Abstract. In order to reduce side effects of survivin-inhibiting anticancer therapies, we determined the expression of the survivin transcripts survivin-wild-type (survivin-wt), survivin-ΔEx3 (ΔEx3) and survivin-2B (2B) in cryo-preserved tumor and non-malignant bladder tissues (18 tumor and 22 non-malignant samples, including 17 autologous tissue pairs) by quantitative PCR. Furthermore, we investigated the biological effects following specific inhibition of the alternative transcripts ΔEx3 and 2B in bladder cancer (BCa) cells. In BCa and non-malignant bladder tissues survivin-wt was the quantitatively dominant transcript followed by ΔEx3 and 2B. The mean mRNA expression of ΔEx3 (0.37 vs. 0.06 zmol/amol GAPDH, respectively) and 2B (0.13 vs. 0.01 zmol/amol GAPDH, respectively) was significantly higher in BCa compared to non-malignant bladder tissues, indicating their accessibility for an expression inhibition in BCa cells. Effective and long-lasting small interfering RNA-mediated inhibition of one alternative survivin transcript caused lower cell growth reduction effects (apoptosis induction, cell cycle arrest, colony formation) compared to simultaneous inhibition of multiple survivin transcripts including survivin-wt. Inhibition of one alternative survivin transcript increased the apoptosis rate by 11% vs. 33-46% when reducing several survivin transcripts. We observed no G2/M arrest or reduction of cell colony formation after inhibiting one alternative survivin transcript. Reduction of cell viability by the chemotherapeutics cisplatin, mitomycin C or gemcitabine was stronger in combination with inhibition of several survivin transcripts than in combination with the reduction of one alternative survivin splice variant. Furthermore, reducing one alternative transcript caused chemosensitization to only one chemotherapeutic agent in contrast to inhibition of several survivin transcripts. Therefore, the alternative survivin transcripts ΔEx3 and 2B do not represent reasonable targets for anticancer, at least BCa, treatment.

Introduction

Survivin (baculoviral IAP repeat-containing protein 5, BIRC5), a potent member of the inhibitor of apoptosis (IAP) gene family, mediates several antiapoptotic functions (1-5). Survivin is also involved in cell cycle progression; as a member of the chromosomal passenger complex it is indispensable to the correct course of cytokinesis (1,2,5-7). As the fourth most common transcript that is overexpressed in numerous tumor entities, survivin is one of the most attractive targets in cancer therapy (1). In many cancers, survivin expression is associated with disease progression and poor prognosis (8-11). Survivin is also involved in cell cycle progression; as a member of the chromosomal passenger complex it is indispensable to the correct course of cytokinesis (1,2,5-7). As the fourth most common transcript that is overexpressed in numerous tumor entities, survivin is one of the most attractive targets in cancer therapy (1). In many cancers, survivin expression is associated with disease progression and poor prognosis (8-11). Survivin is also involved in the development of resistance to apoptosis-inducing agents (5,8). In bladder cancer (BCa), which is, with almost 61500 estimated new cases (12) and 13000 estimated tumor-related deaths in 2006 in the USA (12), one of the most common malignancies, increased survivin mRNA (13) and protein expression (14,15) is associated with increased tumor grade and stage. Furthermore, survivin mRNA detection in urine is a specific marker for BCa (13,16). Like other tumors, BCa develops resistance to conventional therapies, such as chemotherapy (17). Since overexpression of apoptosis inhibitors, such as survivin, is involved in this process, their knock-down presents a useful tool for new anticancer interventions (17,18).

Although the role of survivin-wt is widely understood, little is known about the function and impact of its alternative splice variants, especially in BCa. Until now, four alternatively spliced transcripts of survivin have been described (5,9,19,20).
Two of them, ΔEx3 and 2B, are expressed in different tumor entities (9,21,22). ΔEx3 mRNA lacks exon 3 and has a frame-shift within exon 4, resulting in a 15.6-kDa protein (Fig. 1) (19). An in-frame insertion of a cryptic exon 2B between exons 2 and 3 is found in 2B mRNA, resulting in an 18.6-kDa protein (Fig. 1) (19,23). Different structures of the survivin protein isoforms are responsible for different functions and cellular localization patterns (5,24). The function of alternative survivin splice variants is not yet completely understood. Expression of ΔEx3 decreased apoptosis rates in HepG2 hepatocellular carcinoma cells after methotrexate treatment, whereas the apoptosis rate was increased by 2B expression (19). These findings suggested an antiapoptotic function for ΔEx3 and a non-antiapoptotic or even proapoptotic one for 2B. Therefore, 2B might represent a natural antagonist of anti-apoptotically acting survivin isoforms (19). Heterodimers of survivin-wt and its alternative splice variants are supposed to modulate cellular localization of the wild-type protein, such as localization of the complexes to mitochondria (24). Furthermore, co-expression of survivin-wt and ΔEx3 offered synergistic apoptosis-inhibiting effects in the Daoy medulloblastoma cell line (24). If the function of survivin-wt depends, at least in part, on these heterodimerizations, then inhibition of alternative splice variants may provide a useful therapeutic approach to apoptosis induction. Moreover, since different functions of survivin isoforms may result in different roles in tumor progression, their specific inhibition might cause antiproliferative effects.

Current survivin-inhibiting strategies, such as using small interfering RNAs (siRNAs) or antisense oligodeoxynucleotides (AS-ODN), target different survivin transcripts simultaneously (including survivin-wt). This resulted in remarkable antiproliferative effects, sensitization to apoptosis-inducing treatments as well as tumor growth reduction \textit{in vitro} (8,25,26) and \textit{in vivo} (8,27,28). siRNA- or AS-ODN-mediated survivin knock-down in different BCa cell lines caused increased apoptosis rates and cell cycle arrest, reduced viability and clonogenic survival as well as chemosensitization and radiosensitization (29-31). These observations supported survivin's suitability to be a therapeutic target in anti-BCa treatment.

Nevertheless, survivin expression during the G2/M phase of the cell cycle in normal cells (25) indicates its importance in normal cellular physiology. Furthermore, there is growing evidence showing involvement of survivin in hematopoietic cell maturation (32). Therefore, inhibition of survivin might result in undesirable effects on normal cells. The knock-down of one specific survivin transcript may provide a tool for more selective therapeutic approaches.

Therefore, the aim of this study was to investigate therapeutic efficiency of the specific knock-down of one alternative survivin transcript in comparison to inhibiting different survivin transcripts simultaneously by siRNAs in a suitable \textit{in vitro} BCa model.

\section*{Materials and methods}

\textbf{Tissue samples.} Tumor and non-malignant bladder tissue samples were obtained from radical cystectomy specimens of BCa patients. Twenty-four autologous, paraffin-embedded BCa tissue pairs (1 pT1, 2 pT2, 16 pT3, 5 pT4; all high-grade) were used for immunohistochemical survivin staining. In addition, 18 BCa (1 pT1, 2 pT2, 10 pT3, 5 pT4; all high-grade) and 22 non-malignant cryo-preserved bladder tissue samples (thereof 17 autologous paired samples) of altogether 23 BCa patients were used for mRNA expression analyses. Representative hemalaun and eosin (H&E)-stained sections (4 μm) of each sample were examined by a pathologist to ensure, that the tumor cell count was ≥50% for malignant and ≤5% for non-malignant tissue samples.

\textbf{Cell culture, siRNA and chemotherapeutic treatments.} All cell lines were cultivated under standard conditions (37°C, humidified atmosphere containing 5% CO\textsubscript{2}) in DMEM (Invitrogen, Karlsruhe, Germany), supplemented with 1% non-essential amino acids, 10% fetal calf serum and 1% HEPES. The cell line EJ28 was selected as a human BCa model since it showed a remarkable expression of survivin-wt and its alternative splice variants.

Three days after seeding in appropriate culture plates, cells were transfected with a mixture consisting of 200 nM siRNA (Table I and Fig. 1) and DOTAP (ratio 1:3 w/w; Roche Molecular Biochemicals, Mannheim, Germany), each diluted in serum-free OptiMEM (Invitrogen). This transfection mixture was incubated at 37°C for 4 h, cells were washed with PBS and cultivated in fresh culture medium until further analyses. Anti-survivin-siRNAs were synthesized by Eurogentec (Seraing, Belgium), non-silencing (ns)-siRNA by Qiagen (Hilden, Germany). ns-siRNA-treated cells served as a control for all examinations.

For chemosensitization experiments, 24 h after siRNA transfection chemotherapy drugs (diluted in culture medium) were added at different concentrations, chosen in pre-tests. Mitomycin C (MMC; 0.3 or 0.9 μg/ml) was incubated for 2 h, cisplatin (CDDP; 0.7 or 2.1 μg/ml) and gemcitabine (GEM; 2.5 ng/ml) were incubated for 24 h. Afterwards cells were washed with PBS and cultivated until viability measurements.

\section*{Expression analyses}

\textbf{Immunohistochemistry.} Paraffin sections of autologous BCa tissue pairs (4 μm) were deparaffinized and rehydrated. Endogenous peroxidase activity was blocked with hydrogen peroxide (1.35% in methanol, 10 min) followed by sample pulping (0.01 M citrate buffer, pH 6.0, 15 min, 121°C). After

\begin{table}
\centering
\begin{tabular}{|c|c|c|c|}
\hline
\textbf{WT} & \textbf{exon 1} & \textbf{exon 2} & \textbf{exon 3} & \textbf{exon 4} \\
\hline
\textbf{ΔEx3} & \textbf{exon 1} & \textbf{exon 2} & \textbf{exon 3} & \textbf{exon 4} \\
\hline
\textbf{2B} & \textbf{exon 1} & \textbf{exon 2} & \textbf{exon 3} & \textbf{exon 4} \\
\hline
\end{tabular}
\end{table}

\begin{figure}
\centering
\includegraphics[width=\textwidth]{Figure1.png}
\caption{Figure 1. Scheme of the siRNA target sites within the transcripts survivin-wt, ΔEx3 and 2B.}
\end{figure}
Table I. siRNA constructs, their target mRNAs and sequences of their sense (s) and antisense (as) strands.

<table>
<thead>
<tr>
<th>siRNA construct</th>
<th>Survivin target-mRNA</th>
<th>Target sequence (5’-3’)</th>
</tr>
</thead>
<tbody>
<tr>
<td>siSVV-Bel307 (38)</td>
<td>wt (nt 307–325), 2B (nt 307–325), ΔEx3 (nt 332–350)</td>
<td>GGA GCCU GGA AGG CUG GGA Ctdt as CUC CCC UGC UUC CAG CUC Ctdt</td>
</tr>
<tr>
<td>siSVV284 (30)</td>
<td>wt (nt 358-376), 2B (nt 427-445)</td>
<td>GCA UUC GUC CGG UUG CGC Utdt as AGC GCA ACC GGA AUG Ctdt</td>
</tr>
<tr>
<td>siSVV-D3-332</td>
<td>ΔEx3 (nt 332–350)</td>
<td>GAC GAC CCC AUG CAA AGG Adtdt as UCC UUU GCA UGG GGU CGU Ctdt</td>
</tr>
<tr>
<td>siSVV-2B-405</td>
<td>2B (nt 405–423)</td>
<td>UCA CGA GAG AGG AAC AUA Adtdt as UUA UGU UCC UCU CUC GUG Adtdt</td>
</tr>
<tr>
<td>ns-siRNA</td>
<td>No complementary nucleotide sequence to human mRNAs</td>
<td>UUC UUC GAA CGU GUC ACG Utdt as ACG UGA CAC GUU CGG AGA Adtdt</td>
</tr>
</tbody>
</table>

Data in brackets represent the complementary single-strand motif within the corresponding mRNA sequence (nt, nucleotide position). mRNA sequences were taken from the NCBI database: survivin-wt nm_001168, ΔEx3 nm_001012270 and 2B nm_001012271. siSVV-D3-332 and siSVV-2B-405 were designed by M.K. (Kappler et al., unpublished data). Because of its 100% complementary to human EPR1 (effector cell peptidase receptor 1) mRNA, application of a survivin-wt specific siRNA was not possible.

PBS washing samples were incubated for 1 h with a polyclonal anti-survivin-wt antibody (1:2000; NB 500-201; Novus Biologicals, Littleton, USA) and subsequently stained using Vectastain HRP Kit (Vector, Burlingame, USA) following the manufacturer’s instructions. Staining reaction with 1 mg/ml diaminobenzidine (Dako, Hamburg, Germany) in PBS (+0.02% hydrogen peroxide) was controlled microscopically followed by counterstaining of nuclei with hemalaun and dehydration. For semiquantitative analyses only nuclear survivin staining was examined. Each stained nucleus was classified as survivin-positive, regardless of staining intensity. Survivin staining was evaluated by scoring the percentage of positively stained nuclei in every tissue sample: 1 (0-19%), 2 (20-49%), 3 (50-69%), 4 (70-89%), 5 (90-100%). Three visual fields per sample (magnification x100) were analyzed microscopically and then averaged. These mean values were used for calculating median and mean score of the whole sample collective.

Western blot analyses. Cells (25000 cells per sample) were lysed and proteins were separated on a 12% SDS polyacrylamide gel and blotted onto a nitrocellulose membrane. Immunostaining was performed according to standard protocols, using bovine serum albumin (BSA) diluted in PBS with 0.1% Tween-20 for blocking membrane (5% BSA), dissolving antibodies (0.5% BSA) and washing steps (0.5% BSA). The following antibodies were used: anti-survivin-wt (1:1000; NB 500-201; Novus Biologicals), anti-2B (1:500; ab3729; Abcam, Cambridge, UK), anti-ΔEx3 (1:1000; ab3731; Abcam), HRP-linked anti-mouse immunoglobulin (1:1000; P 0260; Dako) and HRP-linked anti-rabbit immunoglobulin (1:1000; P 0217; Dako). Detection was carried out using the Enhanced Chemiluminescence (ECL) Kit (Amersham, Freiburg, Germany). As a control for equal loading, β-actin was detected by a monoclonal antibody (1:10000; A 5316; Sigma, St. Louis, MO, USA). Protein levels were quantified by densitometry using Intas Gelscan software (Goettingen, Germany) and survivin protein levels were normalized to those of β-actin.

Quantitative real-time PCR. RNA from cells and tissue samples was isolated using standard protocols (Invitrogen, AlphaSpin Cell/Tissue RNA Mini Kit; Invitrogen). RNA from cells and tissue samples was isolated using standard protocols (Invitrogen, AlphaSpin Cell/Tissue RNA Mini Kit; Invitrogen). For further analyses, RNA was reversely transcribed into cDNA (SuperScript™ II RNase H-Reverse Transcriptase Kit; Invitrogen). Transcript levels of the different survivin forms and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) were quantified by quantitative real-time PCR (Roboscreen, Leipzig, Germany). Survivin expression values were calculated in zmol survivin per amol GAPDH. Tissue samples were excluded from analyses of a transcript, if the mean deviation within a dependently repeated quantification of that transcript exceeded 35%.

Biological effects of siRNA treatment

Viability. Cellular viability was examined using cell proliferation reagent WST-1 (Roche Molecular Biochemicals) following the manufacturer’s instructions. For further analyses, cells were washed with PBS and cultivated in fresh medium. Viables were determined 72 and 96 h after siRNA transfection. Viales of siRNA mono-treatments and siRNA/chemotherapeutic combination treatments were normalized to those of ns-siRNA. Viabilities following chemotherapeutic mono-treatments were normalized to those of untreated control.

Colony formation. Cells were harvested 48 h after siRNA transfection, 100 cells were seeded in 6-well plates (3 wells per sample), cultivated and Giemsa stained after 10-12 days of cultivation. Macroscopically visible colonies were counted. Colony formation for each sample was calculated as the mean value of the three examined wells.
Apoptosis and cell cycle. Apoptosis rates were assessed using Annexin V-FITC Apoptosis Detection Kit and cell cycle analyses were performed using Cycle Test Plus DNA Reagent Kit (both BD Biosciences, Heidelberg, Germany) according to the manufacturer’s instructions.

Statistical analyses. Statistical analyses were performed using an unpaired two-sided (homoscedastic) Student’s t-test. Statistical significance was determined at p<0.05.

Results

Survivin expression in human bladder tissue samples

Protein expression. In immunohistochemical analyses both cytoplasmatic and nuclear survivin-wt was detected. Cells of tumor samples showed a staining of the whole nucleus, whereas in non-malignant cells a punctured nuclear staining pattern was noted. All 24 examined BCa samples showed a higher number of stained nuclei and a higher staining intensity compared to autologous non-malignant bladder tissues (Fig. 2). For BCa tissue staining we determined a mean score of 4.8 and a median of 5.0. For corresponding non-malignant bladder tissues the mean score was 2.8 and the median was 2.7. No association was observed between the percentage of nuclear survivin staining and tumor stage.

mRNA expression. Proportional distribution of survivin transcripts ranged from 74-100% survivin-wt, 0-16% ΔEx3 and 0-10% 2B in BCa tissues. In non-malignant bladder tissue the samples’ proportional distribution was 82-100% survivin-wt, 0-18% ΔEx3 and 0-4% 2B. In all examined tumor and non-malignant tissue samples, we observed the following expression patterns: survivin-wt>ΔEx3>2B (when transcripts were detected). In tumor samples survivin-wt was detected in 12/17 cases (71%) in a range of 0.17 to 9.14 zmol/amol GAPDH (Fig. 3). Non-malignant bladder tissue samples showed a survivin-wt expression in 16/21 cases (76%) in a range of 0.07 to 6.05 zmol/amol GAPDH (Fig. 3). ΔEx3 was detected in 8/15 (53%; 0.18-1.41 zmol/amol GAPDH) BCa samples and 5/17 (29%; 0.04-0.68 zmol/amol GAPDH) non-
malignant samples (Fig. 3). We detected 2B in 7/16 (44%; 0.07-0.68 zmol/amol GAPDH) tumor samples and 1/22 (5%; 0.29 zmol/amol GAPDH) non-malignant bladder tissues (Fig. 3). The mean expression was significantly higher in malignant than in non-malignant bladder tissues for all quantified survivin transcripts (Fig. 3). Mean expression of survivin-wt was 3.27 zmol/amol GAPDH in BCa and 0.73 zmol/amol GAPDH in non-malignant bladder tissues. ΔEx3 showed a mean expression of 0.37 zmol/amol GAPDH in tumor and of 0.06 zmol/amol GAPDH in non-malignant samples. 2B was detected with a mean expression of 0.13 zmol/amol GAPDH in BCa and 0.01 zmol/amol GAPDH in non-malignant tissues. Within autologous tissue samples survivin-wt was overexpressed in tumor in 10/17 cases (59%), ΔEx3 in 6/12 cases (50%) and 2B in 7/15 cases (47%).

In vitro inhibition of survivin transcripts and its biological effects
Survivin mRNA expression in BCa cell lines. All tested BCa cell lines expressed survivin-wt and its alternative splice variants ΔEx3 and 2B (Fig. 4). Proportional distribution of these transcripts in the cell lines was similar to that in human BCa tissue samples. ΔEx3 showed a mean expression of 0.37 zmol/amol GAPDH in tumor and of 0.06 zmol/amol GAPDH in non-malignant bladder tissues. 2B was detected with a mean expression of 0.13 zmol/amol GAPDH in BCa and 0.01 zmol/amol GAPDH in non-malignant samples. Within autologous tissue samples survivin-wt was overexpressed in tumor in 10/17 cases (59%), ΔEx3 in 6/12 cases (50%) and 2B in 7/15 cases (47%).

siRNA/chemo-therapeutics combination treatments. Combination treatments of siRNAs and different concentrations of CDDP, MMC or GEM showed statistically significant chemosensitization effects mainly by using siSVV-Bel307 and siSVV284 (Fig. 5). The strongest viability reduction to 52% of that of ns-siRNA/chemotherapeutic therapy was observed for siSVV284/CDDP (2.1 μg/ml) treatment. In contrast, the strongest effect caused by the specific inhibition of an alternative survivin transcript was a viability reduction to 83% of that of the corresponding control (siSVV-2B-405/GEM treatment). Furthermore, siSVV-Bel307 and siSVV284 significantly sensitized EJ28 cells to different chemotherapeutics, in contrast to siSVV-D3-332 and siSVV-2B-405.

Table II. siRNA-mediated knock-down of survivin transcripts survivin-wt, ΔEx3 and 2B in EJ28 cells and their effects on cell proliferation.

<table>
<thead>
<tr>
<th>siRNA construct</th>
<th>mRNA expression</th>
<th>Protein expression</th>
<th>G2/M</th>
<th>Colony formation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>wt</td>
<td>ΔEx3</td>
<td>2B</td>
<td>wt</td>
</tr>
<tr>
<td>siSVV284</td>
<td>-79</td>
<td>+38</td>
<td>-70</td>
<td>-78</td>
</tr>
<tr>
<td>siSVV-D3-332</td>
<td>-16</td>
<td>-26</td>
<td>+43</td>
<td>+10</td>
</tr>
<tr>
<td>siSVV-2B-405</td>
<td>+23</td>
<td>+73</td>
<td>-100</td>
<td>+68</td>
</tr>
</tbody>
</table>

All parameters were measured 48 h (~24 h) after the start of transfection. Data are normalized to those of ns-siRNA-treated cells and declared in % as differences to ns-siRNA-treated control.
in tumor events and cancer therapy is widely unknown. In
order to reduce side effects on normal cells, it appears
reasonable to inhibit only one specific survivin splice variant
instead of multiple ones. In the present study we investigated,
if this approach could provide a suitable therapeutic strategy.

Our investigations showed an overexpression of survivin-
w t at both the mRNA and protein levels in malignant compared
to non-malignant tissue samples obtained from radical cystec-
tomy specimens of BCa patients as found in previous analyses
(1,13,14). So far, expression of alternative survivin transcripts
ΔEx3 and 2B has been characterized in a few tumor entities,
including breast cancer (22,33,34), renal cell carcinoma (35),
medulloblastoma (36), soft tissue sarcomas (37) and colorectal
carcinoma (21). We therefore focussed on these two alternative
splice variants, whereas the other known survivin variants
survivin-3B and survivin-2α were not analyzed (5,9). As
described for the mentioned tumor entities, we found an
expression of ΔEx3 and 2B mRNAs in BCa as well as in
non-malignant bladder tissue samples. Survivin-wt was the
dominant survivin transcript expressed in the examined tissue
samples (in tumor 74-100% of total survivin), whereas ΔEx3
and 2B were detected at significantly lower levels (Fig. 3).

Mean expression of all investigated survivin transcripts
was significantly higher in tumors than in non-malignant
bladder tissues (Fig. 3). Therefore, besides survivin-wt also
its alternative transcripts ΔEx3 and 2B should be accessible
for an siRNA-mediated expression inhibition in BCa. Higher
expression of 2B in tumor samples compared to non-malignant
bladder tissue did not necessarily show an association with
tumor progression. Ryan et al (33) also investigated an over-
expression of 2B in breast cancer compared to normal breast
tissue, but found no relation between 2B expression and
apoptosis. The detection of survivin-wt, ΔEx3 and 2B mRNAs
in non-malignant bladder tissues can be explained by survivin
expression in the G2/M phase of strongly proliferating non-
malignant cells in the urothelium. Since most of the included
BCa samples were histologically classified as pT3/pT4 G3,
we are not able to report an association of survivin splice
variant expression with stage or grade of BCa.

Before performing survivin inhibition experiments, we first
looked for a suitable in vitro model of BCa by investigating
expression of survivin-wt, ΔEx3 and 2B in seven human BCa
cell lines. All examined cell lines expressed these transcripts.
Of these, survivin-wt was the quantitatively dominant transcript
(85-90% dependent on cell line), whereas ΔEx3 (5.8-8.8%) and
2B (3.5-7.7%) were expressed at lower levels. Since EJ28
cells showed the highest expression of all examined transcripts
and a similar proportional transcript distribution as BCa tissues,
they were used as a model system for in vitro survivin knock-
down.

For transfection experiments we used siRNAs targeting
different survivin transcripts simultaneously (siSVV-Bel307,
siSVV-2B84) and also siRNAs specific for one alternative
survivin splice variant (siSVV-D3-332, siSVV-2B-405) and
investigated the biological effects of these treatments in two
independent experiment series. We observed a selective,
effective and long-lasting mRNA reduction to 38% by siSVV-
Bel307 and to 19% by siSVV2B84, resulting in remarkable
antiproliferative effects (Table II and Fig. 5). These findings
and the magnitude of the observed effects confirm previously
published data (30,38-41). Belyanskaya et al (38) reported
a survivin mRNA and protein reduction by 70 and 50%,
Furthermore, when only one alternative survivin transcript was not remarkably change protein levels of survivin isoforms. That inhibition of alternative splice variant mRNAs does detection time (48 h after transfection). It is also supposable, protein isoform. We observed a protein reduction only for only mRNA reduction, but also reduction of the corresponding specific survivin knock-down.

In the present study, we investigated for the first time the effects on cell proliferation caused by specific inhibition of the alternative survivin transcripts ΔEx3 (siSVV-D3-332) and 2B (siSVV-2B-405). Similar to siSVV-Bel307 and siSVV284, we showed a selective reduction of the target mRNA 2B by siSVV-2B-405. siSVV-D3-332 offered an unspecific action by inhibition of survivin-wt (down to 84%) in addition to that of ΔEx3 24 h after the start of transfection (Table II). This construct hybridizes to 10 bases within exon 2 and 11 bases within exon 4, all of which are also present in survivin-wt. This might have been sufficient for inhibiting survivin-wt.

Despite a selective, effective and long-lasting reduction of 2B mRNA (down-regulation of 2B to 0% 24 h after siRNA treatment (Table II) was probably caused by an effective knock-down of this transcript below the detection limit), siSVV-2B-405 treatment resulted only in low antiproliferative effects in EJ28 cells. These observations support the theory of reduced antiapoptotic potential of 2B in BCa cells, as it was described for HepG2 hepatocellular carcinoma cells (19). Proapoptotic action of 2B was also confirmed by its decreased expression in higher tumor grades of colorectal carcinoma (21) and renal cell carcinoma (35), but contradicted by a correlation of 2B expression with poorer prognosis in adult acute myeloid leukaemia (42) as well as by the absence of a relation between 2B expression and apoptosis in breast carcinoma (33). Therefore, the role of survivin splice variants in apoptosis and tumorigenesis or tumor progression has to be further investigated and might even depend on tumor entity or cell line.

siSVV-D3-332 caused lower mRNA inhibition effects compared to the other used constructs (Table II). This finding might explain lower biological effects following treatment with this construct. Formation of secondary structures in the target mRNA could lead to hindrance in siRNA hybridization and therefore, could be responsible for limited mRNA reduction. Specific inhibition of ΔEx3 is possible only at the transition site between exons 2 and 4 (Fig. 1). Therefore, no other siRNA constructs can be designed performing ΔEx3-specific survivin knock-down.

A pre-requisite for an effective target knock-down is not only mRNA reduction, but also reduction of the corresponding protein isoform. We observed a protein reduction only for survivin-wt; an inhibition was not demonstrated for ΔEx3 or 2B. Explanations include the possibility that lifetimes of ΔEx3 protein and of 2B protein were not yet expired at detection time (48 h after transfection). It is also supposable, that inhibition of alternative splice variant mRNAs does not remarkably change protein levels of survivin isoforms. Furthermore, when only one alternative survivin transcript was knocked-down, other transcripts achieved higher expression compared to control (ns-siRNA treated) cells. This might be caused by a mechanism regulating expression of the different survivin transcripts together as described for other alternatively spliced transcripts (43). Absence of protein reduction or mRNA up-regulation could be responsible for the low effects observed following siSVV-D3-332 and siSVV-2B-405 treatments.

However, mRNA inhibition of one specific alternative survivin splice variant (ΔEx3 or 2B) did not result in remarkable antiproliferative effects. These treatments offered less effects in apoptosis induction, cell cycle arrest, reduction of long-term cell proliferation and chemosensitization than treatments with siSVV-Bel307 or siSVV284. These findings were confirmed by the observation that siSVV-2B-405 and siSVV-D3-332 did not radiosensitize human sarcoma cells as effectively as siSVV284 (Kappler et al, unpublished data).

In conclusion, either a simultaneous inhibition of different survivin transcripts including survivin-wt or a reduction of survivin-wt alone was necessary for strong antiproliferative effects caused by siSVV-Bel307 and siSVV284. Since survivin-wt expression is associated with the course of disease, the latter is more plausible. We have shown that the specific inhibition of one alternative survivin transcript is not reasonable for anticancer treatment, at least in BCa therapy.

Acknowledgements

The authors would like to express their gratitude to Katja Robel, Andrea Lohse, Jana Herrmann and Antje Zobjack for their excellent technical assistance, and to Dr Woei-Yun Siow for critically reviewing the manuscript. This study was kindly supported by a grant of the Dr Robert Pfleger-Stiftung (to S.F. and A.M.). M.K. was supported by a grant of Saxony-Anhalt (FKZ: 3584/1104M).

References


