Research

Gene amplification as double minutes or homogeneously staining regions in solid tumors: Origin and structure

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Double minutes (dmin) and homogeneously staining regions (hsr) are the cytogenetic hallmarks of genomic amplification in cancer. Different mechanisms have been proposed to explain their genesis. Recently, our group showed that the *MYC*-containing dmin in leukemia cases arise by excision and amplification (episome model). In the present paper we investigated IO cell lines from solid tumors showing *MYCN* amplification as dmin or hsr. Particularly revealing results were provided by the two subclones of the neuroblastoma cell line STA-NB-IO, one showing dmin-only and the second hsr-only amplification. Both subclones showed a deletion, at 2p24.3, whose extension matched the amplicon extension. Additionally, the amplicon structure of the dmin and hsr forms was identical. This strongly argues that the episome model, already demonstrated in leukemias, applies to solid tumors as well, and that dmin and hsr are two faces of the same coin. The organization of the duplicated segments varied from very simple (no apparent changes from the normal sequence) to very complex. *MYCN* was always overexpressed (significantly overexpressed in three cases). The fusion junctions, always mediated by nonhomologous end joining, occasionally juxtaposed truncated genes in the same transcriptional orientation. Fusion transcripts involving *NBAS* (also known as *NAG*), *FAM49A*, *BC035112* (also known as *NCRNA00276*), and *SMC6* genes were indeed detected, although their role in the context of the tumor is not clear.

[Supplemental material is available online at http://www.genome.org. Sequence data from this study have been submitted to GenBank (http://www.ncbi.nlm.nih.gov/genbank/) under accession nos. HM243501–HM243513, HM243515– HM243529, and HM358636. NimbleGen array CGH data have been submitted to the NCBI Gene Expression Omnibus (http://www.ncbi.nlm.nih.gov/geo/) under accession no. GSE22279. Affymetrix CGH data have been submitted to ArrayExpress (http://www.ebi.ac.uk/microarray-as/ae/) under accession no. E-MEXP-2735.]

Overexpression of cellular oncogenes, a common theme in cancer, can result from gene amplification through the formation of extrachromosomal double minutes (dmin) or intrachromosomal homogeneously staining regions (hsr). Their occurrence has been more frequently reported in solid tumors compared with hematological malignancies. Some oncogene amplifications are associated with specific tumors and usually represent an indicator of poor prognosis (Albertson 2006; Myllykangas et al. 2007).

The breakage-fusion-bridge (BFB) cycle is one of the most popular models invoked to explain intrachromosomal amplification in solid tumors (Hellman et al. 2002; Selvarajah et al. 2006; Reshmi et al. 2007; Vukovic et al. 2007). In this model, as originally proposed by McClintock (1951), multiple cycles of BFB lead to the amplification of genes near the breakage. This feature can be regarded as a specific indication of BFB. A translocation-deletionamplification model has been also demonstrated in some tumors

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(Barr et al. 1996; Roijer et al. 2002; Van Roy et al. 2006). In this model, dmin or hsr derive from the breakpoint region of a translocation event. The "deletion-plus-episome" model, also known as the "episome model," postulates that a DNA segment is excised from an otherwise intact chromosome, circularized, and amplified by mutual recombination to produce dmin or hsr (Carroll et al. 1988). We have recently provided definitive evidence in favor of the episome model for the formation of *MYC*-containing dmin in acute myeloid leukemia (Storlazzi et al. 2006).

In this study we investigated the structure of *MYCN* amplifications, examples of both dmin and hsr, in eight neuroblastoma (NB) and two small cell lung carcinoma (SCLC) cell lines.

Results

The 10 cell lines under study are reported in Table 1. Appropriate fluorescence in situ hybridization (FISH) experiments disclosed that in five cell lines *MYCN* was amplified as dmin, and in the remaining five as hsr (Fig. 1; Table 1). The *MYCN* gene was deleted in one chromosome 2 homolog in both subclones of the STA-NB-10 neuroblastoma cell line (Fig. 1). No *MYCN* deletion was detected in the remaining cell lines, nor were rearrangements at the *MYCN* locus noticed.

Table 1. Tumor cell lines investigated in the	e study
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Cell line	Tumor type	Amplification type	
SK-N-BE	NB	hsr	
STA-NB-3	NB	hsr	
STA-NB-4	NB	dmin	
STA-NB-8	NB	dmin	
STA-NB-10/dmin	NB	dmin	
STA-NB-10/hsr	NB	hsr	
STA-NB-13	NB	dmin	
STA-NB-15	NB	hsr	
GLC8	SCLC	dmin	
GLC14	SCLC	hsr	

(NB) Neuroblastoma, (SCLC) small cell lung carcinoma.

Structure of amplicons

MYCN and the partially overlapping *MYCNOS* were the only genes always involved in the amplification by oligo-array comparative genomic hybridization (CGH) analysis (NimbleGen whole-genome, 385 K platform; average oligo distance ~6.3 kb; Supplemental Figs. 1, 2). FISH with fosmid and BAC probes (listed in Supplemental Table 1) showed that a single hsr was present in two cell lines (SK-N-BE and STA-NB-10/hsr), while two distinct hsr were present in STA-NB-3, STA-NB-15, and GCL14 (Fig. 1). Some amplicons were discontinuous, and the amplification levels of some segments were remarkably different. Notably, in STA-NB-8, FISH provided a clear confirmation of the heterogeneity of amplicon composition suggested by the oligo-CGH analysis (example in Supplemental Fig. 3).

The amplicon structure of the two STA-NB-10 subclones was identical in extent (chr2:15,510–18,630 kb; 3.12 Mb in size), and the identical alternation of amplified and nonamplified segments (Fig. 2) strongly suggested that they were the same amplification. The nonamplified segments internal to the amplicon were shown to be single copies (Fig. 2). All clones mapping inside the 3.12-Mb amplicon of STA-NB-10 failed to yield signals on one chromosome 2 of both subclones (fluorescent in situ hybridization [FISH] examples in Fig. 3). They also failed to detect any signal, in STA-NB-10/dmin, on chromosome 5q at the site of hsr insertion in STA-NB-10/hsr (Fig. 1). Probes mapping at the boundary of this amplicon defined the deletion as spanning, approximately, the region chr2:15,483–18,575 kb (see Fig. 3 and legend).

Sequencing of junction regions

Breakpoint regions were refined by RQ-PCR experiments (Supplemental Table 2A) and by Affymetrix 6.0 SNP array (in STA-NB-8, STA-NB-10/hsr, and STA-NB-15; see Supplemental Table 3). The refinements allowed the setup of normal, long-range, or Vectorette PCR experiments aimed at amplifying, cloning, and sequencing the junction regions. When necessary, nested PCR assays were designed to obtain short, ready-to-sequence amplification products. Thirty-two junctions were successfully sequenced (Supplemental Fig. 4; primers in Supplemental Table 2B) and allowed fine characterization of the internal organization of the amplicons (Fig. 4). The amplicon of the STA-NB-8 cell line was composed of two fragments of different origin (chromosomes 2 and 8), arranged "head to tail." The junction sequence resulting from the circularization of the STA-NB-10/dmin amplicon (Junction I in Fig. 5) defined its boundaries at the sequence level (chr2:15,481,998-18,677,524; 3,195,526 bp in size). An identical junction sequence was obtained from the STA-NB-10/hsr subclone. Additionally, all

the amplification products from the two subclones (about 30, many of them spanning junction regions) yielded bands of identical size (data not shown). Junctions III and VI of both subclones were successfully sequenced and, again, they were identical (Fig. 5). Nonamplified segments internal to the amplicon of the STA-NB-10/hsr, already shown to be single-copy (see above), were found to be homozygous for all the analyzed SNPs (Affymetrix 6.0 SNP array; Supplemental Table 4). Supplemental Figure 5 graphically summarizes the different potential steps of gene amplification in the cell lines, constructed according to the collected data. A bioinformatic analysis of all amplicon breakpoints is reported in the Supplemental material. The analysis did not disclose noteworthy results.

Mapping of hsr insertion sites

All hsr were inserted in chromosomes other than chromosome 2, with the exception of STA-NB-3, in which the hsr mapped on

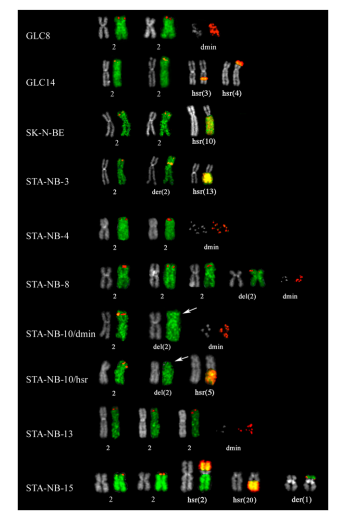


Figure 1. Partial metaphases showing cohybridization FISH experiments using WCP specific for chromosome 2 (green) and RP11-480N14 (chr2:15,999,497–16,004,580), containing the *MYCN* gene (red). GLC8, STA-NB-4, STA-NB-8, STA-NB-10/dmin, and STA-NB-13 showed *MYCN* amplification in form of dmin (example provided at *right*). The remaining cell lines displayed hsr amplification in a single location (SK-N-BE, STA-NB-10/hsr) or in multiple locations (GLC14, STA-NB-3, STA-NB-15). Note the absence of the RP11-480N14 signal on one of the two chromosomes 2 in STA-NB-10 clones (arrows).

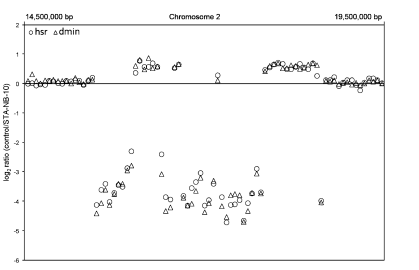


Figure 2. Details of array CGH data (chr2:14,500,000–19,500,000) related to both STA-NB-10/dmin (triangles) and STA-NB-10/hsr (circles) subclones. The pattern of amplified/nonamplified regions perfectly overlaps. Note also that the nonamplified regions, internal to the amplicon, are present in single copy.

chromosome 2, but far away (~61 Mb) from the *MYCN* locus (2p12; *MYCN* is at 2p24.3) (Supplemental Fig. 6). On visual inspection, hsr could be classified as high- (Fig. 6a–c) or low-copy-number (Fig. 6d–f). Occasionally, two different levels of hsr amplification were found in the same cell line, as in STA-NB-3 (Fig. 6a,d). Gajduskova et al. (2007) speculated on the influence of genome position on the level of amplification of the inserted sequences. To investigate this, iterative cohybridization FISH experiments were performed to define the hsr insertion sites (see Supplemental Table 5). The most informative experiments are reported in Figure 6 and in Table 2. We did not detect deletions at the insertion sites, within the limits of FISH resolution.

Fusion genes generated by junctions

Fusion genes are of relevance in cancer (Mitelman et al. 2004). In this study we found several junctions juxtaposing truncated genes in the same transcriptional orientation (see Supplemental material). Reverse transcriptase (RT)-PCR analysis, performed to reveal fusion transcripts, detected two distinct 5'NBAS/3'BC035112 (NBAS also known as NAG; BC035112 also known as NCRNA00276) short chimeric transcripts in cell lines GLC8 and STA-NB-3 (details in Supplemental Fig. 7) that, in silico, did not detect any chimeric NBAS/BC035112 protein (data not shown). In cell line STA-NB-4 we detected another in-frame fusion transcript involving the NBAS gene, composed of the 5' of NBAS, from exons 13-42 (exon 35 is spliced out), and the 3' of FAM49A, exons 3-12 (Supplemental Fig. 6). The transcript was predicted to encode a NBAS/FAM49A chimeric protein of 1511 amino acids, fusing 1185 amino acids of NBAS, containing its Sec39 domain, to 326 amino acids of the FAM49A protein (secretory pathway) (data not shown). In STA-NB-10 subclones we detected a fusion transcript composed of the 5' of FAM49A (exon 1 only) and the 3' of SMC6 (exons 5-13) (accession no. AK292421). This transcript did not encode a chimeric protein, but would encode a 315-amino-acid ORF starting at a codon within exon 5 of SMC6, containing a P-loop NTPase superfamily domain.

Gene expression analysis

Amplicons of the different cell lines showed large size variability, and several genes were coamplified with *MYCN* (Supplemental

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Table 6; Supplemental Fig. 8). We investigated their expression pattern by RQ-PCR (primers reported in Supplemental Table 7). MYCN, amplified in all cell lines, turned out to be always overexpressed, although the increase of transcriptional level was statistically significant only in three NB cell lines (STA-NB-8, STAN-B10, and SK-N-BE). MYCNOS was the only gene always coamplified with MYCN (see Discussion). Expression data of genes with known function are reported in detail in Supplemental Figure 9 and Supplemental Table 8. The overall results indicated that, in addition to MYCN, only GREB1 in STA-NB-8 and RAD51AP2 and SMC6 in STA-NB-10 were significantly overexpressed. Surprisingly, two genes (VSNL1 and GEN1 in STA-NB-10) displayed significant down-regulation despite the gain of copy number. VSNL1 was significantly down-regulated also in

all NB cell lines analyzed (Supplemental Table 8). RQ-PCR experiments were also performed to test the expression of genes interrupted by hsr insertions. The results are described in the Supplemental material and summarized in Supplemental Figure 10 and Supplemental Table 9.

Bioinformatic analysis of hsr insertions

Each hsr insertion site, 400 kb on both sides, was carefully inspected to search for features either shared by all insertion sites or which would allow a differentiation of high- versus low-copy-number hsr (see Supplemental Tables 10–16). The search did not identify any shared features.

Discussion

In the present paper we characterized the structure of *MYCN* amplicons present, in the form of dmin or hsr, in eight NB and two SCLC cell lines. The most interesting result was represented by the perfect matching, in both STA-NB-10 subclones, of the amplicon extension with the extension of the deletion, defined by FISH, found on one chromosome 2. The episomal model has already been hypothesized in solid tumors (Carroll et al. 1988; Corvi et al. 1995). Our finding provided strong evidence of it.

A second revealing result of the present study is the strikingly identical structure of dmin and hsr amplicons of the two STA-NB-10 subclones. This finding suggests that the hsr has evolved from the dmin, as graphically reported in Supplemental Figure 5B. The steps can be summarized as follows: excision, circularization, substantial internal reorganization, amplification, and, in hsr cases, chromosomal integration and further amplification. Theoretically, it can be hypothesized that the excised DNA segment was inserted and amplified as an hsr first. Then, in STA-NB-10/dmin, the hsr extruded to form dmin. The absence, in STA-NB-10/dmin, of any FISH signal at 5q in experiments using probes mapping within the amplicon, and the absence, in STA-NB-10/hsr, of any 5q amplification (CGH array data) imply a perfect and complete extrusion of the hsr to form dmin, which seems unlikely. On the other hand, if hsr represents a potential evolution of dmin, one could expect the simultaneous presence of dmin and hsr in cases in

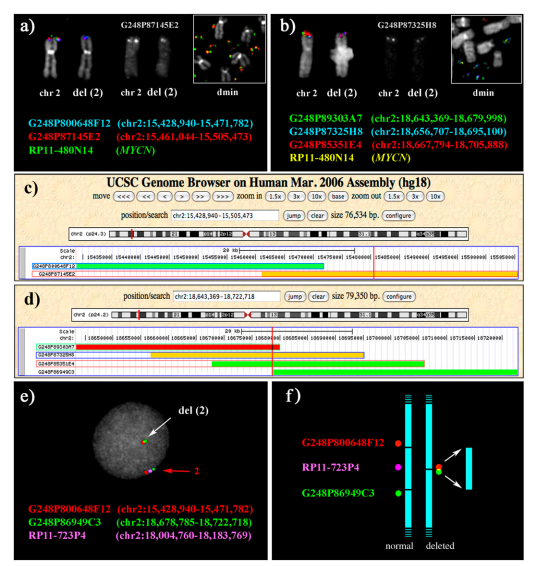


Figure 3. Definition of the boundaries of the deletion on chromosome 2 present in the cell line STA-NB-10/dmin using FISH. (a,b) Partial metaphases showing examples of FISH experiments using fosmid probes spanning the distal (a) and proximal (b) breakpoints of the deleted region, cohybridized with the BAC containing the MYCN gene. (c,d) The position on the human sequence of fosmid clones in a and b, respectively. (Vertical red lines) Amplicon boundaries that were defined by sequencing the amplicon junction (see text and Supplemental Fig. 4). The definition of the deletion breakpoints was obtained as follows. Distal breakpoint. Fosmid G248P87145É2 (chr2:15,461,044-15,505,473; 44.4 kb) gave a signal on del(2) that, by eye, consistently appeared to be half the intensity of the signal on normal chromosome 2 (10 analyzed metaphases; its black and white original signal is separately reported in a). The middle part of the fosmid was therefore assumed to harbor the telomeric break of the deletion at \sim chr2:15,483 kb. G248P87145E2 is present also on the amplicon (a, inset), but the signal intensity is biased by the amplification. The normal intensity on del(2) of fosmid G248P800648F12 (chr2:15,428,940–15,471,782), and the absence of any signal on the dmin, was consistent with the assumption in a. Proximal breakpoint. Fosmid G248P87325H8 (chr2:18,656,707–18,695,100; 38.4 kb) consistently gave a signal of half intensity on del(2) (10 analyzed metaphases; its black and white original signal is separately reported in b). The centromeric break of the deletion was therefore assumed to lay at ~chr2:18,575 kb. Note that sequences of G248P87145E2 are present on the amplicon (b, inset), but, again, the signal intensity is biased by the amplification. The intensity on del(2) of signals yielded by fosmids G248P89303A7, G248P85351E4, and G248P86949C3 was consistent with this hypothesis. The deletion, therefore, spanned approximately the interval chr2:15,483–18,575 kb. Because of growth difficulty of the STA-NB-10/hsr cell line, not all FISH experiments performed on STA-NB-10/dmin were also performed in hsr subclone (see Supplemental Table 1). (e, f) Evidence for the deletion on one of the two copies of chromosome 2. Two fosmid clones flanking the deleted region (red and green) were hybridized with a probe (RP11-723P4, violet) mapping within the deletion but absent in the amplicon (see text). The BAC was chosen in order to avoid the high background due to the amplification. The distance between the red and green probes on normal chromosome 2 (red arrow) is substantially reduced on the del(2) (white arrow), as expected. (f) Rationale of the FISH experiment in e.

which hsr were formed. Some examples of such cases have been reported (Fegan et al. 1995; Yoshimoto et al. 1999; Roijer et al. 2002). It has to be considered, however, that dmin are per se unstable structures. They can be eliminated spontaneously or after addition of certain drugs such as hydroxyurea (Ambros et al. 1997; Narath et al. 2007; Shimizu et al. 2007). Their retention in tumor cells is, very likely, under strong selective pressure, as it has been demonstrated for dihydrofolate reductase dmin originated in response to methotrexate treatment (Martinsson et al. 1982). The selective pressure on dmin, presumably, rapidly drops when an hsr is formed, the latter being intrinsically more stable, and the cell population will soon be composed of a mosaic of dmin- and

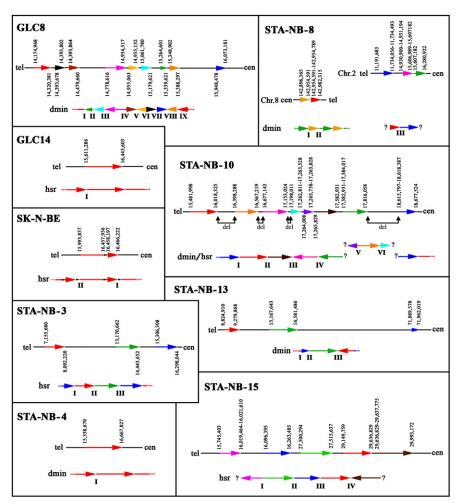


Figure 4. (*Lower* part of each section) Schematic representation of the structure of the amplicon. Different blocks composing the amplicons are represented by arrows of different color. (*Upper* parts) Arrangement of the amplified blocks in the reference sequence (hg18). (Gray line segments) Non-amplified regions; (vertical figures) boundaries of the breaks, at single-base resolution if sequenced (Supplemental Fig. 4), or as a small interval (maximum interval size: 2.5 kb); (?) uncloned fusion regions. In STA-NB-10 the deleted regions (del) internal to the amplified region are indicated. The lengths of the arrows are only approximately proportional the DNA segment length.

hsr-only cells (see Balaban-Malenbaum and Gilbert 1977; Yoshida et al. 1999). The complete loss of dmin can follow. The patient reported by Yoshida et al. (1999) initially showed a hsr/dmin mosaic, then, after therapy, only hsr cells were found. Similar results have been observed also in cultured tumor cells (Quinn et al. 1979; George and Francke 1980; George and Powers 1982). The view that hsr are the result of dmin integration is not new (Coquelle et al. 1998; Bignell et al. 2007; Campbell et al. 2008). Our results on STA-NB-10 subclones add strong support for this view.

STA-NB-10 was the only cell line showing the deletion event. When the excision is postreplicative, the amplified episome, likely conferring a growth advantage, may segregate with either the normal or the deleted chromatid. The deletion, therefore, is not a necessary occurrence. On average, however, it should be found in \sim 50% of postreplicative cases, and up to 100% of non-post-replicative cases. In our recent work on gene amplification in he-matological malignancies, the deletion was detected in 68% (23 out of 34) of the studied cases (Storlazzi et al. 2006). It is unclear why, in the present study, no cell line other than STA-NB-10

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showed the deletion. It is worth noting, however, that, as far as we know, deletions not accompanied by more complex chromosomal events, and corresponding to the amplified sequences, have been reported in solid tumors only by Gibaud et al. (2010).

Amplicon structure

GLC14, SK-N-BE, and STA-NB-4 cell lines showed an amplicon composed of a single fragment; i.e., the excised segment did not undergo internal rearrangement before amplification. The remaining amplicons showed a more complex structure, composed of noncontinuous, non-colinear segments often fused in opposite orientation (Fig. 4). The STA-NB-8 amplicon disclosed coamplification with a sequence from the long arm of chromosome 8 (Fig. 4; Supplemental Fig. 3). Array-CGH analysis disclosed an additional type of complexity. The level of amplification of the different fragments was occasionally substantially different (see STA-NB-8 in Supplemental Fig. 2). Such differences were noted in both dmin and hsr. These findings clearly indicated that these amplicons were heterogeneous and that some events affecting the structure of the amplicon occurred at later stages with respect to the excision event. This heterogeneity has also been reported by Campbell et al. (2008), who assumed that the most frequent rearrangements, estimated by copy-number analysis, occurred early. The FISH experiments on line STA-NB-8, reported in Supplemental Figure 3A, clearly documented a mosaicism. The power of FISH in detecting sequence stretches less than ~ 10 kb in size, however, is relatively low, and we can sup-

pose that the heterogeneity could be higher than suspected. Changes of the amplicon structure could occur at any time during proliferation.

STA-NB-3, STA-NB-13, and STA-NB-15 cell lines showed coamplification of segments located >5 Mb apart from MYCN (up to 55 Mb apart in STA-NB-13) (Fig. 4); in STA-NB-8 the nonsyntenic coamplification involved a ~4-kb DNA stretch located at chr8:142,982,315-142,896,365. These findings suggest that the event that originated the amplicon patchwork was a complex rearrangement in which more than one chromosomal domain participated (Supplemental Fig. 5). The occurrence of coamplification of genes originating from different chromosomes is not a rare event in hematological amplifications (Frascella et al. 2000; Morel et al. 2003; Graux et al. 2004; Martin-Subero et al. 2005; Van Roy et al. 2006). Amplicons with a complex pattern have been reported also in studies utilizing fresh cancer material and analyzed using array-CGH technology (Beheshti et al. 2003; Caren et al. 2008; Fix et al. 2008; Stock et al. 2008). Gibaud et al. (2010) documented, in a glioma case, dmin derived from four different chromosomal domains.

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Proximal chr.2:18,677,505-18,677,54	5 TTCAAGCTGGCTGCGGAAATTTGCATAAGCAATGAGGAGCC
Junction	TTCAAGCTGGCTGCGGAA AT ACAAGAATTCACTTTCATCTT
Distal chr.2:15,481,980-15,482,02	

STA-NB-10/dmin and STA-NB-10/hsr Junction III

STA-NB-10/dmin and STA-NB-10/hsr Junction I

Proximal chr2:17,153,011-17,153,090

STA-NB-10/dmin and STA-NB-10/hsr Junction VI

Proximal chr2:16,567,195-16,567,274

Figure 5. Sequence of Junctions I, III, and VI of STA-NB-10/dmin and STA-NB-10/hsr (see Fig. 4), found identical at the single base-pair level. Note the *AT* microhomology at the junction.

The sequencing of 32 amplicon junctions disclosed 11 microhomologies and 12 anonymous insertions, typical of nonhomologous end joining. Campbell et al. (2008) detected microhomologies in 53% of the joined ends of the acquired rearrangements. Genomic structure, such as segmental duplications, has been shown to affect specific rearrangements in tumors (Barbouti et al. 2004). More specifically, Gibcus et al. (2007) have reported, in their study of 30 patients and eight cell lines showing 11q13 amplification, that amplified regions are flanked by segmental duplications, suggested to play an important role in triggering the initial rearrangement. Many of these amplifications, however, were hypothesized to arise though the BFB mechanism. We did not find any fusion junction joining sequences with high similarity, indicating that, in our cases, mechanisms mimicking nonallelic homologous recombination did not play a role in the genesis of the rearrangements.

The common fragile site FRA2C maps at the band 2p24.2, which encompasses the region chr2:17,000,001–19,100,000, relatively close to the *MYCN* gene (chr2:15,999,497–16,004,580). Some breakpoints (the proximal breakpoint of the STA-NB-10 amplicon, for instance, chr2: 18,677,524) map at 2p24.2. FRA2C, however, was not precisely mapped at sequence level. It is therefore premature to establish a correlation between the FRA2C and excision events.

Recently, the somatically acquired rearrangements of some tumor cell lines showing gene amplification (NCI-H1770 cell line in particular, containing amplified copies of *MYCN*) were investigated by end-sequencing of BAC libraries (Bignell et al. 2007) or by analyzing the end-pairs of short DNA fragments obtained by massive sequencing (Campbell et al. 2008). These studies have the advantage of a precise, genome-wide detection of almost all changes, including those that were acquired in late passages of the cell line, which were probably missed in our study. Very likely, we detected only the more ancestral and more relevant ones, as stressed by Campbell et al. (2008).

Expression of amplified genes, interrupted genes, and fusion genes

MYCN was overexpressed in all cell lines. However the overexpression level reached significance only in three cell lines (Supplemental Table 8A,B). However, there is no doubt that MYCN is the target gene of amplifications of the 2p4.3 domain present in all dmin and hsr amplicons. Amplicon erosion can lead to tumor cell revertance and tumor cell senescence in vitro (Ambros et al. 1997; Narath et al. 2007). The MYCNOS gene partially overlaps with MYCN because it lies on the opposite strand (Armstrong and Krystal 1992). This gene, therefore, was always present in amplicons (Supplemental Table 8A), as reported in similar studies (Scott et al. 2003; Caren et al. 2008; Fix et al. 2008). MYCN and MYCNOS are unique in that they are coregulated in tumor cell lines under basal growth conditions and in response to the differentiating agent retinoic acid (Armstrong and Krystal 1992). We extended the expression analysis to genes whose function was known. Some genes, such as GREB1 in STA-NB-10,

known to be involved in cancer (Sun et al. 2007), or *GEN1* in STA-NB-8, appeared overexpressed although not amplified (Supplemental Table 8A). The present data also indicated that some genes (such as *VSNL1* in NB cell lines, and *PQLC3* in both SCLC and NB cell lines) are consistently down-regulated regardless of their amplification status. *VSNL1* down-regulation has been already reported in squamous carcinoma cells (Gonzalez Guerrico et al. 2005) and in non-small cell lung carcinomas (Fu et al. 2008). Our data suggest that it may have a tumor suppressor role also in NB tumors. The role, in cancer, of *PQLC3*, encoding an integral membrane protein, is presently unknown. Again, its down-regulation in both NB and SCLC tumors may indicate a putative function as a tumor suppressor gene.

Our analysis of the junction points at the sequence level identified four potential fusion genes. For all of them we were able to detect a transcript, and one predicted chimeric protein was identified (NBAS/FAM49A). We did not detect a chimeric protein for the other fusion genes. However, these findings do not exclude a role of a transcript in tumorigenesis (Guastadisegni et al. 2008).

Methods

All sequence data refer to the human genome assembly hg18, March 2006 release.

Cell lines

Cell lines, as an alternative to primary tumors, have been chosen in order to have a higher availability of material to perform the present study. The cell lines investigated are listed in Table 1. SCLC cell lines GLC8 and GLC14 were obtained as previously described (van der Hout et al. 1989) and provided by K. Kok. The NB cell line SK-N-BE (Biedler and Spengler 1976) was provided by G. Della Valle (University of Bologna, Italy). STA-NB cell lines were provided by

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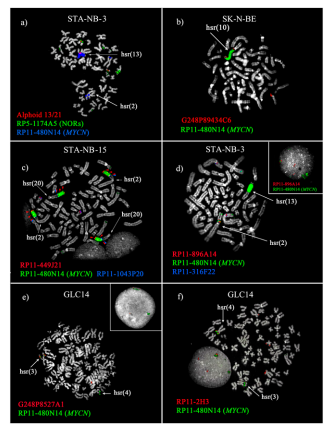


Figure 6. Cohybridization FISH experiments used to characterize hsr insertion sites. *MYCN* was used in all experiments to mark the hsr, and it was cohybridized with contiguous probes flanking the insertion (a, f) or with clones yielding split signals (b-e). (a) In STA-NB-3, the insertion is within the short arm of chromosome 13, between the centromere (alphoid probe, red) and the NOR region (RP5-1174A5). (c) In STA-NB-15, the BAC RP11-449J21 (red) displayed signals distally, proximally, and within the hsr (arrowheads), likely due to a duplication event occurring simultaneously to the hsr insertion process. (e, f) Cell line GLC14 showed two insertion sites. Map positions of all the clones are reported in Supplemental Table 1.

P.F. Ambros, and some of them have been previously described (Ambros et al. 1997; Narath et al. 2007; Stock et al. 2008). STA-NB-10/hsr and STA-NB-10/dmin are subclones of the same cell line, showing MYCN [v-myc myelocytomatosis viral related oncogene, neuroblastoma derived (avian)] amplification in form of hsr or dmin, respectively.

FISH, multicolor-FISH, long-range PCR, Vectorette PCR, sequencing, and RQ-PCR were performed as already described (Storlazzi et al. 2006).

Oligo-array CGH

Whole-genome CGH array analysis was performed to define DNA copy number gains and losses. DNA was extracted from tumor cell lines using standard methods. Tumor DNAs and a NimbleGen human reference sample were labeled and cohybridized on a high-resolution NimbleGen 385 K Whole-Genome Tiling array (build hg18; NCBI Build 36), which has a median probe spacing of 6270 bp. DNA copy number gains and losses were viewed using Roche NimbleGen's SignalMap software. DNA segmentation analysis has been performed using the circular binary segmentation algorithm from Olshen et al. (2004).

Single nucleotide polymorphism (SNP) array CGH

A whole-genome SNP array analysis was performed on STA-NB-8, STA-NB-10/hsr, and STA-NB-15 cell lines using the Affymetrix Genome Wide Human SNP Array 6.0, including over 906,600 single nucleotide polymorphisms (SNPs) and more than 946,000 probes for the detection of copy number variation. Sample preparation, hybridization, and scanning were performed using GeneChip Instrument System hardware according to the manufacturer's specifications (Affymetrix). Quality controls and LOH analysis have been performed using Affymetrix Genotyping Console Software. Copy Number Variant segments have been produced with Partek Genomic Suite Software (Partek Inc.), performing an unpaired analysis using as baseline the 270 HapMap samples and hidden Markov model method.

Gene expression quantification

Gene expression level was determined by RQ-PCR. The cDNA was obtained using the QuantiTect Reverse Transcription Kit (QIAGEN) according to the handbook protocol. The cDNA was amplified using the Applied Biosystems Real-Time PCR System 7300 in the presence of SYBR Green I. The optimization of the RQ-PCR reaction was performed according to the manufacturer's instructions (Applied Biosystems) but scaled down to 10 μ L per reaction. Individual Real-Time PCR was performed in a 96-well plate (Applied Biosystems) containing 1× Platinum SYBR Green qPCR SuperMix-UDG with ROX (Invitrogen), 500 nM ROX Reference (Invitrogen), and 300 nM of sense and antisense primers. All measurements were performed at least in triplicate.

The PCR conditions were as follows: 2 min at 50°C, 10 min at 95°C, followed by 40 cycles of 15 sec at 95°C and 1 min at 60°C for all the primer pairs used. At the end of each reaction, the cycle threshold (Ct) was manually set up at a level that reflects the best kinetic PCR parameters. Primers were designed with an internet-based interface, Primer3 (Rozen and Skaletsky 2000).

The primers were checked for specificity using the BLAT tool in the UCSC Human Genome Browser (http://genome.ucsc.edu/ cgi-bin/hgBlat).

In order to improve the normalization step, 10 housekeeping genes were tested (*ACTB, B2M, GAPDH, HMBS, HPRT1, RPL13A, SDHA, RN28S1, UBC,* and *YWHAZ*) and the three best-performing housekeeping genes (*HPRT1, SDHA,* and *UBC*), selected by applying the geNorm software (Vandesompele et al. 2002), were used. For accurate averaging of the control genes, we used the geometric mean, instead of the arithmetic mean, and the normalized gene of interest (GOI) expression levels were calculated by dividing the raw GOI quantities for each sample by the appropriate geometric mean. We used total brain RNA (Clontech Laboratories, Inc.) and a pool of three normal lung RNA samples as calibrators for NB and SCLC cell lines, respectively.

Finally, statistical significance analysis of data was performed using the Relative expression software tool (REST) (Pfaffl et al. 2002).

Detection of fusion transcripts

RT-PCR experiments were performed in order to detect the occurrence of fusion transcripts, generated by the expression of chimeric genes at amplicon junctions, in cell lines STA-NB-3, STA-NB-4, STA-NB-10, and GLC8. In STA-NB-3 and GLC8, a primer combination of a forward primer specific for exon 41 of *NBAS* with a reverse primer for exon 3 of *BC035112* was used. In STA-NB-4, the same *NBAS* forward primer was placed in combination with a reverse primer for exon 7 of *FAM49A*. In STA-NB-10, a *FAM49A* exon 1 forward primer was used in combination with a *SMC6* exon 7 reverse primer. Fusion products were sequenced and analyzed by

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			11 5			
Cell line	Chromosome	Band	Insertion site interval	Size (bp)	Candidate target gene(s) ^a	
Low-copy-number hsr						
GLC14	Chr 3	3q25.2	chr3:154,013,633– 154,050,461	36,828	P2RY1	
GLC14	Chr 4	4p16.3	chr4:0-354,888	354,888	ZNF genes	
STA-NB-3	Chr 2	2p12	chr2:77,229,736– 77,251,746	22,010	LRRTM4	
High-copy-number hsr						
SKNBE	Chr 10	10p12.31	chr10:21,992,308– 22,029,819	9376	MLLT10	
STA-NB-15	Chr 17	17q25.1	chr17:71,510,913– 71,607,132	96,219	CDK3, EVPL, SRP68, GALR2, ZACN, EXOC7	
STA-NB-3	Chr 13	13p11.2	chr13:8,300,001– 13,500,000	5,200,000	rDNA genes	

Table 2. Overall results of FISH experiments aimed at mapping hsr insertion sites

Insertion site intervals of "low-copy-number" hsr and "high-copy-number" hsr, defined by FISH experiments.

^aGenes located in each interval.

BLAST. In silico translation of the full-length chimeric transcripts was performed using the ORFfinder Tool of the NCBI website (http://blast.ncbi.nlm.nih.gov).

Bioinformatic analysis

The "Self Chain," "Repeat Masker," "RefSeq Genes," "Segmental Duplication," and "Structural Variation" tracks of the UCSC Genome Browser (http://genome.ucsc.edu/cgi-bin/hgTables) were carefully inspected. Additional bioinformatic analysis was locally performed using the Fuzznuc algorithm (http://bioweb2. pasteur.fr/docs/EMBOSS/fuzznuc.html). The 400-kb regions spanning each breakpoint (Supplemental Table 10) were searched for consensus sequences, in order to disclose the frequency of motifs known to be involved in chromosome instability (Supplemental Table 11). Detected motifs are listed in Supplemental Tables 12–16, in BED format (suitable to be charged as UCSC custom Tracks; http://genome.ucsc.edu/goldenPath/help/customTrack. html).

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Gene amplification as double minutes or homogeneously staining regions in solid tumors: Origin and structure

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