatellite instability. An additional peak in tumor DNA compared with nontumor tissue was classified as instability for the marker examined. Instability in BAT 25 or 26 was defined as MSI positive (26).

Results

In the present study, PCR analysis was performed reproducibly and there were no PCR failures, as has been described elsewhere (3). The frequencies of allelic imbalance at the

Figure 1. Well-differentiated adenocarcinoma. (a) As seen under a dissecting microscope and (b) with hematoxylin and eosin staining (H&E). A single moderately differentiated tumor gland (c) under a dissecting microscope and (d) as seen with H&E staining. Single intestinal metaplastic gland (e) as stained by Alcian blue (pH 2.5) under a dissecting microscope and visualized with H&E staining (f). Single non-intestinal metaplastic gland (g).
Figure 2. A detailed view of allelic imbalances on 5q, 17p, 18q, 3p, 4p, 9p, and microsatellite instability in pooled-tumor samples and in corresponding tumor single-gland samples (cases: A-L). P, pooled gland sample; LOH, loss of heterozygosity; N, negative; NI, not informative; NA, not amplified; MSI, microsatellite instability.

Figure 3. A detailed view of allelic imbalances on 5q, 17p, 18q, 3p, 4p, 9p, and microsatellite instability (MSI) in pooled-tumor samples and in corresponding tumor single-gland samples (cases: M-T). P, pooled gland sample; LOH, loss of heterozygosity; N, negative; NI, not informative; NA, not amplified; MSI, microsatellite instability.
chromosomal loci that were studied in the tumor pooled-gland samples are listed in Table II. We defined genotype as the AI pattern of each examined sample.

Concordance of genotype of pooled-gland samples and single-gland samples. We examined the genotype concordance of pooled-gland samples and the predominant genotype in corresponding single-gland samples. In 4 cases there were different AI patterns seen in the tumor pooled-gland samples and in those of the corresponding tumor single-gland samples (samples B, H, O, and S; Figs. 2 and 3). In addition, in another 7 cases (samples D, F, I, J, L, P, and R; Figs. 2 and 3), the predominant genotypes in tumor single-gland samples were different from those of the corresponding pooled-gland samples. Alternatively, in the other 9 cases the genotypes of the tumor pooled-gland samples were consistent with those of the corresponding tumor single-gland samples (Figs. 2 and 3).

In the samples of intestinal metaplastic glands, although no genetic alterations were found in the pooled-gland samples, multiple alterations were frequently detected in single-gland samples (Fig. 4). A representative example of allelic imbalance at each chromosomal locus in a pooled-gland sample and in corresponding intestinal metaplastic single-gland samples is illustrated in Fig. 5 (case C).

In the samples of nonmetaplastic glands, although no genetic alterations were found in the pooled-gland samples, alterations were observed in corresponding single-gland samples (Fig. 6).

Heterogeneous genotypes in individual glands within the same sample. A total of 18 carcinomas (18/20, 90%) consisted of heterogeneous single genotypes (>1 different genotype) within the same tumor. Single glands within the same sample having the same genotype were interpreted as an occurrence of the same subclone. The number of subclones within the same sample was determined and is shown in Table III. Many subclones were identified within the same sample. No more than 7 different genotypes (subclones) in single-tumor glands from the same sample were observed. In addition, multiple subclones within the same sample of not only intestinal metaplastic single glands but also nonmetaplastic single glands were found. The mean numbers of genetic alterations per tumor, intestinal metaplastic, and nonmetaplastic single-gland samples were 4.15, 3.75 and 4.5, respectively.

Two samples (10%) had homogeneous single genotypes within the same tumor (A and N, Figs. 2 and 3). In contrast, MSI did not occur in a homogeneous pattern within the same tumor (G and Q, Figs. 2 and 3). There were no homogeneous patterns observed among either metaplastic or nonmetaplastic single-gland samples (Figs. 4 and 6).

Genotypic pattern of individual glands within the same sample. As shown in our previous study on colorectal carcinoma (3), the genotype of each single-tumor gland was classified as either a major-altered or a minor-altered genotype. The first group, the major-altered genotype, was defined as a single-tumor gland with >1 genetic alteration. In contrast, the minor-altered genotype was defined as a single-tumor gland with none or...
Figure 5. Allelic imbalances on 17p, 5q, 18q, 3p, 4p, and 9p chromosomal loci in an intestinal metaplastic pooled-gland sample and in corresponding intestinal metaplastic single-gland samples (case C). Multiple subclones (genotypes) were seen. Note that although no genetic alterations were detected in the pooled-gland sample, they were frequently found in the paired single intestinal metaplastic gland sample. In intestinal metaplastic single-gland samples, arrows indicate a lost allele at each chromosomal locus. P, pooled gland sample; N, normal.

Figure 6. A detailed view of allelic imbalances on 5q, 17p, 18q, 3p, 4p, 9p, and microsatellite instability in nonmetaplastic pooled-gland samples and in corresponding nonmetaplastic single-gland samples (cases: A, C, D, G, I, L, M, O, and S). Genetic alterations were seen in the single intestinal metaplastic gland samples. The frequency of allelic imbalances in single nonmetaplastic gland samples is low compared with those of single intestinal metaplastic gland samples. P, pooled gland sample; LOH, loss of heterozygosity; N, negative; NI, not informative; NA, not amplified.
We looked for any association between the genotypes of pooled-gland samples (major-altered and minor-altered genotypes) and the genotypic patterns (I-V) of tumor, intestinal metaplastic, and nonmetaplastic single-gland samples; results are shown in Table IV. In 3 tumor samples (Q, R, and S), the genotype in pooled-gland samples was not consistent with the genotypic patterns of tumor single-gland samples. In contrast, in 10 intestinal metaplastic samples (B, C, D, E, I, J, K, N, O, and Q), the genotype of pooled-gland samples in GIM was different from the genotypic patterns of intestinal metaplastic single-gland samples. In addition, in 7 nonmetaplastic samples (C, D, I, J, M, O, and S), the genotype of pooled-gland samples in nonmetaplastic glands was not consistent with the genotype pattern of nonmetaplastic single-gland samples.

### Discussion

In the present study, 12 of 20 tumors showed different predominant genotypes between pooled-gland and single-gland tumor samples. These 12 tumors were primarily classified into 2 groups as an explanation of the discrepancy between pooled-gland and single-tumor gland samples (3). The first explanation is that the AIs in pooled-gland samples at the chromosomal loci we examined may be caused by dilution of minor genotypes within the same tumor (dilution effect). This may represent a mean value of AIs for all collected single-gland tumors. We concluded that the results for 4 tumors (samples: F, I, J, and L) are consistent with this explanation. An alternative explanation may be that the composition of tumors in the pooled-gland samples is different from that observed in the single-gland tumor sample (i.e., different composition effect). This suggests that additional subclones, different from those found in the pooled-gland tumor sample, can exist within the same tumor. According to this explanation, it may be likely that a different subclone or different composition of a single-gland sample will not be detected in the pooled-gland samples (B, D, H, O, P, Q, R, and S).

Such a discrepancy between pooled-gland and single-gland samples was prominent in the nonmetaplastic samples. Although no genetic alterations were detected in the pooled-gland samples, multiple genetic alterations were found in corresponding single-gland samples. In particular, this finding was frequently present in GIM samples.

In the present study, it appears that isolation of single intestinal metaplastic glands can be used to increase the sensitivity of tests for allelic imbalance. This finding suggests

### Table III. The number of subclones within the same sample.

<table>
<thead>
<tr>
<th>Number of subclones within the same tumor</th>
<th>Tumor gland</th>
<th>Intestinal metaplastic gland</th>
<th>Nonmetaplastic gland</th>
</tr>
</thead>
<tbody>
<tr>
<td>Two</td>
<td>3</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Three</td>
<td>3</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>Four</td>
<td>3</td>
<td>7</td>
<td>3</td>
</tr>
<tr>
<td>Five</td>
<td>2</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Six</td>
<td>5</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Seven</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

### Table IV. Frequencies of the genotypic patterns of single-gland samples in gastric tumor, gastric intestinal metaplastic glands, and nonmetaplastic glands according to the genotype of pooled-gland samples.

<table>
<thead>
<tr>
<th>Genotype of pooled-gland samples</th>
<th>Tumor (n-20)</th>
<th>Intestinal metaplastic gland (n-20)</th>
<th>Nonmetaplastic gland (n-10)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>I</td>
<td>II</td>
<td>III</td>
</tr>
<tr>
<td>Major</td>
<td>8</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>Minor</td>
<td>0</td>
<td>2</td>
<td>1</td>
</tr>
</tbody>
</table>
Marked clinical and biological heterogeneity has been noted among human gastric cancers (1-3). However, possible genetic causes of genetic heterogeneity have not been fully investigated. In this study, we used tumor single-gland samples to look for heterogeneous populations within the same tumor. Our findings that 90% of tumors demonstrated heterogeneous composition within the same tumor, show that gastric cancers are genetically highly complex. On average, there were 4.2 genetic alterations per tumor. The high number of genetic alterations per tumor indicates that genetic instability may cause intratumoral heterogeneity, as seen in other human tumors (1,3,27,28), and be an underlying mechanism of gastric carcinogenesis. In addition, our findings suggest that a specific subclone cannot be selected in most gastric cancers during tumor progression. This is an important finding toward understanding the effectiveness of chemotherapy or radiotherapy in gastric cancers, because the existence of heterogeneous populations indicates no single target cell can be defined.

The degree of accumulated LOH (allelic imbalance) has been shown to be of prognostic value in various cancer types including gastric cancer, and a high degree of tumor LOH has been shown to be associated with tumor aggressiveness and a worse prognosis (29). In light of these findings, low and high rates of tumor LOH suggest low and high tumor behavioral aggressiveness, respectively. In some of the present cases, although minor-altered genotype (low LOH) was found in the pooled-gland sample, major-altered genotype (high LOH) was detected in the corresponding tumor single-gland sample. Our findings also indicate that 2 of 5 tumors showing a minor-altered genotype in the pooled-gland sample were classified as type II (composed of major-altered genotype) in the tumor single-gland sample. This finding suggests that highly aggressive subclones may exist within a tumor showing a low frequency of LOH in a pooled-gland sample (2/5, 40%). It is surprising that 40% of pooled-gland samples showing the minor-altered genotype contained a major-altered genotype in the corresponding tumor single-gland samples. On the other hand, in the present study, most of the carcinomas that were examined contained a minor-altered genotype of single tumor glands within the same tumor. One recent study has shown that high-LOH tumors may have a higher rate of response to chemotherapy (29,30). However, in general, the majority of patients with gastrointestinal cancers are thought to obtain a poor response to chemotherapy without survival benefit (29,30). Although the genetic reason for the discrepancy remains unknown, an explanation may be that the minor-altered genotype gland is a supply source to the major-altered genotype gland. Therefore, a minor-altered genotype gland may co-exist with a major-altered genotype gland within the same tumor (3).

In the present study, genotypic pattern for a single-gland sample was classified into 5 groups. Although types I and II were the most frequent genotypic patterns (17/20, 85%), type IV, a tumor genotype showing predominantly minor-altered glands, was relatively rare (3/20, 15%). These findings suggest that single tumor glands with multiple genetic alterations may cause subclonal expansions in different areas within the same tumor, leading to occupation of the whole tumor mass in gastric cancers.

The general strategy of identification of individuals at high risk for progression to cancer offers promising possibilities for cancer prevention, and this approach largely depends on early detection. Therefore, it is important to evaluate GIM, which is generally thought to be a precancerous condition in gastric cancer (12,18). Furthermore, a previous study has shown that GIM is closely associated with Helicobacter pylori infection (31). However, little is known about the genetic events responsible for initiation and progression of gastric cancer (32,33). According to an investigation by Ochiai et al., although $p53$ mutations known to play a key role in human neoplastic progression were identified in GIM, they were detected in only 10 of 756 (1.3%) histological sections (18). This finding indicates that it is difficult to identify such subtle genetic alterations in intestinal metastatic glands. Genetic analysis of a single gland may enable us to identify subtle genetic alterations in GIM. Therefore, we used isolated single glands in the current study to address the issue of whether molecular alterations occur in GIM. In this study, although no genetic alterations were detected in pooled samples of intestinal metastatic glands, alterations were frequently found in the corresponding single intestinal metastatic gland samples. This suggests that intestinal metastatic glands have a markedly heterogeneous composition. This study is the first to identify that expansive microsatellite alterations are seen in samples of intestinal metastatic glands. These data indicate that irreversible genetic changes have already occurred in morphologically non-neoplastic gastric mucosa with intestinal metaplasia, and they support the hypothesis that GIM may be a precursor lesion of gastric cancer.

The present study has demonstrated that multiple genetic alterations are frequently found in nonmetastatic glands. This is a surprising finding and the first study to identify genetic alterations in histologically normal gastric glands. This finding suggests that accumulation of genetic alterations occurs not only in metastatic glands but also in nonmetastatic glands, and that genetic alterations in gastric epithelial cells during chronic gastritis may contribute to an increased risk of gastric cancer (34).

In conclusion, single tumor glands can be useful for investigating genetic alterations in gastric cancers and gastric intestinal metaplasia. The present data indicate that most carcinomas and GIM are genetically heterogeneous. Recently, public policy strategies have been suggested for identification of patients at risk for $H. pylori$-related gastric malignancy (35,36). The thrust of this policy is that eradication of $H. pylori$ infection earlier rather than later in life is anticipated to be more beneficial, because gastric intestinal metaplasia is expected to occur later in life. Our finding that multiple genetic alterations are found in single intestinal metastatic glands may support this opinion.

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References