Atelocollagen-mediated *in vivo* siRNA transfection in ovarian carcinoma is influenced by tumor site, siRNA target and administration route

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Abstract. Ovarian cancer is the leading cause of death from gynecological malignancies worldwide, and innate or acquired chemoresistance of ovarian cancer cells is the major cause of therapeutic failure. It has been demonstrated that the concomitant inhibition of Bcl-x_L and Mcl-1 anti-apoptotic activities is able to trigger apoptosis in chemoresistant ovarian cancer cells. In this context, siRNA-mediated Bcl-x_L and Mcl-1 inhibition constitutes an appealing strategy by which to eliminate chemoresistant cancer cells. However, the safest and most efficient way to vectorize siRNAs *in vivo* is still under debate. In the present study, using *in vivo* bioluminescence imaging, we evaluated the interest of atelocollagen to

vectorize siRNAs by intraperitoneal (i.p.) or intravenous (i.v.) administration in 2 xenografted ovarian cancer models (peritoneal carcinomatosis and subcutaneous tumors in nude mice). Whereas i.p. administration of atelocollagen-vectorized siRNA in the peritoneal carcinomatosis model did not induce any gene downregulation, a 70% transient downregulation of luciferase expression was achieved after i.v. injection of atelocollagen-vectorized siRNA in the subcutaneous (s.c.) model. However, the use of siRNA targeting Bcl-x_L or Mcl-1 did not induce target-specific downregulation *in vivo* in nude mice. Our results therefore show that atelocollagen complex formulation, the administration route, tumor site and the identity of the siRNA target influence the efficiency of atelocollagen-mediated siRNA delivery.

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Abbreviations: HES, hematoxylin, eosin, safran; s.c., subcutaneous; i.p., intra-peritoneal; i.v., intravenous; ASO, antisense oligonucleotide

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Introduction

Ovarian carcinoma therapeutic care remains inefficient owing to the high level of chemoresistance to conventional treatment. With approximately 145,000 new cases each year in developed countries and approximately 100,000 deaths, ovarian carcinoma is the first leading cause of death from gynecological malignancies.

We previously showed that apoptosis control defects are responsible for the strong protection of ovarian cancer cells against cell death (1,2). In particular, we demonstrated that the concomitant inhibition of the anti-apoptotic proteins Bcl-x_L and Mcl-1 either by siRNAs (3,4), or by combining a BH3-mimetic molecule targeting Bcl-x_L with a siRNA or a BH3-mimetic molecule targeting Mcl-1 (5-8) triggers massive apoptotic cell death in chemoresistant ovarian cancer cells in vitro, even in the absence of chemotherapeutic treatment.

This effect is due to the 'primed for death' feature of cancer cells (9). Unlike normal cells, cancer cells contain a high level of the activated apoptosis inducers Bax and Bak, or activator BH3-only proteins such as Bim and Puma, which are sequestered by anti-apoptotic proteins such as Bcl-x_L or Mcl-1 (10). Strategies aimed at the release of Bax/Bak or Bim/Puma from their inhibitors are thus highly attractive in a clinical perspective. In this context, the design of innovative BH3-mimetic molecules able to target Bcl-x_L *in vivo*, such as ABT-263 (Navitoclax), opens up new perspectives for clinical validation of such approaches. However, pharmacological inhibition of Mcl-1 *in vivo* remains problematic despite recent promising advances (8,11-13).

The versatility in the choice of siRNA targets as well as their high efficiency for the downregulation of a specific gene makes the perspective of clinically available siRNA strategies highly attractive. Recent advances in chemistry are now allowing a panel of chemical modifications in siRNAs that avoid unwanted immunostimulatory and off-target effects, thus overcoming some of the major issues preventing the safe and efficient use of siRNAs in clinical practice (14). However, efficient vectorization and delivery of siRNAs *in vivo* remains a major challenge (15), despite the huge amount of research carried out on this topic.

Among the numerous strategies developed for the *in vivo* vectorization of siRNAs (16), atelocollagen constitutes an interesting option. Atelocollagen is a derivative of collagen that has been used for the *in vivo* transfection of nucleic acids such as DNA, antisense oligonucleotides (ASO) and siRNA (17). It is processed from bovine dermal collagen after digestion of pepsine-mediated telopeptides to avoid immunogenicity (18). Atelocollagen is a highly biocompatible and biodegradable compound used in numerous medicinal, surgical and cosmetic applications.

A molecular model has been proposed for atelocollagen/siRNA interactions in which the interaction between atelocollagen, which is positively charged in physiological conditions (19), and the siRNA phosphates promotes the formation of a fibrillar structure containing 5 atelocollagen molecules for 1 siRNA duplex (20). This structure efficiently protects the siRNA from nucleases and is compatible with cellular uptake. In addition, modification of the atelocollagen concentration during complex formation modifies the viscosity of the final product. Low concentration formulations lead to a viscosity close to that of blood and are adapted to intravenous administration. Conversely, higher atelocollagen concentrations lead to a gel-like formation, allowing the continuous delivery of siRNA over an extended period of time after local administration. This plasticity of formulation enables the use of atelocollagen for various administration routes (intravenous, intraperitoneal, intramuscular, intratumoral, peritumoral) (19,21). Atelocollagen has thus been proposed as an adequate vector for the in vivo local or systemic siRNA administration in various tumor models (17,21-24).

Several studies exploring the antitumor effects of atelocollagen-mediated siRNA administration have utilized either intratumoral (24-29) or peritumoral (18,22,30-32) administration. The only study describing atelocollagen delivery of siRNA in a subcutaneous ovarian cancer model used peritumoral administration (32). Using a vasohibin 2 siRNA to target angiogenesis, the authors reported a decrease in tumor growth, tumor dissemination and angiogenesis. However, intratumoral or peritumoral administration is not suitable for treating ovarian carcinoma, in which peritoneal carcinomatosis is sometimes composed of hundreds of disseminated tumor nodes.

The poor overall survival of ovarian carcinoma patients is mainly due to disease recurrence within the peritoneal cavity (33). We thus decided to explore the possibility of siRNA delivery in peritoneal carcinomatosis. In this context, intraperitoneal (i.p.) administration is an attractive option, particularly since the small tumor nodes lack vascularization. Furthermore, local treatment carries a reduced risk of systemic toxicity and could retain some of the properties of peritumoral injections, at least next to the injection site and possibly throughout the peritoneal compartment. Atelocollagen delivery of siRNA via the i.p. route has been used in several studies (33,34), providing an interesting proof-of-concept.

Numerous studies with *in vivo* administration of siRNA-vectorized by atelocollagen involve intravenous (i.v.) administration (24,35,36). It is more feasible in routine clinical use due to the reduced risk for patients compared to i.p. administration as the use of catheters for repeated treatments can lead to serious side-effects (34). Moreover, i.v.-delivered particles likely reach 'in-depth' larger vascularized tumor nodes more easily. The opportunity to deliver siRNAs via systemic administration would make it suitable for several pathologies, including cancer, provided that the chosen site of delivery is vascularized. In the context of ovarian carcinoma, this would mean that peritoneal tumor nodes of only a few millimeters in size would be amenable to siRNA delivery.

As both i.p. and i.v. routes present significant advantages in the context of ovarian carcinoma, we decided to explore the efficiency of atelocollagen-vectorized siRNA delivery in two independent models: a peritoneal carcinomatosis model submitted to i.p. siRNA/vector complex injection, and an s.c. vascularized tumor node model submitted to i.v. siRNA/vector complex injection. To this end, we first used longitudinal imaging studies performed on luciferase-expressing SKOV3 tumors to define optimal administration modalities in both models. Our results evidenced the interest of i.v. administration in an s.c. model of tumor development. We next studied these administration modalities for the siRNA-mediated silencing of Bcl-x_L and Mcl-1 in the same animal models.

Materials and methods

Cell culture and treatment. SKOV3 ovarian carcinoma cell line and its luciferase-expressing counterpart SKOV3-Luc were obtained from the American Type Culture Collection (ATCC; Manassas, VA, USA). They were grown in RPMI-1640 medium supplemented with 2 mM GlutamaxTM, 25 μ M HEPES, 10% fetal calf serum and 33 mM sodium bicarbonate (Fisher Scientific Bioblock, Illkirch, France) and were maintained in a 5% CO₂ humidified atmosphere at 37°C.

In vitro transfection. All siRNAs used in the present study were chemically synthesized and PAGE-purified by Eurogentec (Liège, Belgium) and received as annealed oligonucleotides. siRNA guide sequences targeting the indicated genes were as follows: Mcl-1 siRNA (noted siMcl1), 5'-uguuuagccacaaaggcac-3', Bcl-x_L siRNA (noted siBcl-xL), 5'-ugcgauccgacucaccaau-3', luciferase siRNA (noted siLuc), 5'-ucgaaguacucagcguaag-3' and GFP siRNA (noted siGFP, negative control), 5'-acuuguggccguuuacguc-3'. Exponentially growing cells were seeded (2.5x10⁵ cells/25 cm² flask) the day before transfection to reach 30-50% confluency at the time of transfection. Briefly, the transfecting reagent INTERFERin™ (Polyplus Transfection, Strasbourg, France) was added to siRNA diluted in OptiMEM® serum-free medium (Life Technologies, Saint Aubin, France), and the formation of siRNA/vector complexes was allowed to proceed for 15 min at RT before distribution in culture flasks. The next day, cell medium was changed to remove the remaining transfecting reagent. At the indicated time, cells were trypsinized and washed with cold phosphate-buffered saline (PBS). Cell pellets were used directly or stored at -80°C for later use.

In vitro bioluminescence measurement. Firefly luciferase activity was assayed 72 h after siLuc transfection with the Dual-Luciferase Reporter Assay System (Promega) and measured with a luminometer Centro LB 960 (Berthold, Thoiry, France). Each assay contained 3 technical replicates.

Animal experiments. Female nude mice (Janvier Laboratories, Saint Berthevin, France) 6-weeks of age were injected with human ovarian carcinoma cell lines (SKOV3-Luc or SKOV3) i.p. $(20 \times 10^6 \text{ cells in } 200 \,\mu\text{l PBS})$ or s.c. bilaterally on the flanks $(5x10^6 \text{ cells in } 200 \ \mu\text{l PBS})$. The animals were kept under pathogen-free conditions and fed and watered ad libitum, in cages of 4 to 5 animals (in compliance with recommended area surface/animal), in a dedicated room with a 12h/12h light/dark cycle at a constant temperature of 22°C. A period of 4 weeks was allowed for tumor development before the beginning of the experiments. In vivo experiments were performed in the animal core facilities of the Comprehensive Cancer Centre F. Baclesse in Caen and of the Institute of Advanced Biosciences in Grenoble, both of which are certified by the French Ministry of Higher Education and Research and regularly controlled by the Departmental Management of Population Protection (certification nos. B14 118 003 and C38 516 10001, respectively). Animal experiments were performed according to the current regulations in the respect of animal experiment ethics. Animals were euthanatized at the end points of the experiments, i.e. after a total of 5 weeks of tumoral development. At this stage, s.c. tumors were ~4 mm in size, 33.5 mm³ (V = L x 12 x π /6), whereas for micro-carcinomatosis, nodules were just becoming visible on the peritoneum with the presence of a few nodules (usually 1-5) ~10 mm³ in size in the peritoneal cavity. At this stage no ascites was noted. For euthanasia, animals were anesthetized with a gaseous mix of isoflurane/oxygen (5/95) and thereafter euthanatized with a gaseous mix of oxygen/carbon dioxide (20/80).

In vivo treatments. For local administration, $100 \mu g$ of siRNA complexed with atelocollagen at a final 0.5% concentration was injected i.p. in $400 \mu l$. For systemic administration, $125 \mu g$ of siRNA complexed with atelocollagen at a final concentration of 0.05% was injected i.v. in $200 \mu l$. NaCl injections for control animals were performed with the same volumes.

Local (0.5% Atelocollagen) or systemic (0.05% Atelocollagen) administration kits were purchased from Cosmobio (Koken Co., Ltd., Tokyo, Japan). siRNA/vector complexes were prepared according to the manufacturer's protocol for local use kit, and with a modified protocol, i.e. increase in siRNA concentration in the formulation for the systemic kit, according to the protocol published by Mu *et al* (24). In experiments involving siMcl1 and siBcl-x_L, animals were distributed into 3 subgroups prior to treatment according to the size of the s.c. tumor, i.e. large, medium and small. We thus avoided any possible bias due to unequal average tumor size between groups. Mice from these subgroups were then equally distributed into 4 experimental groups of 5 mice each for a total of 20 mice: vehicle (NaCl), control siRNA (siGFP), Mcl-1 siRNA (siMcl1) and Bcl-x_L siRNA (siBcl-x_L).

Bioluminescence imaging. In vivo imaging was conducted at Optimal (Grenoble, France), a core facility for small animal optical imaging. After 4 weeks of tumor development, mice received an i.p. injection of 150 µg/g luciferin (Promega, Charbonnières-les-Bains, France) for non-invasive bioluminescence imaging (IVIS Kinetic; PerkinElmer, Waltham, MA, USA). Semi-quantitative data of luciferase-positive tumor cell signals were obtained using the manufacturer's software (Living Image; PerkinElmer). Results were expressed as photons/second (photons/s). For animal randomization prior to treatment, mice were separated into 3 subgroups with high, medium and low levels of bioluminescent signals. Mice from each of these subgroups were then distributed into 3 experimental groups for a total of 20 mice: vehicle (NaCl, 4 and 3 mice bearing i.p. or s.c. tumors respectively), control siRNA (siGFP, 3 and 3 mice bearing i.p. or s.c. tumors, respectively) and luciferase siRNA (siLuc, 4 and 3 mice bearing i.p. or s.c. tumors, respectively) treatment. Bioluminescence imaging was performed at day 0, 3 and 7. Results were then expressed as values relative to day 0.

Statistical analysis. In vivo luciferase assays were compared using the non-parametrical Wilcoxon rank-sum test, and p-values <0.05 were considered as significant. Statistical analyses were calculated using the 32-bit R Console software V3.1.0 (R Foundation for Statistical Computing, Vienna, Austria).

Immunohistochemical analysis. Automated immunohistochemistry using a Ventana Discovery XT autostainer was performed on 4 µm-thick paraffin sections. Slides were deparaffinized with EZPrep buffer and epitopes were unmasked by 15 min of high-temperature treatment in CC1 EDTA buffer. Sections were incubated for 40 min at 37°C with an anti Mcl-1 (ab32087; Abcam, Cambridge, MA, USA) or Bcl-x₁ antibody (556361; BD Pharmingen, Franklin Lakes, NJ, USA). Secondary antibody (Omnimap Rabbit; Ventana Medical System Inc., Tucson, AZ, USA) was incubated for 16 min at room temperature. Immunodetection performed without the primary antibody was used as the control. After washes, the staining was performed with DAB and sections were counterstained with hematoxylin using Ventana reagents according to the manufacturer's protocol. Stained slides were then digitized using an Aperio ScanScope slide scanner (Aperio Technologies, Vista, CA, USA).

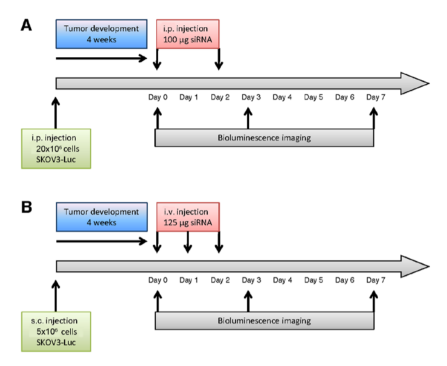


Figure 1. Schedules for *in vivo* siRNA treatments. (A) Mice were injected i.p. with $20x10^6$ SKOV3-Luc cells and tumors were allowed to develop for 4 weeks. Bioluminescence values were acquired at day 0 prior to treatment, day 3 and 7. Atelocollagen (0.5% final) in complex with 100 μ g siRNA was injected i.p. at day 0 and 2. (B) Mice were injected bilaterally s.c. with $5x10^6$ SKOV3-Luc cells in each flank and tumors were allowed to develop for 4 weeks. Bioluminescence values were acquired at day 0 prior to treatment, day 3 and 7. Atelocollagen (0.05% final) in complex with 125μ g siRNA was injected i.p. at days 0, 1 and 2.

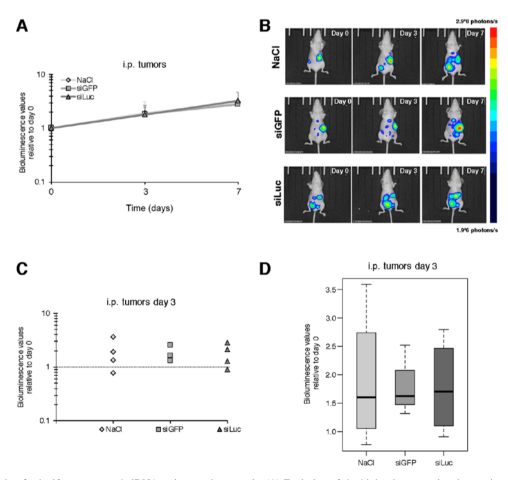


Figure 2. i.p. injection for luciferase-targeted siRNA to i.p. carcinomatosis. (A) Evolution of the bioluminescent signal over time for NaCl- (4 mice), siGFP- (3 mice) and siLuc- (4 mice)-treated groups. Mean values \pm SD for each group of animals relative to bioluminescence values acquired at day 0 for each individual. (B) Representative bioluminescence image for one animal from each group. (C) Bioluminescence values at day 3 for individual mice from each group relative to bioluminescence values acquired at day 0. (D) Box plot of bioluminescence values of each group of animals measured at day 3 relative to bioluminescence values acquired at day 0. No significant difference between groups.

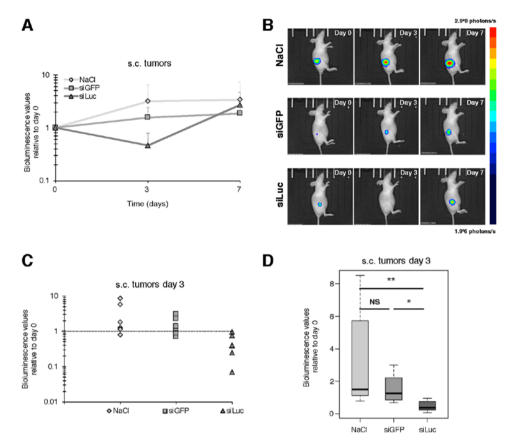


Figure 3. i.v. injection for luciferase-targeted siRNA to subcutaneous tumors. (A) Evolution of the bioluminescence signal over time for NaCl- (3 mice), siGFP- (3 mice) and siLuc (3 mice)-treated groups. Mean values \pm SD for tumors from each group of animals relative to bioluminescence values acquired at day 0 for each tumor. (B) Representative bioluminescence image for one animal from each group. (C) Bioluminescence values at day 3 for individual tumors from each group relative to bioluminescence values acquired at day 0. (D) Box plot of bioluminescence values of each group of animals measured at day 3 relative to bioluminescence values acquired at day 0. The siLuc-treated group was significantly different from the NaCl- (p=0.004) and siGFP (p=0.015)-treated groups (Wilcoxon rank-sum test).

RNA isolation and quantitative reverse transcriptase-PCR (RT-PCR). Total RNA was isolated from the SKOV3 cell line using TRIzol (Qiagen, Courtaboeuf, France) and 1 μ g was reverse-transcribed using Omniscript reverse transcriptase (Qiagen) with random hexamers according to the manufacturer's instructions. cDNA was combined with 10 µmol/l of each forward and reverse primer, 50 µmol/l of the TaqMan® probe and 2X TaqMan® Fast Universal PCR Master Mix (Applied Biosystems, Foster City, CA, USA) in a 20 μl final reaction volume. The following probes were used: for Mcl-1 (Hs00172036_m1), for Bcl-x_I (forward, 5'-TGCGTGGAAAGC GTAGACAA-3' and reverse, 5'-AGGTAAGTGGCCATCCAA GCT-3'; probe, 5'-AGATGCAGGTATTGGTG-3') and GAPDH (Hs9999905_m1). All PCR amplification reactions were carried out in triplicate on an Applied ABI Prism 7500 Fast PCR system (Applied Biosystems). The Mcl-1 and Bcl-x₁ transcripts were quantified relative to GAPDH endogenous reference by the comparative $2^{-\Delta\Delta Ct}$ method and results are expressed as a percentage of untransfected tumors from NaCl-injected mice.

Results

Nude mice were xenografted with SKOV3 ovarian cancer cells constitutively expressing luciferase (SKOV3-Luc) by i.p. or bilateral s.c. injection to induce the development of peritoneal carcinomatosis or s.c. tumors, respectively. The use of *in vivo*

bioluminescence imaging allowed individual longitudinal follow-up of tumor development for each animal. Therefore, by measuring the evolution of bioluminescence signals over time, each tumor constitutes its own control, thus limiting the heterogeneity of response due to inter-animal variation in the rate of tumor development.

Atelocollagen-vectorized siRNA targeting luciferase injected by i.p. route does not decrease bioluminescence in carcinomatosis tumor nodes. In the peritoneal carcinomatosis model, atelocollagen was complexed with siRNA to a 'high' final concentration of atelocollagen (0.5%) to obtain a gel-like compound, according to the manufacturer's protocol. This gel-like formulation was used to allow the continuous release of siRNA in the peritoneal cavity. siRNA/vector complexes or vehicle were injected i.p. at day 0 and day 2. Bioluminescence was measured at day 0 (before treatment), day 3 and day 7. Treatment and the imaging schedules are represented in Fig. 1A.

As expected, an increase in the bioluminescence signal was detected over time from day 0 to 3, and day 7 in the control tumors, corresponding to tumor growth. However, siLuc-treated animals did not display any decrease in bioluminescent signal (178 and 325% of the signal at day 3 and day 7, respectively, relative to day 0) compared to siGFP (182 and 278%) or vehicle (189 and 302%) (Fig. 2 and data not shown).

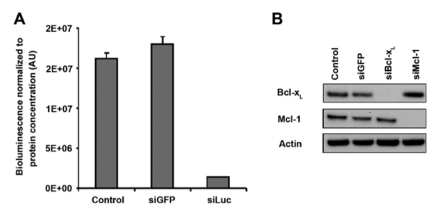


Figure 4. *In vitro* efficiency of the siRNA batches used *in vivo*. (A) Luciferase-targeted siRNAs strongly decrease target expression *in vitro*. Transfection was performed with siLuc in SKOV3-Luc cells with final 20 nM of siRNAs, and target downregulation analyzed 72 h after transfection. Bioluminescent activity was measured in cell lysates. Results are relative to protein concentration in respective cell lysates ± SD. (B) Both Bcl-x_L- and Mcl-1-targeted siRNAs mediate strong downregulation *in vitro* on their respective targets. Transfection was performed with siGFP, siMcl1 or siBcl-x_L in SKOV3 cells with final 20 nM of siRNAs, and target downregulation analyzed 72 h after transfection. Mcl-1 and Bcl-x_L protein expression was assessed by western blotting. Actin was used as a loading control.

Atelocollagen-vectorized siRNA targeting luciferase injected i.v. decreases bioluminescence in s.c. tumors. For the s.c. tumor model, atelocollagen was complexed with siRNA to a 'low' final concentration of atelocollagen (0.05%), so that the siRNA/vector complexes presented a viscosity comparable to blood. SiRNAs - or vehicle - were injected i.v. at day 0, 1 and 2, and bioluminescence was assessed at day 0 (before treatment), day 3 and 7, as described in Fig. 1B. An increase of bioluminescence over time was observed for NaCl and siGFP, consistent with the expected growth of tumors over time (Fig. 3A and B). However, at day 3 the bioluminescence values were significantly lower in siLuc compared to day 0 (47% of day 0). This decrease was even stronger compared to the bioluminescence values of NaCl (319% of day 0) with p=0.004, and vs. siGFP (155% of day 0) with p=0.015 (Fig. 3C and D), which means a decrease in bioluminescence signal of 85 and 70% relative to NaCl and siGFP respectively. In contrast, there was no significant difference at day 7 between groups, with siLuc bioluminescence signal intensity catching up with other groups, supporting that the decrease in bioluminescence intensity observed at day 3 was not the consequence of impaired tumor growth and/or tumor mass reduction for siLuc group (Fig. 3 and data not shown).

Unlike luciferase-directed siRNA, Mcl-1- and Bcl-x_t-directed siRNAs vectorized with atelocollagen do not induce target downregulation in vivo. After defining the successful conditions for in vivo siRNA vectorization with atelocollagen, we sought to downregulate the cancer-related targets of interest, Mcl-1 and Bcl-x_L. We used siRNA sequences which had been validated by our group in previous studies for their ability to downregulate their targets specifically with high efficiency (3,4,6). In addition, the batches of siMcl1 and siBcl-x_L that we used in vivo were checked for their ability to downregulate their respective targets in vitro in SKOV3 cells. The complete disappearance of the protein-specific band was observed in western blotting (Fig. 4A), demonstrating that these siRNAs enable full silencing of their respective targets. These results were comparable to what was obtained upon siLuc transfection in SKOV3-Luc cells, which triggered 89% luminescence inhibition compared to the control conditions (Fig. 4B).

SKOV3 cells were injected into nude mice to develop bilateral s.c. tumors and the above-described protocol and treatment schedule (Fig. 1B) were repeated, with 3 exceptions; i) siLuc condition was removed and replaced with siMcl1 or siBcl-x_L; ii) we discarded the day 7 analysis time point as it did not lead to a decrease in bioluminescence in the above-described experiment; and iii) we used 5 mice/group with a total of 20 mice. Unfortunately, neither RT-qPCR analysis performed on tumor lysates nor IHC performed *in situ* on tumor slices revealed any Mcl-1 or Bcl-x_L downregulation at day 3 after the beginning of the protocol (Fig. 5).

Discussion

Our group has previously shown that the concomitant inhibition of $\operatorname{Bcl-x_L}$ and $\operatorname{Mcl-1}$ is sufficient to trigger apoptosis in ovarian carcinoma cell lines (3), as well as in cell lines from other malignancies (37-39). The use of siRNAs is an attractive option to downregulate $\operatorname{Bcl-2}$ anti-apoptotic family members in ovarian tumors, particularly $\operatorname{Mcl-1}$ for which no pharmacological inhibitor is yet available for clinical use.

The development of safe and efficient tools for siRNA vectorization is a very intensive research topic. More than 50 clinical trials involving siRNAs have been reported (40) but no siRNA-based drug has yet obtained FDA approval and none of the vectorization systems utilized have achieved a consensus (40,41). Furthermore, a large number of different vectorization approaches have been considered for *in vivo* preclinical studies (40,42,43). The choice of a suitable system to vectorize siRNAs *in vivo* is still greatly dependent on the experimental model.

Ovarian carcinoma development can be classified into 4 different tumoral tissue types: primary tumor, peritoneal tumor including micro- and macro-carcinomatosis, distant tumor nodes, and ascites. The primary tumor is usually removed surgically during first-line treatment, and ascites *in vivo* in mice is usually present at a very late stage of tumor development so it cannot be studied for ethical reasons.

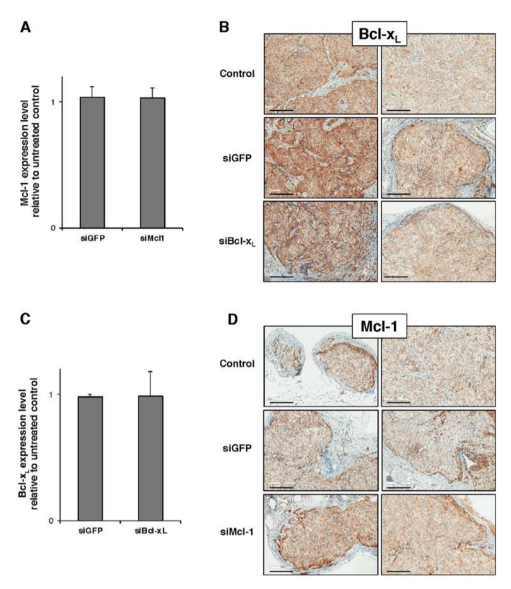


Figure 5. i.v. injection for Mcl-1 or Bcl- x_L -targeted siRNA to subcutaneous tumors. (A and C) RT-qPCR analysis of Mcl-1 and Bcl- x_L mRNA expression level in tumor nodes harvested at day 3. Expression is relative to NaCl tumors for siGFP, siMcl-1 or siBcl- x_L . Data is presented as mean values from 5 tumor nodes from 5 different animals for siMcl-1 and 4 tumor nodes from 4 different animals for siBcl- x_L , \pm SD. (B and D) IHC for Mcl-1 or Bcl- x_L on tumor nodes harvested at day 3. Images are representative of 6 different tumors in each group. Scale bar, 200 μ m.

Micro-carcinomatosis and secondary tumor nodes, which are easily developed in mice, are a good model of the clinically challenging residual disease and recurrence so we targeted them in our siRNA experiments.

The aim of the present study was to evaluate the possibility to vectorize siRNAs in two *in vivo* models of tumoral development in mice mimicking the setting of ovarian carcinoma: the development of carcinomatosis matching the peritoneal dissemination mainly displaying small-sized tumor nodes with low vascularization, and subcutaneous tumors matching distant metastasis with larger tumor nodes displaying more developed vascularization. For the carcinomatosis model with its small islets of tumor cells lacking vascularization, local i.p. injection is attractive. For the distant and/or vascularized metastasis model with i.v. injection, s.c. tumor development is more relevant owing to the unique presence of larger vascularized tumor nodes. We chose to use atelocollagen as a vector since its flexibility (two possible formulations adapted to i.v. and i.p. administration, respectively) enabled us to use it in

both models. In addition, the possibility to use each tumor as its own control, thereby avoiding inter-animal and internodular variability, led us to use a bioluminescent tumoral model to establish a suitable administration regimen.

In the peritoneal carcinomatosis model, we could not evidence any effect of atelocollagen-vectorized siRNAs on the bioluminescence signal in comparison to the control groups. To the best of our knowledge, this formulation with high viscosity has been used only for siRNA delivery with peritumoral injection (18,21,22,30,31), with two exceptions. One study reported atelocollagen-vectorized siRNAs with i.p. injection for intraperitoneal NSCLC tumors (33), but the authors did not give any information either on the atelocollagen concentration or on the amount of siRNA used. Another study demonstrated efficient siRNA delivery and target downregulation on peritoneal tumor nodes in a gastric cancer model after injection of siRNAs-vectorized with atelocollagen at 0.5% together with DharmaFECT 1 transfection reagent (34). Notably, the use of atelocollagen

alone in their model did not allow target downregulation, in accordance with our observations. However, the authors did not report the effects of DharmaFECT 1 transfection reagent in the absence of atelocollagen, so the contribution of atelocollagen to the successful vectorization of siRNAs and silencing of their target could not be estimated.

One of the reasons for the lack of efficacy in this model could be that, unlike with peritumoral or intratumoral injections, the complexes released in the peritoneal cavity are not maintained in contact with the majority of tumor cells, as they cannot cover the entirety of the peritoneal cavity. However, i.p. injection of atelocollagen complexed with ASO was used for delivery to s.c. tumors (44), indicating that small oligonucleotides are at least released efficiently in the peritoneal cavity even with an atelocollagen concentration of 1.8%, which is higher than the one used in the present study, i.e. 0.5%, the concentration recommended in the commercially available kit. A possibility is that once released in the peritoneal cavity, siRNAs, which are more prone to degradation than phosphorothioate-modified ASO, are rapidly excreted and/or degraded without the help of a complementary vector system, before they can reach tumor cells present in this compartment. In order to monitor biodistribution, direct labeling of siRNAs has been used in several instances in vivo. However, the peritoneal compartment is not an organ per se, and such an approach would be difficult to use in our case to monitor the intra-peritoneal release of siRNAs.

Regarding the s.c. tumor model, we first evidenced satisfactory efficiency on the inhibition of luciferase expression as revealed by in vivo bioluminescence (70% decrease in signal vs. control siRNA), which is in line with previous studies in which siRNAs-vectorized with atelocollagen were used (17,23,24,35) on tumors of various origins. This downregulation effect was present at day 3 after 3 daily consecutive injections and was absent at day 7, showing that the effect is transient over time suggesting a loss of the RNA inhibition effect soon after the end of the injections. In several studies describing a long-lasting effect on target inhibition, the protocols included multiple repeated injections (34,45), suggesting that without this, no long-lasting target inhibition can be obtained, which is again in line with our own observations. In addition, when targeting genes involved in cancer progression, successful inhibition by siRNAs prevents further tumoral development and enables tumoral regression. In our luciferase model, however, target inhibition is not supposed to, and indeed does not, impair tumor development. Therefore, the inhibitory effects of transfected siRNAs are likely to be diluted over cell divisions, which is another plausible explanation for a transient decrease in bioluminescence.

Unfortunately, we could not observe the inhibition of target gene expression in s.c. tumors using Mcl-1- or Bcl-x_L-directed siRNAs. The histology of tumors derived from SKOV3 cells (used during Mcl-1 and Bcl-x_L experiments) or SKOV3-Luc cells (used during the bioluminescence experiments) are similar, as confirmed by a certified pathologist (data not shown). Therefore, the use of SKOV3 instead of SKOV3-Luc cells to generate tumors does not likely explain the differences observed between our *in vivo* experiments.

It has been shown that target mRNA half-life influences RNAi efficiency; mRNAs with shorter half-life being more difficult to downregulate (46). Luciferase, Mcl-1 and Bcl-x_L mRNA half-lives are 1.5, 2.5 and 2-3 h, respectively (47,48). These small differences regarding mRNA half-lives are thus quite unlikely to induce any difference in the timing of mRNA downregulation following siRNA injection and do not constitute an explanation for the absence of observed Bcl-x₁ or Mcl-1 downregulation. With regard to protein half-lives, it should be noted that luciferase and Mcl-1 proteins have a 6-fold difference in their respective half-lives [3 h vs. 30 min, respectively (49)]. In addition, the half-life of Bcl-xL protein has been reported to be ~20 h (50). Due to these differences, the effects of siRNA downregulation on Mcl-1 protein should have been measurable earlier than luciferase, whereas Bcl-x₁ protein downregulation should have lasted longer. Overall, the time window in which we should have been able to observe any protein downregulation was larger with Bcl-x_L and Mcl-1. Therefore any difference in the timing of the action of Bcl-x_L or Mcl-1 siRNAs compared to luciferase would have not compromised our ability to observe the effects of siRNA on protein, at least on one of our targets. In fact, these differences in the half-lives of our proteins of interest do not likely explain the lack of observed siRNA activity.

We did consider an increase in dose and/or number of administrations to obtain a more stable and reproducible target downregulation, as well as the use of a panel of alternative siRNA sequences for Mcl-1 and Bcl-x₁. However, in most of the animals we observed a seizure-like behavior shortly after i.v. injection of atelocollagen in complex with each of the 4 siRNAs we used. The mice recovered quickly with no visible sign of physiological or behavioral consequence. It is unlikely that this effect was triggered by contamination of our siRNAs as seizures were observable with 4 different PAGE-purified siRNAs. To the best of our knowledge, this effect with the use of atelocollagen has not been reported elsewhere. Considering previous studies in the literature and the manufacturer's protocol, we could not increase the siRNA concentration in the siRNA/vector complexes. Therefore, as we ruled out increasing the volume or frequency of injections for ethical reasons, we did not push our investigations further with atelocollagen for in vivo siRNA delivery.

In summary, formulation with a high concentration of atelocollagen alone is not suitable for i.p. siRNA transfection in peritoneal nodes. In contrast, a low concentration formulation for i.v. injection is able to deliver siRNAs in s.c. tumors and to induce a strong but transient silencing of the targeted protein activity. In addition, the efficiency of target silencing is very sensitive and depends on the nature of the target. Given that we used the limiting dose for this vector and that the level and duration of silencing are clearly insufficient to explore possible therapeutic effects, we decided not to push our investigations with this vector further for the delivery of siRNAs to ovarian cancer cell xenografts *in vivo* in mice.

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