REVIEW

Progress in the molecular biology of Ewing tumors

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Abstract
Purpose/results/discussion. Rearrangement of the EWS gene with an ETS oncogene by chromosomal translocation is a hallmark of the Ewing family of tumors (EFT). Detectability, incidence, tumor specificity and variability of this aberration have been matters of intense investigation in recent years. A number of related alterations have also been found in other malignancies. The common consequence of these gene rearrangements is the generation of an aberrant transcription factor. In EFT, the ETS partner is responsible for target recognition. However, synergistic and possibly tissue-restricted transcription factors interacting with either the EWS or the ETS portion may influence target selection. Minimal domains of both fusion partners were defined that have proved necessary for the in vitro transformation of murine fibroblasts. These functional studies suggest a role for aberrant transcriptional regulation of transforming target genes by the chimeric transcription factors. Also, fusion of the two unrelated protein domains may affect overall protein conformation and consequently DNA binding specificity. Recent evidence suggests that EWS, when fused to a transcription factor, interacts with different partners than germ-line EWS. Variability in EWS–ETS gene fusions has recently been demonstrated to correlate with clinical outcome. This finding may reflect functional differences of the individual chimeric transcription factors. Alternatively, type and availability of specific recombinases at different time-points of stem cell development or in different stem cell lineages may determine fusion type. Studies on EFT cell lines using EWS–ETS antagonists do suggest a rate-limiting essential role for the gene rearrangement in the self-renewal capacity of EFT cells. The presence of additional aberrations varying in number and type that may account for immortalization and full transformation is postulated. Knowledge about such secondary alterations may provide valuable prognostic markers that could be used for treatment stratification.

Key words: EWS, ETS, IGF1, tumor suppressor, prognosis.

Introduction
Ewing’s sarcoma (ES), being a rare malignant disease affecting bone and soft tissue in children and young adults, was hardly known to people other than pediatric oncologists until the characterization of a chimeric gene product presumed to be causally involved in the generation of this neoplasm. It dramatically gained attention when, from investigating other malignancies, it became apparent that the ES-derived oncoprotein constitutes the prototype of a whole class of aberrant proteins specifically associated with certain tumor types. Consequently, ES may be considered a model system to study malignant conversion on a subclinical level. The discovery of the ES-associated gene rearrangement transiently halted a controversy among pathologists about the existence of distinct categories of ES (i.e. osseous ES, extra-skeletal ES, Askin tumor, peripheral primitive neuroectodermal tumor) because it was found to be expressed in all of them. Clinically, however, there is a need for diversification.

Although more than half of the patients can be cured by multimodal therapy, one third of cases with localized disease and about 80% of patients presenting with metastases succumb to the disease (for a recent review, see Kovar et al.1). Current treatment protocols have largely compensated for classical prognostic markers such as tumor volume and localization of the primary except for the rather unfavorable presence of metastases at diagnosis. It is likely, therefore, that biological differences exist between so far incurable aggressive disease and clinically manageable localized disease inexplicable by the mere presence of the ES-associated gene rearrangement. While Ewing’s tumor research has focused on the clinical exploitability and the function of the ES-specific gene rearrangement since its discovery in 1992, this review will also consider extensively the role of additional molecular aberrations in the search for useful prognostic markers. Neoplastic transformation and metastatic spread is commonly believed to result from a multi-step pro-
cess. In this context, the ES-specific gene rearrangement obviously constitutes a rate-limiting event. According to Knudson’s legendary two-hit hypothesis, at least one additional aberration should be present in an Ewing tumor. It is possible that this second hit is less specific and affects different genes at different times during development of the enigmatic Ewing tumor stem cell, thus defining distinct subcategories of the disease. Consequently, Ewing tumor research is slowly moving towards molecular subclassification and staging.

**Diagnostic tools**

In 1988, the cytogenetic translocation t(11;22) (q24;q12) was described as specifically associated with histopathologically diagnosed ES and peripheral primitive neuroectodermal tumor (pPNET). The presence of this aberration in a largely undifferentiated small round cell tumor of childhood turned out to be a formidable diagnostic marker. However, cytogenetic analysis was restricted to tumor cells with at least limited *in vitro* proliferation potential. The generation of an antibody, HBA71, specifically reacting with the surface glycoprotein encoded by the *MIC2* gene, which was found to be abundantly present in tumor cells carrying a chromosome 22q12 aberration, enlarged the spectrum of diagnostic tools. However, embryonal rhabdomyosarcomas, neuroendocrine tumors and carcinomas occasionally stained positive with HBA71 and, when using a more sensitive antibody (12E7), high level expression of this antigen was also noted in early hematopoietic precursor cells and several lymphomas. The characterization of the ES break-point regions on chromosomes 22 and 11 and the subsequent cloning of a chimeric cDNA resulting from a gene fusion between a novel gene, designated *EWS*, and the ETS transcription factor gene *FLI1* allowed for sensitive detection of tumor cells carrying a 11;22 translocation even in small samples of fresh, frozen or paraffin-embedded material by means of reverse transcriptase polymerase chain reaction (RT-PCR). Subsequently, several alternative fusion partners for *EWS* from the *ETS* oncogene family were identified in ES and pPNET cases. Using the RT-PCR method, combined with genomic analysis of the *EWS* break-point region, on a large series of osseous and extra-skeletal ES (including Askin tumors and pPNET, designated Ewing family of tumors (EFT)) as opposed to several unrelated small round cell tumors, the specificity of the *EWS–ETS* gene rearrangement and the correlation with high MIC2 expression was confirmed. Recently, however, the limitation of this aberration to typical EFT members has been questioned since RT-PCR amplifiable *EWS–FLI1* fusion transcripts have been reported in childhood soft tissue sarcomas with mixed phenotype in some olfactory neuroblastomas which have previously been shown not to express MIC2 and in two cases of classical MIC2-negative neuroblastoma. In the absence of any cytogenetic evidence for a t(11;22) in neuroblastoma the latter finding needs to be independently confirmed. On the other hand, RT-PCR failed to demonstrate the presence of chimeric EWS transcripts in roughly 5% of histologically classified EFT. Sceptics might raise their fingers and recall all the potential pitfalls of using RT-PCR including the method’s inherent susceptibility to cross-contamination as a single tool in the diagnosis of histopathologically ambiguous cases of small round cell tumors. Intriguing questions, i.e. if *EWS–ETS* gene rearrangements can occur outside the EFT and if ‘atypical’ ES exist, can only be assessed by the use of complementary techniques allowing for the visualization of the *EWS* gene rearrangement on a single cell level. It has already been demonstrated that fluorescent *in situ* hybridization (FISH) using cosmids flanking the EFT break-point regions is not restricted to metaphase chromosomes, but is also feasible to detect the gene rearrangement efficiently on interphase nuclei (Hattinger et al., unpublished). Alternatively, antibodies to unique domains of the chimeric gene product could allow the routine pathologist to screen for the *EWS* rearrangement by standard immunohistochemical methods. The author and others have recently obtained preliminary indirect evidence from protein interaction studies that an amino terminal *EWS* domain, which appears to be inaccessible in germ-line EWS, might be specifically exposed on the surface of the chimeric product (see below). One could endeavor, therefore, to generate an agent that distinctly recognizes the altered conformation of the EWS portion present in EWS fusion proteins. The hinge region of EWS–ETS chimeric proteins displays a high degree of variability due to variable break-point locations in the genes contributing to the translocation. So far, five alternative *ETS* family members have been found in *EWS* gene rearrangements. Therefore, antibodies to the linker domain of fused partners would be of only limited use in routine diagnosis. For the analysis of RT-PCR negative ‘atypical’ EFT and for small round cell tumors with a diagnosis other than EFT but RT-PCR positive for an EWS chimeric transcript, it is strongly recommended to confirm the molecular diagnosis by the demonstration of an EWS aberration on either DNA level (FISH or Southern blot) or on RNA level by Northern blotting. Presently, it cannot be excluded that using these approaches, followed by refined cloning procedures, further *ETS* family members will be identified as alternative fusion partners for *EWS* in EFT or non-EFT.
The EWS–ETS gene rearrangement

The EWS gene family

EWS is the prototype of a growing family of putative RNA-binding proteins including TLS (translocated in liposarcoma)/FUS, hTAF_{II}68 (TATA box binding protein associated factor), \textit{hTAF}_{II}68, the small nuclear ribonuclear protein (snRNP)-associated 69-kDa protein, the bovine Pigpen protein and \textit{Drosophila cabeza}/SARFH (sarcoma associated RNA binding fly homologue), that share distinct structural characteristics such as a conserved RNA binding motif flanked by arginine–glycine–glycine (RGG) boxes and a putative zinc-finger domain in the carboxy terminus. This portion is replaced by the DNA binding domain of a transcription factor in the oncogenic EWS and TLS fusion proteins. The amino terminus is rich in glutamine and proline residues. As such, it resembles the activation domain of certain transcription factors such as SP-1. In EWS, this \textit{N}-terminal domain (NTD), which is encoded by the first seven exons, is comprised of 30 copies of a repeated degenerate peptide of 7–12 residues rich in tyrosine, serine, threonine, glycine and glutamine (Fig. 1). TLS was identified as a heterogeneous nuclear ribonucleoprotein (hnRNP) in non-spicosomal complexes on mRNA continuously shuttling between the nucleus and the cytoplasm and SARFH was found to be associated with regions of the \textit{Drosophila} chromatin transcribed by RNA polymerase II. Consistent with a role of EWS family members in gene transcription, hTAF_{II}68, TLS and EWS have been identified in subpopulations of the general transcription factor TF_{II}D. However, recent evidence suggests that the oncogenic derivatives of TLS and EWS are not stably associated with the RNA polymerase II complex and TF_{II}D. Accumulation in nuclear inclusions such as the coiled body and the nucleolus has been reported for Pigpen, the 69-kDa snRNP-associated protein, and, after transcriptional inhibition, for TLS. Such nuclear subcompartments might either constitute the site of normal function of these EWS-related proteins or serve as their reservoir. Interestingly, oncoproteins that contain the amino terminal domain of EWS or TLS are also targeted to the same structure. So far, the functional relevance of this finding is completely unknown.

A role for EWS and its partner genes in determining the EFT phenotype

Figure 2 summarizes all known gene fusions involving either EWS or TLS in human malignancies. In an NIH3T3 transformation study, the type of transcription factor contributing to the chimeric gene product determined cell morphology. This observation might in part explain why only members of the ETS transcription factor family are found in gene fusions with EWS associated with an EFT phenotype. However, while EWS and TLS amino termini appear to be functionally interchangeable...
when fused to the transcription factor CHOP in the \textit{in vitro} model,\textsuperscript{35} as well as in myxoid chondrosarcoma,\textsuperscript{33,34,48} TLS has never been found to replace EWS in EFT. In contrast, fusion of TLS to the ETS family member ERG, which is involved in 10\% of EFT, has been reported for poor prognosis, (t(16;21) positive, acute myeloid leukemia.\textsuperscript{49-51} Rearrange-ment of EWS with other transcription factor genes such as \textit{ATF1}, the Wilms' tumor gene \textit{WT1} and the nuclear receptor \textit{CHN/TEC} have been shown to be associated with malignant melanoma of soft parts, desmoplastic small round cell tumor and myxoid chondrosarcoma, respectively.\textsuperscript{52-55} Thus, it is the specific combination of EWS with a subset of ETS transcription factor genes and/or a particular stem cell in which these genes are sensitive to illegitimate recombination that determine the EFT phenotype. Accessibility to rearrangement by an as yet undefined recombinase might also determine the incidence of EFT. Zucman \textit{et al.} reported recently that sequence analysis of the entire EWS intron 6 region close to the major break-point region in EFT from Caucasian origin revealed a very high density of Alu elements resulting from repeated retroposition during evolution.\textsuperscript{56} The Alu family of short interspersed repetitive DNA elements has previously been demonstrated to be frequently involved in human gene rearrangements.\textsuperscript{57} This region was found to be reduced by 50\% due to deletion in the African population. This inter-ethnic polymorphism in the EWS gene is accompanied by a striking difference in the incidence of EFT between populations of European and African origin.\textsuperscript{58,59} It should be noted that the majority of EWS genomic break-points occur in intron 7 and that intron 6 is, in fact, never directly rearranged in EFT. So far, only three EWS genomic break-points have been sequenced, two in EFT and one in a desmoplastic small round cell tumor,\textsuperscript{60-62} none of which contained Alu elements in the immediate vicinity of the rearrangement sites. Thus, direct proof for the involvement of Alu elements in EWS translocation is not available. In the published cases, the lack of uniformity of sequences affected by the gene rearrangement does not allow the identification of a specific recombinase responsible for the translocation. Chromosome 22 alteration may occur as the only cytogenetically visible aberration in EFT suggesting that the \textit{EWS}–\textit{ETS} gene rearrangement is not the consequence of a general genomic destabilization. However, the frequent involvement of more than two chromosomes in complex chromosome 22 aberrations and evidence for deletion of considerable amounts of sequences from the directly involved genes on the untranscribed counterpart of the derivative chromosome 22\textsuperscript{60} imply a complex mechanism for gene rearrangement in EFT. In addition, while \textit{EWS} and \textit{FLI1} are equally oriented on the long arms of chromosomes 22 and 11 from the centromere to the telomere allowing for simple reciprocal transloca-

The \textit{ETS partner in the EWS fusion gene}\n
The ETS transcription factor family currently counts more than 30 members. It is characterized by the presence of a unique DNA binding domain which is highly conserved from flies to humans and has been first described for the viral oncogene \textit{v-ets} of avian erythroblastosis virus E26 (E twenty-six specific). Several \textit{ETS} subfamilies can be defined on the basis of evolutionary sequence conservation. In ET, 95\% of cases show \textit{EWS} fusion to the \textit{FLI1} (Friend leukemia virus integration site 1)/\textit{ERG} (ETS related gene) subfamily of transcriptional activators.\textsuperscript{16,18} These two gene products share, in addition to almost identical DNA binding domains, a 16-amino acid stretch immediately upstream of the \textit{ETS} domain which is 100\% conserved between \textit{Xenopus} and humans, suggesting an important but as yet unidentified functional role.\textsuperscript{66} This portion is retained in almost all \textit{EWS}–\textit{FLI1} and \textit{EWS}–\textit{ERG}...
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ETS1, GABPa, and FLI1 have recently been demonstrated to bind to the transcription factor PAX5 in vitro. A flanking PAX5 binding site allowed for ETS1 binding to an imperfect ETS recognition motif indicating that cooperativity might change ETS binding specificity to some extent. Some ETS proteins, including ETS1, ET52, FLI1, ERG, GABPa, and TEL, share a further conserved region flanking the amino terminal transactivation domain, referred to as the 'pointed or B-domain'. This structure has been shown to define a specific oligomerization interface. For TEL, it governs homotypic aggregation. No interaction partner has been defined for FLI1 and ERG so far. The conserved fold of the amino terminal domain of ETS proteins is, however, likely to underlie a conserved function. Thus, replacement of this portion by the EWS amino terminus may alter not only quantitatively but also qualitatively the transcriptional activation properties of these two ETS family members by placing them into a different protein context. In addition, several ETS family members have been demonstrated to be a target for the RAS/RAF MAP kinase signaling pathway. For ETS1, ras regulation involves phosphorylation of residues within the amino terminal 'pointed' domain. So far, no link between this signalling pathway and FLI1 or ERG has been reported. Consequently, it remains unclear if fusion to EWS will uncouple FLI1/ERG target gene regulation from extracellular signaling.

In about 1% of ET cases, EWS is rearranged with FEV (fifth Ewing's tumor variant) on chromosome 2. This ETS family member displays 90% identity to FLI1 in the DNA binding domain but the FLI1- and ERG-specific flanking 16 amino acid sequence is missing in this protein. Interestingly, FEV also lacks an amino terminal transactivation domain present in most other ETS family members. Instead, it carries a long C-terminus which, because of the presence of abundant alanine residues, might serve as a putative repressor domain. However, experimental proof for such an activity is not yet available. Since this portion is retained in the EWS fusion, it remains to be established if the gene rearrangement would result in a functional conversion to an activator. Interestingly, germ-line FEV cDNA has been cloned from an EWS–FLI1 expressing Ewing tumor cell line, indicating coexpression of the two genes within the same cell. If EWS–FLI1 and FEV target the same genes and FEV operates as a repressor, it is possible that the EFT gene rearrangement results in the release of these genes from transcriptional inhibition by competitive binding.

So far, three cases of suspected EFT have been reported in which EWS was fused to ETS family genes of a different subclass on chromosomes 7p22 and 17q21, ET71 (ETS translocation variant 1) and E1AF (Adeno virus E1A enhancer binding factor), the putative human homologues of mouse ER81 and

### Table: Tumor-specific rearrangements between an EWS family gene and a transcription factor gene.

| EWS family member | transcription factor (<type>) | neoplasm               |
|-------------------|-------------------------------|------------------------|
| EWS              | FLI1 (ETS)                    | Ewing tumor (85%)      |
| EWS              | ERG (ETS)                     | Ewing tumor (10%)      |
| EWS              | ET51 (ETS)                    | Ewing tumor (<1%)      |
| EWS              | E1AF (ETS)                    | Ewing tumor (<1%)      |
| EWS              | FEV (ETS)                     | myxoid liposarcoma     |
| EWS              | CHOP (bZIP)                   | melanoma of soft parts |
| EWS              | ATF1 (bZIP)                   | desmoplastic small    |
| EWS              | WT1 (Steroid)                 | round cell tumor       |
| EWS              | TEC (Steroid)                 | myxoid chondrosarcoma  |
| TLS              | CHOP (bZIP)                   | myxoid liposarcoma     |
| TLS              | ERG (ETS)                     | acute myeloid leukemia |
| TAF68            | ?                             | ?                      |
| 69kD             | ?                             | ?                      |

Fig. 2. Tumor-specific rearrangements between an EWS family gene and a transcription factor gene.
### Fig. 3. Comparison of DNA-binding domains of ETS proteins rearranged or coexpressed in Ewing's tumors. *: detectable only by RT-PCR; †: identical amino acids; underlined: positions homologous to amino acids making contact to DNA in the PU.1 ETS-family member.
Most notably, the DNA binding domains of these two transcription factors diverge from FLI1, ERG and FEV by 38% including the third $\alpha$-helix that contacts the central core of ETS binding sequences. Do EWS–ETV1 and EWS–E1AF chimeric transcription factors target the same genes as EWS–FLI1, EWS–ERG and EWS–FEV? By a subtractive cloning strategy for genes differentially expressed in EWS–FLI1-transformed and FLI1-transfected untransformed murine fibroblasts (NIH3T3), several potential EWS–FLI1-specific target genes were identified. Among them were the murine homologue of cytochrome P-450 F1, cytokeratine 15, a novel SH2 domain containing protein, EAT2 (EWS–FLI1 activated transcript 2) and the stromelysin gene. Stromelysin is a matrix metalloproteinase involved in metastatic invasion. Previously, E1AF has been demonstrated to regulate the stromelysin gene. In transfection experiments, E1AF was sufficient to confer an invasive phenotype to non-metastatic human breast cancer cells (MCF7) and antisense RNA to E1AF was able to revert it in a squamous cell carcinoma cell line. Stromelysin has also been demonstrated to be activated by ETS2, a member of another ETS subfamily. However, coexpression of ERG, which also strongly binds to the same promoter but is by itself unable to activate it, resulted in inhibition of ETS2-mediated stromelysin gene activation. Thus, different ETS family members, irrespective of their subclass, appear to compete for binding to specific target genes. It is likely, therefore, that target selectivity and the specific mode of gene regulation by the EWS chimeric ETS transcription factors strongly depends on the cellular background. A spectrum of ETS-related gene products coexpressed with EWS–FLI1 in EFT cell lines by amplification of ETS DNA binding domain encoding cDNAs has recently been defined using degenerate primers. Among them, ETS2, E4TF1-60, ELK, ELF1, the putative human homologue of ER71, and a novel gene product ELFR were identified. None of the ETS family members involved in EFT-specific gene rearrangements were found to be expressed in their germ-line configuration. This assay, however, may have missed low level expression of some ETS family members (i.e. FEV). As shown in Fig. 3 comparing the DNA binding domains of the ETS gene products alternatively fused to EWS in EFT with those found to be coexpressed with the chimeric transcription factors, ELK is the most likely ETS-related gene product that might interfere with the EFT fusion proteins in target site selection because the DNA-contacting third $\alpha$-helix of the ETS domain is identical to that of FLI1, ERG and FEV. ELK is one of several alternative ternary complex factors regulating a number of growth factor inducible genes. In fact, EWS–FLI1 can replace ELK within the ternary complex formed on the serum response element of the $c fos$ and the $EGR1$ promoters. In contrast to germ-line FLI1, binding of EWS–FLI1 to the serum response element did not require interaction with the serum response factor SRF. Ternary complex formation by FLI1 and EWS–FLI1 was mediated by a domain preceding the DNA binding domain and present in the majority of EFT-derived EWS–FLI1 fusions that show limited similarity to the ELK1–SRF interaction domain. However, no homologous structure can be identified in ERG and evidence for EWS–ERG involvement in EFT-specific complex formation is not available.

In summary, current knowledge about normal and aberrant ETS proteins suggest a number of interesting candidate target genes for the EFT-specific chimeric transcription factors, potentially involved in the regulation of cell growth, signaling and metastasis, as revealed by the study of heterologous cellular systems such as murine fibroblasts. However, ample evidence exists that ETS transcription factor action is largely context specific. For EFT, the cell of origin remains a matter of speculation. Because of limited neural differentiation potential, EFT is considered as derived from the neuroectoderm. Interestingly, Xenopus FLI1 has been shown to be expressed in a restrictive pattern during embryogenesis evocative of neural crest cell invasion. It is therefore speculated that FLI1 might be involved in neural differentiation in the context of these early stem cells. If so, unscheduled activation of FLI1-responsive genes by the EFT-specific EWS fusion proteins in an undifferentiated cell possibly unrelated to the neural crest might result in limited neural differentiation of the EFT stem cell depending on the degree of determination achieved at the time of gene rearrangement. This model could explain the variable degree of neuroectodermal marker expression in ES and pPNET as well as the occurrence of biphenotypic tumors.

The rate-limiting (first) hit in EFT pathogenesis

EWS–ETS gene rearrangements are the only genetic aberrations that have so far been identified as highly associated with histologically diagnosed EFT. This association is the only available compelling argument that EWS–ETS gene rearrangements can be rate-limiting for tumorigenesis. Although complementary experimental evidence supports this assumption, the mechanism of malignant transformation by these chimeric oncoproteins remains elusive. The best studied biological model for the pathogenic role of inappropriately activated FLI1 is Friend murine leukemia virus (F-MuLV)-induced erythroleukemia. Insertional activation of the FLI1 gene appears to be the first detectable genetic change associated with this disease. The association between the detection of FLI1 rearrangement and clonal outgrowth of erythroleukemia cells suggests
that the activation of this transcription factor may be affecting the self-renewal potential of the infected erythroid progenitors. However, leukemogenesis proceeds in multiple steps and additional aberrations affecting viability of cells (e.g. inactivation of the tumor suppressor gene p53) can be observed in F-MuLV induced erythroleukemia.\textsuperscript{91,92} In contrast to ERG,\textsuperscript{93} normal FLI1 was reported to be unable to transform murine fibroblasts (NIH3T3) while expression of an EWS fusion protein resulted in pronounced anchorage-independent clonogenicity of NIH3T3 cells.\textsuperscript{94,95} However, rat embryo fibroblasts and some murine fibroblast subclones were resistant to EWS–FLI1-mediated transformation. These findings again suggest that the oncogenic potential of normal and aberrant FLI1/ERG ETS subfamily members may depend on a cell type-specific availability of relevant synergetic factors and possibly on the presence of additional aberrations. NIH3T3 transfection studies with various recombinant EWS–FLI1 deletion mutants revealed a dependence of transformation on both the EWS portion and the ETS domain.\textsuperscript{95} However, optimal transactivation potential mediated by the 30 EWS amino terminal degenerate repeats included in almost all EFT-derived fusion proteins was dispensable for maximal focus formation of transfected NIH3T3 in soft agar. The minimal EWS domain required to transform murine fibroblasts was delineated to the first 82 amino acids.\textsuperscript{94} Recently, evidence obtained shows that within the EWS–FLI1 fusion protein, but not within germ-line EWS, this peptide directly contacts a component of the RNA polymerase II complex, RPB7, and that this interaction is sufficient to drive EWS–FLI1-mediated reporter gene transactivation.\textsuperscript{125} Interactions of full-length EWS with the general transcription factor TFIIID, an essential component of the transcriptional preinitiation complex, were absent from EWS fusion proteins.\textsuperscript{126} It is therefore possible that fusion of the EWS amino terminus to the FLI1 DNA binding domain alters the protein conformation and directly recruits RNA polymerase II to FLI1 target genes. Since RPB7 displays similarities to prokaryotic sigma factors, it might be involved in EFT-specific target site selection. Further protein–protein interactions presumably occurring downstream of the 82 amino acids might be required for efficient gene regulation within the EFT context. In addition, using the yeast two-hybrid protein interaction trap, further candidate proteins not directly related to transcription regulation were identified that interacted with the 82 amino acids long minimal transformation domain. These interactions await detailed characterization.

Since no tissue of EFT origin has been identified so far, transformation studies of authentic EFT stem cells cannot be performed. Alternatively, several investigators have used EWS–FLI1 antagonists (antisense RNA expression vectors, antisense oligonucleotides, dominant negative proteins) to modulate expression of the chimeric oncoprotein in EFT cell lines.\textsuperscript{86,96–98} These studies revealed a growth inhibitory and anti-tumorigenic effect of these agents. Reduction in cell growth appeared to result from cell cycle arrest and not from reduced tumor cell viability. Recently, EWS–FLI1-mediated transformation of murine fibroblasts was demonstrated to require the presence of a functional insulin-like growth factor-1 (IGF1) receptor.\textsuperscript{99} Interestingly, consistent expression of IGF1 and its receptor was previously reported for EFT and IGF1 was demonstrated to act as a potent growth factor for EFT cells in the absence of serum.\textsuperscript{90–93} This cytokine appears to regulate negatively several mechanisms of programmed cell death at a far downstream step.\textsuperscript{104} It has been shown that inhibition of the IGF1 autoregulatory circuit by anti-IGF1 receptor antibodies resulted in increased apoptosis and reduced tumorigenicity of EFT cells.\textsuperscript{107} Taken together, the EWS–ETS gene rearrangement appears to be involved in the aberrant self-renewal capacity of EFT cells but might not be sufficient to guarantee survival of initiated tumor cells. However, as demonstrated recently, there might still be some role for FLI1, ERG and their EWS fusions to play in the protection from stress (i.e. calcium ionophore and serum deprivation)-induced cell death.\textsuperscript{105}

The second hit

Assuming that the EWS–ETS gene rearrangement is able to initiate EFT pathogenesis but is not sufficient to generate malignant transformation, the presence of additional mutations must be postulated. These aberrations might not necessarily be tumor specific but may display inter-individual variation that could account for variations in EFT phenotype as well as in clinical behavior. Also, they may determine differentiation capacity, invasive potential and treatment resistance. Consequently, while from a clinical point of view the EWS–ETS gene rearrangement provides a valuable diagnostic tumor marker, knowledge about the nature of additional aberrations in EFT may assist subclassification and provide prognostic tools. Since studies on facultative genetic anomalies in EFT have been largely neglected since the discovery of the EWS–ETS gene rearrangements, enhanced efforts to define the multitude of additional aberrations are warranted for the benefit of patients.

Clues from cytogenetics

In three independent reviews of cytogenetically informative EFT cases, non-random numerical and structural chromosomal aberrations were reported to occur with variable frequencies in addition to the
tumor specific t(11;22)(q24;q12)\textsuperscript{106–107} (Hattinger \textit{et al.}, unpublished). These include trisomy 8 in about 50\% of cases frequently coupled with trisomy 12 occurring in roughly 20\%, and a derivative chromosome 16 as a result of an unbalanced t(1;16) in 18\% of EFT. In rarer cases, other aneuploidies have been identified. Structural chromosome 1 aberrations that either result in gains of chromosome 1q21–22 or relative losses of the short arm of chromosome 1, a frequent alteration in neuroblastoma and other neuroectodermal tumors, have also been observed in EFT (Hattinger \textit{et al.}, unpublished). Interestingly, this chromosomal region harbours a gene encoding a protein (p73) that is structurally and functionally related to the tumor suppressor p53, a transcription factor involved in the regulation of cell growth and apoptosis, and frequently inactivated during the progression of many tumors.\textsuperscript{108,109} While research currently focuses on the role of p73 for neuroblastoma pathogenesis, its relevance for a subset of EFT is something that has to be explored.

In general, excluding chromosome 22q12 translocations, numerical chromosome changes are the most frequent cytogenetic findings in EFT. These are likely to affect gene dosage. However, no candidate genes that could promote \textit{EWS–ETS}-initiated EFT pathogenesis when expressed at aberrant levels have been identified so far. Also, information on genes affected by the recurrent chromosome 16 and 1 structural alterations is not available yet. Frequently, these cytogenetic alterations occur in only a subpopulation of neoplastic cells within the tumor suggesting that they may be associated with late stages of tumor progression.

\textit{The role of non-specific cancer genes}

In the absence of recurrent candidate progression-associated genetic alterations identifiable in EFT by the means of cytogenetics, work has focused on the analysis of mutations generally associated with a broad range of human malignancies. Genes investigated during the last years include the oncogenes \textit{ras}, \textit{cmyc} and \textit{MDM2}, the tumor suppressors p53, p16, and \textit{Rb}, the metastasis-associated splice variants of the CD44 adhesion molecule, and the tumor-specific metastasis suppressor gene \textit{nm23H1}.\textsuperscript{110–113,127} None of the studied oncogenes was found to be altered by mutation or in expression although occasional low level amplification of \textit{MDM2} has been reported in an independent study on a similar sized cohort of EFT patients.\textsuperscript{111} Neither mutation nor differences in expression levels of the nucleotide diphosphate-kinase \textit{nm23H1} were observed irrespective of the disease extension.\textsuperscript{112} Only standard \textit{CD44} expression was detectable in EFT.\textsuperscript{127} In contrast, homozygous deletions of the p16 tumor suppressor was identified in about one-third of primary EFT samples. This finding was surprising since chromosomal aberrations of band 9q21 containing the \textit{p16} gene were not reported before suggesting a high-frequency of microdeletions. Expression studies on EFT cell lines suggested that the frequency of \textit{p16} inactivation might be even higher since post-transcriptional gene silencing was observed in several cases.\textsuperscript{113} p16 acts as an inhibitor of the cyclin D1/cyclin-dependent kinase 4 (CDK4) complex that inactivates the cell cycle inhibitor pRb by phosphorylation. Inactivation of p16 should compromise the G1 cell cycle checkpoint. Over-expression of either cyclin D1 or CDK4, or loss of pRb function, is believed to mediate a similar effect.\textsuperscript{114} In fact, we observed frequent cyclin D1 over-expression as well as variable CDK4 abundance in EFT cell lines and loss of Rb in one case. However, expression data from primary tumor material are not available yet. In addition, low level CDK4 amplification was previously reported for two of 30 EFT samples.\textsuperscript{111}

In virally induced malignancy, G1 check-point control is frequently compromised by concomitant inactivation of the pRb and p53 pathways. In addition, F-MuLV-induced erythroleukemia involves not only the activation of the \textit{FLI1} oncogene as a rate-limiting step but also mutation of p53. The author and others have, therefore, investigated the status and expression of p53 and related genes in EFT.\textsuperscript{110,115} The frequency of p53 mutations in primary tumors was found to be lower than 10\% as opposed to a mean frequency of 40–60\% in most human malignancies. In contrast, in about half of EFT cell lines, the p53 gene was mutated and showed loss of heterozygosity. Comparison of p53 gene status between cell lines and the respective primary tumors of origin, when available, demonstrated that the observed increase in mutation frequency was due to selection and was not acquired during \textit{in vitro} expansion of tumor cells. This result suggested that p53 mutation might release EFT cells from some \textit{in vivo} growth or survival factor dependency. Transient transfection and over-expression of wildtype p53 in cell lines with endogenous mutant or wildtype gene status demonstrated frequent but variable reduction in apoptotic responsiveness,\textsuperscript{116} suggesting the presence of some as yet unidentified cell-protective mechanism in EFT cell lines. Preliminary expression analysis of members of the cell death regulatory \textit{Bcl}2 gene family did not reveal any significant variations between individual EFT cell lines (our unpublished observations). Previously, high levels and activity of poly(ADP-ribose) polymerase, a nuclear enzyme that participates in DNA replication, repair and the triggering of apoptosis induced by DNA strand breaks, have been reported for some EFT cell lines.\textsuperscript{117} Sensitivity of EFT cell lines to DNA damaging agents (etoposide, actinomycin D, X-rays) varied considerably in a manner independent from p53 responsiveness and endogenous p53 gene status suggesting that complete
mutational or partial inhibition of the p53 apoptosis pathway in EFT cell lines plays a role different from radio- and chemosensitivity. However, the physiological signals that stimulate p53-dependent cell death have not been defined so far.

**Molecular markers of prognosis**

As a result of variable break-point localization in the involved genes, EWS–ETS gene products vary considerably in size. Most fusions include EWS exons 1 to 7 (89%) and FLI1 exons 6 to 9 (54%). EWS/FLI1 exon 7/6 fusions (type 1) predominate independent of the disease extension (51%). In about one-third of EFT, FLI1 exon 5 is included into the chimeric gene product, most frequently joined to EWS exon 7 (type 2) (27%). In rare cases, the chromosomal translocation results in the inclusion of EWS exons 9 (1%) or 9 plus 10 (10%) or FLI1 exon 4 (1%). In about 3% of cases, FLI1 exon 6 or exon 6 plus 7 are missing from the gene fusion. This variability has prompted us to investigate a possible prognostic impact of the gene fusion type. The study, performed on 55 patients with localized disease and 30 patients with metastases at diagnosis, treated according to the European Intergroup Coordinated Ewing’s Sarcoma Studies (CESS 86 and EICESS 92), revealed a significantly better outcome for patients with localized disease carrying a type 1 EWS–FLI1 expressing tumor as compared to non-type 1 cases. A recent update after a median observation time of 3.12 years confirmed this result (Zoubek et al., unpublished). In addition, an independent American study performed on a similar sized cohort of patients after a median follow up of 31 months, using a similar treatment regimen, not only supported our findings but also identified the EWS–ETS gene fusion type as a prognostic marker independent from the presence of metastases at diagnosis in a multi-variate analysis. About 55% of the ‘non-type 1’ group in the two studies were comprised of type 2 gene fusions. Because of the low incidence of the individual ‘other-gene’ fusion types, no distinction has been made between various non-type 1 subgroups so far. In the absence of a biological explanation for the observed prognostic differences, large collaborative prospective studies are warranted to highlight the specific chimeric molecules and the protein domains associated with adverse patients’ outcome.

Still, about 20% of patients with localized tumors and more than half of the patients with metastases succumb from the disease despite the expression of a type 1 EWS–FLI1 gene fusion, suggesting the existence of additional adverse factors. Comparison of p16 gene status and clinical course of 23 EFT patients analyzed so far suggested an adverse prognosis associated with this aberration independent from the extension of the disease. These results, which have not been subjected to statistical analysis, must be considered as preliminary since patients’ numbers in the study were small and the median observation period did not exceed 2 years. Retrospective immunohistochemical analysis of biopsy material from a large number of patients will help to clarify the prognostic relevance of a disrupted pRb cell cycle regulatory pathway in EFT. Also, mutation of p53 might be linked to an adverse outcome, since none of the three EFT patients from our series carrying such an aberration survived. However, because of the rarity of this alteration it cannot serve as a useful prognostic marker.

A prognostic relevance for EFT of the observed numerical and structural cytogenetic changes has not been demonstrated with confidence due to low sample numbers in the studies performed so far. Most recently, deletion at 1p36, occurring in 6/22 localized EFT, was discussed as being associated with unfavorable outcome in this group (Hattinger et al., unpublished).

In the absence of reliable molecular markers to predict outcome in EFT, the presence of clinically overt metastases at diagnosis is commonly considered as the only prognostic criterion that is used for treatment stratification. The EWS–ETS gene rearrangement as a tumor cell specific marker detectable by the highly sensitive RT-PCR method provides a powerful means for the detection of minute numbers of circulating tumor cells that may be the source of clinically occult micrometastases. However, except shortly after surgical intervention, mobilization of PCR detectable amounts of tumor cells (>1/10⁶) into the bloodstream has rarely been observed. In contrast, tumor cells were detected at diagnosis by this method in the bone marrow of 30% of patients with localized disease, 50% of cases with isolated lung metastases and all patients with bone metastases. In a preliminary series of 23 patients lacking clinically overt dissemination, RT-PCR screening for bone marrow involvement did not allow the prediction of early relapse after a median observation time of 30 months. It is, however, necessary to recall several factors that may affect tumor cell detection in the bone marrow by RT-PCR: (1) tumor cell infiltration may be focal and bone marrow aspiration may miss these sites, (2) bone marrow aspirates may contain variable amounts of diluting blood resulting in insufficient sensitivity, (3) primary EFT cells may differ in vitality, although diluted tumor cells from cell lines can be detected in blood samples even after 48 h at 4°C, and (4) bone marrow infiltrating tumor cells may be in a resting state and express lower levels of chimeric EWS RNA than proliferating tumor cells. RT-PCR measures RNA quantity rather than tumor cell abundance. In a recent study, up to 10-fold variations in the content of chimeric EWS–ETS tran-
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scripts between individual EFT cell lines have been reported. Moreover, germ-line EWS expression in T-cells has been demonstrated to depend on the proliferative activity. Since the EFT-specific chromosomal rearrangement places the chimeric gene under the control of the EWS regulatory sequences, it is possible that EWS–ETS gene expression may also vary. Consequently, detectability of EWS–ETS chimeric transcripts does not necessarily reflect true tumor cell content. Even if RT-PCR studies fail to demonstrate significance of positive blood or bone marrow screening results for relapse in patients with localized disease, the question of prognostic relevance of tumor cell infiltration remains unsolved. To assess this problem, immunohistochemical studies may prove to be superior to RT-PCR. While MIC2 may serve as a valuable surface marker in tumor diagnosis, its exploitability for tumor cell detection in hematopoietic tissue is limited. The recently discovered expression of gastrin-releasing peptide (GRP) by all EFT cell lines and about half of the primary tumors tested (Lawlor et al., unpublished) may provide a marker that, in conjunction with MIC2, may allow the identification of EFT cells in blood and bone marrow with increased specificity. The study of EWS–FLI1 transcriptional targets may result in the identification of tumor cell-restricted immunohistochemical markers. In order to detect positively staining cells with very low abundance on a routine basis, automated microscopic screening and consequently sophisticated technical equipment is warranted.

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