Abstract. Uterine leiomyomas were shown to be clonal lesions, but the relationship among different tumor nodules in multiple uterine leiomyomas remains unresolved. In this study, X-chromosomal inactivation patterns of these tumor nodules were shown by allelic polymorphism analysis through polymerase-chain reaction at the phosphoglycerate kinase and androgen receptor loci following pretreatment with the methylation-sensitive restriction enzyme HpaII or HhaI. A total number of 113 cases of uterine leiomyomas were examined. Monoclonality was demonstrated in all of the 315 nodules from 76 informative cases. The inter-nodular relationship was evaluated in 55 multiple cases with 294 tumor nodules. Different inactivation patterns were observed in 20 cases, demonstrating a multicentric origin, while an identical inactivated allele was found in all or most of the nodules in the rest of the cases, indicating a common clonal origin. The occurrence of the unicentric cases appeared to be associated with an elevated mitotic activity. Seven nodules from a multinodular case with a morphology indicative of mitotically active leiomyoma were shown to carry the identical inactivated allele, which demonstrates their unicellular origin and malignant nature. In addition, the same androgen receptor gene alteration was identified in two discrete leiomyoma nodules from a uterus. These results approve the monoclonality of uterine leiomyomas and demonstrate the presence of unicentric multiple leiomyomas.

Introduction

Uterine smooth muscle neoplasm is one of the most common tumors in females, whose prevalence among adults ranges from 20% (1) to 70% (2,3). It is frequently benign. However, tumors showing histologic features intermediate between clearly benign leiomyomas and leiomyosarcomas, namely mitotically active leiomyomas (4-7), smooth muscle tumors of uncertain malignant potential (8,9) or atypical leiomyoma (10), are not rare, and their clinical behavior cannot be predicted reliably (11,12). The distinction between benign and malignant behavior is further blurred by the presence, even though infrequent, of intravenous leiomyomatosis (IVL) (13,14), benign metastasizing leiomyoma (BML) (14-18) and disseminated peritoneal leiomyomatosis (DPL) (19,20).

There is often more than one tumor observed in cases of uterine leiomyoma. It remains to be settled as to whether all of these tumors are fully independent (multicentric) or most of the nodules originate from a parent tumor by cell migration or spreading (unicentric). While most of the reports favor the former possibility (21-28), the data from our group, obtained in 15 multiple leiomyomas using X-chromosomal inactivation analysis, have indicated that, in some cases, different tumor nodules may share a common clonal origin (29). Actually, identical genetic alterations and X-chromosomal inactivation patterns were also observed incidentally in different leiomyomas from the same uterus by other authors (27,30,31). A better understanding of the relationship between multiple tumor nodules may be helpful for distinguishing the borderline neoplasms from the clearly benign leiomyomas. In the current study, monoclonality was revealed in all of the 318 smooth muscle tumor nodules from 76 cases, and the relationship between different nodules was evaluated in 55 multi-nodular cases with 297 lesions by their X-chromosomal inactivation patterns.

Materials and methods

Samples. Fresh tissue samples were obtained by 113 hysterectomies for uterine corpus neoplasms in Tangdu Hospital, Xi'an from February 1998 to August 2001. The study protocol
was approved by the Medical Ethics Commission of the Fourth Military Medical University in Xi’an. Sampling was finished within 40 min after the uterus was taken from patients. All tumors were excised. Four specimens of apparently normal smooth muscle tissue (5 mm³) adjacent to each of the tumor nodules were used as controls. The tissues were stored at -70°C.

Histological and immunohistochemical examinations. Each sample was examined histologically for the parenchyma/mesenchyma ratio on 4-μm paraffin sections stained with hematoxylin and cosin. Samples with admixed mesenchymal cells >20% of the total nucleated cells were excluded from the assay. All cases were examined separately by three pathologists (Q. Su, S.-F. Wang and W. Zhang) according to the reported criteria (3-5,11,12,32). The multiple leiomyomas were examined for the histologic variants including atypical (bizarre), myxoid types and those with pronounced hyaline degeneration (32). Mitotic figures were counted within 50 randomly selected high-power fields (HPF), and mitotic indices were calculated and expressed as number of mitotic figures/10 HPF.

For the samples suspected for leiomyosarcoma, immunostaining was also conducted, as described previously (33), to identify their differentiation pathways. Following deparaffinization and rehydration, serial sections were treated with 0.3% of hydrogen peroxide in 80% methanol for 30 min, blocked using 3% of preimmune goat serum in 10 mM phosphate-buffered saline (pH 7.4), and incubated overnight at 4°C with mouse monoclonal antibodies against smooth muscle actin, desmin, CD10 and vimentin (Dako A/S, Glostrup, Denmark). The reactions were demonstrated using a peroxidase-labeled streptavidin-biotin kit (K0675; Dako A/S) and visualized in a buffer containing 0.5 mg/ml 3,3'-diaminobenzidine and 0.01% hydrogen peroxide for 15 min. The sections were lightly counterstained with hematoxylin, dehydrated, and mounted in neutral resin medium.

Principles of X-chromosomal inactivation analysis. There are two X chromosomes within each somatic cell in females, one from the father (Xp) and another from the mother (Xm). Both PGK and AR genes are located in X chromosomes, showing readily demonstrable polymorphism in different frequencies. The former carries a BstXI restriction-polymorphic site, reflecting the G/A single-nucleotide polymorphism (SNP) at exon 1 (34), while the latter is polymorphic at the CAG short-tandem repeat (STR) (35). These allelic differences enable us to discriminate Xp from Xm.

During embryogenesis, one of the X chromosomes in each somatic cell in females is inactivated through methylation, while the genetic activity of the second X chromosome is preserved throughout life. As the inactivation is random, in all females somatic cells can be divided into two types, one with the inactivated paternal PGK and AR alleles, named Type A, and another with the inactivated maternal alleles, termed Type B. The inactivation is rather stable, and the inactivated allele of a cell can be transmitted to its daughter cells including those undergoing neoplastic transformation (36). The clonality status of the somatic female cells, therefore, can be determined based on the X-chromosomal inactivation mosaicism (37). Loss of the X-chromosomal inactivation mosaicism, therefore, can be demonstrated by the allelic imbalance using SNP analysis at the PGK locus through BstXI digestion or STR length estimation at exon 1 of the AR gene on denaturing polyacrylamide gels.

As depicted in Fig. 1, each inactivated allele at Xp or Xm, both for PGK and AR loci, should be detected at the same frequency in uterine smooth muscle tumors. Hence, occurrence of tumors composed of transformed type A cells and those of type B cells should be at similar incidences, if the tumors themselves are fully independent from each other, and frequencies of coincidental occurrence of type A or that of type B representation in two, three, four and six independent tumors should be 25%, 12.5%, 6.25%, 3.13% and 1.56%, respectively. For the multiple lesions, therefore, it should be safe to regard five or more tumor nodules carrying an identical inactivated allele as unicentric, i.e., sharing a common parent clone. It is also reasonable to consider that three or more tumor nodules with identical inactivation patterns as demonstrated at both PGK and AR loci are most probably of unicentric origin.

Primers used to amplify PGK and AR genes. Two primer pairs were used for amplifying PGK gene exon 1 (29) (PGK1A: 5'-CTG TTC CTG CCC GCG CGG TGT TCC GCA TTC -3'; PGK1B: 5'-ACG CCT GTG ACG TAA GCT CGT CAG GCC TCC -3'; and PGK2A: 5'-AGC TGG ACG TTA AAG GGA AGC GGC TGG TTA -3'; PGK2B: 5'-TAC TCC TGA AGT TAA ATC AAC ATC CTC TGG -3'). Two primer pairs were for amplifying AR gene exon 1 (29) (AR1A (221+): 5'-GAG GAG CTT TCC AGA ATG TG -3'; AR1B (454+): 5'-CAT GGG CTT GGG GAG A -3'; and AR2A (230+): 5'-TCC AGA ATC TGT TCC AGA GC -3'; AR2B (446-): 5'-TGG GGA GAA CCA TCC TCA CC -3').

Pretreatment with HpaII and amplification for the PGK gene. One-mm thick tissue specimens, as large as 5 mm³ or through the largest cross-section, were minced separately. Genomic DNA was isolated using an extraction kit (QIAamp, Qiagen GmbH, Hilden, Germany) as described previously (29).
DNA extracts from tumors and the non-neoplastic smooth muscle tissues, 10 μl each, were incubated with 5 U of HpaII (Promega, Madison, WI, USA) at 37°C for 12 h in a volume of 20 μl containing 0.2 μl of 10 mg/ml bovine serum albumin (BSA) and 2 μl of 10X reaction buffer, and heated at 95°C for 5 min to inactivate the enzyme. The digested DNA samples, 5 μl each, were then subjected to nested polymerase-chain reaction (PCR). The reaction mixture was 50 μl in volume, containing 4 μl of 10 mM dNTP (Gibco BRL, Life Technologies, Inc., Gaithersburg, MD, USA), primers PGK1A and PGK1B (0.4 pmol each), 5 μl of 10X buffer, 1.5 μl of 50 mM MgCl2, and 2.5 μl of Taq DNA polymerase (Gibco BRL). The amplification was conducted using a PT-200 thermocycler (MJ Research, Inc., Watertown, MA, USA) for 35 cycles (94°C, 40 sec; 58°C, 50 sec; 72°C, 1 min). The PCR mixture (1 μl) was then seeded to the second reaction mixture with PGK2A and PGK2B as primers. The reaction was conducted for 35 cycles (94°C, 40 sec; 56°C, 50 sec; 72°C, 1 min).

Detection of BstXI restriction-polymorphism of the PGK products. The PCR products, 10 μl each, were incubated at 47°C for 8-10 h, in a mixture of 20 μl in volume containing 5 U of BstXI (Promega), 0.2 μl of 10 mg/ml BSA and 2 μl of 10X reaction buffer. The digested products were then resolved in 2% (g/ml) agarose gel containing ethidium bromide (0.2 μg/ml), and visualized under ultraviolet light. Images were recorded using an image-acquisition system (UVP, Inc, Upland, CA, USA). Sizes of products were determined by a 100-bp DNA ladder (Gibco BRL), fluorescence intensities of the products at the positions of 530 bp and 433 bp were assessed using the analysis software LabWork 3.0 (UVP).

Pretreatment with HhaI or HpaII and amplification for the AR gene. Genomic DNA extracts, 10 μl each, were incubated with 10 U of HhaI (Promega) at 37°C for 3 h in a volume of 20 μl containing 0.2 μl of 10 mg/ml BSA and 2 μl of 10X reaction buffer, and heated at 65°C for 20 min to inactivate the enzyme. HpaII-pretreatment, as described above, was also used in some samples to ensure reliability of the reaction, as inconsistent results have been observed regarding HhaI- and HpaII-pretreatment (38). The digested DNA samples, 5 μl each, were then subjected to nested PCR, with AR1A and AR1B as primers for the first round, AR2A and AR2B as primers for the second round, and 56°C as the annealing temperature for both reactions.

Resolving AR products by denaturing polyacrylamide gel electrophoresis. The efficacy of the PCR for the AR gene was assessed by electrophoresis on 2.0% agarose gels. The amplification products, 3 μl each, were mixed with the same volume of sample buffer (99% formamide, 1 mg/ml bromophenol blue, and 1 mg/ml xylene cyanol), and then resolved on the 10% polyacrylamide gel containing 8.0 M urea at 80 V for 8 h using the Hoefer MiniVE apparatus (Amersham-Pharmacia Biotech, Inc., San Francisco, CA, USA). For those with allelic variations as small as one or two CAG repeats, the size difference could not be clearly visualized on the 8-cm long gel, and then the samples were loaded onto a 0.75-mm thick gel as long as 26 cm and resolved using the Hoefer SE660 apparatus (Amersham Bioscience Corp., San Francisco, CA, USA) for 26 h at a voltage of 150 V. Gels were fixed for 4 min in 10% ethanol containing 0.5% acetic acid, treated with 10% nitric acid for 4 min, washed in deionized water for 4 min, and then impregnated in 0.6% silver nitrate for 45 min. After rinsing the gel in deionized water for 5 min, the reaction was visualized in a solution containing 3% sodium carbonate and 0.1% formaldehyde for 5 min. After being soaked in 10% acetic acid for 5 min, the gel was washed in deionized water for 5 min. The reaction was recorded using the image-acquisition system and an ordinary optical camera. Sizes of products were determined using 100-bp and 10-bp DNA ladders (Gibco BRL).

Cloning and sequencing of PCR products. Selected amplification products of PGK, all migrating at 530 bp or at 433 bp, were cloned and sequenced to verify the specificity of the BstXI cleavage reaction. Representative PCR products for AR exon 1 were also analyzed by sequencing in order to correlate their migration behaviors and CAG repeat numbers. All aberrant amplification products were cloned and sequenced to identify the underlying structural alterations.

Evaluation of data and statistical analysis. For both reactions for the PGK and AR genes, a reduction of ≥50% in their signal intensities for the products of either allele, as compared to those obtained using the samples not treated with HpaII or HhaI, is regarded as loss of X-chromosome inactivation mosaicism (29). Each reaction was repeated at least once to guarantee the reproducibility. Moreover, parallel assays were made using non-neoplastic smooth muscle samples of comparable sizes from four different sites of the same uterus as controls. In addition, adjacent smooth muscle tissues, ranging from 1 to 10 mm2 in size and as thick as 1 mm, from 6 uteri were also subjected to the assay at PGK locus in order to exclude possible interference of clonal patches to the evaluation of data as encountered by other authors (39-41).

Statistical analyses were carried out using the software package SPSS 10.0 (SPSS, Inc, Chicago, IL, USA). The mitotic indices of different types of multiple leiomyomas were expressed as median values. Their difference was determined by the two-sided Wilcoxon rank-sum test (42). Relationships between the types and histologic variants were described by t-test and Pearson (χ2) tests. P<0.05 was regarded as being statistically significant.

Results

Non-neoplastic smooth muscle tissues from the 113 uteri were subjected to PCR. A PGK-specific product was obtained in all the cases, digestion with BstXI showing its polymorphism in 34 (30%). Frequently, the band migrating at 530 bp was stronger than the lower one at 433 bp, with their ratio being approximately 3:1 (Figs. 2 and 3), as noted previously (29). Among the 34 cases with polymorphism at the PGK locus, 6 (18%) showed marked allelic skewing (29,43), and were excluded from the following assay.

In order to approve the specificity of PCR and the accuracy of BstXI digestion, amplification products from cases 77 and 78 were cloned and their sequences were determined. Both of the samples were shown on gels to be homozygous for the
BstXI site, giving rise only to a band of 530 bp for the former case and 433 bp for the latter. Sequencing data showed that both products were from exon 1 of the PGK gene, being in accordance with the data reported previously (44,45). As demonstrated by BstXI digestion, a difference was found at position 438 from the 5’-end between these 2 samples, with adenosine at this position for case 77, and guanosine at this position for case 78.

In total, 101 (89%) of the 113 cases examined were found to be length-polymorphic at exon 1 of the AR gene, showing two products migrating between positions of 190- and 240-bp DNA markers on the denaturing polyacrylamide gel (Figs. 2 and 4). Sixty-three (62%) of the polymorphic cases, with 287 tumors, were suitable for the subsequent assay, and 38 (38%) were excluded for the following reasons: i) the allelic length-variation was too small for precise assessment in 33 cases; ii) marked skewing between two AR alleles in 4 cases; and iii) structural alteration in one of the AR alleles identified in one case. However, the mutation in the latter case (case 79), with two tumor nodules, provided another parameter to describe their inter-nodular relationship, as detailed below.

Combining these two reactions, 76 of the 113 samples (67%) were suitable for X-chromosomal inactivation analysis, whereas only 28 (25%) or 63 (56%), respectively, were suitable when either the assay for the PGK gene or that for the AR locus were used.

Among the 76 informative cases examined, 21 were found to carry a single tumor nodule, and 55 showed multiple leiomyomas with 2 (20 cases) or >2 (35 cases) tumor nodules. A total number of 318 tumors were examined using the PGK or/and AR assays, with 315 being informative. Three tumors from case 46, with massive hyalinization, were excluded from the analysis. Loss, or a pronounced reduction, of one band was observed for all the samples after the pretreatment with HpaII or HhaI, reflecting their monoclonality (Figs. 2-4).

Reproducibility of the results was ensured by taking more than one sample from different areas within tumors >1 cm in diameter. Consistent results were obtained for all of the cases examined (Fig. 2). In order to exclude possible pitfalls in the interpretation of the results by ignoring clonal patches, 221 samples of the adjacent smooth muscle tissue of different sizes were taken from six uteri and subjected to the assay for PGK locus. The tissues with 10-mm² (n=6) and 5-mm² (n=42)

<table>
<thead>
<tr>
<th>Tumor nodules (n)</th>
<th>Cases examined (n)</th>
<th>Case codes (n)</th>
<th>Cases with identical allele inactivated in all (or most) nodules (n)</th>
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cross-sections did not show identifiable allelic differences in
the signal intensity ratio by HpaII-pretreatment. However,
imbalance of the X-chromosomal inactivation was observed
in 3-, 2-, and 1-mm² samples at frequencies of 14% (6/42),
19% (8/42) and 28% (25/89), respectively, its occurrence being
closely associated with the sample size examined (F=5.807,
P<0.01). The data showed that clonal patches are indeed present
in the uterine leiomyogenic tissues. The patch sizes should be
much smaller than 5 mm² in cross-section. Therefore, patching
cannot interfere with the evaluation of data with the procedures
employed in this study (Fig. 5).

A total number of 297 tumor nodules were found in the
55 multiple leiomyomas examined (5.4 nodules per case in
average). The relationship among different tumor nodules in
multiple cases, as revealed by their inactivated PGK and AR
alleles, appeared to be associated with the nodule numbers
(Tables I and II). For the cases with >10 tumor nodules (cases
32-50), a unicentric origin will be considered if all tumor nodules share
the same inactivated PGK and/or AR alleles. As listed in
Table I, the 19 cases examined were divided into three types
according to their inter-nodular relationships: i) the multicentric
type (cases 39, 43-45, and 50), with the two alleles randomly
inactivated among different lesions as listed in Table II; ii) the
unicentric type (cases 32-35, 40-42 and 46), with all lesions
sharing the same inactivated allele as shown in Fig. 6; and iii)
the concomitant occurrence of the unicentric and multicentric
types in 6 cases (cases 36-38, and 47-49; Table II), with only
one nodule carrying a different inactivated allele as shown in
Fig. 7.

Among the unicentric multiple cases, case 46 was a 43-
year-old woman when admitted to the hospital. Multiple
masses were detected in her uterus and a hysterectomy was
performed. Grossly, 10 apparently discrete nodules sizing
from 15 to 30 mm were found in the uterine corpus, with
myxoid appearance in some areas. Microscopic examination
showed features characteristic of the smooth muscle tumor.
This was confirmed by its immunoreactivities for vimentin.

<table>
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<tr>
<th>Case codes</th>
<th>Nodules (n)</th>
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<th>AR alleles at</th>
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-, data non-informative due to the lack of polymorphism or marked allelic skewing.

For the cases with 4 to 10 tumor nodules (cases 32-50), a
unicentric origin will be considered if all tumor nodules share
the same inactivated PGK and/or AR alleles. As listed in
Table I, the 19 cases examined were divided into three types
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types in 6 cases (cases 36-38, and 47-49; Table II), with only
one nodule carrying a different inactivated allele as shown in
Fig. 7.
smooth-muscle actin and desmin, and its negativity for CD10. Focal myxoid change and massive hyaline degeneration were observed, with mild nuclear atypia in focal areas (Fig. 8A). A mitotic index of 14/10 HPF (56/40 HPF) was found in more cellular areas. Mitotically active leiomyoma was proposed for the case. Then, the sample was subjected to the X-chromosome inactivation assays. Among these ten nodules, PCR was successfully conducted in 7, the others failed because of massive hyalinization (Fig. 8B). As shown in Fig. 9, similar inactivated alleles were found in all of these nodules, taking the sensitivity of the PGK clonality assay as showing allelic imbalance with the examined sample containing >50% of monoclonal cells (29), i.e. with the ratio of two cell types >3:1.

Figure 5. Tissue size requested for the clonality analysis based on X-chromosomal inactivation mosaicism as a negative control, in consideration of clonal patches in uterine myometrium. The patches composed of type A cells (white) and those of type B cells (gray) are distributed randomly, each size is ~2 mm in diameter according to the data from the current study. The frame, 5 mm², reflects the sample size requested for the clonality assays in the study, covering 20 patches, clearly demonstrating the mosaicism in non-neoplastic tissues. However, for the samples of 1 mm², as represented by the smallest square, 7 of the 25 squares (28%) fall into the area occupied completely (A, B, D and G) or mostly (C, E and F) by the same cell type. With areas of 2 and 3 mm² considered, the frequencies decrease to 12.5% (2/16) and 11% (1/9), respectively, taking the sensitivity of the PGK clonality assay as showing allelic imbalance with the examined sample containing >50% of monoclonal cells (29), i.e. with the ratio of two cell types >3:1.

Figure 6. Multiple leiomyomas of the unicentric type, cases 35 and 41 with 5 (T1-T5) and 6 tumor nodules (T1-T6), respectively, demonstrated by X-chromosomal inactivation analysis at both PGK and AR loci for the former case, and only at the PGK locus for the latter. Identical PGK and AR alleles were preserved through the pretreatment (−, before; +, after) with the restriction enzymes HpaII and HhaI, respectively. NT, non-neoplastic smooth muscle tissue.

Figure 7. A multiple uterine leiomyoma of the mixed type (case 51), with the PGK allele, recognizable by BstXI, preserved through the pretreatment with HpaII (−, before; +, after) in only one tumor nodule (T1), the other allele was preserved in all of the remaining 8 nodules (T2-T9). NT, non-neoplastic smooth muscle tissue.

smooth-muscle actin and desmin, and its negativity for CD10. Focal myxoid change and massive hyaline degeneration were observed, with mild nuclear atypia in focal areas (Fig. 8A). A mitotic index of 14/10 HPF (56/40 HPF) was found in more cellular areas. Mitotically active leiomyoma was proposed for the case. Then, the sample was subjected to the X-chromosome inactivation assays. Among these ten nodules, PCR was successfully conducted in 7, the others failed because of massive hyalinization (Fig. 8B). As shown in Fig. 9, similar inactivated alleles were found in all of these nodules.
samples, demonstrating a common clonal origin for the 7 nodules examined. The multi-nodular appearance simply reflected intramural spreading of the tumor cells, and the aggressiveness of this tumor. Focal coagulative necrosis (Fig. 8C) and highly cellular areas (Fig. 8D) were found by re-sampling and further examination. A final diagnosis of leiomyosarcoma was made.

Among the 20 cases with two tumor nodules and suitable for the X-chromosomal inactivation analysis, 14 (70%) showed the same inactivated PGK or AR allele (Table I). Three of them were examined by both assays at the PGK and AR loci, with concordant results demonstrated in two and different data obtained in one. Of the 11 cases with three tumor nodules, four (46%) showed different X-chromosomal inactivation patterns (Fig. 4), and 7 (64%) were found to carry the same inactivated allele, with consistent results obtained for both the PGK and AR loci in one of them. It is evident that the identical inactivation pattern in three discrete lesions from the same uterus frequently, if not always, reflects a common parental cell origin.

Considering the limitations in evaluating the inter-nodular relationships in cases with only a few tumor nodules, the presence of an identical gene mutation or the same chromosomal alteration, which occurs otherwise rarely, would be strong evidence for a unicentric origin of different tumors in the same patient. Fortunately, it was encountered in a multiple case (case 79) with two grossly discrete tumors. Loss of heterozygosity (LOH) was observed in these two tumor nodules, the larger product being undetectable. An aberrant amplification product was found in both samples, migrating at the position corresponding to that of the 190-bp DNA marker (Fig. 10). The products from tumors and adjacent smooth muscle tissues were cloned and sequenced. Those from the smooth muscle tissue showed homology to the AR gene as compared to the reported sequence (46-48), the CAG repeat number of the smaller and larger products being 21 and 23, respectively. A deletion mutation within the STR was detected in the aberrant products from both tumors, with the CAG repeat number being reduced to 11.

Forty-nine cases of multiple leiomyomas were examined for their mitotic activity and histologic variants, with one (case 46)
study, we examined 315 leiomyomatous nodules from 76 uterine leiomyomas were examined for their X-chromosomal inactivation analysis at the PGK and AR loci, and provided strong evidence for the view.

Considering possible interference with the clonal cell clusters occurring under physiological and pathological conditions (39-41,48), we demonstrated the patches in the non-neoplastic smooth muscle tissue. Their sizes were shown to be <5 mm², most probably ranging from 2 to 3 mm² in cross-section, as estimated by comparing the observed frequencies of X-chromosomal inactivation mosaicism in the samples and its probabilities in a representative, albeit simplified, model as depicted in Fig. 5. The data clearly demonstrated the reliability of the assay with the control samples covering an area of 5 mm².

While most of the uterine leiomyomas appear benign, some rare cases (1-3/1000) can progress to leiomyosarcoma (49,50). Different chromosomal alterations were identified in both types of lesions by comparative genomic hybridization (51). In addition, there are more cases showing histologic features intermediate between them (4-10). These lesions often cause diagnostic problems, because their clinical behavior cannot be predicted with certainty (11,12).

The relationship between different tumor nodules in multiple uterine leiomyomas remains a matter of debate. Recent observations on three leiomyogenic proliferative diseases related to the multiple uterine leiomyoma, including DPL, IVL and BML, have demonstrated the common clonal origin of different lesions and apparently associated the unicentric feature to their clinical aggressiveness. DPL has been shown to be neoplastic. An identical inactivated X chromosome was found in all of its extraterine leiomyogenic proliferative nodules and the intramural leiomyoma (20). Consistent cytogenetic alterations were found in most of the nodules examined as well (19,20). These data reveal the disseminating or metastatic feature of the neoplasm. A study by Quade et al (52) on IVL demonstrated the same X-chromosomal inactivation pattern in different lesions within and outside the uterus. Chromosome translocation (12,14) (q15;q24), a frequent cytogenetic alteration in uterine leiomyoma (53), was also observed in all of these nodules. The same X-chromosomal inactivation pattern was demonstrated in uterine tumors and all pulmonary tumorlets from the BML cases (17,54), supporting the view that pulmonary lesions are metastatic (15,16). In consideration of these findings, the relationship among different tumor nodules in the multiple leiomyomas may also reflect their biological behavior.

While most authors favor a multicentric origin of different tumors in multiple uterine leiomyomas based on cytogenetic (22,23,25) and X-chromosomal inactivation analyses (21,23,24,26-28), macroscopically discrete tumors occurring in the same uterus were incidentally observed to share the same inactivated PGK or AR allele and identical cytogenetic alterations by a few groups in a limited number of cases (19,27,29-31). In the present study, 55 cases of multiple leiomyomas were examined for their X-chromosomal inactivation patterns. Random inactivation patterns were revealed among different tumor nodules from 20 (36%) of these cases, indicating their independent, multicentric origins. In 29 (53%) of them, however, the identical inactivated allele was found in all tumor nodules from the same uterus. Obviously, the occurrence of the same X-chromosomal inactivation pattern

![Image](image.png)

Figure 10: Loss of heterozygosity at AR exon 1 in both of the tumor nodules (T1, T2) from case 79. Allelic length polymorphism at the AR locus was visualized on the denaturing polyacrylamide gel (upper) in the non-neoplastic smooth muscle tissue (NT). Loss of band A (arrowheads) and occurrence of an aberrant product (arrow) migrating at the position corresponding to the 190-bp DNA marker (M) were seen in samples from both tumor nodules. Sequencing data were shown in the frame (low). A 36-base deletion, represented by dash lines, was found within the short-tandem repeat region of AR2B. Positions of primer pair AR2A/AR2B are underlined.

excluded from the test for its obvious malignant behavior as described above. The mitotic counts were not >2/10 HPF in 48 cases and reached 2.6/10 HPF in one. Median indices for the unicentric/mixed-type cases and the multicentric-type tumors were 0.6/10 HPF and 0.2/10 HPF, respectively. The mitotic activity appeared significantly elevated (Z=2.424, P=0.015) in the former compared to the latter group. No association was observed between mitotic index and the numbers of tumors (P>0.05). In addition, no statistical significance was found between the inter-nodular relationship and the histologic changes, including nuclear pleomorphism (χ²=1.918, P>0.05), myxoid (χ²=1.367, P>0.05) and hyaline degeneration (χ²=0.453, P>0.05).

Follow-up was conducted for the 55 multiple cases, ranging from 64 to 112 months (78 months in average). Metastases were found in case 46 with the multi-nodular leiomyosarcoma occurrence of an aberrant product (arrow) migrating at the position corresponding to the 190-bp DNA marker (M) were seen in samples from both tumor nodules. Sequencing data were shown in the frame (low). A 36-base deletion, represented by dash lines, was found within the short-tandem repeat region of AR2B. Positions of primer pair AR2A/AR2B are underlined.

Discussion

It has been proposed that uterine leiomyomas are monoclonal in their cell composition (23,24,29,37,48). Hence, the development of this neoplasm is considered a process of clonal expansion of an altered leiomyogenic cell. In the present study, we examined 315 leiomyomatous nodules from 76 cases using X-chromosomal inactivation analysis at the PGK and AR loci, and provided strong evidence for the view.

While most of the uterine leiomyomas appear benign, some rare cases (1-3/1000) can progress to leiomyosarcoma (49,50). Different chromosomal alterations were identified in both types of lesions by comparative genomic hybridization (51). In addition, there are more cases showing histologic features intermediate between them (4-10). These lesions often cause diagnostic problems, because their clinical behavior cannot be predicted with certainty (11,12).

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in different tumors may be a fortuitous event in some of these cases. This is particularly true for those with two tumors. In the latter cases, however, presence of the identical AR gene mutations in different tumors adds a piece of strong evidence for the unicentric origin, as the incidence of this genetic event is <0.7% (2/287) and the probability for its coincidental occurrence in two independent tumors from the same uterus is <0.005% (0.7% x 0.7%).

Among the 55 multiple cases examined, 6 (11%) showed an inactivated allele in one tumor and another inactivated allele in all of the remaining nodules, comprising a mixed type. In these cases, one of two preexisting genotypically different, independent tumors may acquire invading or migrating capacity and hence give rise to several daughter nodules with identical genotype.

The results of our study, for the first time, provide strong evidence for the presence of unicentric multiple uterine leiomyomas, even though the definite incidence remains to be elucidated. The results indicate that, at least in some of the cases, most tumor nodules developed from one parent tumor cell clone by migration or spreading of the neoplastic cells, which means that some of the unicentric multiple leiomyomas are potentially aggressive. This is further evidenced by the elevated mitotic activity observed in the unicentric multiple leiomyomas.

It remains to be settled whether the mitotically active leiomyoma is more aggressive than leiomyomas with low mitotic activity. In the present study, identical X-chromosomal inactivation patterns at both PGK and AR loci were found in seven discrete tumors examined from a case (case 46) otherwise considered a mitotically active leiomyoma. Our X-chromosomal inactivation analysis data revealed the invading capacity of the tumor. Focal coagulative necrosis was detected by re-sampling, and a diagnosis of leiomyosarcoma was made for the case. The subsequent follow-up proved its malignant behavior. Therefore, X-chromosomal inactivation analysis is of diagnostic value for distinction of the leiomyosarcoma from other multinodular smooth muscle neoplasms with elevated mitotic activity.

In the uterus, those with mitotic figures <5/10 HPF are considered non-aggressive, benign lesions (55), while most of the tumors falling into the category of leiomyosarcoma are highly malignant (56,57). Actually, little is known about the invading capacity of uterine leiomyomas, though it may be a more representative parameter for behavior evaluation of the smooth muscle tumors. Between the highly malignant and benign entities of the spectrum, there should be some lesions showing local aggressiveness and classified into benign uterine leiomyoma based on the current criteria. Some of the cases may correspond to the mitotically active leiomyoma, but the relevance remains to be established between the elevation of mitotic activity and acquisition of invading capacity. The intramural spreading usually does not result in extraterine dissemination or metastasis possibly due to their dependence on hormonal stimuli and anatomical features of the uterus. The occurrence of unicentric multiple leiomyomas, as described above, may reflect the local aggressiveness. There have been several reports describing the presence of some extraterine smooth muscle tumors associated with multiple uterine leiomyomas (58-62). We think that some of the multiple uterine tumors are unicentric, and that these extrauterine lesions may represent dissemination or metastasis. In this context it is interesting to note that a majority of the clinically aggressive leiomyogenic neoplasms, including IVL (14), BML (15-18,54,63-66) and DLP (19,20,67,68), emerge as multiple uterine lesions. Further study is needed for clarification of the clinical behaviors of this tumor type by long-term follow-up surveys.

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