Molecular Analysis of 11q13 Breakpoints in Multiple Myeloma

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The t(11;14)(q13;q32) chromosomal translocation, which is the hallmark of mantle cell lymphoma (MCL), is found in approximately 30% of multiple myeloma (MM) tumors with a 14q32 translocation. Although the overexpression of cyclin D1 has been found to be correlated with MM cell lines carrying the t(11;14), rearrangements of the BCL-1/cyclin D1 regions frequently involved in MCL rarely occur in MM cell lines or primary tumors. To test whether specific 11q13 breakpoint clusters may occur in MM, we investigated a representative panel of primary tumors by means of Southern blot analysis using probes derived from MM-associated 11q13 breakpoints. To this end, we first cloned the breakpoints and respective germ-line regions from a primary tumor and the U266 cell line, as well as the germ-line region from the KMS-12 cell line. DNA from 50 primary tumors was tested using a large panel of probes, but a rearrangement was detected in only one case using the KMS-12 breakpoint probe. Our results confirm previous findings that the 11q13 breakpoints in MM are scattered throughout the 11q13 region encompassing the cyclin D1 gene, thus suggesting the absence of 11q13 breakpoint clusters in MM.

CHROMOSOMAL translocations affecting the immunoglobulin heavy chain (IGH) locus on 14q32 represent the mechanism of activation of a number of proto-oncogenes in B-cell lymphoid neoplasms.1 The t(11;14)(q13;q32) chromosomal translocation is associated with approximately 70% to 90% of mantle cell lymphomas (MCL)2,3 and leads to the overexpression of the cyclin D1 gene.4,5 The breakpoints on chromosome 11q13 were initially found clustered in a 1-kb region, named the major translocation cluster (MTC), of the BCL-1 locus. Further investigations have shown that breakpoints may occur telomeric of the MTC in a region of about 120 kb between the MTC and the cyclin D1 loci.3 Molecular analyses, including fluorescence in situ hybridization (FISH), have also demonstrated that breakpoints may occur either centromeric or telomeric of this 120 kb region in some cases.4,5,9

Multiple myeloma (MM) is a malignant proliferation of bone marrow plasma cells that is characterized by a wide spectrum of clinical entities and whose molecular pathogenesis is still largely unknown.10-11 Cytogenetic analyses in MM are limited and difficult mainly because of the low proliferation rate of malignant plasma cells. However, in about 20% to 40% of tumors with an abnormal karyotype, a 14q− marker has been reported.12,13 This marker is generally the consequence of translocation events involving the IGH locus on chromosome 14q32. Interestingly, in almost 30% of cases with cytogenetically detectable 14q−, the marker is the result of a t(11;14)(q13;

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MATERIALS AND METHODS

Pathological samples. Bone marrow or peripheral blood samples from 50 MM patients investigated by Southern blot analysis were collected during the course of standard diagnostic procedures. The diagnosis and clinical staging of MM was made according to the criteria described by Durie and Salmon.17 These samples came from a larger series of 88 previously investigated primary MM tumors.18 Forty patients were at first diagnosis: six in stage I (indolent phase), 22 in stage II, and 12 in stage III; five patients were evaluated at clinical relapse and five were affected by plasma cell leukemia (four at diagnosis and one at relapse). Twenty-one of the patients were male and 29 female; their median age was 61 years (range, 42 to 80). Monoclonal component was as follows: IgG (35 patients), IgA (11), kA chain 30/16; k chain (3) λ chain (1). No conventional cytogenetic analyses (G-banding) of these samples were available.

The previously reported tumor LB41118 was derived from a 69-year-old male patient affected by IgGk-type plasma cell leukemia with 2 months survival; no karyotype was available in this case. Case AC97 was a 74-year-old female with a κ-type MM in clinical stage III that had
of comigration between the DNA fragments containing constant switch-mediated chromosomal translocations was based on the absence of any rearrangement of the 

IGH alleles from case LB411 and the U266 cell line was kindly provided by Dr T. Otsuki (Okayama, Japan).

DNA preparation and Southern blot analysis. Mononuclear cell suspensions with more than 95% viability were prepared from the pathological samples by means of Ficoll-Hypaque gradient centrifugation; the percentage of malignant plasma cells identified by immunocyto-morphologic analyses was between 22% and 98%. DNA from pathological samples and cell lines was purified by extraction with phenol-chloroform and ethanol precipitation.21 A total of 10 µg of genomic DNA was digested with BamHI, EcoRI, or HindIII restriction enzymes, electrophoresed in a 0.7% agarose gel, and then denatured, neutralized, and transferred to nylon filters (Amer- sham International, Amersham, UK). The filters were hybridized to 32P-labeled probes according to the manufacturer’s specifications, washed in 0.5 × SSC (NaCl/Na citrate) at 65°C, and autoradiographed using an intensifying screen at −80°C.21 IGH gene rearrangement was analyzed using previously described probes.22-24 The probes used for the rearrangement analysis of the BCL-1/cyclin D1 locus were MTC, p94, and cyclin D1 cDNA.2,3,6

Molecular cloning. The identification of a t(11;14)(q13;q32) chromosomal translocation in an unknown case (LB411) and the U266 cell line was made possible following a Southern blot approach recently reported by us,15 which allows the identification of putative switch-translocated IGH alleles on the basis of the absence of any linkage between different IGH regions. A quite similar approach has been previously reported by others.25 We reasoned that, as a result of immunoglobulin gene recombination during maturation of B cells, the rearranged joining-switch-constant IGH regions are generally contained on a novel BamHI restriction fragment, and that a translocation event involving the switch region should therefore generate a rearranged BamHI fragment containing the 3′ constant region, but not the 5′ joining sequences of the IGH gene. In our Southern blot assay, the DNA was digested with BamHI restriction enzyme, and the filters subsequently hybridized with the JH, Cµ, Cα1, and Cy1 probes. The identification of rearranged IGH alleles as potential candidates for switch-mediated chromosomal translocations was based on the absence of coniguration between the DNA fragments containing constant IGH regions and those positive for the JH probe. Recombinant phage clones containing translocated Co-rearranged IGH alleles from case LB411 and the U266 cell line were obtained by complete digestion of genomic DNA with BamHI, the subsequent ligation of gel-purified fragments into λEMBL3 phage vectors (Stratagene, La Jolla, CA) and screening with the Cs1 probe. The germ-line regions of chromosome 11 were isolated by screening a genomic library of human placenta DNA (Clontech, San Diego, CA) using probes derived from recombinant clones. The normal 11q13 region encompassing the breakpoint in KMS-12 was cloned by screening a phage genomic library with a 193-bp fragment specific for the KMS-12 breakpoint region, which was obtained by polymerase chain reaction (PCR) amplification of genomic DNA using a pair of previously reported primers.16 The library screening and plaque isolation were performed according to established procedures.21 The inserts were analyzed using restriction enzyme mapping and then subcloned into plasmid vector pGEM3 (Promega, Madison, WI).

DNA sequencing. DNA sequence analysis was performed on restriction fragments cloned into pGEM3 plasmid (Promega) by “dideoxy” chain-termination analysis using the Sequenase sequencing kit (USB, Cleveland, OH).

cDNA amplification. The synthesis of the first strand cDNA was performed as previously described.18 PCR amplifications were made by diluting 5 µL of first-strand cDNA from each individual case into a 50-µL PCR mixture. A 196-bp fragment encompassing exons 3 and 4 of cyclin D1 gene6 was amplified using the following primers: sense 5′-AAGACATCATCCGCAAACAC-3′; antisense 5′-TACACCTGTGACGGTCGA-3′. Thirty cycles of amplification were performed at 94°C for 30 seconds, 60°C for 30 seconds, and 72°C for 30 seconds.

FISH. The chromosome preparations were hybridized in situ with probes labeled with biotin or directly with the fluorochrome, Cy3 (Amersham, Little Chalfont, UK) by nick translation, as described21 with minor modifications.26,27 Briefly, 200 ng of labeled probe were used for each experiment, and hybridization was performed at 37°C in 2 × SSC, 50% (vol/vol) formamide, 10% (wt/vol) dextran sulphate, 5 µg Cot1 DNA (Boehringer, Mannheim, Germany), and 3 µg of sonicated salmon sperm DNA in a volume of 10 µL. Posthybridization washing was at 60°C in 0.1 × SSC (three times). Biotin-labeled DNA was detected using fluorescein isothiocyanate (FITC)-conjugated avidin (Vector Laboratories, Burlingame, CA). The chromosomes were identified by simultaneous 4′,6-diamidino-2-phenylindole dihydrochloride (DAPI) staining, which produces a Q-banding pattern. Chromosome 11 and 14 painting probes were obtained by Ala-PCR amplification of the somatic cell hybrids retaining only the human chromosome 11 or 14.28 Digital images were obtained using a Leica DMR epifluorescence microscope equipped with a CCD camera (Cohu, Inc, San Diego, CA). FITC-avidin, Cy3, and DAPI fluorescence signals were detected using specific filters and recorded separately as gray scale images.
oring and image merging were performed using Adobe Photoshop software (Adobe Systems, Mountain View, CA).

Probes for FISH analysis. The probes used are described in Fig 1. The cosmid clones representative of the cyclin D1 and FGF3 loci were isolated by screening a cosmid library of human placenta (Clontech) with probes specific for cyclin D1 or FGF3/int-2 cDNA.29 The FGF4/int-1 locus was investigated using an 8-kb genomic fragment containing the 5′ of the gene.30 The cosmid clones I4, R4B, and pHS-11 encompassing the MTC region have been previously described.9 All of the YACs were obtained from the human CEPH library (YAC Screening Center, DIBIT, Milan, Italy); YAC clones 961-H-2 (800 kb) and 744-F-12 (330 Kb) have been previously reported15 and are both negative for the MTC region; YAC clone 877-D-8 (1080 kb) was identified by the presence of the D11S911 marker, telomeric of the GARP locus on 11q14.9,32 The IGH locus was analyzed using a phage clone containing the 18-kb ColI BamHI germ-line fragment.23

RESULTS
Cloning of a t(11;14)(q13;q32) in a MM primary tumor. We have recently reported the identification of IGH rearranged alleles as potential candidates for switch-mediated chromosomal translocations in 21 of 88 cases of primary MM tumors investigated.18 To confirm the presence of chromosomal breakpoints in these alleles, we cloned the rearranged fragments from three cases (two Col and one Cµ rearrangement). In two of these tumors, molecular cloning and FISH analyses showed the presence of a novel t(4;14)(p16.3;q32) chromosomal translocation.18 In the third MM tumor (case LB411), a t(11;14)(q13;q32) chromosomal translocation was identified by cloning a 10-kb BamHI ColI rearranged fragment (Fig 2) that was apparently negative for JH sequences.18 This fragment was isolated as described in Materials and Methods and used as a probe in the FISH analysis of metaphase spreads from mitogen-stimulated normal peripheral blood lymphocytes. This clone hybridized to both chromosome 14q32 and chromosome 11q13, thus indicating that the rearrangement was a result of a t(11;14)(q13;q32) chromosomal translocation (data not shown). About 25 kb of the normal genomic region surrounding the breakpoint on chromosome 11q13 were cloned by isolating two overlapping phages from a normal genomic library (Fig 2). None of this genomic region hybridized with probes specific for the BCL-1 locus (MTC, p94, and full-length cyclin D1 cDNA), thus confirming the absence of rearrangements in these regions observed in Southern blot analysis of LB411 DNA (data not shown). Subsequently, this region was hybridized with a set of 11q13 genomic clones (see Fig 1); cosmids R4B and pHS-11 were both positive, thus indicating that the breakpoint in case LB411 occurred approximately 40 to 45 kb telomeric of the MTC region and consequently 65 to 70 kb centromeric of the cyclin D1 gene. Interestingly, reverse transcriptase (RT)-PCR

Fig 2. Molecular cloning of the chromosomal breakpoint from case LB411. (A) Schematic representation of the breakpoint and the respective 11q13 germ-line region. From the top: a diagram of the BCL-1/cyclin D1 locus where the cosmid clones I4, R4B, pHS11 are located is shown; the 11q13 germ-line region and the probes used for the Southern and Northern blot analyses are shown as solid lines. The vertical arrow indicates the breakpoint position. In the t(11;14) breakpoint clone, chromosome 14 is indicated by open boxes with black or stippled boxes representing different IGH regions and chromosome 11 is shown as a solid line. Restriction enzyme symbols: B, BamHI; R, EcoRI; H, HindIII; X, XhoI. (B) The nucleotide sequence of the breakpoint region and its alignment with 11q13 and 14q32 germ-line sequences are shown.
analysis showed that this case was overexpressing cyclin D1 (data not shown).

Cloning and mapping of the 11q13 breakpoint in MM cell line U266. We investigated cyclin D1 expression in a panel of MM-derived cell lines (U266, KMM1, JNJ3, OPM2, Karpas, IM9, Sultan, LP-1) for which karyotypic analyses have been reported to be negative for the t(11;14)(q13;q32) chromosomal translocation. Interestingly, Northern blot analysis demonstrated a high level of cyclin D1 expression in U266 cell line (Fig 3A). On the basis of this evidence, we looked for the presence of an illegitimate IGH recombination in U266 DNA by Southern blot analysis. As shown in Fig 3B, hybridization of BamHI-digested DNA with the Cca1 probe detected a rearranged fragment that did not comigrate with the rearranged JH allele. A phage library was constructed using BamHI-digested DNA from the U266 cell line and screened with the Cca1 probe. The rearranged Cca1 fragment was isolated and tested on normal metaphases by FISH analysis, which showed hybridization with chromosomes 14q32 and 11q13 (data not shown) that was consistent with the presence of a t(11;14)(q13;q32) chromosomal translocation. We next cloned about 20 kb of the normal genomic region surrounding the breakpoint by isolating a recombinant phage from a normal genomic library (Fig 3C). This genomic region did not hybridize with MTC, p94, or full-length cyclin D1 cDNA and with any of the available 11q13 clones (see Fig 1).

To define the location of the U266 11q13 breakpoint more precisely, we performed FISH analyses on metaphases spreads. As shown in Fig 4A, the chromosome 11 painting probe hybridized with two chromosomes on U266 metaphases: an apparently normal chromosome 11 and a structurally altered chromosome consisting of the 11pter→11q13-14 region and extra material of unknown origin. Hybridizations with probes specific for the MTC region, and the cyclin D1, FGF4, and FGF3 loci, indicated that all of these regions are contained in the putative 11q13 region of this abnormal chromosome (data not shown).
Hybridization with the chromosome 14 painting probe clearly detected two different chromosomes: an apparently normal chromosome 14 and an abnormal chromosome with a large amount of chromosome 14 material on its long arm. (Fig 4B). Thus, no detectable exchange of material between chromosomes 11 and 14 was observed with painting probes hybridization: a finding consistent with the absence of a cytogenetically detectable t(11;14)(q13;q32) in U266 cell line. Interestingly, the Cα1 clone was clearly detected at 11q13-14 on the abnormal chromosome 11 (colocalized with each of the 11q13 probes tested above; Fig 4C and data not shown), as well as in the telomeric region of both chromosomes recognized by the 14 painting probe (Fig 4C). These findings indicated the presence of a t(11;14) and suggested that the breakpoint on 11q13 may be located telomeric of the FGF3 locus. Interestingly, the YAC clone 877-D-8 specific for a region telomeric of the GARP locus on 11q14 hybridized to the abnormal chromosome 11 in U266 metaphases, thus suggesting that this locus is still retained. Triple-hybridization experiments using MTC (green), Cα1 (red), and 877-D-8 (green) probes showed that the Cα region is apparently located between the MTC and GARP loci (Fig 4C). These findings indicate that the t(11;14) translocation in U266 is the result of a complex chromosome rearrangement, as is also suggested by the presence of extra material on the abnormal chromosome 11. The FISH experiments performed in an attempt to elucidate the origin of the extra material on this chromosome gave very complex results; however, chromosomes 3 and 4 are apparently involved (data not shown).

Mapping of 11q13 breakpoints in MM tumors with a cytogenetically detectable t(11;14)(q13;q32) chromosome translocation. We used FISH analysis to investigate the approximate location of the 11q13 breakpoints in the cell line KMS-12 and a primary MM tumor (AC97), both carrying a t(11;14)(q13;q32) chromosomal translocation. While this work was in progress, Chesi et al16 reported the cloning of a t(11;14) breakpoint in the KMS-12; more recently, Vaandrager et al33 have reported that this breakpoint is located approximately 215 kb centromeric of the MTC region and is juxtaposed to the Cγ2 of the IGH locus. Therefore, only part of our results will be presented. In particular, double-color FISH on KMS-12 metaphase spreads with the painting probes specific
for chromosomes 11 and 14 showed the presence of five putative 14 (der) chromosomes containing material from chromosome 11 (Fig 5A1). FISH analyses further demonstrated that the MTC, cyclin D1, FGF4, and FGF3 loci were all contained in these chromosomes and apparently juxtaposed to sequences recognized by the Cox1 probe (Fig 5A2 and data not shown). The YAC clones 961-H-2 and 744-F-12 located centromeric of the MTC region (see Fig 1) did not hybridize with the 14 (der) chromosomes (Fig 5A3 and data not shown), but with two small and structurally-altered chromosomes recognized by the painting and centromeric-specific probes of chromosome 11 (Fig 5A1, 3, and data not shown). These results suggest that the breakpoint is centromeric of the MTC region and are consistent with previously reported data.33

Case AC97 carried a t(11;14)(q13;q32) chromosomal translocation (see Materials and Methods; data not shown) without any evidence of the involvement of the BCL-1/cyclin D1 regions at Southern blot analysis (data not shown). No data concerning cyclin D1 expression were available in this case. Two-color FISH analysis in interphase nuclei with probes of the 11q13 region showed that the breakpoint in this case is apparently located between MTC and the cyclin D1 locus. In particular, colocalization of signals (red and green) were detected on interphase nuclei when YAC 744-F-12 was used together with the pH511 clone (Fig 5B) and when the cyclin D1 cosm id was cohybridized with the FGF3-specific clone (data not shown), but disassociation of signals was observed when the pH511 and cyclin D1 cosmids were used (Fig 5B).

Lack of clusters of 11q13 breakpoints in MM. Because the t(11;14)(q13;q32) translocation is recurrently found in MM, we investigated by Southern blot whether any of the cloned 11q13 breakpoints were involved in the translocation in other MM cases using the probes described in Fig 6, which are specific for case LB411 and cell lines U266 and KMS-12. We tested DNA from 50 MM primary tumors for which karyotype data were not available. Furthermore, all of these cases have been found to be negative for rearrangements of the MTC and p94 regions and cyclin D1 locus (data not shown). Southern blot analysis was performed on BamHI DNA digests and, when possible, on EcoRI and HindIII digests. Rearrangements were detected in only one tumor (patient LB104) and involved the 11q13 region where the breakpoint of the KMS-12 cell line is located (Fig 6). Case LB104 was a 72-year-old female patient affected by an IgA l-type MM in clinical stage IIA at diagnosis and with 32 months survival. Cyclin D1 expression was evaluated by RT-PCR in only 11 of the 50 MM patients (including case LB104). Specific amplified fragments were detected in case LB104 and in an IgGa-type MM tumor (case LB413) from a 75-year-old female patient in clinical stage IIIB at diagnosis with 13 months survival (data not shown).

**DISCUSSION**

Conventional cytogenetic analyses of MM are generally a difficult task because of the low proliferation rate of malignant plasma cells. Abnormal karyotypes have been reported only in about 40% of MM and at a higher frequency in plasma cell leukemia; interestingly, in about 20% to 40% of MM patients with an abnormal karyotype a 14q- marker is observed.12,13 In about 30% of the cases, the 14q- marker originates through a t(11;14)(q13;q32) chromosomal translocation, which is mainly associated with MCL and involves the cyclin D1 gene on 11q13.12 However, rearrangements of the BCL-1/cyclin D1 regions involved in MCL have been rarely demonstrated in
cases of MM carrying such a translocation (mainly cell lines) or in unselected tumors.\cite{9,14,16} The overexpression of cyclin D1 has been observed in the majority of MM-derived cell lines carrying the t(11;14), thus suggesting that this translocation may lead to the deregulation of this gene in MM, as it does in MCL.\cite{7,9,16} The main aim of the present study was to investigate whether specific regions on 11q13 are associated with 11q13 breakpoints in MM. To this end, we cloned and mapped different 11q13 breakpoints, isolated the corresponding germ-line regions, and evaluated in Southern blot a representative panel of MM primary tumors for genomic rearrangements.

In case LB411, the breakpoint was located between MTC and cyclin D1, approximately 40 kb telomeric of the MTC. In case AC97 carrying a t(11;14), FISH analysis allowed us to map the breakpoint in this region, in which the breakpoints from the MM cell lines XG1 and SK-MM2 have also been found to be located.\cite{9,16} In the U266 cell line, the breakpoint is telomeric of the cyclin D1 locus between the FGF3 and GARP loci; however this rearrangement is complex because the Co1 region is located to the abnormal chromosome 11, and apparently the GARP locus is still retained on this chromosome. As far as we know, this is the second MM tumor in which the 11q13 breakpoint has been mapped telomeric of cyclin D1. Raynaud et al\cite{9} reported a similar finding in the XG2 cell line; however, the cyclin D1 gene was not found to be expressed in this cell line. Given the large distance between FGF3 and GARP, it can be suggested that the breakpoint in U266 may be closer to the FGF3 gene, at a distance that can affect cyclin D1 expression. In the KMS-12 cell line, the breakpoint is centromeric of the MTC.

During the course of this study, Vaandrager et al\cite{133} reported that the breakpoint in this cell line is located 215 kb centromeric of the MTC region; Southern blot analysis showed that the breakpoint in our case LB104 appears to be located in the same region. A breakpoint centromeric of the MTC locus has been previously reported in the XG5 MM cell line.\cite{9} Taken together, these data confirm the notion that 11q13 breakpoints in MM are scattered along the 11q13 (see scheme in Fig 7) and suggest that they may occur either centromeric of the MTC, between the MTC and cyclin D1 loci, or telomeric of the cyclin D1. Interestingly, we found by RT-PCR that the presence of 11q13 breakpoints in tumors LB411 and LB104 correlates with cyclin D1 overexpression (data not shown).

The availability of 11q13 probes specific for breakpoints in MM would make it possible to investigate by Southern blot analysis whether distinct breakpoint clusters may occur in MM. As far as we know, this is the first study regarding the screening of MM tumors using such an approach; however, our analysis with probes from three distinct 11q13 regions representative of MM breakpoints detected a rearrangement in only one case, further confirming the notion that 11q13 rearrangements associated with MM are scattered over a large region encompassing the cyclin D1 gene. These findings suggest that other technical approaches, such as FISH on interphase nuclei,\cite{4} are needed to assess the frequency and involvement of 11q13 breakpoints in MM.

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