Urothelial cancer cell response to combination therapy of gemcitabine and TRAIL

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Abstract. High-risk superficial urothelial carcinoma of the bladder (UCB) is commonly treated with intravesical bacillus Calmette-Guerin (BCG), but with significant side effects. We recently showed that the tumour necrosis factor-related apoptosis-inducing ligand (TRAIL) exhibited high therapeutic potential against UCB cells and with only limited toxic effects in normal cells. However, many cancer cells are refractory to TRAIL during monotherapy. Therefore, our experimental aim was to develop combinatorial approaches with other pro-apoptotic agents to reactivate apoptosis in resistant phenotypes. We demonstrate that UCB cells varied in their response to TRAIL, and the effect was caspase-dependent (reduced or abrogated by pre-incubation of cells with caspase-inhibitor peptides). In contrast wortmannin, a PI3K/Akt inhibitor, enhanced the TRAIL effect. Furthermore, combination therapy of TRAIL with low dose gemcitabine markedly enhanced UCB cell response (except in the TRAIL-resistant HT1376 cell line). The enhanced response was both time- and concentration-dependent and asymptotic at gemcitabine concentration >1 µmol/l. To define the mechanisms underlying gemcitabine-augmented TRAIL action, we evaluated the expression of several proteins regulating the apoptotic pathway. Gemcitabine-augmented TRAIL effect was associated with inhibition of the Bcl-2 protein (intrinsic signalling) along with activation of the caspase (extrinsic) cascade. The combined maximal stimulation of both the intrinsic and extrinsic signalling pathways also appeared to overcome the survival (PI3K/Akt) pathway as evident by the lack of response to wortmannin. Our solid-spheroid model showed that TRAIL and gemcitabine selectively caused UCB cells to undergo apoptosis without affecting normal cells, and both appeared to penetrate deeply enough to allow for combination intravesical therapy.

Introduction

The majority (80%) of urothelial carcinomas (UC) are non-invasive. Recurrence of superficial UC of the bladder (UCB) is frequent (as high as 80%) even after complete transurethral tumour resection [TURBT] (1). Bacillus Calmette-Guerin (BCG) is considered the most effective adjuvant intravesical treatment, and appears to induce tumour regression by host immune stimulation via tumour necrosis factor (TNF)-related apoptosis-inducing ligand [TRAIL] (2,3). Systemic and intravesical cytotoxic chemotherapy is used to treat and prevent seeding at the time of TURBT, but BCG cannot be used immediately after TURBT or used systemically for invasive UC because of infectious complications. UC resistance to both BCG and cytotoxic chemotherapy can be induced. At a molecular level there is mechanistic rationale for combining these agents to target both the intrinsic and extrinsic apoptotic pathways. Unlike BCG, TRAIL can be combined with cytotoxic chemotherapy; hence we explored this further in an effort to identify a combinatorial treatment option that can selectively destroy only the transformed cells with minimal risk to normal tissues or the host.

A major mode of resistance to antitumour therapy is loss of sensitivity to apoptosis induction (4). Apoptosis is a cellular death program that controls tissue homeostasis (5,6), and is tightly regulated by a variety of proapoptotic and antiapoptotic proteins; with the relative balance of the opposing activities determining cell fate. TRAIL, a member of TNF family, closely related to Fas-Ligand and TNF-α, induces apoptosis in a wide variety of cancer cell lines and tissues. However, unlike Fas-Ligand and TNF-α, TRAIL spares normal cells even though most express TRAIL receptors at significant levels (7). Upon ligation to its death receptors (DR4 and DR5), TRAIL not only stimulates apoptosis via the formation of the Death Inducing Signalling Complex (DISC) that contains procaspase-8 and Fas Associated Death Domain (FADD) (8), but also activates NF-kB, which regulates the expression of survival factors such as members of the inhibitors of apoptosis and the Bcl-2 families. Although TRAIL is quick at inducing apoptosis in many cancer cell lines, high concentrations of it are necessary to induce apoptosis in other cell lines with low-sensitivity. Cytotoxic chemotherapy and ionizing radiation can up regulate DR expression on cancer cells (9,10). In addition, cytotoxic chemotherapy can trigger the mitochondrial (intrinsic) pathway.
(4,11). Thus, combinations with chemotherapeutic agents may be necessary to sensitize UCB cells.

Gemcitabine [dFdC] is a nucleoside analog with broad spectrum activity against a variety of solid tumours. The cytotoxic effects of this agent are due to the ability of its diphosphate derivative to inhibit DNA synthesis; and the direct incorporation of gemcitabine into DNA, where it inhibits replication (12). Although clinical effectiveness has been demonstrated in the systemic treatment of advanced UC, both as monotherapy (13) and in combination with other agents (14), its use in combination therapy with TRAIL in UC has not been explored. The purpose of the study was to determine whether gemcitabine could sensitize UCB cells to TRAIL and thus potentiate TRAIL-induced apoptosis in those cells that exhibit TRAIL resistance.

Materials and methods

Cell culture, spheroid preparation and reagents. Six established human bladder UC cell lines were used in the study: HT-1376 (ATCC, Manassas, VA, USA); UMCU-14 (from Dr. H.B. Grossman, MD Anderson, Houston, TX, USA); 253J, T24, (from Dr. M. Gleave, University of British Columbia, Vancouver, BC, Canada); RT112 and MGH-U3 (from Dr Y. Fradet, University Laval, QC, Canada). All cell lines were cultured at 37°C with 5% CO2 (standard conditions) in RPMI-1640 or MEM medium supplemented with 10% fetal calf serum FCS, 10 mM HEPES (RPMI only), sodium pyruvate, non-essential amino acids, 2 mM L-glutamine, 100 U/ml penicillin, and 100 µg/ml streptomycin and 0.25 µg/ml amphotericin B. TRAIL-resistant (UC14-T1000) cells were obtained through stepwise exposure of the parental cells (UMUC-14) to 1000 ng/ml TRAIL (PeproTech Inc., Rocky Hill, NJ, USA). Gemcitabine was obtained from Eli Lilly (Indianapolis, IN, USA), reconstituted in sterile phosphate buffered saline and stored in aliquots at -20°C. A liquid overlay (semisolid) culture technique was used to produce multicellular spheroid culture (MCS) as previously described (15). Cells were seeded at a concentration of 3x10^5 cells/well in 96-well plates for tumour cells, and 6x10^3 cells/well for fibroblasts in 200 µl of medium onto 1% solid agarose (50 µl). After 3 days of culture, fibroblast MCS were transferred 1:1 with a pipette tip into a well with a tumour MCS. Resulting co-culture MCS (co-MCS) were ready for experimentation after 3-4 days. As previously reported (15), fibroblasts provide the optimal substrate to generate an in vitro co-cultured spheroid model with proper cellular orientation and cooperative paracrine growth with the malignant cell lines.

Bcl-XL phosphorothioate oligonucleotides. Antisense oligonucleotide to Bcl-2 (Bcl-XL) chemically modified with phosphorothioates (ASO-Bcl2; from Dr. M Gleave and Oncogenex) was delivered into cells in the form of complexes with cationic lipid Lipofectin reagent (Invitrogen, Burlington, ON). ASO-Bcl2 and Lipofectin® reagent were mixed with 350 µl of Opti-MEM in an Eppendorf tube. As well, 7 µl of Lipofectin were diluted into 23 µl of Opti-MEM in a second set of tubes and incubated at room temperature for 30 min. Oligonucleotides and the Lipofectin/Opti-MEM mixtures were combined (ASO-LIPO complex) and incubated for additional 20 min. The mixture was then allowed to dilute with complete growth medium to achieve final concentration of 1000 nM. Cells were incubated for 6 h at 37°C after which equal volumes of two times concentration of antibiotic-free, FCS-supplemented media were added, and cells were then cultured for additional 20 h before being treated with gemcitabine (10 µM).

Dose-response relationship for gemcitabine-induced cell death. The growth inhibitory effects of TRAIL on UC cells during combination therapy with gemcitabine were tested in similar fashion to our recent paper on TRAIL dose-dependent UC cell death (16). Cells were seeded in 96-well plates at a density of 0.4 to 0.7x10^5 cells/well and cultured overnight to adhere to plate. Cells were then pulse-treated with varying concentrations of gemcitabine (10^-4 to 10^-1 M) without or with TRAIL (100 ng/ml) for 3 h, after which the therapeutic medium was replaced with fresh drug-free media. After 24, 48 or 72 h of culture (depending on experiment), cell survival was quantified using the tetrazolium dye 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT; Sigma-Aldrich, St. Louis, MO, USA). To determine the effect of caspase-3, -8, -9 and PI3K inhibitors on tumour cell growth, cells in 96-well plates were pretreated for 2 h with all (2 times concentrated) 100 µl 20 nM wortmannin (to inhibit pro-survival growth factor signalling via PI3K/Akt pathway), or 40 µM caspase inhibitory peptide IETD-fmk (for caspase-8) or 20 µM DEVD-fmk (for caspase-3), or 20 µM caspase inhibitory peptide Z-LEHD-fmk (for caspase-9) (Clontech, Mountain View, CA). This was then followed by addition to the wells of (2 times concentrated) 100 µl of 20 µM gemcitabine or 200 ng/ml TRAIL or their combination (final concentrations were 10 µM and 100 ng/ml, respectively) and cells cultured for 48 h before MTT assay. For all experiments, cell death or growth inhibition was calculated as a percentage of the untreated controls. Each data point represents the average value from three experiments each of which was replicated at least four times.

Co-culture spheroid viability testing. Procedure for differentiating between the two cell populations of UC and fibroblasts, and their orientation in the co-MCS has been described (17), with the fibroblast occupying the inner core of the co-MCS. Untreated, gemcitabine- and/or TRAIL-treated co-MCS were labelled at 24 or 48 h post-treatment with nucleic acid stains SYTO® 16 and propidium iodide (PI) for live/dead assay as previously described (17). Multiple fluorescent images of stained spheroids were captured using fluorescein isothiocyanate and Cy3 filters on a Zeiss Axioplan upright digital imaging microscope equipped with a SenSys cooled CCD camera (Photometrics, Tucson, AZ) using a Zeiss infinity corrected EC Plan-Neofluar 10x objective. All imaging parameters were identical, and acquired images were merged and processed with Metamorph® and Adobe Photoshop® CS2.

Determination of apoptotic cells by Annexin-V/PI and flow cytometry. Untreated or cells treated with gemcitabine and/or TRAIL were incubated with 80 µg/ml Hoechst 33342 (C32H42C7N4O8) for 1 h at 37°C under standard conditions, followed by Annexin-V- FITC (BD Biosciences, San Jose, CA) for additional 20 min. Hoechst, stains the nuclei of live cells whereas membranes of apoptotic cells, but not of live cells are permeable to Annexin-V. The cell-dye suspension was fixed in 1% paraformaldehyde (PFA; in PBS), and image acquisition
of 3-4 non-overlapping fields per well of a 24-well plate was performed on the Zeiss fluorescence microscope. Annexin-V and Hoechst-positive cells were counted after setting a threshold using Metamorph® software. The percentage of apoptotic cells (apoptotic ratio) was calculated as the ratio of the Annexin-stained cells to total cell count (Annexin + Hoechst stained).

For FACS analysis, after incubating with the various agents in vitro cells were collected by gentle trypsinization, washed with ice-cold PBS, and then 1x10^6 cells were fixed in 70% ethanol for 2 h at 4°C. The pellet was resuspended in 1 ml staining solution containing 20-µg/ml PI and 375-µg/ml RNase A. Stained cells were incubated at room temperature for 30 min (in the dark), and the cellular DNA content analyzed by flow cytometry. The relative percentage of DNA content in the sub-G1 phase was then quantitated and used as an estimate of cells undergoing apoptosis (18).

**Immunoblotting.** Following in vitro treatment with gemcitabine and/or TRAIL, cells were lysed in buffer containing 50 mM HEPES [pH 8.0], 150 mM NaCl, 1% Triton X-100, 5 mM EDTA, 5 mM EGTA, 20 mM sodium pyrophosphate, 1 mM sodium orthovanadate, 10% glycerol, 1 mM phenylmethylsulfonyl fluoride, and 4 µg/ml each of leupeptin, aprotinin and pepstatin A. Protein amounts of postnuclear supernatants were assayed using BCA (Pierce, Rockford, IL). Aliquots containing 40 µg of protein were then separated on 4-15% SDS-polyacrylamide gels and electrophoretically transferred to nitrocellulose membranes (Bio-Rad, Hercules, CA). After blocking, membranes were incubated with primary antibodies directed against Bcl-2 (Upstate, Charlottesville, VA), Bid, PARP, and caspases-3, -8 and -9 (Cell Signaling) followed by their respective horseradish-peroxidase-conjugated secondary antibodies (Jackson ImmunoResearch, West Grove, PA). Immunoreactive proteins were detected by enhanced chemiluminescence and autoradiography using HyperFilm-ECL (Amersham, Buckinghamshire, UK). Stripped blots were reprobed with
anti-tubulin or mouse anti-β-actin IgG (Sigma) to establish equivalent loading.

Statistical analysis. Data are provided as means ± SEM. The in vitro cytotoxic effects of gemcitabine and/or TRAIL were analyzed using an ANOVA model. Association between the expression levels of anti-apoptotic proteins and percent cell death was analyzed using the Spearman rank correlation. All statistical tests were two-sided, and the level of significance was set at p<0.05. All analyses were performed using GraphPad Prism®.

Results

Differential sensitivity of UCB cells to gemcitabine and TRAIL treatment. We exposed UCB cell lines to escalating concentrations of gemcitabine (from 0.01 to 100 µM) for 4 h (a relevant clinical intravesical dwell time) and measured cell death by MTT. As depicted in Fig. 1A, gemcitabine cytotoxicity on UC was time-dependent and increased dramatically from 24 to 72 h. At 24 h post treatment, UC cells displayed minimal response at all gemcitabine concentrations. However, when cells were cultured for 72 h there was marked increase in cell death (p<0.05), even at the low gemcitabine concentrations suggesting that both concentration and temporal effects are important in eliciting gemcitabine effect. The time x concentration interaction was further examined in cells treated with gemcitabine for 3 consecutive days (4 h/day; multiple dosing) compared with single dosing above. We observed that the single-dose gemcitabine was as effective as the multiple doses at inducing UC cell death (data not shown), further confirming the time-dependence of gemcitabine effect.

Gemcitabine was previously reported to sensitize pancreatic cancer cells to TRAIL-mediated killing (19). Therefore, in combination experiments we pulse-treated UC cells with escalating concentrations of gemcitabine for one day followed by TRAIL [100 ng/ml; optimal concentration determined in prior studies (16)] treatment for an additional day (Fig. 1B). Treatment with TRAIL as a single agent significantly increased cell deaths by 30-95% in all cell lines (except in HT-1376) relative to untreated control. We have recently reported that these same cell lines displayed differential responses to TRAIL, with UMUC-14 and MGH-U3 being sensitive; RT112, 253J and T24 partially resistant; and HT-1376 being resistant to TRAIL (16). Thus our current data is consistent with this prior observation. In combination experiments, partially TRAIL-resistant cells treated with gemcitabine followed by TRAIL showed an enhancement in cell death (p<0.05) that is beyond that seen for TRAIL alone. Combination of gemcitabine with TRAIL did not improve TRAIL sensitivity in the resistant HT-1376 cell line. However, in UMUC-14 cells made partially TRAIL-resistant through stepwise exposure to 1000 ng/ml TRAIL (UC14-T1000), gemcitabine presensitization reactivated TRAIL sensitivity in these cells (Fig. 2A). Epifluorescent microscopy (Hoeschst 33258 / Annexin V staining) and FACS analysis showed that gemcitabine inhibited the growth of UC cells in a time-dependent manner (increased apoptosis at 2 days post-treatment vs one day; Fig. 2A), and that UC cells underwent gemcitabine induced TRAIL-mediated apoptosis as a method of cell death. In the presence of TRAIL or gemcitabine alone there was increased accumulation of sub-G1 fraction in all cell lines except HT-1376 when compared with the control (Fig. 2B, top panel). Combination of the two agents increased the sub-G1 fraction in the partially TRAIL resistant cells more than individual effect alone.
Next we examined whether the two agents can selectively produce cytotoxicity in a co-MCS setting while sparing normal cells. The MCS model provides \textit{in vitro} simulation of tumour micro-environment and has been used extensively in prior studies of resistance of tumour cell layers to chemotherapy. Pure and co-MCS (MGH-U3 and CRL-1120 fibroblasts) were exposed to gemcitabine (10 µM) alone or with TRAIL (100 ng/ml), and cell death and shedding (MCS dissolution) were monitored (Fig. 3). In homogeneous spheroids (comprised exclusively of MGH-U3), PI uptake by gemcitabine- or TRAIL-treated cells increased dramatically from 24 to 48 h; an indication of increased cell death. Similarly, in heterogeneous co-MCS (MGH-U3/CRL-1120) the PI uptake was time-dependent for the peripheral MGH-U3 cells whereas CRL-1120 fibroblasts took up SYTO 16 dye, indicating tumour-specific gemcitabine-initiated cell death and dissolution. Furthermore, combination of the two agents produced marked UC cell killing and extensive dissolution compared with either agent alone. There were no cytotoxic effects observed on the fibroblasts. Collectively these results suggest that the combination of TRAIL and gemcitabine caused rapid cell death, and selectively cause UC cells to undergo apoptosis in co-MCS without affecting normal fibroblasts.

Apoptosis response of UC cells to gemcitabine/TRAIL is dependent on suppression of Bcl-2 and activation of the...
caspase pathway. Next we examined changes in the expression of key proteins that are centrally involved in regulating apoptosis. Most death modulators function by acting through the Bcl-2 family. Although other members of this family stimulate programmed cell death, bcl-2 is the prototypic antiapoptotic gene (20), and its expression has been correlated with apoptotic responses of a variety of neoplastic cell lines. Gemcitabine alone decreased Bcl-2 expression in the partially-TRAIL resistant cell lines, and the effect was both time and concentration dependent (Fig. 4A-C). TRAIL alone had little or no effect on Bcl-2 expression in the same partially TRAIL-resistant cell lines. However, when TRAIL was combined with gemcitabine the decrease in Bcl-2 expression was more pronounced than the gemcitabine effect alone. Further analysis showed that Bcl-2 level was inversely correlated with gemcitabine cytotoxicity (percent cell death) in UC cell lines consistent with Bcl-2 being prognostic of tumour recurrence and progression (21-23).

Figure 4. Effect of gemcitabine and TRAIL on Bcl-2 expression. (A) Shows arbitrary densitometry values, respectively for Bcl-2 protein expression in RT-112 cells after exposure to varying concentrations of gemcitabine for the time indicated. Data are the means ± SEM from three separate experiments; (B) shows immunodetectable Bcl-2 protein in gemcitabine-treated RT112 and T24 cell lines harvested at 24 or 72 h post-treatment; and (C) expression levels of Bcl-2 protein in TRAIL-resistant (HT-1376) and partially TRAIL-resistant (RT112, T24, 253J) UCB cell lines after combination therapy with gemcitabine and TRAIL. Cells were treated with the agents for 4 h and then cultured in drug-free medium for 3 days before harvesting for Western blot analysis. β-actin was used as loading control. (D) Partially TRAIL responsive cell lines (RT112, T24 and 253J) were transfected with antisense oligonucleotide targeting Bcl-2 (ASO-Bcl2) or MisMatch control (MM). Twenty-four hours after transfection cells were treated with gemcitabine (10 µM for 4 h) and then cultured for 2 days in drug-free medium followed by MTT cell proliferation assay. Western blot analysis showed that ASO-Bcl2 reduced Bcl-2 expression in the treated cells. a-d represent significant differences, p<0.05.
apoptotic resistance and/or improved tumour cell response to the cytotoxic agent. We transiently transfected 3 partially-TRAIL-resistant cell lines (RT112, T24, and 253J) with antisense oligonucleotide to Bcl-2 (ASO-Bcl2) or mis-match control (MM) and cultured them in the absence or presence of gemcitabine (10 µM). Western blot analysis showed that ASO-Bcl2 treatment greatly reduced Bcl-2 protein expression (Fig. 4D) along with increased cell death compared with the MM control. Combination of ASO-Bcl2 and gemcitabine significantly increased cell death compared with either gemcitabine or ASO-Bcl2 alone.

We also analyzed caspase-mediated Bid activation in partially TRAIL resistant and TRAIL-sensitive cells. Gemcitabine effect on Bid expression was somewhat variable and did not show strong relationship with treatment. However, we detected greatly reduced levels of Bid (suggesting consumption of Bid protein by caspase-8 during activation) after treatment with TRAIL alone [similar to our recent observation

Figure 5. Effect of gemcitabine and its combination with TRAIL on apoptotic protein expression in UCB cells. Cells were exposed to 10 µM gemcitabine for 4 h and then incubated in drug-free media for additional 20 h followed by TRAIL treatment (100 ng/ml for 4 h). Cells were harvested immediately following TRAIL treatment and lysates examined by Western blot analysis using a panel of antibodies directed against Bid, PARP, caspase-8, -3 and -9. Equal protein loading of samples was verified by the level of β-actin in stripped blots (25 mM glycine, pH 2.0, 1% SDS).
(16] or in combination with gemcitabine (Fig. 5A and B). We did not observe an active fragment of Bid (tBid) in any of the treatments. Also gemcitabine-augmented TRAIL-induced caspase-8, -9 and -3 activation was enhanced in the cell lines (except HT1376). Further, we observed enhanced PARP cleavage in response to gemcitabine and during combination therapy with TRAIL. These results could mean that both the extrinsic and intrinsic pathways were involved in the apoptotic cell death initiated by gemcitabine/TRAIL. On the other hand there was little or no activation of Bid, PARP, pro-caspase-8, -3 and -9 in the TRAIL-resistant HT-1376 cell line (Fig. 5B); suggesting there could be a defect upstream of caspase activation, which may result in failure to adequately activate and initiate apoptotic signalling in this cell line. To further investigate the involvement of the caspases in gemcitabine- and TRAIL-induced cell death, we pre-treated cells with caspase inhibitor peptides before exposing them to gemcitabine and/or TRAIL. Addition of individual caspase inhibitor peptides or their combination significantly inhibited TRAIL-induced cell death (Fig. 6), supporting that the observed TRAIL effects were directly dependent on the activation of these caspases. However, inhibition of the caspases was not effective in diminishing gemcitabine cytotoxicity. Also, inhibition of the pro-survival function via wortmannin inhibition of PI3K/Akt had very little to no effect on TRAIL- or gemcitabine-induced cell death or when gemcitabine was combined with TRAIL (Fig. 6).

**Discussion**

UC response to BCG is believed to be mediated by TRAIL. TRAIL induces apoptosis in a variety of tumour cells with minimal toxicity to normal cells. Resistance to TRAIL-mediated apoptosis by some tumour cells however limits its widespread application. Several factors have been implicated in this resistance. Amongst them is the low expression of agonistic TRAIL receptors DR4 and DR5, but there does not appear to be a common mechanism for all cancer cells (15-17). Recently we showed that the TRAIL-induced apoptosis is mediated via activation of caspase-3 and -8 in UC cells (16), which is often a function of ligand-dependent trimerization of DR4 and DR5 and the efficiency by which TRAIL ligation to the DRs induces protein association in the caspase cascade. In some experiments we observed that UC cell lines that are less responsive to TRAIL express DR4 and DR5 receptors at levels comparable to the TRAIL-sensitive ones (data not shown). That is, the apoptotic sensitivity of the UC cells did not appear to be related to the DR expression patterns alone (27-29). Thus, our current observation that TRAIL induced apoptosis in a caspase-
dependent fashion may suggest that resistance to TRAIL may depend more on the level of intracellular signalling molecules rather than differences in receptor expression.

Given our prior data (15-17) we have focused on identifying effective therapeutic strategies for overcoming resistance within heterogeneous and adaptive solid tumour cell population, with the view that multiple apoptotic pathways (the extrinsic and intrinsic) would need to be targeted. Recent studies in the literature have indicated that sequential treatment of tumour cells with combination of TRAIL and chemotherapy can overcome resistance to TRAIL-mediated apoptosis, and in particular significant synergism between TRAIL and gemcitabine action has been demonstrated in other cancer cell lines (27,30,31). In the current studies we investigated the utility of combining gemcitabine with TRAIL for the treatment of localized (non-muscle invasive) and possibly advanced (invasive) bladder cancer. In the normally TRAIL-resistant UC (HT1376) cells, pretreatment with gemcitabine did not overcome resistance of the cells to TRAIL-induced apoptosis. This is discordant with the observation by Seol et al (32) that showed pretreatment of TRAIL-resistant A549 cells with gemcitabine enhanced TRAIL-induced apoptosis, which was mediated not only via caspase-dependent and mitochondrial pathway but also by upregulating DR5 expression. Consistent with other studies (32,33) however, we showed that gemcitabine induced apoptosis in the partially-TRAIL-resistant bladder UC cells, and its anti-proliferative effects were concentration-dependent. Also, we observed significant enhancement of cytotoxicity in nearly all UC cells through rapid induction of apoptosis when co-treated with gemcitabine and TRAIL. These observations, like other similar observations in the literature (19,32,34,35), could be of clinical importance in that TRAIL could serve to sensitize cancer cells to chemotherapeutic agents or vice versa and thus enable dose reduction and therefore reduce side effects (36,37). Also concentrations of gemcitabine required to sensitize UC cells to TRAIL-mediated apoptosis are well within a range that is clinically achieved in patients, and the dose required would be expected to lower potential toxicity while enhancing efficacy. Thus, given its previously reported systemic activity as a single agent (13,38), and high body clearance rate (39), gemcitabine is a logical choice for both systemic and intravesical therapy; and several reports have confirmed its feasibility and tolerability for such application (40-42). For any intravesical application however, penetration of the molecule is crucial. Our current data and those from our previous studies (15,17,43) demonstrate that gemcitabine can penetrate well in vitro into co-MCS UCB models with evidence of selective toxicity. We also have demonstrated here that TRAIL appears to have similar robust penetration to allow for potential intravesical administration of this agent alone or in combination with gemcitabine.

The presence of anti-apoptotic proteins such as members of the Bcl-2 family can also contribute to TRAIL resistance. Depending on cell type TRAIL apoptotic effect can be amplified by suppression of antiapoptotic molecules and/or upregulation of pro-apoptotic molecules. Importantly, we showed that during the first 24-72 h of gemcitabine-TRAIL therapy there was robust downregulation of Bcl-2, accompanied by increased apoptosis (and cell death) in the UC cell lines investigated. In addition gemcitabine induced apoptosis with caspase-3, -8, -9, and PARP cleavage. These observations suggest that gemcitabine sensitizing effects in partially TRAIL-resistant cell lines might not be solely Bcl-2 dependent but could also be due to other mechanisms, including death receptor pathway and mitochondrial damage. In TRAIL-resistant cells (HT1376) gemcitabine or its combination with TRAIL had no effect on Bcl-2 level or TRAIL apoptosis. This could be due to the fact that activation of the TRAIL receptor can lead to activation of nuclear factor-κB which may subsequently mediate proliferation, invasion and metastasis (47). In addition, the HT1376 cell line appears to have a lower level of apoptotic proteins. Also post gemcitabine sensitization, TRAIL can induce a rapid negative-feedback loop that could downregulate TRAIL receptor expression thus allowing tumour cells to escape TRAIL mediated effect (28). In addition, we observed in this and other studies (unpublished data), that TRAIL can trigger the intrinsic pathway via Bid and that this pathway may be hampered or impaired in TRAIL-resistant UC cells. The Bid activation is cross-talk between the receptor-mediated pathway and mitochondria pathway that normally occurs when active caspase-8 cleaves Bid. This then further supports the presence of alternative pathways that may resist or bypass apoptosis, or a situation where other anti-apoptotic proteins may substitute for Bcl-2 (after its downregulation) resulting in chemoresistance. Our data are, however, discordant with other observations where treatment of TRAIL-resistant cancer cells with gemcitabine enhanced TRAIL effect (29,32). This enhancement was only observed in cells with partial resistance.

The gemcitabine effect on activation of the caspases could potentially permit combination therapy with TRAIL being administered after Bcl-2 downregulation with gemcitabine. By selectively inhibiting the caspase cascade with the two inhibitory peptides, our results demonstrated what could potentially happen with such a combination therapy and the near absolute need of these molecules for TRAIL-mediated apoptosis in UC cells. Apoptosis induced by TRAIL was dependent upon caspase activation as indicated by the effect of the caspase inhibitor peptides. However, inhibition of the caspases was ineffective in reducing gemcitabine-induced cytotoxicity. This observation suggests that these proteins are critical effectors of extrinsic apoptosis, whereas gemcitabine can activate caspase-3 by caspase-9 or independently through another mechanism either of which is sufficient for induction of apoptosis. In our studies pretreatment with gemcitabine enhanced TRAIL-mediated apoptosis suggesting that reduction of Bcl-2 had a more dominant impact than the extrinsic caspase pathway. Alternatively, gemcitabine may increase DR expression which we could not detect (data not shown) (54). As demonstrated, deficiency of one of the caspases examined can potentially result in compensatory activation of alternative (redundant) death activating pathways. As such the caspases should be the key rate limiting step as they are consumed in their activation and were thus chosen for targeting in these studies. However, the redundancy in the apoptotic pathways allows the process to proceed through upstream Bid or other mechanisms not yet elucidated. Bcl-2 appears to be a key or dominant regulator of both the intrinsic and extrinsic pathway in UCB.

In conclusion, we show here that treatment with TRAIL activates the caspase cascade to induce apoptosis in the UC
cell lines. We also showed that gemcitabine enhanced the sensitivity of UC cells to TRAIL-induced apoptosis, and this was achieved in part through down-regulation of Bcl-2, and activation of the proteins (caspases, DISC) involved in the extrinsic (type II) death signalling cascade. The combined maximal stimulation of both the type I and type II pathways appears to overcome the survival pathway of PI3K-Akt pathway as evident by the lack of response to wortmannin. Furthermore, our data suggest that gemcitabine and TRAIL both appear to be able to penetrate deeply enough to allow for combination intravesical therapy in addition to systemic therapy. Thus, further preclinical and clinical studies are required to assess this therapeutic strategy for UC.

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