

Preclinical and 'near-patient' models for the evaluation of experimental therapy in prostate and bladder cancer Merbel. A.F. van de

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Cationic Amphiphilic Drugs as Potential Anti-Cancer Therapy for Bladder Cancer

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Abstract

More effective therapy for patients with either muscle-invasive or high-risk non-muscle invasive urothelial carcinoma of the bladder (UCB) is an unmet clinical need. For this, drug repositioning of clinically-approved drugs represents an interesting approach. By repurposing existing drugs, alternative anti-cancer therapies can be introduced in the clinic relatively fast, because the safety and dosing of these clinically-approved pharmacological agents are generally well known. Cationic amphiphilic drugs (CADs) dose-dependently decreased the viability of a panel of human UCB lines in vitro. CADs induced lysosomal puncta formation, a hallmark of lysosomal leakage. Intravesical instillation of the CAD penfluridol in an orthotopic mouse xenograft model of human UCB resulted in significantly reduced intravesical tumor growth and metastatic progression. Furthermore, treatment of patient-derived ex-vivo cultured human UCB tissue caused significant partial or complete anti-tumor responses in 97% of the explanted tumor tissues. In conclusion, penfluridol represents a promising treatment option for bladder cancer patients and warrants further clinical evaluation.

Introduction

Urothelial carcinoma of the bladder (UCB) is the fifth most common cancer in the Western world (1). Despite the prevalence and high economic costs of UCB, this cancer is still relatively understudied (2). UCB presents either as non-muscle-invasive (NMIBC) or muscle-invasive carcinoma (MIBC), and many histopathological and molecular subgroups have been defined (3). NMIBC patients undergo transurethral resection of the bladder tumor (TURBT), followed by adjuvant intravesical instillations with Bacillus Calmette-Guérin (BCG) in patients at high risk of recurrence and progression. However, patients do not always tolerate BCG and the risk of relapse in BCG-treated patients varies between 30-40% (4). Neoadjuvant cisplatin-based chemotherapy and radical surgery is the recommended treatment option in non-metastatic MIBC patients. Nevertheless, the 5-year survival rate is only a mere 50% (5). For patients with metastatic UCB, systemic cisplatin-based chemotherapy is the standard of care (6). Taken together, more effective therapies for high-risk NMIBC and advanced or metastatic UCB patients are warranted.

By repurposing existing drugs, alternative anti-cancer therapies can be introduced in the clinic relatively fast, because the safety and dosing of these clinically-approved pharmacological agents are generally well known. The risk of clinical failure due to (serious) adverse effects or reactions, therefore, strongly reduced (7). Consequently, the time needed for bringing the drug from bench-to-bedside is reduced (8). Upon meta-analysis, the overall risk-ratio of cancer incidence was reduced amongst schizophrenia patients, although findings remain controversial (9, 10). Strikingly, the incidence of UCB in patients with schizophrenia is significantly reduced when adjusted for differences in smoking prevalence (9). Strikingly, a class of commonly used antidepressants, antihistamines and antipsychotics- the so-called Cationic Amphiphilic Drugs (CADs)- were previously found to preferentially induce cell death in transformed cells (11). The most potent CADs, including penfluridol, display preferential cytotoxicity towards transformed cells in vitro and potent anti-tumor activity even as single agents in murine tumor models (amongst others in breast, colon, pancreatic and lung carcinoma (11-17).

We hypothesized that CADs display anti-tumor activity in human UCB. We evaluated and compared the potential anti-tumor efficacy of multiple CADs in human UCB cell cultures in vitro, in a preclinical orthotopic human UCB growth and progression model, and in cultured, patient-derived UCB tissue ex-vivo.

Materials and Methods

In vitro

Human UCB cell lines were cultured as described in **Supplementary table 1** (18, 19).

Viability

Cells were seeded as described previously (19). Cells were seeded to reach subconfluency after 24h and subsequently treated for 2h with a dose-range. After replacing the medium, cells were incubated at 37°C for an additional 24, 48 or 72h.

Live cell count

Cell death was measured after 15 min propidium iodide (dead cells;0.2 µg/mL) and Hoechst-33342 staining (live cells;2.5µg/mL) at 37 °C employing Celígo® Cytometer according to the manufacturer's manual.

Apoptosis and Necrosis

Cells were seeded in a 96-wells plate at a density of 10,000 single-cells per well with 3 technical replicates. After 24h, detection reagent (RealTime-Glo™ Assay) was added. Subsequently, a dose-range of penfluridol was added and luminescence and fluorescence were measured using SpectraMax iD3.

Clonogenicity

Clonogenic assay was performed as described previously (19).

Lysosomal membrane permeabilization

60,000 cells were seeded on PET-coated chamber slides and left for 24h to adhere. Cells were treated with a dose-range of penfluridol for 2h. After 24h, lysosomal membrane permeabilization was detected by staining paraformaldehyde-fixed cells with LGALS1 and LAMP-1 (**Supplementary table 1**). Confocal images were taken with TCS SP8.

Orthotopic in vivo xenograft model

Female BALB/c nude mice (8-weeks old) were housed in ventilated cages under sterile conditions according to Dutch guidelines (evaluated by the Leiden Ethical Animal Welfare committee; DEC_14212; **Supplementary table 1**). Human luciferase-expressing UM-UC3-luc2 cells were inoculated into the bladder as described previously (18) (**Supplementary information**).

For both inoculation of healthy mice and the xenograft model, mice were intravesically treated weekly with either vehicle or penfluridol solution ($100\mu M$ equivalent to $130\,\mu g/kg/w$). The bladder was emptied by mild abdominal massage. Subsequently, $50\,\mu l$ solution was inserted via an angiocatheter (24G) and the urethra was closed for 1h using a suture. Thereafter, followed by emptying of the bladder (18). Bioluminescence imaging was performed using the IVIS Lumina (18). Quantification of bioluminescent signals was performed by the Living Image® (Xenogen) software. Values were expressed as RLUs in photons/second. Numbers of metastases per animal were counted by eye from a dorsal and ventral view.

Ex-vivo cultured UCB tissues

Tumor tissues from patients diagnosed with various stages of UCB were obtained during transurethral resection of the bladder upon informed consent (evaluated by the Erasmus Medical Ethical Committee MEC-2014-553). Explanted tumor tissue slices (TS) were sectioned and cultured as previously described (20, 21) (**Supplementary information**).

Scoring and inclusion criteria

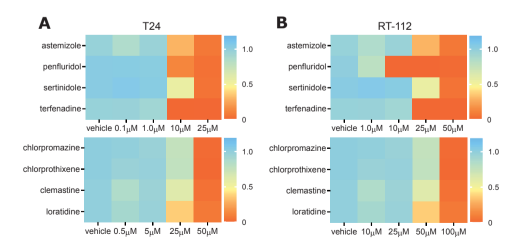
Upon fixation, paraffin embedding and sectioning, TS were stained for H&E, apoptosis (c-CASP-3), proliferation (PCNA) and cytokeratins (KRT) (20). Stained sections were scanned with a slide scanner and evaluated with Caseviewer. Immunohistochemical evaluations were based on visual estimations from complete histologic tissue sections. Confocal images were taken from representative regions. Criteria for the inclusion of UCB tissue are 1) diagnosis of UCB in TS is confirmed by the pathology report; 2) majority of directly fixed and/or vehicle-treated TS contain tumor cells; 3) size of obtained material is sufficient to perform at least vehicle and 1 CAD dosage. First, H&E stained TS were scored for overall quality (score 0: good quality; 1: poor tissue integrity and quality in >50% of TS area; 4: tissue completely degraded or fragmented). Then, sections were scored for the presence of apoptosis (c-CASP-3+ tumor cells) and fragmented cytokeratin (loss of cell integrity) (score 0: sporadic; 1: multiple clusters). Finally, TS were scored for presence of PCNA (score 1:<50% of KRT+ cells displayed nuclear PCNA). Total score: cumulative individual scores (**Supplementary figure 4**).

Statistics

Analysis was performed using GraphPad Prism 7.0 and Excel. For in vitro assays, one-way ANOVA and Bonferroni post-hoc test. For in vivo: Mann-Whitney-U tests. For ex-vivo: Chi-squared test for categorical data with more than two categories. p<0.05;**p<0.01;***p<0.001.

Results

Cationic Amphiphilic drugs (CADs) induce lysosomal-dependent cell death in vitro To identify CADs that display anti-tumor effects, 8 CADs were selected -at least in part- based on a library screen on lung cancer and on the NCI drugs sensitivity database (22). These CADs were initially tested for their effect on viability of T24 and RT-112 human UCB cells in vitro. The CADs penfluridol, astemizole and terfenadine induced the strongest anti-tumor effects upon short-term exposure, (Figure 1A-B; Supplementary figure 1A-D). Subsequently, short-term exposure to a dose-range of these CADs also led to a dose-dependent inhibition in viability of a panel of UCB cells (Figure 1C-E; Supplementary figure 1E-G). Viability was decreased when a confluent layer of UCB cells was exposed to a dose-range of penfluridol (Supplementary figure 1H), Furthermore, penfluridol significantly decreased the clonogenicity of multiple UCB cell lines (Figure 1F-G; Supplementary figure 2A-D). Since we observed strongest anti-tumor potential with penfluridol this compound was used in other disease models in vivo and exvivo. Penfluridol treatment resulted in redistribution of phosphatidylserine from the internal to external membrane surface, an early indicator of apoptosis. This was followed by lysis of the cell as measured with a DNA binding dye (Figure 2A-B). For the higher dosage (100 µM), only a small increase in apoptosis was observed, and already loss of membrane integrity was observed within 2 hrs. Furthermore, lysosomal LGALS1 puncta formation, a hallmark of lysosomal leakage, was found in penfluridol treated cells (Figure 2C) (11, 12, 23, 24).



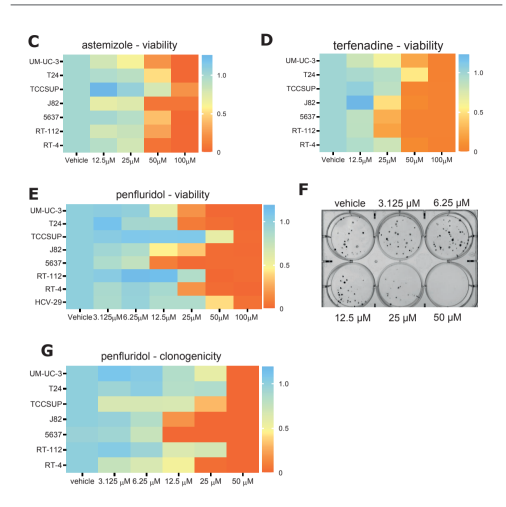
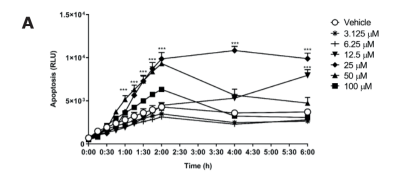


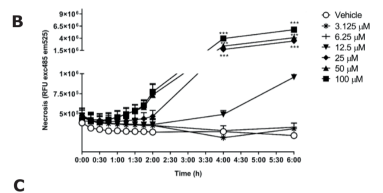
Figure 1 CADs reduce viability and clonogenicity in a panel of human bladder cancer cells.

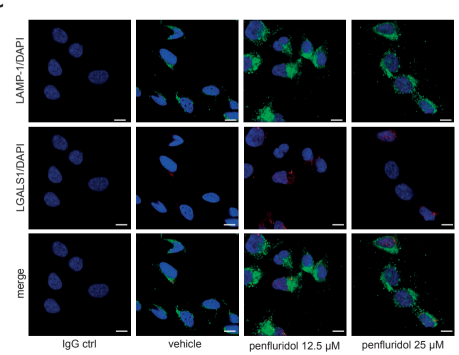
Assessment of the viability of T24 (\mathbf{A}) and RT-112 (\mathbf{B}) cells after treatment with 8 different CADs for 40h (n=3; 6 replicates each). Panels (\mathbf{C} - \mathbf{E}) represent dose-response experiments with astemizole, terfenadine and penfluridol respectively on the viability of multiple subconfluent bladder cancer cells. Viability was measured 48h after a 2h treatment. Mean normalized to vehicle treated cells. (n=3; 6 replicates each). Clonogenic assay: multiple bladder cancer cells were treated for 2h with a dose range of penfluridol. (\mathbf{F}) Representative image of a clonogenic assay UM-UC-3 cells. (\mathbf{G}) The number of colonies was measured using Image J after 10-14 days of culture. (n=3; 3 replicates each).

Penfluridol inhibits UCB growth and metastasis in a preclinical orthotopic xenograft model

In order to evaluate the anti-tumor effects of penfluridol in a preclinical xenograft mouse model, firefly luciferase2-expressing human UM-UC-3luc2 cells were inoculated in the bladder of female immunodeficient mice (BALB/c nu/nu) (18). Four days after tumor cell inoculation, tumor burden was assessed by wholebody bioluminescence imaging (BLI) and properly inoculated mice were equally distributed among two groups with similar median tumor burden (Supplementary figure 3A-B) (18). At this stage, bladder tumor cells have not yet invaded the muscle layer of the murine bladder. Subsequently the mice were treated by weekly intravesical instillations of penfluridol or vehicle solution. Penfluridol significantly reduced total tumor burden by 63% (Figure 3A-B). No significant changes were observed in body weight (Supplementary figure 3C). At day 29, mice were sacrificed and size and weight of the bladders of penfluridol-treated mice were diminished compared with vehicle-treated animals (Supplementary figure 3D-E). As expected, histological examination revealed large orthotopically-growing tumors in vehicle-treated mice with invasion of the tumors into the surrounding connective tissue and muscle layers (Figure 3C). In contrast, in penfluridoltreated mice no tumor invasion into the bladder muscle layer was observed and large necrotic regions in tumor tissues at the luminal side of the bladder were detected (Figure 3D). All vehicle-treated mice developed metastatic disease with an average of 6.8 metastases/mouse, while only 33% of the penfluridoltreated animals developed distant metastases with an average 4.3/mouse (Figure 3E). Metastases were observed in the lungs, reproductive system, liver, intestinal or mesenteric lymph nodes, pancreas, bone and spleen lymph nodes (Supplementary figure 3F). Both the number of metastases/mouse and metastatic tumor burden were significantly decreased in penfluridol-treated mice (Supplementary figure F-G).





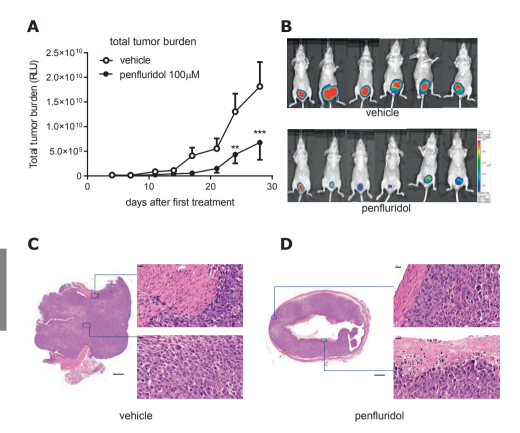


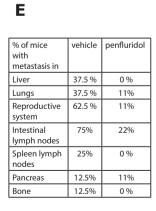
◆ Figure 2 Penfluridol induces cell death in human bladder cancer cells.

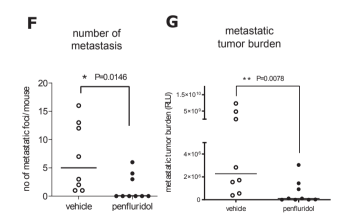
(A) Real-time apoptosis (luminescence measurement of phosphatidylserine on outer leaflet of cell membranes with Annexin V) and (B) cell death (fluorescence measurement of a DNA binding dye) after administration of a dose range of penfluridol. Data presented are mean +/- SE. (n=3; 6 replicates each). One-way ANOVA. *p<0.05; **p<0.01; ***p<0.001. (C) Penfluridol targets lysosomal structures in UM-UC-3 cells (n=3, 3 replicates each). Representative images of LAMP-1 (green) LGALS1 (red)/ DAPI (blue) stained UM-UC-3 cells treated for 2h with a dose range of penfluridol.

Anti-neoplastic response of Penfluridol in ex-vivo cultured, patient-derived cancer tissue slices

Next, we examined the anti-neoplastic effect of penfluridol in our 'near-patient' model, i.e. ex-vivo cultured patient-derived UCB tissue slices (20). Tumor tissues from 57 patients diagnosed with different stages of bladder carcinoma were obtained during transurethral resection of the bladder (TURB) procedures. Tumor tissues from 39/57 patients (68%) were included according to a pre-defined set of tissue quality criteria (see **Materials and methods section 2.3.1; Table 1**). Tumor tissues were treated with vehicle solution or a dose-range of penfluridol (n=18) for 3 days. When tumor biopsy material was limited (n=21), explanted tumor tissues were cultured in presence of vehicle solution or 100 µM penfluridol for 3 days. Culturing of the tumor tissue slices under control conditions only marginally affected tissue integrity and viability or not at all, as indicated by marginal differences in immunohistochemical localization of fragmented cytokeratin (indicative of cancer cell debris), apoptotic cells (c-CASP-3) and proliferating cells (PCNA) compared to directly fixed tissue (scoring method in **Supplementary figure 4** and results in **Supplementary figure 5**).







◆ Figure 3 The anti-tumor effects of penfluridol in an orthotopic murine xenograft model with stable firefly luciferase-2 UM-UC-3 human bladder cancer cells.

Female BALB/c nu/nu mice were inoculated with firefly luciferase2 labelled UM-UC-3 bladder cancer cells and intravesically treated with vehicle (n=8) or penfluridol (n=9); 100 μ M once weekly; equivalent to 130 μ g/kg/w). (**A**) quantification of the whole-body total tumor burden in real time (relative light units (RLU). Data are presented as mean +/- SE). Mann-Whitney-U test. *p<0.05; **p<0.01; ***p<0.001. (**B**) representative bioluminescent images of vehicle and penfluridol treated mice at day 29. Representative images of vehicle (**C**) and penfluridol (**D**) treated bladders stained with Hematoxylin & Eosin. Scale bar 500 and 20 μ m respectively. (**E**) The percentage of mice with detectable metastases in the listed organs. (**F**) number of metastatic foci per mouse per experimental group. (**G**) metastatic tumor burden per experimental group (RLU). Mann-Whitney-U test. *p<0.05; **p<0.01; ***p<0.001.

In almost all patients (35/39), strong changes were observed after treatment with penfluridol, ranging from complete loss of tumor to significant reduction of the tumor (**Figure 4A**). An overall reduction in the number of UCB cells, proliferating cells (PCNA), increased numbers of apoptotic cells (c-CASP-3) and significantly increased amounts of fragmented cytokeratins were observed (P value=4,36424E-11, X2 test). In tissue slices obtained from only three patients, partial anti-neoplastic responses were registered (e.g. an induction of apoptosis). In tissue slices obtained from one patient (1/39), no response to the treatment was found. The effect of a dose-range of penfluridol revealed that, starting from 12.5 μ M, major anti-neoplastic changes were observed (**Figure 4C-D**).

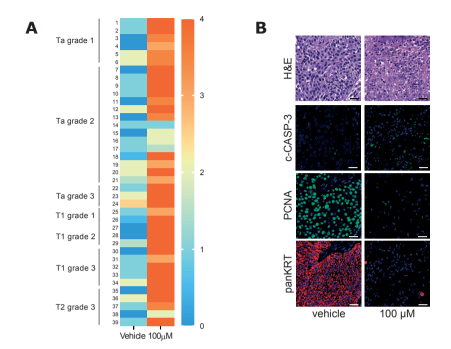
► Table 1 Patient characteristics.

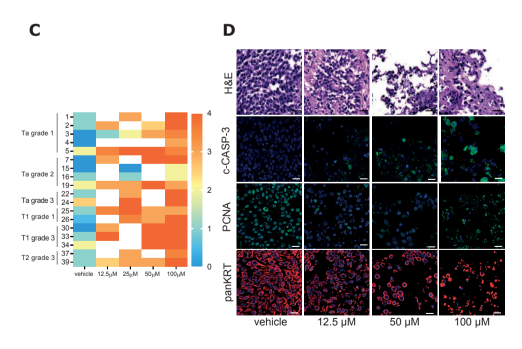
Disease stage, grade, presence of CIS and detrusor was determined by a pathologist. Patient #40 is a patient diagnosed with prostate carcinoma Gleason Tumor tissues from 57 patients diagnosed with various stages of urothelial carcinoma of the bladder were obtained during transurethral resection of the bladder (TURBT upon informed consent, MEC-2014-553). Tumor tissues from 39 patients were included according to the tissue quality inclusion criteria. grade 3+4 with no history of UCB. NA: not applicable. BCG: Bacillus Calmette-Guérin.

patient	stage	grade	tumor	recurrence	cis	detrusor	previous	previous treatment	gender	age
1	Та	G1	primary	NA	unknown	yes	no	modality NA	male	75
2	Та	G1	primary	NA	unknown	yes	no	NA	male	71
3	Та	G1	recurrence	>1 / year	unknown	yes	unknown	unknown	male	72
4	Та	G1	primary	NA	unknown	yes	no	NA	female	69
2	Та	G1	primary	NA	unknown	no	no	NA	male	62
9	Та	G1	recurrence	>1 /year	unknown	no	yes	nephroureterectomies	male	63
7	Та	G2	primary	NA	unknown	yes	no	NA	male	92
8	Та	G2	primary	NA	по	yes	no	NA	male	67
6	Та	G2	recurrence	>1 /year	no	yes	yes	TURBT	male	98
10	Та	62	recurrence	>1 /year	no	yes	yes	chemotherapy	male	26
11	Ta	62	recurrence	>1 /year	ou	yes	yes	BCG	male	87
12	Та	62	recurrence	>1 /year	no	yes	yes	nephroureterectomies	male	62
13	Та	G2	primary	NA	no	unknown	no	NA	male	39
14	Ta	62	recurrence	>1 /year	no	yes	yes	BCG	male	59
15	Ta	G2	primary	NA	no	yes	no	NA	female	69
16	Та	G2	recurrence	>1 /year	no	yes	yes	chemotherapy	female	53
17	Та	G2	recurrence	>1 /year	no	yes	yes	Chemohyperthermia	male	57
18	Та	G2	primary	NA	no	yes	no	NA	male	79
19	Та	G2	primary	NA	no	yes	no	NA	male	73
20	Ta	G2	recurrence	<1 / year	no	yes	yes	chemotherapy	male	75

► Table 1 Patient characteristics Contiuned.

patient no	stage	grade	tumor	recurrence frequency	cis present	detrusor present	previous treatment	previous treatment modality	gender	age
	Та	G2	recurrence	>1 /year	no	yes	yes	BCG	female	29
	Ta	G2	recurrence	unknown	no	yes	yes	chemoradiation	male	84
	Ta	63	primary	NA	no	yes	no	NA	male	75
	Ta	63	recurrence	>1 /year	no	ou	yes	BCG	male	92
	Та	63	primary	NA	unknown	no	no	NA	male	26
	Та	unknown	primary	<1 / year	no	unknown	no	NA	female	53
	T1	G2	recurrence	<1 / year	unknown	no	no	NA	male	98
	T1	G2	recurrence	>1 /year	no	yes	yes	BCG	male	74
	T1	G2	primary	NA	yes	yes	no	NA	male	29
	T1	G2	primary	NA	no	yes	no	NA	male	55
	T1	63	primary	NA	no	yes	yes	TURBT	male	99
	T1	63	recurrence	>1 /year	no	yes	yes	BCG	male	52
	T1	63	primary	NA	no	no	no	NA	female	29
	T1	63	primary	NA	unknown	yes	no	NA	male	81
	T2	63	primary	NA	no	yes	no	NA	male	74
	T2	63	primary	NA	yes	yes	no	NA	male	28
	T2	63	recurrence	>1 /year	yes	yes	no	NA	male	72
	Т2	63	primary	NA	unknown	yes	no	NA	male	99
	T2	highgrade	primary	NA	unknown	yes	no	NA	male	64
	T3	GI3+4	primary	NA	NA	NA	yes	LHRH agonist	male	9





◄ Figure 4 Ex-vivo treatment of cultured human bladder cancer slices with penfluridol.

Heat maps showing median cumulative scores of the bladder cancer slices per condition per patient. Multiple explanted tissue slices were cultured for each condition. Cumulative score is based on histological evaluation of entire tissue slice (TS; see Supplementary figure 4). TS were scored in 4 categories: 1) overall quality based on H&E staining, 2) presence of apoptotic cells (cleaved-caspase 3+(c-CASP-3+)/ Keratin+ (KRT+) cells) and 3) cancer cell debris (fragmented Keratin), and 4) nuclear proliferation based on proliferation cell nuclear antiqen (PCNA). TS received a score of either 0 or 1 for each category. For category 1, TS received a score of 1 for >50% of tissue was fragmented/degraded. For category 2 and 3, TS received a score of 1 when multiple clusters were observed. For category 4, TS received a score of 1 when <50% of KRT+ cells in the TS displayed nuclear PCNA. Cumulative score was calculated as the sum of the 4 categories, median cumulative scores are shown. (see also **Supplementary figure 4**) Explanted tumor slices were ex-vivo cultured with vehicle solution or penfluridol (100 µM for A-B and a dose range for C-D) for 3 days. Chi-squared test for categorical data with more than two categories. P value=4,36424E-11. (B) Representative images of bladder cancer slices obtained from a patient diagnosed with NMIBC stage T1 grade 2 (#28) after 3 days in the presence of vehicle solution or penfluridol (100 μM). Complete loss of tumor cells was observed in 5 patient-derived tumor explants (#9,28,29,31 and 35). Significant reduction was found in 30 patient-derived tumor explants (#1-8, 10-15, 17, 20, 22-27, 30, 32-34 and 36-39). A partial response was observed in 3 patient-derived tumor explants (#18,19 and 21). No response was observed in one patient-derived tumor explant (#16). (D) Representative images of bladder cancer tissue slices obtained from a patient diagnosed with NMIBC stage T1 grade 2 (#9) after 3 days in the presence of vehicle solution or a dose range of penfluridol. c-CASP-3 and PCNA: green, panKRT red and DAPI blue, Scalebar 25 um.

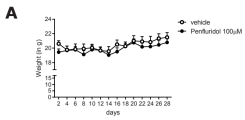
Penfluridol has no effect on normal urothelium.

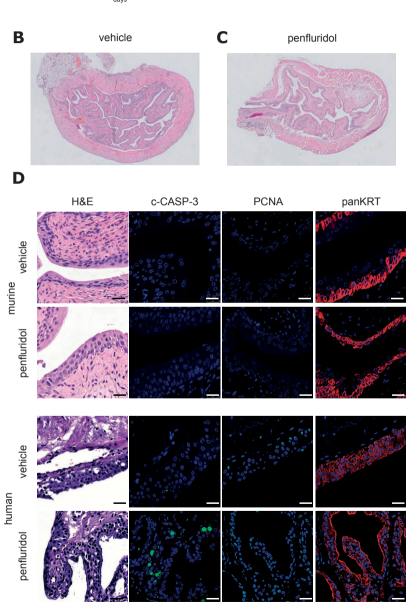
Intravesical treatment represents a potential alternative route of administration of CADs for the treatment of localized bladder cancer. In order to evaluate safety of penfluridol instillations on normal urothelium, a dose-range of penfluridol was administered to non-tumor bearing mice by weekly intravesical treatment (**Figure 5A**). Histological analyses revealed no detectable changes of normal murine urothelium after multiple doses of intravesical penfluridol (**Figure 5B-D**). Moreover, while penfluridol treatment of ex-vivo cultured patient-derived UCB tissues caused significant anti-tumor effects, adjacent 'normal' human urothelium was not or only marginally affected (**Supplementary figure 6A-B**). No changes were observed in epithelial tissue integrity, the proliferation index (PCNA), viable epithelial cell numbers or fragmented KRT, despite a mildly increased apoptotic response (**Supplementary figure 6A**).

Because matched normal urothelium samples are scarce, we assessed the effect of penfluridol on 'normal' urothelium of a prostate cancer patient with no history of bladder carcinoma. Again, no detectable tissue integrity changes were observed in the penfluridol-treated human urothelium versus vehicle treated urothelium. Limited c-CASP-3+ cells were only observed in the penfluridol-treated normal urothelium layers but no detectable changes of urothelium tissue integrity were found (**Figure 5E**).

► Figure 5 Evaluation of the effect of penfluridol on normal murine or human urothelium integrity.

(A) Average body weight (mean +/- SE) of non-tumor bearing female BALB/c nu/nu mice treated intravesically with vehicle (n=5) or penfluridol solution (n=5;100 μ M once weekly for 4 weeks; equivalent to 130 μ g/kg/w). One-way ANOVA. *p<0.05; **p<0.01; ***p<0.001. Representative overview images of H&E stained murine bladders of vehicle-treated (B) and 100 μ M penfluridol-treated (C) mice. (D) Representative images of vehicle- and penfluridol-treated, non-tumor bearing murine bladders. c-CASP-3 and PCNA: green, panKRT red and DAPI blue. Scalebar H&E 20 μ m, IF staining 25 μ m. (E) Representative images of non-transformed 'normal' urothelium in tissue slices obtained from a patient with prostate cancer with no history of bladder carcinoma, treated either with vehicle solution or 100 μ M penfluridol for 3 days (#40). c-CASP-3 and PCNA: green, panKRT red and DAPI blue. Scalebar 25 μ m.





Discussion

In this study, we provide compelling evidence that CADs display significant antitumor effects in multiple preclinical bladder cancer models, i.e. cultured human UCB cells in vitro, an orthotopic preclinical xenograft model of UCB growth and metastasis in vivo, and patient-derived ex-vivo cultured UCB tissues. Screening of eight different CADs in a panel of cultured human UCB cell lines identified penfluridol as a potent candidate anti-cancer agent. We observed that the antitumor properties of penfluridol in human UCB may rely on rapid destabilization of lysosomal structures in human UCB cells. Our data in UCB are in line with the observed anti-neoplastic effects of CADs in other solid tumors (11-17). Other antineoplastic mechanisms of this class of compounds may also play a role, including anti-migratory and invasion effects, cell cycle effects, changes in cholesterol homeostasis, autophagy, immune modulation or non-homologous end-joining (reviewed in (25-27)).

CADs have been shown to induce lysosome-dependent cell death mediated by lysosomal cathepsin proteases leaking from the destabilized lysosomes to the cytosol in other tumors (28-30). Selectivity of the CADs for transformed cells is due to the lower constitutional acid sphingomyelin levels in their lysosomes compared to healthy cells. CADs reduce ASM levels to the point of lysosomal membrane permeabilization which leads to lysosome-dependent cell death specifically in transformed cells (28, 29, 31). Selectivity of CADs for transformed bladder epithelial cells is further substantiated by the observed lack of adverse effects of CAD treatment on normal urothelium, which was not (or only marginally) affected after intra-vesical administration of a same dose of penfluridol in a nontumor bearing in vivo model and administration of penfluridol to ex-vivo cultured normal human bladder tissue. In our preclinical disease model of orthotopically growing human UCB, weekly intravesical instillation with penfluridol significantly reduced tumor progression, invasion and formation of distant metastases. Our preclinical data demonstrate that effective delivery of penfluridol can be achieved by instillation in the bladder thus mimicking the clinical use of standard-of-care pharmacological agents mitomycin C or BCG. This way potential adverse systemic side effects of this class of antipsychotic drugs may be largely avoided.

For UCB patients that present with a disease confined to the mucosa (stage Ta, CIS) or submucosa (stage T1) treatment consists of transuretheral resection of the bladder (TURB) followed by an immediate postoperative intravesical instillation with chemotherapy. In patients who are at intermediate- or high-risk of recurrence, maintenance treatment with intravesical chemo- or immunotherapy is advocated in the guidelines. In general, chemotherapy with mitomycin C or epirubicin for 1 year (4 weekly instillations followed by 6-8 monthly instillations) is indicated in intermediate-risk patients, whereas high-risk patients are treated with 1 to 3 years of Bacillus Calmette-Guérin (BCG) (32). Patients in whom NMIBC recurs after TURB and BCG treatment, and who have BCG unresponsive disease, are unlikely to respond to further BCG therapy and may need a cystectomy. The alternative treatment option is lifelong cystoscopic surveillance and frequent TURB re-operations are necessary. Obvious downsides of radical cystectomy are that it is a major surgical procedure with substantial cost and a negative impact on the quality of life of patients. If feasible, patients often prefer a bladder sparing treatment and new intravesical therapies are being developed in clinical trials (33).

Despite these new developments, only Valrubucin -a doxorubucin-analog- was FDA-approved for BCG-refractory CIS (34). Based on the growing body of evidence and the previously established safety profile of CADs as FDA-approved drugs, penfluridol appears to be a promising anti-cancer compound, especially when systemic exposure is reduced by intravesical administration to patients with (recurrent) high risk bladder cancer. Taken together, our current findings emphasize the need for a clinical trial that will address safety and anti-tumor efficacy aspects of intravesical penfluridol in organ-confined bladder cancer.

Conclusions

Cationic amphiphilic drugs like penfluridol significantly reduced UCB cell viability. Weekly bladder instillations of penfluridol resulted in significant decreased tumor growth, progression and metastasis in vivo. Treatment of cultured patient-derived UCB tissue slices with penfluridol resulted in significant anti-tumor responses ranging from complete loss of tumor to significant reduction. Normal urothelium, when treated with penfluridol was not at all or only marginally affected upon instillation of CADs in a preclinical UCB in vivo model and cultured normal human bladder tissue slices ex-vivo.

Taken together, penfluridol is a promising anti-cancer agent in human UCB and clinical studies are required to determine if this FDA-approved antipsychotic agent can provide a benefit to patients with UCB.

Financial support

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Supplementary table

► Supplementary table 1 key resources

Key reagents and resources used in the manuscript with their respective suppliers, Research Resource Identifiers (RRID) and additional information. Cells were routinely cultured in a humidified incubator at 37°C and 5% CO2 and were regularly (once every 2 months) tested for mycoplasm by RT-PCR. Supplements were from Life Technologies, Gibco. For used antibodies, respective dilutions were added and the indicated time in the pressure cooker (PC) when antigen retrieval was performed by cooking the slides in unmasking solution (Vector Labs, H-3300); N.A. not applicable.

Supplementary table 1 key resources.

Supplements were from Life Technologies, Gibco. For used antibodies, respective dilutions were added and the indicated time in the pressure cooker (PC) Cells were routinely cultured in a humidified incubator at 37°C and \$5% CO2 and were regularly (once every 2 months) tested for mycoplasm by RT-PCR. Key reagents and resources used in the manuscript with their respective suppliers, Research Resource Identifiers (RRID) and additional information. when antigen retrieval was performed by cooking the slides in unmasking solution (Vector Labs, H-3300); N.A. not applicable.

		Experimental models: cell lines	lels: cell lines		
Cell line	Source	Identifier	Medium	Supplier	Supplements
			Eagle's		
		0 × ++0 0 0 × ++0 0 0 × ++0 0 0 0 × 0 ×	minimal		10% FBS , 100 units/
UM-UC-3	ATCC	AICC Cat# CRE-1/49,	essential	ATCC, 30-2003	ml penicillin, 50 µg/ml
		NAID.CVCE_1703	medium		streptomycin
			(EMEM)		
			Dulbecco's		, of TD 000
C	() ()	ATCC Cat# HTB-1,	Modified	Life technologies,	10% FBS , 100 uffits/
J82	AICC	RRID:CVCL_0359	Eagle medium	Gibco, 31966-021	mi penicilin, 50 µg/mi
			(DMEM)		srreptornycin
		# +*** J J J L V			10% FBS , 100 units/
T24	ATCC	AICC Cat# 1115-4,	RPMI 1640	Lonza, BE12-167F	ml penicillin, 50 µg/ml
		RKID: CVC F_0334			streptomycin, GLUTAMAX

Supplementary table 1 Continued.

		Experimental models: cell lines	els: cell lines		
Cell line	Source	Identifier	Medium	Supplier	Supplements
			Eagle's		
		3 GFT #+~C CCEV	minimal		10% FBS , 100 units/
TCCSUP	ATCC	AICC Cat# 1118-3,	essential	ATCC, 30-2003	ml penicillin, 50 µg/ml
		RAID.CVCL_1/30	medium		streptomycin
			(EMEM)		
	Cell lines		Dulbecco's		/otion: 001 303 %001
PT 113	service.	CLS Cat# 300324/p10280_	Modified	Life technologies,	IU% rbs , IUU uliits/
NI_112	https://	RT-112, RRID:CVCL_1670	Eagle medium	Gibco, 31966-021	riii peinciiiii, oo pg/iiii
	clsgmbh.de		(DMEM)		scieptoliiyciii
		0-8TH #+c7 77TV			10% FBS , 100 units/
5637	ATCC	PPID: CVC 0126	RPMI 1640	Lonza, BE12-167F	ml penicillin, 50 µg/ml
					streptomycin, GLUTAMAX
		C GTU #+~O OOTV		Thermo Fisher	10% FBS , 100 units/
RT-4	ATCC	PRID: CVCI 0036	McCoy's 5A	Scientific Invitrogen	ml penicillin, 50 µg/ml
		RAID.CVCE_0030		36600-021	streptomycin, GLUTAMAX

Supplementary table 1 Continued.

		Experimental models: organisms and strains	ganisms and stra	ins	
Cell line	Source	Identifier	Gender	Ethical committee	Ethical protocol
				Committee on the Ethics	
Female balb c nude mice	Cildres River	RRID:		of Animal Experiments	7
(CAnN.Cg-Foxn1nu/Crl)	CCF 002702	MGI:2160479	מפופ	of Leiden University, The	DEC_14212
	SCK_003/92			Netherlands	
		Experimental models: biological samples	biological sample:	S	
Biological sample	Source	Identifier	Gender	Ethical committee	Ethical protocol
Human bladder samples	Erasmus Medical				
transurethral resection of	Center bladder cancer	Coded samples			MEC-2014-553
the bladder (TURB)	biobank				

Supplementary table 1 Continued.

		Antibodies		
Antibody	Source	Identifier	Dilution	PC
Mouse-anti human Proliferating cell nuclear antigen (PCNA)	Sigma-Aldrich, P8825	RRID: AB_477413	1:2000	40′
Rabbit anti-human Pan cytokeratin (panKRT)	Life technologies, 180059	RRID: AB_10942225	1:250	40′
Guinea Pig anti-murine Pan cytokeratin (panKRT)	Origene, BP5069	RRID: AB_979529	1:250	40′
Goat anti-human Collagen I Southerenbiotech, 1310-01	Southerenbiotech, 1310-01	RRID: AB_2753206	1:1000	40′
Rabbit anti-human Cleaved caspase-3 (c-CASP-3)	Cell signaling, 9661L	RRID: AB_2341188	1:500	40′
Mouse anti-human Keratin 18 (KRT18)	Dako, M7010	RRID:AB_2133299	1:500	40′
Mouse anti-human LAMP-1	Abcam ab25630	RRID:AB_470708	1:20	N.A.

Supplementary table 1 Continued.

		Antibodies		
Antibody	Source	Identifier	Dilution	PC
Rabbit anti-human Galectin-1 (LGALS1)	AbCaM ab25138	, RRID:AB_2136615	1:2000	N.A.
Donkey-anti Mouse Alexa Fluor 488	Life technologies, R37114	RRID:AB_2556542	1:250	N.A.
Donkey-anti Rabbit Alexa Fluor 488	Life technologies, A-21206	RRID:AB_141708	1:250	N.A.
Donkey-anti Mouse Alexa Fluor 555	Life technologies, A-31570	RRID:AB_2536180	1:250	N.A.
Donkey-anti Rabbit Alexa Fluor 555	Life technologies, A-31572	RRID:AB_162543	1:250	N.A.
Donkey-anti Goat Alexa Fluor 647	Life technologies, A-31572	RRID:AB_141844	1:250	N.A.
Goat-anti Guinea Pig Alexa Fluor 647	Life technologies, A-21450	RRID:AB_2735091	1:250	N.A.

Supplementary table 1 Continued.

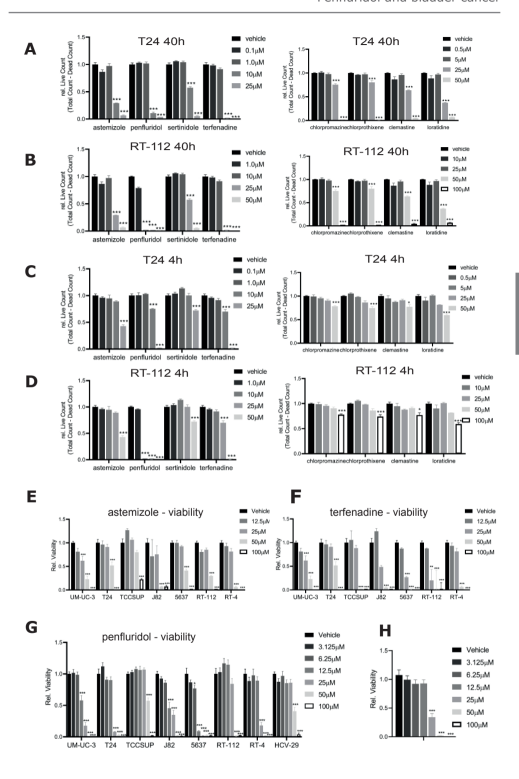
	Chemicals	
Chemical	Source	Identifier
	Sigma-Aldrich	D2271
loniin di	RRID:SCR_008988	F35/1
مامدنسمهم	Sigma-Aldrich	19861
Astellizole	RRID:SCR_008988	A2001
Ciberofic	Sigma-Aldrich	T0652
ים ופוופ	RRID:SCR_008988	19032
;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;	Sigma-Aldrich	28033
ספורווומסוע	RRID:SCR_008988	2/00/2
, id	Sigma-Aldrich	C1671
מוואלוווא	RRID:SCR_008988	C10/1
3	Sigma-Aldrich	00130
CITIOT PLOTINGE	RRID:SCR_008988	00130
Compotino	Sigma-Aldrich	SMI 0445
Circiliastille	RRID:SCR_008988	SML0443
4470	Sigma-Aldrich	19661
Lorataume	RRID:SCR_008988	L9004

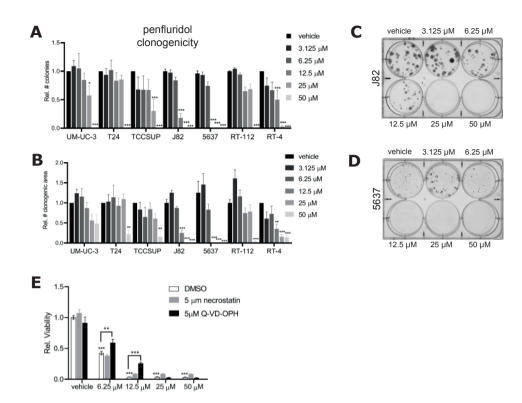
Supplementary table 1 Continued.

	Critical commercial assays	
Assay	Source	Identifier
CellTiter 96® AQueous		
One Solution Cell	Promega RRID:SCR_006724	G3581
Proliferation Assay		
RealTime-Glo [™] Annexin		
V Apoptosis and Necrosis	Promega RRID:SCR_006724	JA1012
Assay		
	Critical equipment and software	
Assay	Source	Identifier
Celígo® Imaging	Nevcelom Bioscience	
Cytometer		
SpextraMax iD3	MolecularDevices	
Image J version 1.48v	https://imagej.net/	RRID:SCR_003070
	Leica Microsystems	
Colliocal ICS_SPo	RRID:SCR_008960	1
	Critical equipment and software	
Assay	Source	Identifier
IVIS Lumina Imaging	۸۵۱ مصورت کی فار سرمارات	
System	Calipel LifeSciences, USA	1
MIDI slidescanner	Pannoramic	1
Caseviewer	3DHistech	RRID:SCR_017654

▶ Supplementary figure 1 CADs reduced viability in a panel of human bladder cancer cells.

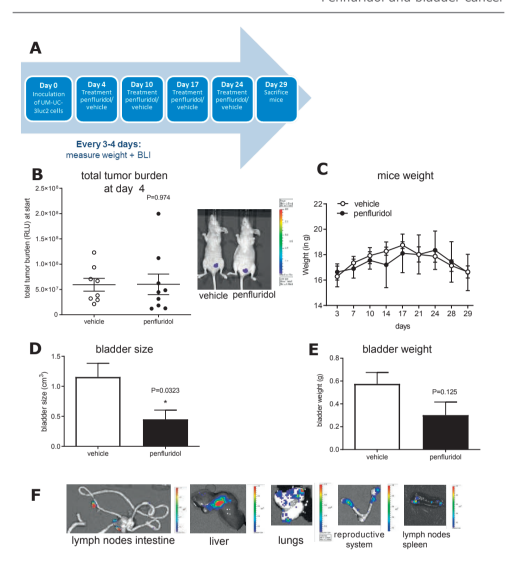
Assessment of the viability of T24 ($\bf A$) and RT-112 ($\bf B$) cells after treatment with a dose range of 8 different CADs for 40h. Assessment of the viability of T24 ($\bf C$) and RT-112 ($\bf D$) cells after 4hrs treatment. CADs were replaced by regular growth medium after 4h and 40h later live cell count was quantified in a Celigo(R) cytometer as the total cell count (Hoechst-positive cells) - dead cell count (Propidium Iodide-positive cells). Mean +/- SE normalized to vehicle treated cells. One-way ANOVA *p<0.05; **p<0.01; ***p<0.001 Multiple subconfluent UCB cells were treated for 2h with a dose range of either astemizole ($\bf E$), terfenadine ($\bf F$) or penfluridol ($\bf G$). Viability was measured using MTT assay after 48h. Mean +/- SE normalized to respective vehicle treated UCB cells. ($\bf H$) Confluent UM-UC-3 cells were treated for 2h with a dose range of penfluridol. Viability was measured using MTT assay after 48h. Mean +/- SE normalized to vehicle treated cells. ($\bf n=3$; 6 replicates each). One-way ANOVA *p<0.05; **p<0.01; ***p<0.001.





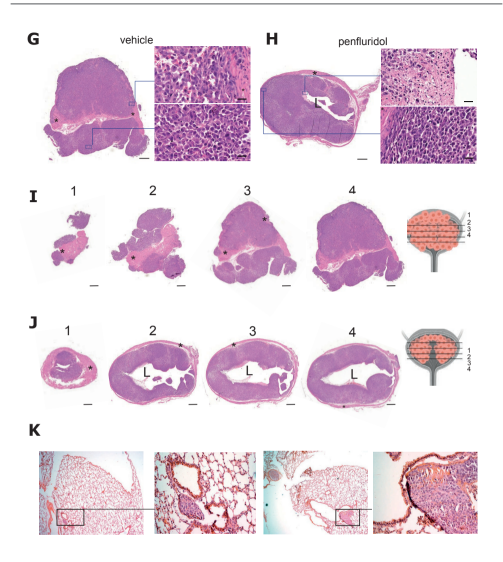
Supplementary figure 2 CADs reduced viability and clonogenicity in a panel of human bladder cancer cells.

Clonogenic assay: multiple bladder cancer cells were treated for 2h with a dose range of penfluridol and subsequently after 10-14 days, (\mathbf{A}) number of colonies and (\mathbf{B}) clonogenic area were measured using Image J. Representative images of clonogenic assay of respectively J82 (\mathbf{C}) and 5637 (\mathbf{D}) cells. (n=3; 3 replicates each). One-Way ANOVA. *p<0.05; **p<0.01; ***p<0.001. (\mathbf{E}) Mitochondrial activity was measured after 24h in response to treatment with a dose range of penfluridol in combination with either necroptosis inhibitor NS1 or pan-caspase inhibitor Q-VD-OPh. Mean+/- SE normalized to vehicle treated UM-UC-3 cells (n=3; 6 replicates each). One-way ANOVA. *p<0.05; **p<0.01; ***p<0.001.



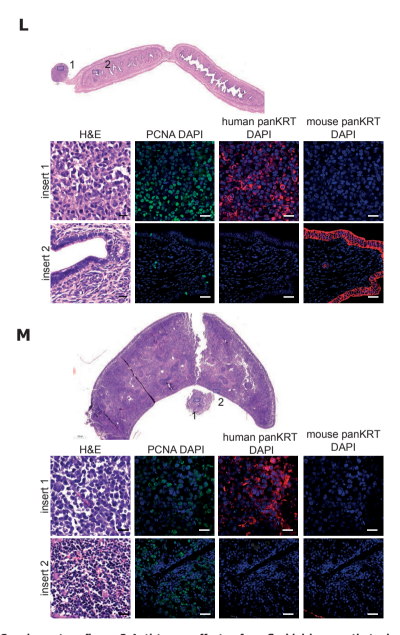
Supplementary figure 3 Anti-tumor effects of penfluridol in an orthotopic murine xenograft model with stable firefly luciferase-2 UM-UC-3 human bladder cancer cells.

Female BALB/C nu/nu mice were inoculated with UM-UC-3 luciferase 2 bladder cancer cells and intravesically treated with vehicle (n=8) or penfluridol (n=9; 100 μ M once weekly; equivalent to 130 μ g/kg/w). (A) Schematic overview of the experiment. (B) Mice were divided into 2 groups with equal tumor burden based on total tumor burden; representative BLI images of mice are shown. (C) Weight of the mice over time. (D) Bladder size in cm3 at day 29. (E) Bladder weight in g at day 29. (F) Representative BLI images of metastasis in intestinal lymph nodes, liver, spleen, reproductive system, and lungs at day 29. Mann-Whitney-U test. *p<0.05; **p<0.01; ***p<0.001.



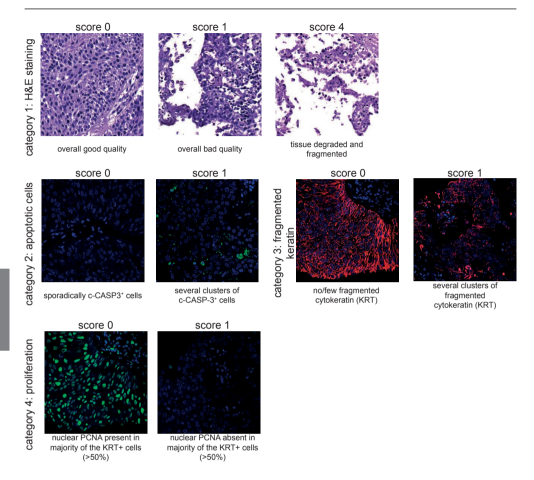
Supplementary figure 3 Anti-tumor effects of penfluridol in an orthotopic murine xenograft model with stable firefly luciferase-2 UM-UC-3 human bladder cancer cells.

Representative images of vehicle (**G**) and penfluridol (**H**) treated bladders stained with Hematoxylin & Eosin with respective magnifications. Scale bar 500 and 20 μ m respectively. (**I**) Representative images of vehicle treated bladders stained with Hematoxylin & Eosin at different planes of sectioning displaying muscle-invasion. Scale bar 500 μ m. Schematic overview of the bladders is shown next to the different planes (created with Biorender.com).(**J**) Representative images of penfluridol treated bladders stained with Hematoxylin & Eosin at different planes of sectioning. Scale bar 500 μ m. (**K**) Representative Hematoxylin & Eosin images of a metastasis in a lung. L: lumen; * muscle layer.



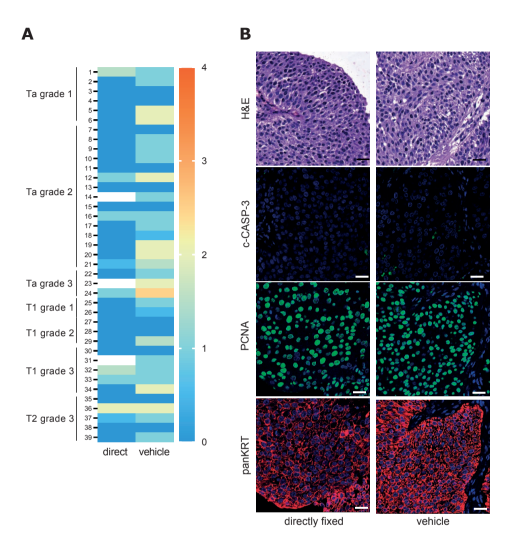
Supplementary figure 3 Anti-tumor effects of penfluridol in an orthotopic murine xenograft model with stable firefly luciferase-2 UM-UC-3 human bladder cancer cells.

(L) Representative image of a metastasis in a lymph node (adjacent to spleen). (M) Representative image of a metastasis in the reproductive system. Scale bar for overview 500mm. PCNA: green, panKRT (human or mouse): red and DAPI blue. Scalebar 25 μ m.



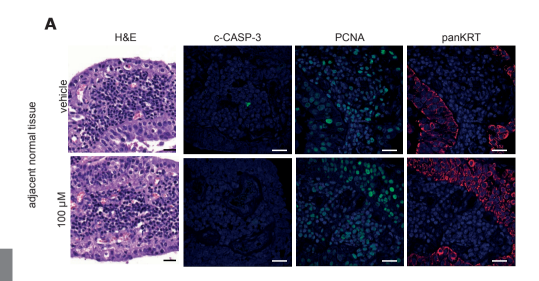
Supplementary figure 4 Scoring of ex-vivo cultured human bladder cancer slices.

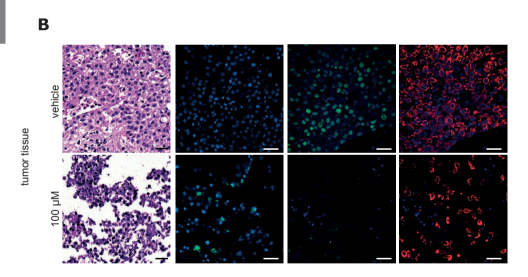
Multiple explanted tissue slices were cultured for each condition. Cumulative score is based on histological evaluation of entire tissue slice. Representative images are shown. First, overall quality of the TS was scored based on H&E staining (category 1). TS received a score of either 0, 1 or 4. TS received a score of 1 when overall quality was poor in >50% of the H&E stained TS. The TS scored the maximum score of 4 when tissue was completely degraded or fragmented. When TS did not already directly receive a score of 4 for overall quality based on the H&E stained tissue, sections were subsequently scored for 3 additional categories: 2) presence of cleaved-caspase 3+(c-CASP-3+)/Keratin+ (KRT+) cells, 3) fragmented KRT, and 4) nuclear proliferation based on proliferation cell nuclear antigen (PCNA). TS received a score of either 0 or 1 for each category. For category 2 and 3, TS received a score of 1 when multiple clusters were observed. For category 4, TS received a score of 1 when <50% of KRT+ cells in the TS displayed nuclear PCNA. Cumulative score was calculated as the sum of the 4 categories, median cumulative scores are shown. c-CASP-3 and PCNA: green, panKRT red and DAPI blue.



Supplementary figure 5 Ex-vivo treatment of cultured human bladder cancer slices with penfluridol: directly fixed vs cultured tissue.

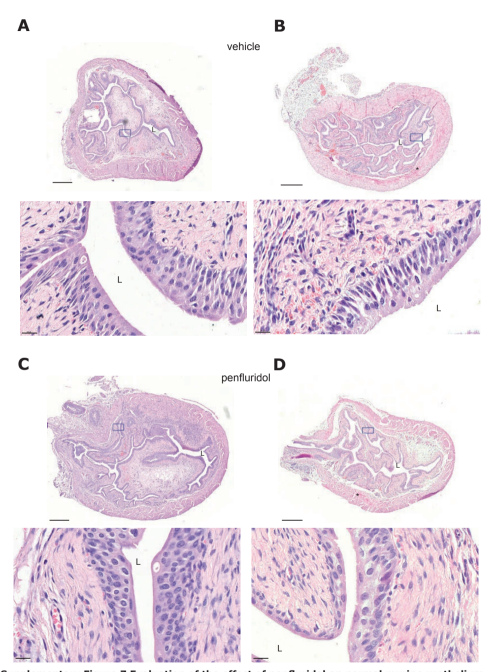
(A) heat map showing the median scores of the bladder cancer slices per patient (sum of the scores based on PCNA, c-CASP-3, fragmented KRT and H&E staining). Bladder cancer slices from directly fixed and ex vivo cultured in vehicle solution for 3 days. (B) representative images of bladder cancer tissue obtained from a patient diagnosed with NMIBC stage T1 grade 2 (#28) directly fixed or cultured for 3 days in the presence of vehicle solution. c-CASP-3 and PCNA: green, panKRT red and DAPI blue. Scalebar 25 µm.





Supplementary figure 6 Evaluation of the effect of penfluridol on normal human urothelium.

(A) Representative images of urothelium in tissue slices obtained from adjacent normal tissue from a bladder cancer patient with NMIBC T1 grade 2 (#29). (B) Representative images of tissue slices from tumor tissue from this patient (#29). H&E, c-CASP-3 and PCNA: green, panKRT red and DAPI blue. Scalebar 25 µm.



Supplementary Figure 7 Evaluation of the effect of penfluridol on normal murine urothelium. (A) Representative overview images of H&E stained non-tumor bearing murine bladders of vehicle-treated (A-B) and 100 μ M penfluridol-treated (C-D) mice. L: lumen; * muscle layer

Supplementary information

Cationic Amphiphilic Drugs

Penfluridol, astemizole and terfenadine were obtained from Sigma-Aldrich®. Penfluridol (P3371) and terfenadine (T9652) were diluted in Absolute EtOH (EMSURE®) to a stock solution of 50 mM. Likewise, astemizole (A2861), sertinidole (S8072), chlorprothixene (C1671), chlorpotmazine (C8138), clemastine (SML0445) and loratadine (L9664) were diluted in dimethyl sulfoxide (DMSO; Sigma-Aldrich®) to a stock solution of 50 mM (**Supplementary table 2**). Serial dilutions were established by diluting these stock solutions in the cell-specific medium of the various cell lines (**Supplementary table 1**).

In vitro Viability assay

Cells were seeded in a 96-wells plate at a concentration of 1500 single cells per well with 6 technical replicates. Cells were either treated after 24h, or cells were allowed to grow until confluency and subsequently treated for 2h with a dose range. After replacing the medium, cells were incubated at 37°C for an additional 24, 48 or 72h. Subsequently, 20µl of MTS reagent was added and after 2 hours, the OD was measured at 490nm using the VersaMax ELISA Microplate Reader. Apoptotic nuclear condensation was evaluated in Hoechst-33342 stained cells using Olympus IX81 microscope with a 20 \times Olympus objective, Scan^R automated acquisition software (version 2.3.0.5) and analysis with ImageJ (version 1.48v).

In vitro clonogenicity

100-single cells were seeded in a 6-wells plate. After 24h, cells were treated with a dose-range of penfluridol for 2h. After replacing medium, cells were incubated at 37°C. After ~14 days, cells were fixed with 4%PFA and stained for 10 minutes with 5% crystal-violet. The number of colonies and colony area were calculated with ImageJ software.

In vitro Lysosomal membrane permeabilization

60000 cells were seeded on PET-coated chamber slides and left for 24h to adhere. Cells were treated with a dose-range of penfluridol for 2h. After 24h lysosomal membrane permeabilization was detected by staining paraformaldehyde-fixed cells with LGALS1 and LAMP-1 primary antibodies (**Supplementary table 1**) at 4°C. DAPI staining was used to visualized the nuclei. Next, the slides were stained for 1.5h at RT with DaM 488 secondary antibody and mounted with prolong Gold antifade. Confocal images were taken with the Confocal TCS_SP8 (Leica).

Near-patient ex vivo bladder cancer cultures

Tumor tissues from patients diagnosed with various stages of UCB were obtained during transurethral resection of the bladder upon informed consent (MEC-2014-553). Explanted tumor tissue slices (TS) were sectioned and cultured as previously described (20, 21). Bladder cancer material was transported in EMEM medium at RT, transferred into Penicillin Streptomycin solution for 10 minutes and cut into consecutive tissue slices of approximately 500-1000 μ m thick (20). The tissues slices were placed on nitrocellulose filter inserts (6 well filter inserts, pore size of 3 μ m, Corning Costar) and cultured with EMEM, supplemented with 10% FCS and 1% Pen-Strep. The tissue was cultured in an oxygenated and sealed system. After 24 hours, the tissue slices were treated with either vehicle solution or penfluridol (depending on the amount of tissue either 100 μ M or a dose range). At 3 days post treatment, the tissue was fixed in 4% PFA for 1 hour at RT and transferred into a plastic cassette and stored in 70% ethanol at 4°C prior to paraffin embedding.

Histology

Haematoxylin and eosin staining (H&E) was performed according to a standard protocol to assess general histology. For immunofluorescence staining, deparaffinization was performed via incubation in Histoclear (National Diagnostics, HS-200) and rehydrated by incubation in a series of decreasing concentrations of ethanol. Antigen retrieval was performed by cooking the slides in unmasking solution (Vector Labs, H-3300) in the pressure cooker. After blocking with 1% BSA (Sigma-Aldrich, A7906) for 30 minutes, the slides were incubated overnight with primary antibody at 4°C. Next, the slides were stained for 1.5 hours at RT with secondary antibody and mounted with prolong Gold antifade (Molecular probes, Thermofisher Scientific, P36930) (**Supplementary table 1**).

Orthotopic inoculation tumour cells

- -Anaesthetize mouse with isoflurane (see below), induce miction by administering mild abdominal pressure.
- -Administer Temgesic according protocol. (see below)
- -Place the mouse ventrally on a cardboard piece
- -Tape the tail to the cardboard
- Remove the catheter (Jelco I.V. Catheter Radiopaque 24G, ¾G, Smiths medical ref 4053) from the needle and put some instilla gel (Farco-PHARMA GmbH Koln, Germany) on the catheter.
- Hold the urethra with a pair of tweezers and gently insert the catheter into the urethra. If you feel some resistance, pull back and try a different angle. The catheter should enter the urethra easily.

- Insert 1 cm of the catheter and tape the catheter on top of the tail.
- Rinse the bladder with 1 ml PBS (Check if the liquid has been equilibrated to room or body temperature)
- -Fill a syringe with HCl (0.1M)and place the needle on it. Inject approx. 100 ul into the bladder
- Stir, using a back and forth motion of the syringe for 15" (1 stroke/sec) and remove the HCl.
- Inject 100 ul KOH (0.1M) to neutralize the HCl.
- Stir, using a back and forth motion of the syringe for $15^{\prime\prime}$ (1 stroke/sec) and remove the KOH
- Inject cell suspension (5*10^6 in 35 μl PBS)
- Close the urethra by binding it off using a suture (Prolene Ethicon 6-0 13mm 3/8c). Let someone remove the catheter while binding off.
- Remove tape
- Keep the mouse anaesthetized with his lower body lifted slightly up for 2 hours
- Cut suture open before ending the anaesthesia.

Administration of Analgesia (Temgesic)

- Prepare temgesic solution (dosage: 0.1 mg/kg buprenorphine s.c. every 8 hours as needed)
- Check if the liquid has been equilibrated to room or body temperature.
- Fill the syringe with the substance that needs to be injected. Make sure there are no air bubbles in the syringe and that the needle is filled with fluid.
- Take the animal from out of the cage.
- Hold the scruff of the animal between thumb and forefinger and restrain the tail.
- Disinfect the injection place with a cotton swab wetted with ethanol 70%.
- Carefully insert the needle, between thumb and forefinger and into the center of the triangular shaped section of the scruff. Keep the needle and syringe parallel to the mouse's head and back.
- Carefully and slowly insert the substance by pushing down the plunger. You can feel the bulging of the substance between your fingers.
- Carefully remove the needle from the scruff of the animal.
- Check the injection site for liquid flowing back or blood loss. Do not return the mouse to its cage unless the injection site is clean and dry. Place the animal back into the cage. Assure yourself of the proper/ normal movement of the animal
- Check your animal for signs of pain or discomfort at least 6-8 hours after initial dose to determine if there is the need for additional analgesia

Intravesical Instillation

- Anaesthesize mouse with isoflurane (see below)
- Administer Temgesic according to protocol. (see below)
- Place the mouse ventrally on a cardboard piece
- Remove the catheter (Jelco I.V. Catheter Radiopaque 24G, ¾G, Smiths medical ref 4053) from the needle and put some instilla gel (Farco-PHARMA GmbH Koln, Germany) on the catheter.
- Prior to the rinsing, the bladder will be emptied by mild abdominal massage, to induce a better contact between compound and urothelial cells.
- Hold the urethra with a pair of tweezers and gently insert the catheter into the urethra. If you feel some resistance, pull back and try a different angle. The catheter should enter the urethra easily.
- Insert 1 cm of the catheter and tape the catheter on top of the tail.
- Rinse the bladder with 1 ml PBS (Check if the liquid has been equilibrated to room or body temperature)
- Tie a suture around the urethra, but do not pull tight yet.
- Insert the needle and inject 50 µl of the compound.
- Tighten the suture, while a second person, first removes the needle. Directly followed by the catheter.
- During 1 hour, the mice are regularly turned to contact the whole bladder surface with the compound.
- After 1 hours the suture will be removed and the bladder will be emptied by mild abdominal compression.
- Mice will be daily checked for weight, behavioural changes and blood in urine.

Anaesthesia

- Isoflurane: induction phase: 2 4 %, maintenance phase: 0.25 2%
- Airflow: induction phase: 0.8 L/min, maintenance phase: 0.4 L/min
- When operation takes longer than 15 min. the airflow (from DL meter) needs to be enriched with O2 till 30-40%.

Bioluminescence measurements

Outgrowth of spread of the tumour cells was monitored (number of measurements depending on the experiment) by whole body bioluminescent imaging (BLI) using an intensified charge-coupled device (CCD) video camera in a light-tight specimen box of the *in vivo* Imaging System (IVIS Lumina). Animals were anesthetized with isoflurane and were given D-luciferin (15 mg/ml in D-PBS; 25 μ l/10 g body weight) by intraperitoneal injection. Mice were placed into the light-tight camera box with continuous exposure to isoflurane. Imaging time ranged from 10 s to

1 min (generally 30s), depending on the tumor model, 5 min after D-luciferin injection. Three mice were imaged each time from a dorsal and a ventral view. The photons emitted from the bioluminescent tumours or cells were detected by the IVIS camera system, integrated, digitized and displayed. When saturated pictures are taken, first the F-stop is adjusted (diaphragm). If the image is still saturated, the image time can be adjusted.

Intraperitoneal injections (luciferin)

- Check if the liquid has been equilibrated to room or body temperature.
- Fill the syringe with the substance that needs to be injected. Make sure that there are no air bubbles in the syringe and that the needle is filled with fluid.
- Take the animal from out the cage
- Restrain the animal. Hold the animal with the belly facing up, tilting the animal in a head down position.
- Disinfect the injection place with a gauze wetted with ethanol 70%.
- Insert the needle, with the bevel facing up, into the animal's lower right abdominal quadrant, avoiding the abdominal midline so that you will not inject into the urinary bladder
- Carefully retract the plunger to check if you haven't hit a blood vessel
- With no blood in the syringe gently inject the substance.
- Carefully remove the needle from the animal
- Check the injection site for liquid flowing back or blood loss.
- Place the animal back into the cage. Check for proper/ normal movement of the animal.

Cervical dislocation

- Anaesthetize the mouse with isoflurane.
- Place the thumb and first finger of the other hand against the back of the neck at the base of the skull
- To produce the dislocation, quickly push forward and downward with the hand or object restraining the head, while pulling backward with the hand holding the tail base
- The effectiveness of dislocation can be verified by feeling for a separation of cervical tissues. When the spinal cord is severed, a 2-4 mm space will be palpable between the occipital condyles and the first cervical vertebra. Occasionally, however, the dislocation occurs between thoracic vertebrae. Check closely to confirm respiratory arrest and when possible, verify by palpation, that there is no heart beat.

Quantification of signals

Quantification of signals was performed by the Living Image® (Xenogen) software. Values were expressed as RLUs in photons/second. Numbers of metastases per animal were counted by eye from a dorsal and ventral view.