Stable interference of EWS–FLI1 in an Ewing sarcoma cell line impairs IGF-1/IGF-1R signalling and reveals TOPK as a new target

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BACKGROUND: Ewing sarcoma is a paradigm of solid tumour-bearing chromosomal translocations resulting in fusion proteins that act as deregulated transcription factors. Ewing sarcoma translocations fuse the EWS gene with an ETS transcription factor, mainly FLI1. Most of the EWS–FLI1 target genes still remain unknown and many have been identified in heterologous model systems.

METHODS: We have developed a stable RNA interference model knocking down EWS–FLI1 in the Ewing sarcoma cell line TC71.

RESULTS: EWS–FLI1 inhibition induced apoptosis, reduced cell migratory and tumourigenic capacities, and caused reduction in tumour growth. IGF-1 was downregulated and the IGF-1/IGF-1R signalling pathway was impaired. PBK/TOPK (T-LAK cell-originated protein kinase) expression was decreased because of EWS–FLI1 inhibition. We showed that TOPK is a new target gene of EWS–FLI1.

TOPK inhibition prompted a decrease in the proliferation rate and a dramatic change in the cell’s ability to grow in coalescence.

CONCLUSION: This is the first report of TOPK activity in Ewing sarcoma and suggests a significant role of this MAPK-like protein kinase in the Ewing sarcoma biology.

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Ewing sarcoma is an aggressive neoplasm of the bone and soft tissues of children and young adults. Ewing sarcoma is characterised by the presence of chimeric proteins joining the NH2-terminal domain of the RNA-binding protein EWS with the DNA-binding portion of an ETS transcription factor, FLI1, in 90% of the cases (Delattre et al, 1992; Sorensen et al, 1994; May et al, 1997). There is strong evidence for the oncogenic potential of EWS–ETS fusions (May et al, 1997). EWS–ETS chimeric proteins act as deregulated transcription factors, both activating and repressing the expression of target proteins (Prieur et al, 2004). Several of these have already been identified, including transcription, apoptotic, or signal transduction factors (Dauphinot et al, 2001; Amsellem et al, 2005; Mendiola et al, 2006). The proliferation and survival of Ewing sarcoma is also determined by autocrine and paracrine activation of growth factor receptors and their ligands, such as IGF-1 (Scotlandi et al, 1998).

The cellular context contributes to the phenotype because the introduction of EWS–ETS fusions into different cellular models resulted in diverse outcomes ranging from the induction of cell-cycle arrest or apoptosis to dedifferentiation (Thompson et al, 1999; Deneen and Denny, 2001; Lessnick et al, 2002; Zwerner et al, 2003). The use of RNA interference in the Ewing sarcoma cell lines allows for a detailed molecular analysis of the targeted gene fusion and its downstream targets in the tumour context itself, precluding the use of heterologous model systems.

We have used a stable RNA interference system knocking down EWS–FLI1 in the Ewing sarcoma cell line, TC71, to establish a long-term model in which we can analyse in depth the molecular behaviour of the fusion protein. Our shRNAi model disclosed TOPK, a kinase involved in cell proliferation and motility, as a new target of EWS–FLI1. TOPK expression was downregulated because of fusion interference. EWS–FLI1 inhibition decreased tumourigenic potential, reduced the tumour volume in a xenograft model,
impaired migration, and rendered cells more sensitive to both apoptosis and the action of inhibitors of the IGF-1/IGF-1R pathway, showing the interaction between these two key players in Ewing sarcoma tumourigenesis.

MATERIALS AND METHODS

Cell lines and drugs

The Ewing sarcoma cell line, TC71, was maintained as described earlier (Martins et al, 2006). NVP-AEW541 was obtained from Novartis Pharma AG (Basel, Switzerland). PD-98059 and LY-294002 were purchased from Stressgen (Victoria, BC, Canada), and G418 was acquired from Invitrogen (Carlsbad, CA, USA). Cell lines were tested for the absence of mycoplasma using the VerorGeM kit (Minerva Biolabs, Berlin, Germany).

shRNAi interference

An shRNAi approach was used based on the pSUPER-neo-GFP plasmid (Oligoengine, Seattle, WA, USA) (Brummelkamp et al, 2002). The shRNAi sequences were designed to target the fusion point between EWS and FLI1 following the manufacturer’s instructions in order to avoid affecting the wild-type partners. The targeted sequences of EWS – FLI1 were 5’-CAGAGGGACGACGCTACGGGCGACGACGAC-3’ for 7-6 shRNAi and 5’-GGGCGACGACGACGACGACG-3’ for 7-6 M shRNAi. The presence of the insert within the recombinant pSUPER vector was confirmed by sequencing (ABI Prism 3100 Genetic Analyzer, Applied Biosystems, Foster City, CA, USA). T771 cells were electroporated (125 V, 800 mA, 40 – 45 ms) using Gene Pulser II (Bio-Rad, Hercules, CA, USA), seeded, and allowed to recover for 24 h. G418 selection (500 μg ml⁻¹) was carried out for 3 weeks, and individual selected clones were grown for 2 additional weeks and split into several plates to create a stock of the early and late stages (eight cellular passages later).

Validated TOPK siRNA oligos were purchased from Ambion (Austin, TX, USA). (sense sequence: 5’-GGUGAAAGUUGUCUAGAUAUGt-3’; antisense sequence: 5’-CAUUAAGACAGCUUUUCACUCc-3’). TOPK siRNA oligos were transfected using jetPEI (Polyplus-transfection, Illkirch, France).

Western blot

Western blots were performed as described earlier (Martins et al, 2006). EWS– FLI1 expression was determined using the anti-FLI1 antibody (Santa Cruz BT, Santa Cruz, CA, USA). Other antibodies used were anti-phospho-p44/p42 MAPK (Thr202/Tyr204), anti-p44/p42 MAPK, anti-phospho- Akt (Ser473), anti-AKT, anti-p57/Kip2, and anti-Caspase 9 (all from Cell Signaling, Danvers, MA, USA), anti-DAX1 (a gift from Dr Javier Alonso-IIB, Madrid, Spain), anti-PARP/Caspase 9 (Serotec Germany, Duesseldorf, Germany), anti-Kip2, and anti-Caspase 9 (all from Cell Signaling, Danvers, MA, USA), anti-GAPDH, anti-IGF-1R and anti-NKX2.2 (Santa Cruz BT), anti-actin antibody (Santa Cruz BT, Santa Cruz, CA, USA). The cells were treated with the NVP-AEW541 for 15 min and with the inhibitors for 2 h, before IGF-1 stimulation (50 ng ml⁻¹) for 15 min in serum-free conditions.

Quantitative real-time PCR

Quantitative real-time PCR was performed as described earlier (Martins et al, 2008). Primers for the selected genes were designed using primer3 software (MIT, Cambridge, MA, USA) (Supplementary Table 4)
hierarchical clustering analysis and to clearly distinguish the shRNAi clone samples from the others. Genes with a P-value < 0.05 were considered candidate genes.

**Chromatin immunoprecipitation (ChIP)**

The bioinformatics program, TFM-explorer (http://bioinfo.lifl.fr/TFME/), was used for searching EWS–FLI1 putative binding sites in the TOPK promoter and introns. ChIP was performed using the Magna ChIP G Chromatin Immunoprecipitation kit (Millipore, Billerica, MA, USA) according to the manufacturer’s instructions. The antibodies used for each sample were 1 μg of IgG-negative control, 1 μg anti-RNA polymerase II antibody-positive control (provided by the kit), 5.0 μg of EWS goat polyclonal IgG (Santa Cruz BT) – antibody of interest, no antibody–ChIP reaction containing pre-cleared chromatin, and DNA–protein complexes before immunoprecipitation, named ‘Input DNA’. As EWS contains a RNA-binding domain, the immunoprecipitated products corresponded only to the fusion protein (DNA-binding domain of FLI1) and not to wild-type FLI1. Cell extracts were sonicated four times with a 12 s constant burst (100 J energy, 20% amplitude) (SONICS, Newtown, CT, USA), following by 30–40 s rest on ice between each pulse. Purified DNA was subjected to PCR. The control human PCR primers used for ChIP were those described by Zupanska et al. (2007). The primers used for probable TOPK-recognised sequences are detailed in Supplementary Table 5. The PCR program consisted of 38 cycles of amplification for 20 s at 94°C, 30 s at 58°C, and 30 s at 72°C. An earlier denaturing step of 3 min at 94°C and a final extension step of 2 min were added.

**In vivo studies**

Four to five-week old female NOD/SCID mice (Charles River, Barcelona, Spain) were used, following the Spanish and European Union guidelines (RD 1201/05 and 86/609/CEE, respectively). The study was approved earlier by the Bioethics Committee of our institution (CB-A4). Cell suspensions, containing 5 × 10⁶ alive cells in 0.2 ml of 1:1 cellular medium (Matrigel Matrix (BD), were injected s.c. into the right flank of the mice. Cells were counted using a Neubauer chamber (VWR) and cellular viability was checked by trypan blue staining (Sigma). Mice were randomised into three controls (TC71wt, early mock, and late mock) and two treated groups (early and late shRNAi clone). Tumours were measured every 5 days with a caliper, and the diameters were recorded. Tumour volume was calculated as described earlier (Martins et al., 2008) by the formula \( a^2 b/6 \), in which \( a \) is the smallest diameter and \( b \) the biggest one. Mice were killed by anaesthesia overdosing 4 weeks after the cells injection, and tumours were collected for histopathology analysis.

**Statistics**

For in vivo studies, one-way ANOVA for independent samples was performed using the SPSS 15.0 software (SPSS Inc., Chicago, IL, USA) and mice with a tumour volume higher than 2.5 cm³ were excluded from the analyses. For comparisons between shRNAi and mock (early and late stages), we computed two-sided t-tests of independent samples for each condition.

**RESULTS**

**shRNAi permitted a sustained knockdown of EWS–FLI1**

To analyse the effects of a stable suppression of EWS–FLI1 expression in its native context, we used a pSUPER vector that directs intracellular synthesis of siRNA-like transcripts. Two different and specific interfering shRNAi (named 7-6 and 7-6 M, respectively), both targeting the fusion point of type 1 EWS–FLI1 mRNA isoform (7-6), were tested in the Ewing sarcoma cell line, TC71 (Figure 1A). The loss-of-function phenotype was assessed in selected clones in early and late stages (eight cellular passages) to confirm the maintenance of the EWS–FLI1 ‘knockdown’, although a drop in the interference effect was detected (Figure 1B). Wild-type FLI1 was detected only in the multiple myeloma cell line MM1S, and its expression was not affected by the EWS–FLI1 shRNAi constructs. Expression of wild-type EWS protein was not changed by EWS–FLI1 RNAi, confirming the specificity of our shRNAi designs (Figure 1C). The effect of RNA interference was also measured at the mRNA level in the selected shRNAi clones resulting in 42–54% inhibition (P < 0.01), and also at later passages (P < 0.05) (Figure 1D). These results confirmed the validity and the stable nature of the shRNAi model through the course of cellular passages.

**Stable reduction of EWS–FLI1 prompted an increase in the apoptotic fraction, and a reduction of the migratory capacity and tumourigenicity**

Up to 40 different TC71 shRNAi clones were obtained. We selected the one, shRNAi clone 6, showing the most pronounced decrease in EWS–FLI1 expression, to further characterise and evaluate its molecular and cellular phenotype.

We observed a marked increase in the apoptotic index of up to 70% in both, early and late stages. Western blotting analysis confirmed the processing of caspase 9 in the interfered clone in both stages, although a slight activation could also be seen in the mock sample, which seemed insufficient for promoting apoptosis (Figure 2A). PARP was also processed in the shRNAi clone, confirming caspase cascade activation (data not shown).

To determine whether EWS–FLI1 knockdown affects the ability of TC71 cells to migrate, we performed a transwell experiment. The migratory capacity of shRNAi cells was found to be impaired, with around a 30% decrease in both stages (Figure 2B).

The tumourigenicity of the shRNAi clone was tested using a soft agar growth assay. The shRNAi cells were largely restricted in their tumourigenicity, giving rise to fewer and smaller colonies with a decrease of around 35–40% (P < 0.0003) compared with the mock, which was maintained through the course of cellular passages (Figure 2C). Similar results were obtained with the shRNAi clone M2 (data not shown).

Analogous results regarding cell-cycle distribution and apoptotic index were obtained using other shRNAi clones in which interference of EWS–FLI1 expression was less dramatic. There was a direct relationship between the extent of interference of the fusion transcript and the degree of change observed, thus confirming that the effects appearing after EWS–FLI1 inhibition were due to its specific suppression of expression and not due to unspecific clonal ‘off-target’ effects (Figure 2D).

The growth rate of these interfered cells was reduced, as measured by MTT assay, to around 30% in the early passage (P < 0.05). However, this inhibitory effect was lost in the late stage (Supplementary Figure 1A). Assessment of the cell-cycle pattern of the shRNAi clone showed no changes in cell-cycle distribution (Supplementary Figure 1B).

**EWS–FLI1 interference reduced tumour growth**

We injected in immunodeficient mice cells from the shRNAi clone, mock, and TC71wt, both in the early and late stages. Tumours appeared at the same time in all the animals, but mice harbouring cells from the shRNAi clone in both temporal stages developed tumours with a marked volume reduction (Figures 3A and B). Most of the animals injected with the early and late shRNAi clone cells developed tumours of < 0.5 and 0.3 cm³, respectively, with no tumour reaching 1 cm³ in any case. However, tumours in control
groups (injected with wild-type, early mock cells, and late mock cells) were markedly larger, most of them close to 1 cm³ or even larger (P < 0.05). Moreover, five animals (two from the wild-type, two from the early mock, and one from the late mock) developed tumours between 2.5 and 4.9 cm³ and were not included in the statistical analyses.

**Figure 1**  Targeting of EWS–FLI1 through shRNAi for stable protein and mRNA knockdown. (A) Diagram of wild-type EWS, wild-type FLI1, and EWS–FLI1 fusion type 1. The illustration of EWS–FLI1 cDNA and the location of the two shRNAi designs used, 7-6 and 7-6 M, both against fusion break-point. (B) Analysis by western blotting showed that the EWS–FLI1 expression was reduced in several TC71 shRNAi clones. The EWS–FLI1 reduction is maintained partially in some shRNAi clones through the course of cellular passages (early and late stages). EWS–FLI1 68 kDa, FLI1 51 kDa, and actin 42 kDa. 3, 4, 5, 6: TC71 shRNAi clones transfected with shRNAi design type 7-6; M2: TC71 shRNAi clone transfected with shRNAi design type 7-6 M. MM1S: multiple myeloma cell line. Positive control for FLI1. NB: neuroblastoma cell line SK-N-JD. Negative control for EWS–FLI1. Actin is shown as a loading control. (C) Analysis by western blotting showed that the EWS–FLI1 shRNAi designs are specific and did not alter the EWS expression in the early and late stages. The shRNAi clone corresponds to the TC71 shRNAi clone 6. Actin is shown as a loading control. (D) The shRNAi reduction in the EWS–FLI1 mRNA level assessed through qRT–PCR (SYBR green probes) in the early (T0) and late passage (T8). The shRNAi clone corresponds to the TC71 shRNAi clone 6. GAPDH was used as an internal control. Columns, mean of triplicates of three different replicates; bars, s.d.

**EWS–FLI1 interference downregulated IGF-1 and sensitised Ewing sarcoma cells to inhibitors of the IGF-1/IGF-1R signal transduction pathway**

We next analysed the effects of EWS–FLI1 interference on the gene expression signature of the TC71 Ewing sarcoma cells. RNA
from the shRNAi clone, mock, and TC71wt from the early and late stages was isolated and used to perform duplicate hybridisation experiments with the Affymetrix HG-U133A platform. EWS–FLI1 activated and repressed genes in similar numbers. Several of the genes identified were previously reported to be repressed (Supplementary Table 1) or induced (Supplementary Tables 2 and 3) by EWS–FLI1, including p57/Kip2, NXXK2.2, and DAX1-NR0B1. Their expression was validated, confirming the original microarray data (Supplementary Figures 2A–C, respectively).

As IGF-1 was pointed as a relevant downregulated gene, we decided to focus on it. Western blot and qRT–PCR analyses confirmed IGF-1 downregulation (P < 0.05) in the shRNAi clone

Figure 2 The long-term reduction in EWS–FLI1 provoked an increase in the apoptotic fraction, reduced migration capacity, and oncogenic transformation. There is a direct relationship between the extent of the interference of the fusion transcript and the degree of change observed. (A) Induction of apoptosis by reduction in the expression of EWS–FLI1 using specific shRNAi in the TC71 Ewing sarcoma cell line in an early and late stage. The shRNAi clone corresponds to the TC71 shRNAi clone 6. The cells were seeded on 24 well plates, 24 h after apoptotic index was determined by FACS using an annexin V detection kit assay. The means ± standard deviations (error bars) of four independent experiments are shown. FACS Apoptotic Analysis Figures show the variances cell populations. A, alive fraction; EA, early apoptosis; LAN, late apoptosis/necrosis; N, necrosis. X axis: FL3-H Annexin V; y axis: FL1-H PI. Panel below shows the activation of Caspase 9 in the shRNAi clone in both stages by western Blot. (B) The shRNAi clone cells were less able to migrate through the matrix of Transwell in both stages, early and late. The shRNAi clone corresponds to the TC71 shRNAi clone 6. A total of 100,000 alive cells were counted using a Neubauer chamber and seeded over the transwell in 24 well-format plates. After 16 h, cells of the upper part of the transwell were scrapped, cells migrated to the bottom part of the transwell were fixed, stained with blue violet solution and discoulered with acetic acid. Absorbance was measured at 570 nm. Photographs of migrated cells taken with a Nikon 10 x objective. Columns, mean of replicates of three different experiments; bars, s.d. (C) The diminished expression of EWS–FLI impaired the number and size of colonies in the shRNAi clone. The shRNAi clone corresponds to the TC71 shRNAi clone 6. Colonies were photographed after 2 weeks. Experiments were performed three times with several replicates. (D) The EWS–FLI1 shRNAi extension determined the degree of the changes that were not because of unspecific clonal 'off-target' effects. Analysis by western blotting showed that the EWS–FLI1 expression was reduced in the early stage in both shRNAi clones, 5 and M2, although only shRNAi clone M2 maintained partially EWS–FLI1 reduction in the late stage. Actin is shown as a loading control. The increase in the apoptotic index was directly correlated to the degree of EWS–FLI1 interference. The shRNAi clone number 5 showed no change in the apoptotic index compared with shRNAi clone M2 that had a more pronounced inhibition of EWS–FLI1 and showed a increase in its apoptotic fraction. The cells were seeded on 24 well plates, 24 h after apoptotic index was determined by FACS using an Annexin V detection kit assay. The means ± standard deviations (error bars) of four independent experiments are shown. Columns, mean of triplicates of three separate replicates; bars, s.d.
during the course of the cellular passages compared with TC71wt and mock cells, although a decrease in IGF-1 expression could be detected in the late stage of the mock because of unknown reasons (Figures 4A and B). No change in IGF-1R expression was observed by WB (Figure 4B) or IHC (data not shown). We hypothesised that the shRNAi clone would be more sensitive to inhibitors of this signal transduction pathway. NVP-AEW541 is an inhibitor of the IGF-1 pathway (Garcia-Echeverría et al., 2004). PD98059 and LY294002 inhibit MEK and PI3-K, respectively, which are key genes in the IGF-1/IGF-1R pathway. We studied the influence of NVP-AEW541, PD98059, and LY294002 administration on AKT, p-AKT, MAPK42/44, and p-MAPK 42/44 levels by WB, and evaluated the induction of apoptosis and repression of proliferation. Knockdown of the EWS–FLI1 fusion protein rendered the shRNAi clone more sensitive to the action of NVP-AEW541, PD98059, and LY294002 administration on AKT, p-AKT, MAPK42/44, and p-MAPK 42/44 levels by WB, and evaluated the induction of apoptosis and repression of proliferation.

TOPK is a new target of EWS–FLI1 involved in proliferation

We focused our attention on a new potential target gene of EWS–FLI1, namely TOPK/PBK (T-lymphokine-activated killer cell–originated protein kinase). Our gene expression analyses indicated TOPK as a target gene downregulated as a consequence of EWS–FLI1 inhibition (score value /C0 6.78 in Genetrix analysis). qRT–PCR and western blot analysis confirmed a decrease in TOPK at the mRNA (P < 0.05) (Figure 5A) and protein levels (Figure 5A) which was more intense in the early stage. These results pointed out TOPK as a candidate EWS–FLI1 target gene. To confirm this fact we looked for EWS–FLI1-binding sites in the TOPK proximal promoter and introns using the bioinformatics program TFM-explorer. Several consensus binding sites were found confirming TOPK as a putative interacting target of the fusion gene (TOPK proximal promoter location (−37:−248), P-value = 2.32e−02; TOPK intron1 location (−2197:−2740), P-value = 7.23e−03). Immunoprecipitation of chromatin with anti-EWS antibody from mock yielded amplification of the TOPK intron 1 selected region (Figure 5B). Immunoprecipitation with IgG or with anti-RNA polymerase II resulted in no amplification of the TOPK promoter proximal region, intron 1 and intron 4.

Figure 3 Tumour growth evolution. The EWS–FLI1 shRNAi reduced tumoural growth in vivo. (A) NOD-Scid mice were s.c. injected with 5 × 10^6 TC71 cells. The treated mice (early and late shRNAi clone) showed smaller tumours than the control groups (TC71wt, late and early mock) since the third week after cells injection. At the end of the study these differences were statistically significant (P < 0.05). (B) Visual and histopathological evaluation of mice tumours. All of the tumours showed the same histopathological pattern: a large area of necrotic tissue, an area of proliferating cells, and a layer between them with cells in apoptosis.
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We inhibited TOPK expression in TC71wt cells using siRNA oligos. Seventy-two hours after transfection a reduction in both TOPK RNA and protein levels was observed (Figure 5C).

A noteworthy effect of TOPK inhibition was a mild decrease in proliferation rate as measured by MTT (Figure 5D) compared with TC71 wt Mock shRNAi clone. The shRNAi clone corresponds to the TC71 shRNAi clone 6. Figure 4 EWS–FLI1 shRNAi impaired the IGF-1/IGF-1R survival pathway and its downstream targets. (A) Histogram representation of IGF-1 transcript level knockdown in the shRNAi clone as assessed by qRT–PCR normalised to GAPDH. The shRNAi clone corresponds to the TC71 shRNAi clone 6. SYBR probes were used. All experiments were performed in triplicate in the early and late stages. Column, mean of triplicates of three different experiments; bars, s.d. (B) The IGF-1 protein level is reduced in the shRNAi clone as assessed through western blot whereas the IGF-1R protein level is not changed. The shRNAi clone corresponds to the TC71 shRNAi clone 6. (C) IC50 of proliferation measured by MTT assay after the NVP-AEW541, LY294002, or PD98059 treatment (72 h of incubation). The shRNAi clone showed more sensitivity to the action of inhibitors of the IGF-1/IGF-1R pathway being statistically significant for AEW571 and PD98059 (P<0.05). The shRNAi clone corresponds to the TC71 shRNAi clone 6. (D) Apoptotic index after the NVP-AEW541, LY294002, or PD98059 treatment (72 h of incubation). Apoptosis could be further induced in the shRNAi clone under all conditions. The means ± standard deviations (error bars) of four independent experiments are shown. The shRNAi clone corresponds to the TC71 shRNAi clone 6. P<0.05 values are considered as significant. (E) Effects of NVP-AEW541 combined with inhibitors LY294002 and PD98059 on the activation of the IGF-1/IGF-1R signalling pathway. The AKT and MAPK42/44 phosphorylation were diminished in the shRNAi clone under IGF-1 stimulation whereas TC71wt and mock cells maintained the active phosphorylation status in both kinases. All conditions were treated with AEW571 during 15 min and with the inhibitors for 2 h, before IGF-1 stimulation (50 ng ml–1) during 15 min (serum-free conditions). The shRNAi clone corresponds to the TC71 shRNAi clone 6.

Figure 4

We inhibited TOPK expression in TC71wt cells using siRNA oligos. Seventy-two hours after transfection a reduction in both TOPK RNA and protein levels was observed (Figure 5C).
TC71wt and negative siRNA control. A dramatic change in the growth pattern was detected after knocking down TOPK. TOPK siRNA cells grew in clumps and were not able to merge as compared with TC71wt cells (Figure 5D). The decrease in TOPK levels did not affect the cell-cycle pattern and did not cause an increase in apoptotic rate (data not shown). Transfection of EWS–FLI1 type 1 mRNA in the shRNAi clone restored EWS–FLI1 levels; TOPK protein expression was also significantly restored (Figure 5C). In addition, AKT levels also decreased, thus pointing to a possible connection between both protein kinases (Figure 5E). When the expression of TOPK was diminished an increase in the transcript level of p27/Kip1 was detected (Figure 5E). TOPK expression could be detected through qRT–PCR in nine Ewing sarcoma cell lines analysed regardless of the type of EWS–ETS fusion (Figure 5F).

DISCUSSION

Ewing sarcoma fusion proteins act as aberrant transcription factors by interfering with different signalling pathways (Prieur et al, 2004). Identification of EWS–FLI1 targets represents a key aspect in the understanding of the molecular behaviour of Ewing sarcoma. RNA interference has been shown to be a powerful tool that allows targeting chimeric oncoproteins and gaining a deeper knowledge of tumour biology in its own context (Sharp, 1999; Tijsserman et al, 2002).

Ectopic expression models for the study of gene fusions are disadvantaged as they do not account for the true cellular context. A wide variety of effects have been documented after EWS–FLI1 induction in heterologous models, ranging from dedifferentiation to growth arrest (Kovar, 2005).

shRNAi studies to suppress EWS–FLI1 in the Ewing sarcoma cells have been carried out using different systems (Siligian et al, 2005; Kinsey et al, 2006; Owen and Lessnick, 2006; Carrillo et al, 2007; Stegmaier et al, 2007). Our shRNAi approach, based on the pSUPER vector, targets at the junction point of EWS–FLI1 type 1 fusion transcripts in a reliable model, the Ewing sarcoma cell line, TC71. The effects due to EWS–FLI1 interference were observed in different clones and were related to the degree of fusion inhibition, indicating a direct and specific correlation. We reported a significant reduction in EWS–FLI1 expression, which was maintained through several cellular passages, confirming the validity of our approach, although a partial loss of RNA interference was detected through the course of cellular passages as also described by other group (Smith et al, 2006) in the Ewing sarcoma cell line A673.

The main consequence observed in our model because of the silencing of EWS–FLI1 was a strong induction of apoptosis, correlating with caspase 9 pathway activation and PARP processing. EWS–FLI1 interference has been documented to increase remarkably the number of apoptotic cells (Prieur et al, 2004). A decrease in the anchorage-independent capacity to grow has also been reported (Smith et al, 2006). The ability of cells to grow in semisolid media was also affected in our shRNAi model. Disruption of EWS–FLI1 provoked the appearance of fewer and smaller colonies mainly in the early stage. EWS–FLI1 reduction also impaired migration capacity. CCN3 (NOV) has been described as an important mediator of migration and invasion (Benini et al, 2005) and its expression was downregulated in our model; this could explain loss of function of interfered cells although it is not possible to rule out the contribution of early apoptosis to that observation.

NOD/SCID mice injected with shRNAi cells in both temporal stages developed tumours with a significant reduction in tumour volume with respect to mock and TC71wt. shRNAi cells had restricted their tumourigenicity and migratory capacity, which could explain the tumour growth reduction observed. Tumoural growth reduction in nude xenograft models has been described in several studies either using an shRNAi model (Smith et al, 2006) or through the inhibition of EWS–FLI1 targets such as IGF-1/IGF-1R (Scotlandi et al, 1998), VEGF (Guan et al, 2005), NR0B1/DAX1 (Kinsey et al, 2006), and cholecytokinin (Carrillo et al, 2007).

The strong decrease in IGF-1 gene expression observed after EWS–FLI1 silencing was analysed. The IGF-1/IGF-1R signal transduction pathway is over-expressed in many sarcomas (Sekyi-Otu et al, 1995) and has great relevance for the biology of Ewing sarcoma (Scotlandi et al, 1996). Blocking IGF-1/IGF-1R has shown its effectiveness in restricting tumour growth (Martin et al, 2006). Our shRNAi clone showed an increased sensitivity to the action of IGF-1/IGF-1R pathway inhibitors NVP-AEW541, LY294002, and PD98059, and a reduction in the phosphorylation levels of AKT and MAPK 42/44. IGF-1R protein status is not affected, and therefore the effects observed were due to IGF-1 downregulation. This observation underlines the fact that IGF-1/IGF-1R signalling activation and EWS–FLI1 fusion are two key molecular features in Ewing sarcoma and suggests the need for blocking both of them in order to achieve therapeutic benefits.

Understanding the oncogenic properties of EWS–FLI1 requires identification of the downstream targets. The aim of our gene expression analysis was to identify potentially regulatory genes, whose transcript levels change in response to oncoprotein inhibition. Well-known EWS–FLI1 targets were among the genes identified, including NR0B1/DAX1, ryzin, p57/Kip2, and NKK2.2 (Dauphinot et al, 2001; Amsellem et al, 2005; Mendiola et al, 2006; Smith et al, 2006), confirming the validation of our stable shRNAi model.

We have mainly focused our attention on a new target gene of EWS–FLI1, TOPK/PBK. TOPK, T-lymphokine-activated killer cell-originated protein kinase, is a 36-kDa protein located on 8p21.2 belonging to the Ser/Thr protein kinase family. It was cloned as a new MAPK-like protein kinase, highly conserved among vertebrates, specifically expressed in the testis, placenta, activated lymphoid cells, and neural lineages (Abe et al, 2000; Dougherty et al, 2005). TOPK has been involved in haematological malignancies (Simons-Evelyn et al, 2001; Nandi et al, 2004), in breast cancer (Park et al, 2006), in preventing apoptosis in a melanoma cell line (Zykova et al, 2006), as a proliferative agent related to DNA damage response through p38 MAPK, c-Jun-NH2-kinase, and H-Ras signalling (Ayllon and O'Connor, 2007; Oh et al, 2007), or as an enhancer of Cdk1/cyclin B1-dependent phosphorylation of PRC1 and promoter of cytokinesis (Abe et al, 2007). Our gene expression analysis pointed out TOPK as a target gene downregulated because of EWS–FLI1 inhibition. TOPK reduction at protein and mRNA levels was confirmed through WB and qRT–PCR, respectively, and was maintained during cellular passages.

Using an in silico approach, we analysed the TOPK promoter and introns in order to find EWS–FLI1 binding sites. EWS–FLI1 and some ETS family members, such as wild-type FLI1, require a 9-bp consensus sequence harbouring a GGAA ‘core’. A 9-bp sequence, GAAGGAAGT, was found in the TOPK intron 1, which showed limited similarity to the high-affinity ETS-binding consensus (ACCGGAAGT) (Gangwal and Lessnick, 2008). It has been shown in promyelocytic leukaemia cells that the transcriptional control of TOPK promoter is mostly because of binding of transcription factors E2F and CREB/ATF to two distinct binding sites within it (Nandi and Rapoport, 2006). As a proliferative agent related to DNA damage response through p38 MAPK, c-Jun-NH2-kinase, and H-Ras signalling (Ayllon and O'Connor, 2007; Oh et al, 2007), or as an enhancer of Cdk1/cyclin B1-dependent phosphorylation of PRC1 and promoter of cytokinesis (Abe et al, 2007). Our gene expression analysis pointed out TOPK as a target gene downregulated because of EWS–FLI1 inhibition. TOPK reduction at protein and mRNA levels was confirmed through WB and qRT–PCR, respectively, and was maintained during cellular passages.
inhibition; TOPK-interfered cells seemed to be disabled in their ability to reach confluence compared with TC71wt cells. The same observation was described earlier when TOPK was knocked down using siRNA oligos in the prostate carcinoma cell line, DU145 (Ayllon and O’Connor, 2007). The main consequence observed because of TOPK inhibition was a 35% reduction in the proliferation rate. The results were concordant with the significant suppression of cell growth caused by TOPK interference in human breast cancer and colorectal cell lines (Park et al., 2006; Zhu et al., 2007). Surprisingly, TOPK silencing raised AKT levels. When EWS–FLI1 expression was restored in the shRNAi clone, the TOPK level also recovered and AKT level decreased to the level observed in TC71wt, thus proving that AKT increase was due to the TOPK knockdown and not due to EWS–FLI1 interference. AKT is a relevant gene in the Ewing sarcoma biology that is involved in proliferation and survival, and the therapeutical benefit of its inhibition is clear (Martins et al., 2006); further research to clarify the relationship between TOPK and AKT needs to be carried out. Neither cell-cycle pattern nor apoptotic index was modified after TOPK silencing, as was the case in the MCF-10A breast epithelial cell line (Ayllon and O’Connor, 2007), although the levels of p27/Kip1, a gene described in Ewing sarcoma by its expression level inversely correlated with that of EWS–FLI1 (Matsunobu et al., 2006), were increased.

In this study, we have developed a stable RNAi model system. Our shRNAi model maintained the EWS–FLI1 knockdown
through the course of the cellular passages, showing that it is a valid one for assessing the long-term effects of EWS–FLI1 interference. EWS–FLI1 interference caused a decrease in IGF-1 levels and rendered cells more sensitive to the action of inhibitors of the IGF-1/IGF-1R pathway, pointing to the importance of this survival pathway and the necessity to block it for therapeutic benefits. Our model could be a valuable tool for confirming targets already described and for discovering new ones such as TOPK, a kinase involved in proliferation and motility. This is the first time that TOPK expression is described in Ewing sarcoma, and the changes observed after TOPK interference were concurrent with those observed in other cellular models, confirming the relevant role of this gene in the Ewing sarcoma biology.

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