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Research Article

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EWS-*erg* and EWS-Flil Fusion Transcripts in Ewing's Sarcoma and Primitive Neuroectodermal Tumors with Variant Translocations

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Abstract

We have determined the frequency of EWS fusion transcripts in a series of primary Ewing's sarcomas and peripheral primitive neuroectodermal tumors and cell lines. Type 1 and 2 EWS-Flil fusions were demonstrated in 8 cell lines and 14 patient samples. Five patients with cytogenetically characterized rearrangements of chromosome 22 that did not involve chromosome 11 were included in these studies. A novel EWS-Flil in-frame isoform fusing EWS to exon 8 of Flil was isolated from a tumor with a variant t(12;22;22)(q14;p1;q12) translocation. Three in-frame isoforms of a novel hybrid transcript derived from the fusion of EWS with the ETS domain of the human *erg* gene were identified in patient samples and a cell line with cytogenetically unidentified or cryptic translocations involving chromosomes 21 and 22. Interphase analysis by fluorescent in situ suppression hybridization using two overlapping *erg* yeast artificial chromosome clones demonstrated disruption of the *erg* gene on chromosome 21 in a patient sample with monosomy 22. Our results provide new information about the involvement of EWS in small round cell tumors involving exchange of its putative RNA-binding domain with DNA-binding domains derived from different members of the ETS family of transcription factors. These studies emphasize the utility of reverse transcriptase PCR analysis and fluorescent in situ hybridization as additional diagnostic tools for differential diagnosis among small round cell tumors. (*J. Clin. Invest.* 1994. 94:489–496.) **Key words:** small round cell tumors • chromosomal translocations • yeast artificial chromosomes • fluorescent in situ hybridization • interphase cytogenetics

Introduction

Ewing's sarcoma (ES)¹ is a highly malignant tumor of bone and soft tissues that most often affects young adolescents (1).

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1. **Abbreviations used in this paper:** AT, Askin's tumor; ES, Ewing's sarcoma; FISH, fluorescent in situ suppression hybridization; PN, peripheral neuroepithelioma; PNET, primitive neuroectodermal tumors; RT-PCR, reverse transcription PCR; SRCT, small round cell tumors; YAC, yeast artificial chromosomes.

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Approximately 83% of all cases are associated with a t(11;22)(q24;q12) reciprocal translocation (2–4). Cytogenetically identical translocations have been described in other small round cell tumors (SRCT) such as primitive neuroectodermal tumor (PNET), peripheral neuroepithelioma (PN), and Askin's tumor (AT) (5). The recent cloning of the EWS and Flil genes disrupted by the t(11;22)(q24;q12) reciprocal translocation characteristic of these SRCT has provided new insight into possible mechanisms of tumorigenesis in solid tumors (6). Further, the cloning of hybrid fusion transcripts between EWS and the ATF-1 transcription factor gene in malignant melanoma of soft parts with a balanced t(12;22)(q13;q12) translocation (7), the fusion of TLS (translocated in liposarcoma) gene with CHOP gene (member of the c/EBP family of transcriptional activators) as a result of the t(12;16)(q13;p11) translocation in myxoid liposarcoma (8), and the fusion of PAX3 gene to a member of the forkhead gene family in alveolar rhabdomyosarcomas with a t(2;13)(q35;q14) translocation (9, 10) suggest a common mechanism of activation of transcription factors that is responsible for development of different tumor types.

Human Flil (11) is a member of the ETS family of transcription factors that exhibit homology within a conserved DNA-binding domain, termed the 3'-ETS domain (12). EWS (6) is a gene of unknown function with extensive sequence similarity to TLS (8). Both belong to a new subclass of RNA-binding proteins that contain a conserved 80-amino acid domain in their carboxyl terminus (13). The ES- and PNET-associated t(11;22)(q24;q12) translocation results in a hybrid mRNA transcribed from the derivative chromosome 22, in which the 3' DNA-binding domain of Flil replaces the 3' RNA-binding domain of EWS (6). Hybrid EWS-Flil constructs induce malignant transformation when introduced into murine fibroblasts (14). EWS-Flil fusion genes are heterogeneous since three different breakpoints in Flil and two breakpoints in EWS have been described so far, resulting in four unique fusion sequences (6, 14). The incidence of the different fusion transcripts in a large number of patients has not yet been determined. Variant and complex translocations have been reported in 9% of all ES and PNET cases (4, 15). In most instances they involve chromosome 22q12, suggesting that the fusion gene on the derivative chromosome 22 is related to tumorigenesis. Molecular studies published to date demonstrated that all ES and PNET with typical t(11;22)(q24;q12) translocations and one ES cell line with a complex t(11;22;14) translocation contained EWS-Flil fusion transcripts (6, 14, 16). Rearrangements of the EWS and Flil genes were also found in two tumors with variant t(7;22) and t(14;22) translocations and in one tumor with a normal karyotype (17). These findings are similar to the cases of chronic myeloid leukemia in which cytogenetic analysis does not demonstrate the typical t(9;22)(q34;q11) translocation yet contains bcr-abl chimeric transcripts (18).

Table I. Pathological and Cytogenetic Characteristics of the Cell Lines and Tumors

	Age/sex	Diagnosis	Partial karyotype	Fusion
Cell line (reference)				
TC-71 (5)	22/M	ES of bone	t(11;22)(q24;q12)	Type 1
6647 (5)	14/F	ES of bone	t(11;22)(q24;q12),22q+	Type 2
SK-ES-1	18/M	ES of bone	t(11;22)(q24;q12)	Type 2
RD-ES	19/M	ES of bone	22q-	Type 2
TC-32 (5)	17/F	PN	t(11;22)(q24;q12)	Type 2
LAP-35 (23)	12/F	PNET of bone	t(11;22)(q24;q12)	Type 2
SK-PN-LI (24)	3/M	PNET of bone	t(21;22)(q22;q12)	Type 9e
SK-PN-DW (24)	17/M	PNET of soft parts	-11; t(11;22)(q24;q12)	Type 1
SK-N-MC (5)	14/F	AT of chest wall	t(2;11;22;21)(q32;q24;q12;p11)	Type 1
U2OS (25)	15/F	Osteosarcoma	Complex abnormalities	None
SK-N-SH	4/F	Neuroblastoma	der(22)t(22;?)(q13;?)	None
Tumor				
T89-135	17/M	SRCT consistent with ES	-22	Type 1e
T91-124	11/M	PNET	der(21)t(21;?)(q22;?) ?der(22)t(22;?)(q12;2)	Type 1e
T92-60	11/F	ES of bone	der(22)t(22;?)(q12;?)	Type 3e
T93-98	12/F	SRCT consistent with alveolar rhabdomyosarcoma	-22 der(16)t(16;22)(q?23;q11-12) der(22)t(22;?)(q11-12;?)	None
T93-101	14/F	ES	t(11;22)(q24;q12)	Type 2
T93-113	13/M	SRCT consistent with PNET	t(12;22;22)(q14;p1;q12)	Type 8

Variants of specific chromosomal translocations in multiple leukemia subtypes have been reported. The cloning of some of these rare variant translocations has led to the discovery of new genes that share sequence homology and/or common motifs with the gene disrupted by the "typical" translocation. A notable example is the group of translocations that fuse the *MLL* gene on 11q23 (19–21) to genes from a variety of different chromosomes (22). These genes share extensive sequence homology and presumably confer similar biological activities to the chimeric protein. Cloning of the genes involved in variant translocations in solid tumors should also result in an increased understanding of the mechanisms involved in the development of these malignancies.

In this study, we have investigated the frequency and specificity of EWS alterations in a series of cell lines and tumors obtained from patients with SRCT. Cell lines and tumor specimens with typical t(11;22) translocations and with rearrangements of chromosome 22 in the presence of normal chromosomes 11 were characterized cytogenetically and studied with molecular techniques. Novel transcripts involving EWS and the ETS family-related *erg* or *Fli1* genes were observed, expanding the repertoire of possible gene rearrangements involved in the pathogenesis of ES and PNET.

Methods

Cell lines and patient samples. 11 tumor cell lines were analyzed in this study. Their clinical features at diagnosis and partial karyotypes are listed in Table I (23–25). Cell lines RD-ES, SK-ES, SK-N-MC, SK-N-SH, and U2OS were obtained from the American Type Culture Collection (Rockville, MD). Six SRCT cases were cytogenetically characterized at The Children's Hospital of Philadelphia, according to standard methods (26). Cytogenetic abnormalities of the relevant chromosomes are listed in Table I. 17 patient samples of cytologically, histologically, and immunohistochemically confirmed SRCTs were not characterized

cytogenetically and were obtained from the Istituti Ortopedici Rizzoli, Bologna, Italy. Surgical tumor samples were cryopreserved before their use in these studies. Informed consent was obtained according to institutional guidelines.

Reverse transcription PCR (RT-PCR) analysis of EWS-*Fli1* and EWS-*erg* fusions. Total RNA was isolated from frozen tumor fragments and from cell lines in culture as described (27). 1 µg of total RNA was reverse transcribed with an oligo-d(T)₁₆ primer using the GeneAmp RNA-PCR kit (Perkin-Elmer Corp., Norwalk, CT). PCR amplification was performed with 2.5 U of Taq DNA polymerase, 2 mM MgCl₂, 1× PCR buffer II (Perkin-Elmer Corp.), 200 µM deoxynucleotide triphosphate, and 0.15 µM each 5' and 3' primers. Oligonucleotide 22.1 (7) or 22.3 (6) were used as 5' (forward) EWS primers. Primer 11.3 (6) was the reverse primer on exon 9 of the human *Fli1* gene (Zucman, J., T. Melot, B. Plougastel, L. Selleri, M. Giovannini, G. A. Evans, O. Delattre, and G. Thomas, manuscript submitted for publication). The reverse primer for the human *erg* gene (28) 21.9, 5'-ATGAGAAGG-CATATGGCTGG-3', was used to specifically amplify *erg* moieties. The amplification profile consisted of denaturation at 94°C for 30 s, primer annealing at 65°C for 60 s, and extension at 72°C for 2 min, for a total of 30 cycles on an automated heat block (GeneAmp PCR system 9600; Perkin Elmer-Cetus Instruments, Norwalk, CT). Positive and negative controls were carried through all steps. As a positive control for the integrity of RNA isolated from each tumor and cell line, PCR amplification of the β -actin gene was performed using forward primer β A1 (nucleotides 2563–2582 of GenBank M10277) in combination with reverse primer β A2 (nucleotides 2970–2989 of GenBank M10277), which produce a 314-bp product.

Direct sequencing of PCR products. PCR products were purified and concentrated on a Microcon 100 (Amicon, Beverly, MA). Primer 22.3 (6) was used for sequencing of EWS-*Fli1* and EWS-*erg* PCR products using a Taq DyeDeoxy terminator cycle sequencing kit (Applied Biosystems, Inc., Foster City, CA). Reactions were run and analyzed on a fluorescent DNA sequencer (373A; Applied Biosystems, Inc.).

Multicolor fluorescent in situ hybridization with *erg* yeast artificial chromosomes (YACs). Metaphase spreads and interphase nuclei were

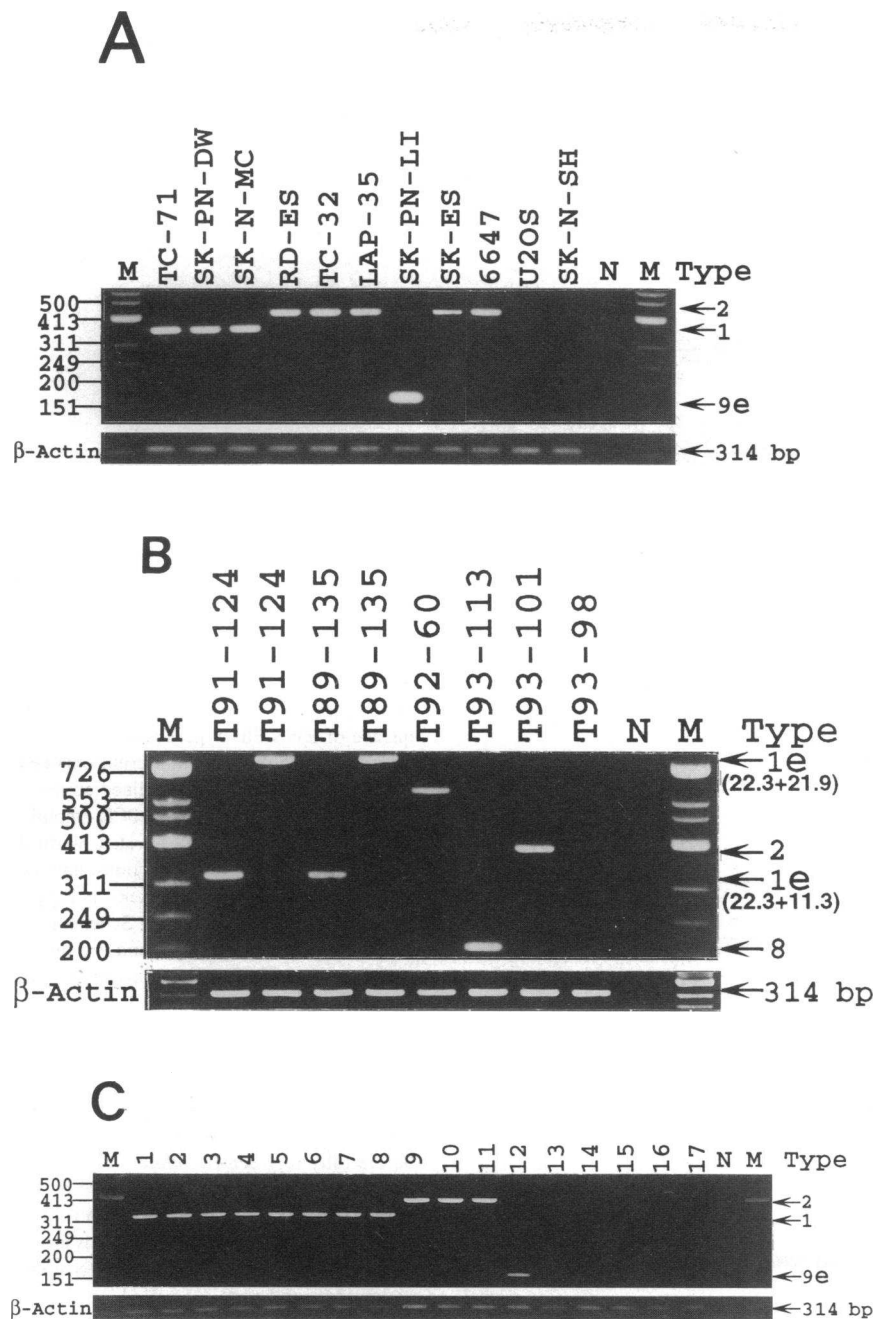


Figure 1. RT-PCR analysis of EWS-Flil and EWS-erg chimeric transcript expression in SRCT. (A) Cell lines. PCR analysis was performed on oligo-dT reverse-transcribed total RNA samples (1 µg) with primers 22.1 (EWS exon 7) and 11.3 (Flil exon 9). Amplification products were separated by electrophoresis on 2% agarose gels and visualized by ethidium bromide staining. RNAs derived from t(11;22)-negative U2OS and SK-N-SH were used as negative controls. N, no template; M, molecular weight standards (ϕX174/HinfIII). The fusion types are indicated on the right side, and their sizes with primers 22.1 and 11.3 or 21.3 are: type 1, 355 bp; type 2, 421 bp; and type 9e, 181 bp. The 314-bp fragment shown below is amplified using b-actin primers for quality control of the RNAs. (B) Cytogenetically characterized patient samples. Samples were amplified using primers 22.3 and 11.3. EWS-erg type 1 fusions isolated from patients T91-124 and T89-135 (lanes 3 and 5) were amplified using primer 21.9 that is specific for the erg gene and does not amplify EWS-Flil fusions. The sizes of the fusion types indicated on the right are: type 1e, 328 bp (primers 22.3 and 11.3) and 735 bp (primers 22.3 and 21.9); type 2, 394 bp (primers 22.3 and 11.3); type 3e, 580 bp (primers 22.3 and 11.3); and type 8, 202 bp (primers 22.3 and 11.3). N, no template; M, molecular weight standards (ϕX174/HinfIII). (C) Cytogenetically uncharacterized samples. PCR analysis was performed on 17 cytologically, histologically, and immunohistochemically confirmed SRCTs. Included were eight ESs (lanes 1–5 and 10–12), three metastatic ESs (lanes 6–8), one PNET (lane 9), two small cell osteosarcomas (lanes 13 and 14), and three non-Hodgkin lymphomas of bone (lanes 15–17). Samples were amplified using primers 22.3 and 11.3, and the sizes of the fusion types are indicated in B. N, no template; M, molecular weight standards (ϕX174/HinfIII).

prepared as described previously (25). Overlapping YAC clones A125B12 and B19C12 from the Washington University human genomic YAC library were shown previously to be colinear and to contain the human erg gene (29). To confirm that the YAC strains A125B12 and B19C12 contained the desired human insert, PCR analysis with two sets of primers corresponding to the 5' and 3' regions of the human erg gene was performed. Primers for the erg 5' region were erg5S (5'-CCTCTCGGTTATTCCAGGATC-3') (forward) and erg5A (5'-GTCCGGGACAGTCTGAATCAT-3') (reverse). Primers for the erg 3' region were erg3S (5'-TTCAAGATGACGGATCCCGAC-3') (forward) and primer 21.9 (reverse). Fluorescent in situ suppression hybridization (FISSH) analysis using YAC clones A125B12 and B19C12 was carried out as reported (30). For two-color FISH, YAC A125B12 DNA was labeled with biotin-11-dUTP (ENZO Diagnostics, Syosset, NY), and YAC B19C12 DNA was labeled with digoxigenin-11-dUTP (Boehringer Mannheim Corp., Indianapolis, IN) by random priming. Hybridization and posthybridization washes of the slides were carried

out as described (25). After posthybridization washes, slides hybridized only to biotin-labeled probes were incubated in avidin-fluorescein as described (25). Slides hybridized to both biotin- and digoxigenin-labeled probes were amplified and detected as described (31).

Results

Identification of distinct EWS-Flil and EWS-erg fusion isoforms in ES and PNET without t(11;22)(q24;q12) translocation. To define the chromosomal translocations and/or gene fusions in cases with variant rearrangements, we performed molecular analysis on a variety of cell lines and tumor material. RD-ES and SK-PN-LI cell lines (Table I) were analyzed by RT-PCR. The PCR amplification product generated using reverse-transcribed RNA from cell line RD-ES as a template and primers

A

**EWS-Flil1
Type 8**

P T S Y P P O T G S Y S Q A P S Q Y S Q
CCCCTAGTTACCCACCCCAACTGGATCCTACAGCCAAGCTCCAAGTCAATATAGCCAA
 22.1
 Q S S S Y G Q Q N P Y Q I L G P T S S R
CAGAGCAGCAGCTACGGGCAAGAGATTCGGTATCAGATCCTGGGCCCGACCAGCAGTCCG
 L A N P G S G Q I Q L W Q F L L E L L S
 CTAGCCAAACCTGGAAGCGGGCAGATCCAGCTGTGGCAATTCTCTGGAGCTGCTCTCC
 D S A N A S C I T W E G T N G E
 GACAGCGCCAAACGCCAGCTGTATCACCTGGAGGGGACCAACGGGGAGT
 11.3

**EWS-erg
Type 1e
& 9e**

P T S Y P P O T G S Y S Q A P S Q Y S Q
CCCCTAGTTACCCACCCCAACTGGATCCTACAGCCAAGCTCCAAGTCAATATAGCCAA
 22.1
 Q S S S Y G Q Q N P Y Q I L G P T S S R
CAGAGCAGCAGCTACGGGCAAGAGATTTACCATATGAGCCCCCAGGAGATCAGCCTGG
 (Type 1e) 22.3
 T G H G H P T P Q S K A A Q P S P S T V
 ACCGGTCACGGCCACCCACGCCCGAGTCGAAAGCTGCTCAACCATCTCTTCCACAGTG
 P K T E D Q R P Q L D P Y Q I L G P T S
 CCCAAAATGAAGACCGCTCCTCAGTTAGATCCTTATCAGATTCCTTGGACCAACAAGT
 S R L A N P G S G Q I Q L W Q F L L E L
 AGCCGCTTGCATAATCCAGGCAGTGGCCAGATCCAGCTTTGGCAGTTCCTCTGGAGCTC
 (Type 9e)
 L S D S S N S S C I T W E G T N G E F K
 CTGTGCGCAGCTCCAACTCCAGCTGCATCACCTGGAAAGGCCAACCGGGAGTTCAGG
 M T D P D E V A R R W G E R K S K P N M
 ATGACGGATCCCGACGAGGTGGCCGGCGCTGGGGAGAGCGGAAGAGCAACCCCAACATG
 N Y D K L S R A L R Y Y Y D K N I M T K
 AACTACGATAAGCTCAGCCGCGCCCTCCGTTACTACTATGACAGAAGACATCATGACCAAG
 V H G K R Y A Y K F D F H G I A Q A L Q
 GTCCATGGGAAGCGCTACGCCCTACAAGTTTCGACTTCACCGGATCGCCAGGCGCTCCAG
 P H P P E S S L Y K Y P S D L P Y M G S
 CCCACCCCCCGGAGTCATCTCTGTACAAGTACCCCTCAGACCTCCCGTACATGGGCTCC
 Y H A H P Q K M N F V A P H P P A L P V
 TATCACGCCCAACCCACAGAAGATGAACCTTTGTGGCGCCCAACCTCCAGCCCTCCCGCTG
 T S S S F F A A P N P Y W N S P T G G I
 ACATCTCCAGTTTCTTGTGCTGCCCAAAACCCATACTGGAATTACCAACTGGGGGTATA
 Y P N T R L P T S H M P S H
 TACCCCAACACTAGGCTCCCC**ACAGCCATATGCCTTCTCAT**
 21.9

**EWS-erg
Type 3e**

M D E G P D L D L D L P Y E P P R R
 ...ATGGATGAAGGACCGATCTTGATCTAGTTTACCATATGAGCCCCCAGGAGA...

Figure 2. Molecular analysis of EWS-*erg* and EWS-Flil1 chimeric mRNAs. (A) Nucleotide sequence and predicted amino acid sequence of EWS-Flil1 type 8 and EWS-*erg* type 1e, 3e, and 9e fusion transcripts obtained by RT-PCR. Sequences underlined correspond to the primers used for PCR amplification and their names are indicated. Vertical bars indicate the different junctions, and the contribution of the EWS sequence is highlighted in bold characters. (B) Schematic representation of EWS, Flil1, and *erg* functional domains and their locations on the non-rearranged proteins and the fusion proteins translated from the cDNAs represented in (A). The RNA-binding domain of EWS (RNA BD) is replaced by the ETS domains (ETS D) of Flil1 or *erg*, depending on the fusion type. Fusion proteins type 8 and type 9e are fully represented, while type 1e and type 3e can be deduced from type 9e by the insertion of 59 additional *erg* amino acids. In type 3e, 84 additional amino acids are derived from the EWS sequence. Interrupted codons are indicated.

22.1 and 11.3 was 421 bp in size (Fig. 1 A). Nucleotide sequence analysis of the hybrid transcript revealed in-frame junctions between EWS codon 265 and Flil1 codon 197, which is identical to type 2 fusion described previously (6). Amplification of SK-PN-LI cDNAs resulted in a product of 181 bp derived from the fusion of EWS with the human *erg* gene, located on chromosome 21 (Fig. 1 A). Although not reported previously (24), a t(21;22)(q22;q12) translocation was observed in karyotypes prepared from the SK-PN-LI cell line. Basic local alignment search tool (32) analysis (BLAST) of the GenBank database with primer 11.3 nucleotide sequence revealed 90% homology with the region extending from nucleotide 842 to 861 of the published *erg* sequence (28). Therefore, primer 11.3 can amplify both EWS-Flil1 and EWS-*erg* fusion transcripts. Nucleotide sequence analysis of the fragment revealed an in-frame fusion occurring between the first base (A) of codon 265

of EWS (Ser) and the G in the second position of codon 192 of *erg* (Gly), creating a new AGC codon for a serine residue (Fig. 2, A and B). This region of the *erg* gene shows 98% homology with the exon 9 region of Flil1, and we will therefore refer to this fusion as type 9e. Six different *erg* isoforms have been described, generated by alternative splicing events (28, 33, 34). The *erg* moieties cloned in this study are in a region common to all the described *erg* isoforms. The *erg*-1 cDNA sequence was therefore used for nucleotide and codon numeration (GenBank database accession number M21535). Oligo-dT-primed cDNAs from five SRCT patients that cytogenetically do not contain the typical t(11;22)(q24;q12) translocation (Table I) were amplified using the combination of primers 22.3 and 11.3 (Fig. 1 B). Tumors from patients T91-124 and T89-135 showed a 328-bp amplification product consistent with an EWS-Flil1 type 1 fusion. Surprisingly, however, sequence

B

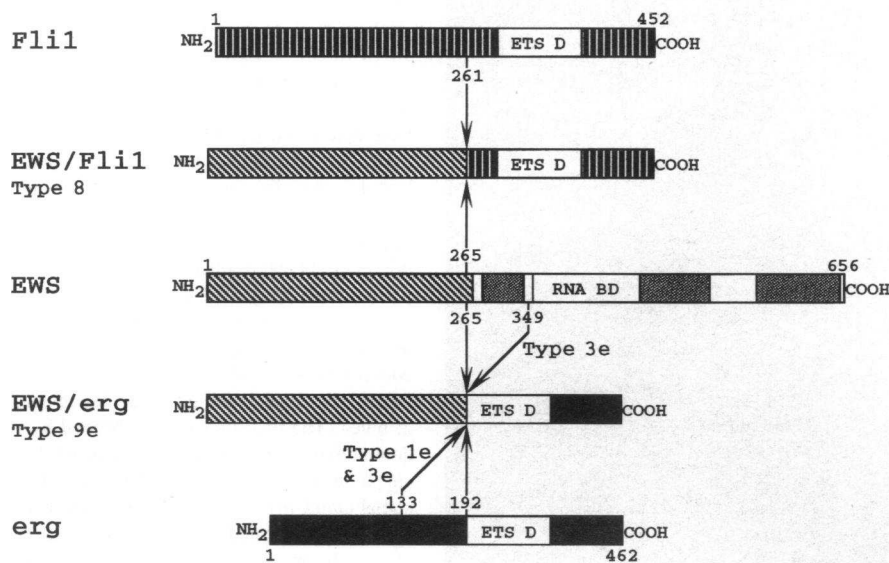


Figure 2. Continued

analysis revealed a novel in-frame EWS-*erg* fusion occurring between the first base (A) of codon 265 of EWS (Ser) and the A in the second position of codon 133 of *erg* (Asp), creating a new AAT codon for an asparagine residue (Fig. 2, A and B). Because of this similarity in size with EWS-Fli1 fusion type 1, we define this fusion as type 1e. To improve the quality and specificity of the PCR amplification of EWS-*erg* fusions, a new reverse primer, 21.9, was designed. Primer 21.9 is in the region immediately downstream of the *erg* ETS domain (nucleotides 1249–1268 of GenBank M21535) that does not show homology with Fli1. As expected, amplification of T91-124 and T89-135 cDNAs using primers 22.3 and 21.9 resulted in single amplification products of 735 bp (Fig. 1 B, lanes 3 and 5). The same combination of primers failed to detect the EWS-Fli1 transcripts present in SK-PN-DW and RD-ES cell lines (data not shown), confirming the predicted specificity of primer 21.9 for the human *erg* gene. Reexamination of karyotypes from case T91-124 demonstrated a translocation involving 21q22 and a possible translocation involving 22q12. Case T89-135 was missing one chromosome 22, and no obvious rearrangements of chromosome 21 were seen. To confirm that the rearranged *erg* in fact resulted in disruption of the gene on chromosome 21, we performed interphase FISH analysis with two overlapping YAC clones encompassing the *erg* gene (29). PCR analysis demonstrated that YAC clone A125B12 contains the 3' region of *erg* and not the 5' and, conversely, YAC B19C12 contains the 5' and not the 3' region (data not shown, see Methods for details on primer sequences). Together these YACs span ~430 kb of genomic DNA on human chromosome 21q22.2 (29). After FISH, 46% of the nuclei from tumor sample T89-135 contained three hybridization signals (Fig. 3), one yellow signal due to the overlap of the two YACs on the normal chromosome 21, one green signal corresponding to YAC B19C12 (5' *erg* region), and one red signal corresponding to the translocated 3' *erg* region contained in YAC A125B12. The remaining nuclei displayed two yellow hybridization signals indicating two normal *erg* genes and probably represent normal diploid stromal

cells infiltrating the tumor biopsy. A third, distinct type of EWS-*erg* fusion was found in tumor T92-60. This tumor demonstrated what appeared to be the derivative 22 from a t(11;22)(q24;q12) translocation. A single fragment of 580 bp was detected after 30 cycles of amplification with primers 22.3 and 11.3 (Fig. 1 B). Sequence analysis of the region encompassing the fusion breakpoint revealed an in-frame junction between the first base (G) of codon 349 of EWS (Gly) and the A in the second position of codon 133 of *erg* (Asp), creating a new GAT codon for an asparagine residue (Fig. 2, A and B). EWS codon 349 was reported to be interrupted by fusion with Fli1 codon 219. This was classified as type 3 (6). Because of the striking similarities of the two fusion mechanisms, we refer to this new EWS-*erg* fusion as type 3e. An EWS-Fli1 fusion isoform was isolated in tumor T93-113. Cytogenetic studies from this tumor demonstrated a variant t(12;22;22) translocation in the presence of two apparently normal chromosomes 11. A representative karyotype is shown in Fig. 4. One der(22) is similar to the der(22) seen as a result of the typical t(11;22)(q24;q12) translocation. RT-PCR amplification with primers 22.3 and 11.3 (Fig. 1 B) produced a 202-bp fragment resulting from an in-frame junction between the A in position one of EWS codon 265 (Ser) and the A in position 2 of Fli1 codon 261 (D), the first codon of Fli1 exon 8 (Fig. 2, A and B). A cryptic translocation involving chromosomes 11 and 22 was thus present in this tumor. We termed this EWS-Fli1 fusion as type 8. Despite amplification of the β -actin gene (Fig. 1 B), tumor T93-98 did not demonstrate EWS-Fli1 or EWS-*erg* fusions by RT-PCR analysis. The diagnosis for this patient was undifferentiated sarcoma consistent with alveolar rhabdomyosarcoma. Cytogenetic studies demonstrated translocations involving both chromosomes 22, although neither resembled the der(22) of a t(11;22)(q24;q12) translocation. The presumed 22q11-12 breakpoints are proximal to EWS and suggest the presence of another gene on chromosome 22 involved in the development of solid tumors.

Incidence of the various fusion transcripts in ES/PNET

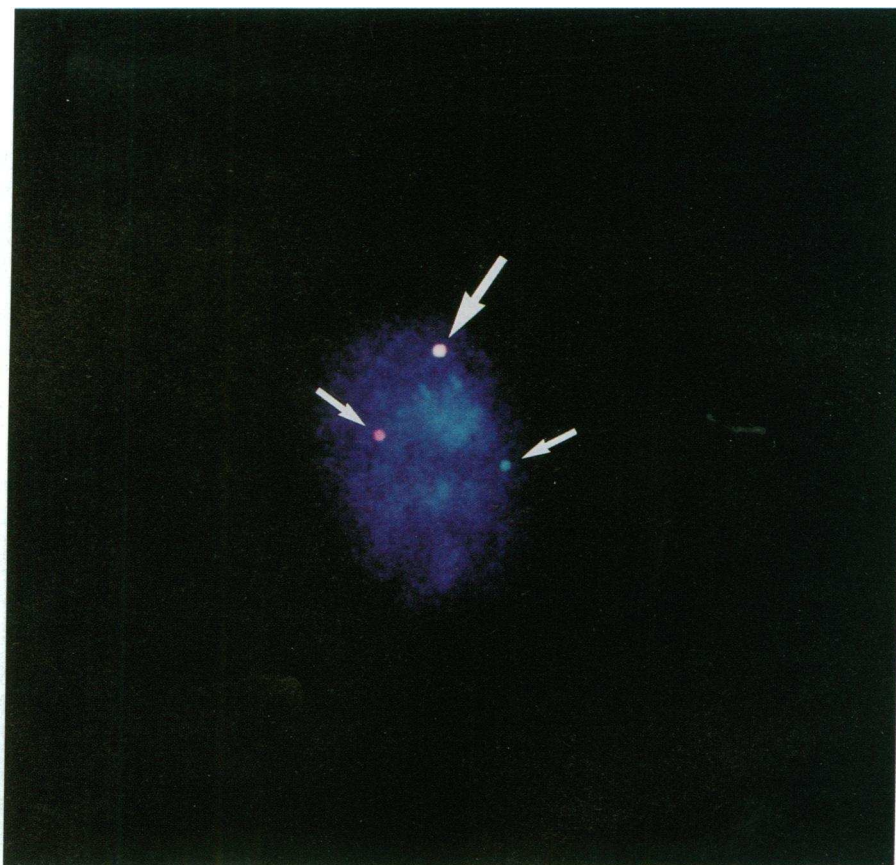


Figure 3. FISH with *erg* YAC clones. Interphase nucleus from ES tumor T89-135 after two-color FISH with YACs A125B12 and B19C12, biotin- and digoxigenin-labeled, respectively. Hybridization signals were detected using Texas red avidin and fluorescein-anti-digoxigenin. Data were collected, and images were reconstructed using a confocal microscope (MRC-600; Bio-Rad, Cambridge, MA). Separate images of each fluorochrome were collected and digitally superimposed (image one: red and green). Delineation of the nucleus was obtained by blue pseudocoloring of a third image collected with the 610-nm filter to visualize the nucleus stained by diluted propidium iodide (image 2: blue). The final image was obtained by double-exposure photography of images 1 and 2 on the computer screen. The yellow signal (*thick arrow*) corresponds to the normal, nontranslocated *erg* allele and is generated by the proximity of green/red hybridization signals of the two overlapping YAC clones. Disruption of the *erg* gene with the translocation is visualized by separation of the red (*left*) and green (*right*) signals (*thin arrows*) corresponding to YAC A125B12 (*erg* 3' region: ETS domain) and B19C12 (*erg* 5' region), respectively.

carrying the typical $t(11;22)$ translocation and in cytogenetically uncharacterized tumors. To study the incidence of the different fusion transcripts and their occurrence in tumors with different phenotypes and cytogenetic abnormalities, we extended our RT-PCR analysis to 6 cell lines and 1 patient sample (T93-101) with typical $t(11;22)(q24;q12)$ translocations, 1 cell line with a complex $t(2;11;22;21)$ translocation, and 17

tumor samples with cytologically, histologically, and immunohistochemically confirmed SRCT for which cytogenetic data were not available. An EWS-Flil type 1 chimeric transcript was found in eight unkaryotyped Ewing's tumors (Fig. 1 C) and in three cell lines (Fig. 1 A). Of the cell lines, one was derived from ES (TC-71), one from a PNET (SK-PN-DW), and one from an AT (SK-N-MC) carrying a complex $t(2;11;22;21)$

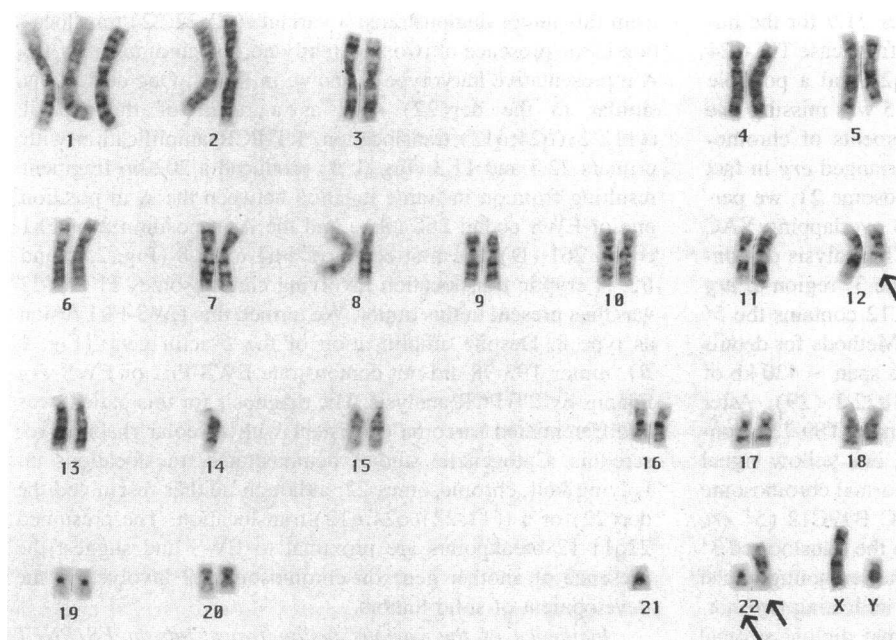


Figure 4. Representative karyotype from case T93-113. Arrows point to the der(12) and the two derivative chromosomes 22. The chromosome 22 on the left is similar to the der(22) seen in a typical $t(11;22)(q24;q12)$ translocation. The der(22) on the right contains the long arm of chromosome 12 ($q14 \rightarrow qter$) translocated to 22p. The der(12) contains the long arm of chromosome 22 ($q12 \rightarrow qter$). The chromosomes 11 appear to be normal. Loss of a chromosome 14 and 21 are random, and there is also an $i(1q)$.

translocation. Karyotypes prepared from the SK-PN-DW cell line demonstrated a typical t(11;22)(q24;q12) translocation, as well as the monosomy 11 which had been reported previously (24). An EWS-Flil Type 2 transcript was found in 3 of the 17 tumors (Fig. 1 C): 2 Ewing's Sarcomas and 1 PNET of the bone, in 4 cell lines (Fig. 1 A): 1 was derived from a PN (TC32), 1 from a PNET of bone (LAP-35), 2 from ES (SK-ES; 6647), and in ES patient T93-101 (Fig. 1 B). 1 additional ES among the 17 SRCT demonstrated a type 9e EWS-*erg* fusion (Fig. 1 C). Our RT-PCR assay did not reveal the presence of the type 4 fusion in PN cell line TC-32, which had been described previously (14). Despite amplification of the β -actin gene (Fig. 1 C). RNA from five unkaryotyped tumor samples did not result in amplification of fusion transcripts. On the basis of cytologic, histologic, and immunohistochemical analyses, three tumors had been diagnosed as non-Hodgkin's lymphoma of bone and two as small cell osteogenic sarcoma.

Discussion

Molecular studies of primary tumor specimens from ES and PNET patients as well as established cell lines demonstrate that the majority of tumors contain a typical t(11;22)(q24;q12) translocation, resulting in a type 1 or type 2 EWS-Flil fusion transcript (6, 17). At present, there does not appear to be any correlation with tumor phenotype, and the relationship of the molecular findings to clinical patient outcome is unknown. In this report, we describe five tumor specimens which were selected for study based on the cytogenetic finding of a rearranged or missing chromosome 22, in the presence of normal chromosomes 11. In one case with a variant t(12;22;22)(q14;p1;q12) translocation, we identified a novel EWS-Flil fusion (type 8). This tumor therefore had a cryptic or masked translocation involving chromosomes 11 and 22 and is similar to what has been seen in chronic myeloid leukemia (35). Three of these tumors, as well as one cell line, and an additional SRCT in which cytogenetic analysis was not performed do not have an EWS-Flil fusion, but contain one of three EWS-*erg* fusion transcripts. In each case, the NH₂-terminal domain of EWS is fused to the ETS DNA-binding domain of the *erg* gene. ERG is a member of the ETS family of genes which share highest homology with Flil (11, 12) and was found to be rearranged in human myeloid leukemia with t(16;21) translocation (36).

In recent studies (Zucman, J., T. Melot, B. Plougastel, L. Salleri, M. Giovannini, G. A. Evans, O. Delattre, and G. Thomas, manuscript submitted for publication), we determined the genomic organization and exon structure of the Flil gene and showed that it spans 120 kb of genomic DNA and is encoded by nine exons. Flil exon 9 encodes the ETS domain and begins with a conserved glycine codon that is split by the 8th intron/9th exon junction between nucleotides in position one and two (G/GA). Although the genomic structure and exon organization of the human *erg* gene have not been determined yet, the striking amino acid sequence identities and the high degree of conservation of intron/exon boundaries found for the other members of ETS family for which the genomic structure is known (Flil, ETS1 and ETS2) imply that *erg* has an exon structure similar to that of Flil. Furthermore, analysis of the EWS-*erg* fusions we have isolated indicates two potential exon/intron boundaries. Flil codon 219 (asparagine), which is split in EWS-Flil type 1 and 3 fusions (6), is interrupted by an intron/exon junction. In the EWS-*erg* fusion types 1e and 3e, we found *erg* codon 133 (asparagine) to be split between the G

in the first position and the A in the second position, indicating a potential intron/exon junction. Another intron/exon junction could interrupt codon 192 of *erg* (glycine) between the G in the first position and the G in the second position which was found to be fused to EWS in EWS-*erg* fusion type 9e. This fusion isoform, connecting the NH₂-terminal domain of EWS directly to the *erg* ETS domain, has not been described for Flil and may indicate that the ETS domain itself is sufficient to induce transformation. In fact, similar studies (37–39) have demonstrated that the EWS/Flil chimeric protein is a transcriptional activator, suggesting that it deregulates the target genes of Flil.

The mapping of marker D21S60 to the *erg* YAC A125B12 (29) gives us additional information regarding the genomic organization of *erg*. In an integration map of human chromosome 21, D21S60 is located centromeric to the *erg* gene (40). In addition to our PCR results, this evidence strongly suggests that the *erg* gene is transcribed in a telomere to centromere direction. Since a simple balanced translocation would juxtapose the 5' region of EWS to the 5' end of *erg* and not the 3' end, this putative *erg* orientation suggests that a productive EWS-*erg* fusion requires a complex chromosomal rearrangement. Additional FISH studies with chromosome-specific painting probes and probes encompassing or flanking the translocation breakpoints will be necessary to precisely define the dynamics of these chromosomal rearrangements.

Another rationale for this study was to analyze the incidence of the different fusion types in a spectrum of tumors with known phenotype. The breakpoints resulting in EWS/Flil type 1 and type 2 fusions appear to be, by far, the most common, and the occurrence of either type 1 or 2 does not seem to be correlated to whether the tumors are considered to be ESs or PNETs. Although we have analyzed a significant number of tumors, analysis of a larger series is necessary to determine whether the occurrence of distinct fusion types is a random event. Considering the number of novel fusion transcripts we have identified in a relatively small number of cases, we predict that additional studies will probably reveal other fusion isoforms occurring between EWS-Flil, EWS-*erg*, and fusion of EWS with genes on other chromosomes. Further biological studies are needed to determine the role of the fusion transcripts in the development of these tumors. At present, RT-PCR and FISH clearly provide a sensitive means by which to identify the molecular events involved in the standard, variant, and masked translocations in ES and PNET.

Note added in proof. Since submission of this article two independent groups (Zucman, J., T. Melot, C. Desmaza, J. Ghysdael, B. Plougastel, M. Peter, J. M. Zucker, T. J. Triche, D. Sheer, C. Turc-Carel, et al. 1993. *EMBO [Eur. Mol. Biol. Organ.] J.* 12:4481–4487) (Sorensen, P. H. B., S.-L. Lessnick, D. Lopez-Terrada, X. F. Liu, T. J. Triche, and C. T. Denny. 1994. *Nat. Genet.* 6:146–151) have reported similar EWS-*erg* fusion transcripts.

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