Polyclonal nature of diffuse proliferation of interstitial cells of Cajal in patients with familial and multiple gastrointestinal stromal tumours

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Gastrointestinal stromal tumours (GISTs) are the most common mesenchymal tumours in the human digestive tract. Most GISTs express a receptor tyrosine kinase, KIT. 3,4 KIT is encoded by the c-KIT gene 5 and its ligand is stem cell factor (SCF). 1,3 GISTs frequently have a mutation of the c-KIT gene which results in constitutive activation of KIT without stimulation by SCF. 6 As the development of interstitial cells of Cajal (ICCs) is dependent on SCF-KIT interaction and since ICCs and GISTs both express KIT and CD34, 7,8 we considered that GISTs may originate from ICCs.

Recently, we and others found some families with multiple GISTs and a germline gain of function mutation of the c-KIT gene. 9–13 In addition to the development of multiple GISTs, family members with the mutation showed diffuse proliferation of spindle shaped cells between the circular and longitudinal muscular layers—that is, the myenteric plexus of the small intestine. 13–15 As the proliferative spindle shaped cells expressed both KIT and CD34, they were also considered to originate from ICCs. Although we regarded diffuse ICC proliferation as hyperplasia and the macroscopically apparent mass lesions as GISTs, the clonal nature of them remained unclear.

To examine the clonality of human samples, random inactivation of one of the two female X chromosomes (lyonisation) is most commonly used. 16–18 Inactivation occurs by methylation during early embryogenesis in each somatic cell, and is stably transmitted to all progeny cells. It also persists during neoplastic transformation. Therefore, normal tissues or non-neoplastic hyperplastic lesions composed of uniform cells having either maternally or paternally derived inactivated X chromosomes demonstrated a monoclonal pattern. Currently, the polymorphism of the human androgen receptor (HUMARA) locus, which is located on the X chromosome, is most frequently used to examine clonality as it has a highly polymorphic tandem repeat. 19,20 Approximately 90% of females are heterozygous for the number of CAG trinucleotide repeats. 17 Moreover, methylation sensitive restriction enzyme sites of HpaII and HhaI are within the HUMARA locus. 17

In the present study, we examined the clonality of diffuse ICC proliferation in familial and multiple GIST cases using polymorphism of the HUMARA locus. To obtain pure populations of cells from the diffusely proliferative lesions, the laser capture microdissection (LCM) method was used. 21–23 GISTs showed a monoclonal pattern but diffuse ICC proliferation displayed a polyclonal pattern as did normal mucosal tissues.

MATERIALS AND METHODS

Samples

Tissues were obtained from three female patients with multiple GISTs and a germline mutation of the c-KIT gene during operation. Case No 1 was a daughter of case No 2, and they had a c-KIT gene mutation at exon 11, as described previously. 9 Case No 3 had no familial relationship with case Nos 1 and 2, and had a c-KIT gene mutation at exon 17, as described previously. 11 In each case, some GIST tissues were resected, including macroscopically normal adjacent intestine, and were fixed in formalin and embedded in paraffin. Histological diagnosis was made by haematoxylin and eosin staining and
immunohistochemistry of KIT and CD34. Immunohistochemistry was performed as described previously.21 In the present study, we regarded non-mass forming proliferation (<3 mm in diameter) of spindle shaped cells which were double positive for KIT and CD34 as diffuse ICC proliferation, while apparent mass forming lesions (>5 mm in diameter) which were double positive for KIT and CD34 as GISTs. The adjacent sections were used for DNA extraction.

**DNA extraction**

DNA samples were extracted from formalin fixed paraffin embedded tissues according to a previous method.22 Normal intestinal mucosa, diffuse ICC proliferation, and GISTs were microdissected from 4μm sections stained with haematoxylin and eosin using an LCM system (LM200; Arcturus Engineering, Mountain View, California, USA). Representative histology before and after microdissection is shown in fig 1. In the section of large GISTs, three samples were extracted from three distinct sites, respectively. A total volume of 20μl of PK buffer containing 1.0% proteinase K, 10 mM Tris HCl (pH 8.0), 1 mM ethylenediamine tetraacetic acid, and 1% Tween 20 was mounted on a microdissected specimen attached to a piece of LCM Transfer Film (Arcturus Engineering). Then, a 0.5 μl micro test tube (Eppendorf-Netheler-Hinz-GmbH, Hamburg, Germany) was placed over it. After 16 hours of incubation at 37°C, proteinase K was inactivated by heating at 95°C for 10 minutes. The lysis mixture was centrifuged for five minutes to remove undigested tissue fragments.

**Clonality assessment**

We used a quantitative fluorescent polymerase chain reaction (PCR) procedure that enables accurate measurement of peak intensities of each allele, as described previously.23 The HUMARA gene includes a polymorphic ((CAG)n) repeat located at 3′ of the methylation sensitive HhaI restriction enzyme sites.24 The PCR assay used primers, the product of which spanned both the HhaI sites and the ((CAG)n) polymorphism. Variations in length of the ((CAG)n) repeats on the paternal and maternal X chromosomes yield HUMARA alleles of different lengths. Methylation of the HhaI sites distinguishes the active (non-methylated) from the inactive (methylated) X chromosome. It is only the undigested inactive methylated allele that is subsequently amplified by PCR.25 Each DNA sample (7.5μl) was digested overnight at 37°C in a 2.5 μl reaction mixture containing 0.5 μl of 16 units of HhaI (Toyobo, Osaka, Japan), 1 μl of concentrated (×10) TA buffer (330 mM Tris acetate, 660 mM KOAc, 100 mM MgOAc, 5 mM dithiothreitol), and 1 μl of concentrated (×10) bovine serum albumin. For each case, a control sample without DNA was run simultaneously to rule out contamination of DNA in the reaction mixture. The restriction enzyme was then inactivated by heating at 95°C for 10 minutes.

For PCR, 1 μl of each digested DNA sample was added to 24 μl of a PCR reaction mixture containing 2.5 μl of concentrated (×10) PCR buffer,1.5 μl of deoxyribonucleoside triphosphate (200 μM), 0.2 μl of primers 1A and 1B (10 μM each), 0.1 μl of Taq polymerase (Roche Diagnostics GmbH, Mannheim, Germany), and 19.7μl of deionised H2O. The DNA samples were amplified using a sandwiched primer approach.26 The first step was performed using outer primers 1A (5′-GCT GTG AAG GTT GCT GGT CCT CAT-3′) and 1B (5′-GGT CCA AGA CCT ACC GAG GAG C-3′). The second step was performed with inner primers 2A (5′-TCC AGA TTC TGT TCC AGA GGG TGC-3′) and 2B (5′-ATG GGG AGG GTG GGG AGA ACC ATC CTC-3′). Primer 2A was labelled at the 5′ end with 6-carboxyfluorescein. Initial denaturation was performed for 10 minutes at 95°C, followed by 35 cycles of one minute at 95°C, one minute at 60°C, and one minute at 72°C. In the final cycle, extension at 72°C was prolonged for 10 minutes. The second step PCR profile was the same as the first step PCR.

Following the second step amplification, 5 μl of the PCR product was assessed using 2.0% agarose gel electrophoresis to confirm amplification of the HUMARA target. After amplification, 1 μl of the PCR products was mixed with 12 μl of a Template Suppression Reagent (Applied Biosystems, Foster City, California, USA) and 0.5 μl of internal size standards (GENESCAN-500 (TAMRA), Applied Biosystems). The mixture was denatured at 95°C for two minutes, and analysed through a DNA Sequencing Polymer (Applied Biosystems) with an ABI PRISM Genetic Analyser (Applied Biosystems), according to a previous method.23 Data were analysed using Genescan 310 Software (Applied Biosystems).

**Data interpretation**

Amplification of each HUMARA allele usually generated a set of multiple peaks, including one major peak and a few associated peaks of lesser intensity, as described previously.23 Clonality assessment was based on the major peak generated from each allele. Patients were considered heterogeneous when PCR amplification of undigested DNA showed two major peaks of almost equal intensity. This suggested that maternal and paternal X chromosomes have HUMARA alleles of different molecular weights. PCR products showing a single major peak suggested that maternal and paternal X chromosomes have HUMARA alleles of the same molecular weight. Such patients were considered to be homozygous for the HUMARA gene, and thus uninformative for the analysis.

Samples were considered to be polyclonal when PCR amplification of digested DNA showed two major peaks similar to that of normal tissue from the same organ. PCR products showing only one of the two major peaks were considered to be monoclonal.

**RESULTS**

Specimens were obtained from three women. Immunohistochemistry of KIT and CD34 was carried out on sections with normal intestinal mucosa, diffuse ICC proliferation, and/or GIST tissues of each patient. Almost all cells of GIST or diffuse ICC proliferation tissues were double positive for KIT and CD34. Adjacent sections were stained with haematoxylin and eosin, then normal intestinal mucosa, diffuse ICC proliferation and GIST tissues of each patient were carefully removed by LCM (fig 1). In each case, two GISTs from different sites were studied. DNA was extracted from each sample, and the portion of the HUMARA locus containing the trinucleotide repeats was amplified by PCR and the length and intensity of the PCR products were analysed.

In case No 1, all of the PCR products using undigested and digested DNA from normal intestinal mucosa, diffuse ICC proliferation, and two GISTs showed only one major peak with an allelic size of 246 bases (fig 2A1–A4), indicating that the patient was homozygous for the number of trinucleotide repeats and uninformative for the analysis.

In case No 2, the mother of case No 1, the PCR products using undigested DNA from normal intestinal mucosa, diffuse ICC proliferation, and two GISTs showed two major peaks with allelic sizes of 231 and 246 bases, indicating that the patient was heterozygous for the number of trinucleotide repeats and informative for the analysis. After digestion by HhaI, the PCR product derived either from normal intestinal mucosa or from diffuse ICC proliferation demonstrated two major peaks (fig 2B1–B2). On the other hand, the PCR product derived either from a small GIST or from a large GIST showed only one allele with a lower molecular weight (allelic size of 231 bases) (fig 2B3–B4). All of the PCR products from three distinct sites inside the same large GIST demonstrated disappearance of the same allele with higher molecular weight.

In case No 3, the PCR products using undigested DNA from normal intestinal mucosa, diffuse ICC proliferation, and two GISTs showed two major peaks with allelic sizes of 240 and...
252 bases. After digestion by HhaI, the PCR product derived either from the normal intestinal mucosa or from diffuse ICC proliferation showed two major peaks (fig 2C1–C2). The PCR product derived from a large GIST showed only one allele with a lower molecular weight (allelic size of 240 bases) (fig 2C3). Again, all of the PCR products from three distinct sites inside the same large GIST demonstrated disappearance of the same allele with a higher molecular weight. On the other hand, the PCR product derived from a small GIST showed the other allele with a higher molecular weight (allelic size of 252 bases) (fig 2C4).

**DISCUSSION**

In this study, we examined the clonal nature of diffuse ICC proliferation observed in three female patients with a germline mutation of the c-KIT gene. One case was uninformative because the HUMARA locus was homozygous. In two informative cases, PCR products derived from diffuse ICC proliferation showed the two major peaks as those derived from the intestinal mucosa. In other words, diffuse ICC proliferation is polyclonal, suggesting that the lesion is hyperplastic but not neoplastic. On the other hand, the PCR product derived from a small GIST showed the other allele with a higher molecular weight (allelic size of 252 bases) (fig 2C4).

In rare cases, X chromosome inactivation in normal tissue does not occur randomly, and is responsible for a false monoclonal pattern of inactivation in normal tissue. Therefore, interpretation of the clonal pattern of a lesion must always be performed with the knowledge of the X chromosome inactivation pattern of a normal tissue from the same organ. In the present study, we demonstrated that normal tissues from the intestine were apparently polyclonal, and therefore our clonality analysis can be considered reliable.

PCR products from three distinct sites inside the same large GIST showed disappearance of the same peak in case Nos 2 and 3. Although PCR products from a small GIST and a large GIST in case No 2 demonstrated disappearance of the same peak, those in case No 3 showed disappearance of different peaks. The results indicated that multiple GISTs observed in patients with a germline KIT mutation were not metastatic lesions from one GIST but developed independently.
Although monoclonality is considered to be a hallmark of neoplasia, the relationship between monoclonality/polyclonality and neoplasia/hyperplasia is currently controversial, especially in some familial syndromes. For example, in multiple endocrine neoplasia type 2A, each focus of C cell proliferation had been considered to be hyperplastic, but Diaz-Cano et al reported that the lesion was monoclonal intraepithelial neoplasia. Moreover, Merritt et al reported that Min (multiple intestinal neoplasia) mice frequently had intestinal adenomas showing a polyclonal structure. In the case of neurofibromatosis type 1, the clonality of single neurofibroma is controversial. Some reports showed that each benign neurofibroma was monoclonal and neurofibrosarcoma probably developing from benign neurofibroma was monoclonal, while other reports demonstrated that both benign neurofibromas may be indistinguishable for the development of GISTs and diffuse ICC hyperplasia. Further research is needed for the elucidation of the relationship between diffuse ICC hyperplasia and the development of GISTs in patients with familial and multiple GISTs.

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