A modified random oligonucleotide-based combination therapy for adjuvant treatment of pancreatic ductal adenocarcinoma

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Abstract. Anti-cancer therapy in pancreatic ductal adenocarcinoma (PDAC) is mostly based on surgical removal or palliative therapy using antimetabolites, like 5'-fluorouracil or gemcitabine. Adjuvant treatment using these chemotherapeutics has recently proven a beneficial concept, though general survival rates are still poor. Most recently, combination therapy of gemcitabine with other targeted drugs was evaluated in clinical trials. We present here a study performed in a mouse orthotopic xenotransplant model of PDAC, using an oligonucleotide-based approach. We have shown previously that antisense oligonucleotides against p53 reduce the weight of orthotopic pancreatic tumours in immune-deficient mice. We further characterised terminal modifications of phosphorothioate oligonucleotides in vitro and found a random, unrelated control sequence carrying a D,L-α-tocopherol modification at the 5' and 3' ends to be most efficient in induction of cell death in PancTu-1 cells. Modified random oligonucleotide (MRON) were thus further tested in vivo. MRON showed a reduction of tumour weight in established primary orthotopic tumours in SCID/bg mice. In a surgically adapted pre-clinical model, where primary tumours were resected and animals received adjuvant treatment, MRON was very efficient in suppression of relapse and metastasis, when combined with gemcitabine. While the exact molecular mechanism of MRON activity still needs to be elucidated, the compound showed a remarkable preference for uptake into tumour cells in vivo.

Introduction

Ductal adenocarcinoma of the pancreas is currently the fifth leading cause of cancer-related deaths in the United States (1). Despite all efforts, which have been made to improve the efficacy of treatment, the prognosis of pancreatic adenocarcinoma remains poor. The only curative option consists of radical surgical resection with lymphadenectomy, by which a 5-year survival rate of 6-20% (2-6) and, rarely, up to 30% (7) can be achieved. But even if no detectable residual tumour is left behind (R0), there is a high rate of local recurrence, which accounts for >70% of relapses (8). Therefore, the need for new therapeutic strategies is evident even for an adjuvant situation. One such new strategy would be to address genetic alterations associated with pancreatic adenocarcinoma. Mutation of the p53 tumour suppressor gene is one of the most frequent genetic alterations during development of malignancies (9,10). Thus, an approach to address the expression and function of mutated p53, also representing gain of function mutations (11), in pancreatic adenocarcinoma using antisense oligonucleotides represents such a new therapeutic strategy. Antisense p53-based therapy has been used in the treatment of certain leukaemia (12). We have investigated a well-studied p53 antisense sequence, directed against exon 10 of the p53 gene, and have shown therapeutic effects in a murine orthotopic xenotransplant model (13). When studying the effectiveness of p53 antisense therapy, different terminal modifications of antisense oligonucleotides were designed to increase stability and cellular uptake. The differently modified p53 antisense oligodeoxynucleotides (ODN) were compared to scrambled p53 antisense sequences and an unrelated random control sequence. Tests were performed in vitro for impact on DNA synthesis and induction of apoptosis in PancTu-1 cells. Upon comparison of the results, the unrelated random sequence, carrying a double D,L-α-tocopherol terminal modification, called MRON, proved to be most efficient and was thus investigated in more detail using a murine orthotopic xenotransplant model in vivo.

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Abbreviations: MRON, modified random oligonucleotide; PDAC, pancreatic ductal adenocarcinoma; ODN, oligodeoxynucleotide; NF-κB, nuclear factor-κB; SCID/bg, severe combined immuno-deficiency/beige mutation; scr, scrambled

Key words: oligonucleotide therapy, adjuvant treatment, pancreatic ductal adenocarcinoma
Materials and methods

Laboratory animals. Four-week-old female SCID beige (SCID/bg) mice weighing 14–19 g were obtained from Charles River Laboratories (Sulsel, Germany). The mice were allowed to become acclimatized for 1 week in a sterile environment where bedding, food and water were autoclaved. Animal experiments and care were in accordance with the guidelines of the institutional authorities (no. 252-722-41.121-7).

Cell culture. The cells used in this study were PancTu-1 (14,15) human pancreatic adenocarcinoma cells. They were routinely cultured in RPMI-1640 medium (Invitrogen, Germany), supplemented with 10% heat-inactivated fetal calf serum (FCS, PAN-Systems, Aidenbach, Germany), 2 mM glutamine, and 1 mM sodium pyruvate (Invitrogen, Germany) (standard culture medium). The cells were maintained in monolayer culture at 37°C in a humid atmosphere with 5% CO₂.

DNA-synthesis (methyl-[^3H]-thymidine-incorporation assay). Cells were plated in 96-well culture dishes (Nunc, Wiesbaden, Germany) at a density of 10,000 cells per well in standard culture medium. After 24 h, the medium was replaced with medium containing only 0.5% FCS (starvation medium) and incubated for another 24 h. Cells were then treated with 2.5 μM ODN in medium containing only 0.5% FCS (starvation medium) and incubated for another 24 h. Cells were then treated with 2.5 mM or 5 μM ODN in medium with 0.5% FCS without transfection reagents for 24 or 48 h. During the last 3 h of this period, a total amount of 7.4 kBq methyl-[^3H]-thymidine per well was added. Cells were plated in 96-well culture dishes at a density of 10,000 cells per well in standard culture medium. After 24 h medium was replaced 96-well culture dishes at a density of 10,000 cells per well in DNA-synthesis (methyl-[^3H]-thymidine-incorporation assay). Cells were plated in 96-well culture dishes (Nunc, Wiesbaden, Germany) at a density of 10,000 cells per well in standard culture medium. After 24 h, the medium was replaced with medium containing only 0.5% FCS (starvation medium) and incubated for another 24 h. Cells were then treated with 2.5 μM or 5 μM ODN in medium with 0.5% FCS without transfection reagents for 24 or 48 h. Cells were collected using a cell-harvester (Skatron, Lier, Norway) and incorporated radioactivity was determined by liquid scintillation counting.

Electrophoretic mobility shift assay (EMSA). Nuclear extracts from MRON-stimulated PancTu-1 cells were prepared essentially as described previously (17).

Morphological analysis of in vivo distribution. For analysis of systemic distribution of MRON, 3 mg/kg Cy3-labelled MRON was administered i.p. to 2 tumour-bearing mice which were sacrificed 3 and 24 h after injection. Organ samples were frozen in liquid nitrogen and cryosections of 5-10 μm were prepared and post-fixed for 1 h in 2.5% paraformaldehyde. For reduction of autofluorescence, sections were treated with 1% sudan black in 70% ethanol and subsequently subjected to counterstain with bis-benzimide and Oregon Green 488-labelled WGA as described previously (13). Analysis of tissue sections was carried out using a Zeiss Axioptot equipped with an Axioscan digital camera (Carl Zeiss Jena, Jena, Germany).

Orthotopic xenograft model. The human pancreatic adenocarcinoma cell line, PancTu-1, forms a highly invasive pancreatic adenocarcinoma when injected orthotopically into SCID/bg mice, as has also been described previously (13,18). For orthotopic injection, PancTu-1 cells were trypsinized and suspended in serum-free RPMI-1640 medium and stored on ice. General anaesthesia was induced using fentanyl (0.05 mg/kg), midazolam (1 mg), medetomidin (0.5 mg/kg). A median laparotomy was performed and the pancreas identified. One million tumour cells (volume 30 μl) were injected into the body of the pancreas. The abdominal wall was closed using Vicryl 6/0 (Ethicon, Germany). The recovery was carefully supervised with red light re-warming and volume substitution. Tumour formation and growth was allowed for 8 days before the start of conservative therapy or tumour resection.

Conservative therapy of orthotopic tumours. Tumour-bearing mice were randomly assigned to either one of the different treatment or control groups. The mice in the treatment groups either received daily i.p. injection of the ODN or injections followed an interval scheme (alternating 3 days with and without treatment). Single doses varied from 1 mg/kg body weight (BW) per day up to 3 mg/kg BW per day. Total doses ranged from 24 mg/kg up to 72 mg/kg. The control group received a daily injection of 500 μl PBS intra-peritoneally. Animals were sacrificed after 24 days of treatment (32 days after tumour cell inoculation). The tumours in the pancreas were removed and weighed, length, width and height were measured and the tumour volume was calculated using the formula recommended in a meta-analysis by Tomayko and Reynolds (19). Furthermore, organs were investigated for formation of metastases macroscopically and by histology of selected sections.

Adjuvant therapy of resected animals. Tumour resection was carried out essentially as described previously (20). Animals were allowed to recover from resection for 2 days before start of adjuvant therapy. Control animals received PBS i.p. MRON only was applied at 3 mg/kg BW at 3 day intervals. Gemcitabine only was applied i.p. at 120 mg/kg BW twice a week. Combined therapy of MRON and gemcitabine was 3 mg/kg BW MRON at 3 day intervals with gemcitabine at 120 mg/ kg BW injected the day after MRON treatment.
In vitro data were analysed for homogeneity of variance (ANOVA) using the Levene test. In case of heterogeneity of variance, post-hoc test Dunnett T3 was employed for analysis of significance. In vivo data were subjected to Student’s t-test.

Results

Efficiency of inhibition of DNA synthesis by modified oligodeoxynucleotides depends on terminal modifications. To test for influence of terminal modification of phosphorothioate ODNs, four different sets of terminal modifications were investigated. For comparison, a previously described p53 anti-sense 15mer (13) was compared to scrambled p53 antisense ODN. These sequences were modified by 3’ hexaethylene glycol, 5’-D,L-α-tocopherol modification; scr, scrambled-p53-control sequence; PT, 3’ polyethylene glycol 1500, 5’-D,L-α-tocopherol; and TT, 5’3’-D,L-α-tocopherol modification. To test for sequence independence, an unrelated random oligonucleotide was designed and used in its double-tocopherol (TT) modified variant, MRON. The effects on DNA synthesis were analysed by [3H]-thymidine-incorporation assays, which were performed in PancTu-1 cells at concentrations of 2.5 and 5.0 μM for 24 and 48 h. As shown in Fig. 1A, p53-HT-ODN reduced DNA synthesis less efficiently than p53-PT-ODN and p53-TT-ODN. Remarkably, for scrambled-p53-sequences (scr-p53-sequences), notable effects on DNA synthesis were seen, a significant reduction of DNA synthesis as compared to untreated controls (p>0.02, post-hoc test Dunnett T3) was found for all ODNs except scr-p53-HT and scr-p53-PT ODN at 2.5 μM for 24 h. The highest efficiency after 24 h at a 2.5-μM concentration was found for MRON. After 24-h treatment with a 5.0-μM ODN concentration, the inhibition was more pronounced for all different modifications as well as for all tested sequences, as seen in Fig. 1C. This dose-dependent effect was even higher than the time-dependent effect of a 2.5-μM concentration of ODN after 48 h, as seen in Fig. 1B. Prolonged incubation appears to strengthen inhibitory effects for PT and TT modifications of anti-p53-ODNs. Surprisingly, MRON was the most efficient in inhibition of DNA synthesis among all ODNs tested. Besides MRON, efficiency of inhibition of DNA synthesis was significantly higher for p53 antisense ODNs compared to scrambled-p53-sequences (p>0.001, Dunnett), except for 24-h 2.5-μM HT modification and 48-h 5.0-μM PT modification.
Induction of apoptosis by modified oligodeoxynucleotides is time- and dose-dependent. Induction of apoptosis was analysed by means of DNA fragmentation as determined by JAM-assay. A significant impact was documented for antisense p53 carrying PT and TT modifications and MRON compared to untreated controls (p>0.05, TukeyHSD/Dunnett T3). Prolonged incubation with a 2.5-μM concentration for 48 h enhanced DNA fragmentation for double-tocopherol modified ODNs but appeared to be unrelated to sequence, with antisense-p53-TT as efficient as MRON (approximately 40% reduction of intact DNA) (Fig. 2C). Treatment with 5.0-μM ODN for 48 h showed the most efficient impact on DNA stability, although the effect appeared to be sequence independent, as scrambled p53-ODNs are as potent as p53 antisense ODNs. HT-modified ODNs showed no significant induction of DNA fragmentation at low doses. Even at high concentrations, only 48 h of incubation yielded a significant effect. PT modification of ODNs always resulted in significant effects compared to controls, but only at high concentrations and 48 h of incubation were significant differences found between antisense and scrambled sequence (p>0.005, Dunnnett T3). In all settings, MRON showed the highest efficiency of all tested modified ODNs.

Figure 2. Effect of terminal modification of oligodeoxynucleotides (ODN) on DNA fragmentation in PancTu-1 cells in vitro. Antisense p53 oligodeoxynucleotides (ODN) and scrambled p53-ODN bearing different terminal modifications were incubated at 2.5- and 5.0-μM concentrations for 24 and 48 h. Data are presented as % of intact DNA (*no significance between antisense and scrambled control sequence, #no significance against untreated control, for all other p-values see Results).

MRON activates NF-κB in vitro. To test for a mechanism of MRON activity, we performed gel shift analysis (EMSA) of PancTu-1 cells stimulated with MRON at concentrations of 1.0, 2.5 and 5.0 μM for different times. Fig. 3 shows a representative analysis revealing an induction of NF-κB after 1 h of incubation with 1.0-μM MRON. After 3 and 6 h, NF-κB activity returns to control levels. When cells were incubated with 2.5-μM MRON, the induction of NF-κB was stronger and remained above control levels even after 6 h.

Figure 3. Gel shift analysis of NF-κB activity in cells stimulated with MRON. PancTu-1 cells were treated with 1.0- and 2.5-μM MRON for 1-, 3-, and 6-h treatment with 5.0-μM MRON was carried out for 6 and 24 h.
Induction of NF-κB by incubation with 5.0-μM MRON was visible after 6 h and returned to control levels after 24 h. Rapid, dose-dependent induction of NF-κB activity within 1 h of treatment and subsequent decrease of NF-κB activity points towards a specific effect targeting NF-κB in PancTu-1 cells, arguing for a receptor-mediated mechanism of MRON activity in PancTu-1 cells.

MRON shows a preference for tumour cells in vivo. To test for in vivo uptake of MRON into cells, tumour-bearing mice were injected with Cy3-labelled MRON, and sacrificed after 3 and 24 h of treatment. After 3 h, Cy3-MRON was found enriched in all organs investigated; intraperitoneal organs, such as the colon and spleen, and extraperitoneal organs, such as the lung and heart (data not shown).

Closer investigation of the liver, as shown in Fig. 4A and B, revealed that non-parenchymal cells were heavily labelled with Cy3-MRON after 3 h of treatment, while hepatocytes showed only diffuse uptake (Fig. 4A). After 24 h, Cy3-label was reduced in hepatocytes and less intense in non-parenchymal liver cells (Fig. 4B). Orthotopic tumours in the pancreas were identified by WGA-staining, as WGA has a strong affinity to the plasma membrane of tumour cells, while it binds to zymogen granules in pancreatic acinar cells. Fig. 4C shows tumour cells as outlined in green with strong labelling of Cy3-MRON throughout the cell and the nucleus. The inset shows an acinus with WGA-stained zymogen granules (green), surrounded by Cy3-MRON (red) labelled interstitial cells.

Effects of MRON in a murine orthotopic xenotransplant model. Effectivity of MRON in vitro as an anti-tumour agent was then tested in vivo in a murine orthotopic xenotransplant model. PancTu-1 cells were injected orthotopically into the pancreas of SCID/bg mice and tumour formation was allowed for 8 days prior to the start of therapy. In a first experiment, the efficacy of MRON was tested with regard to different doses and application regimes over a period of 3 weeks on...
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In vivo testing of MRON on established orthotopic tumours, resembling conservative or palliative therapy in humans, revealed a dose-dependent reduction of tumour size. At a dosage of 3 mg/kg BW, the pancreatic tumour weight showed a higher homogeneity of distribution around the mean, which argues for the effectivity of MRON at this dosage in vivo. At higher concentrations of 6 mg/kg BW, no further effect was seen regarding reduction of tumour size and the animals did not tolerate higher dosages as well. However, at lower concentrations of 1 mg/kg BW, there was a marked heterogeneity of tumour size with a higher mean than untreated controls. This finding points towards the dose-dependent differential activity of MRON. A slight growth-promoting effect was also observed in vitro, when MRON was used at the low dosage of 1 μM (data not shown). Application of MRON at 3 mg/kg BW in adjuvant therapy employing our model of tumour resection (20) resulted in 2 animals without local recurrence and no metastases. Adjuvant treatment with gemcitabine resulted in local recurrence in 3 animals and liver metastasis in 1 animal. Adjuvant treatment with a combination of MRON and gemcitabine showed no local recurrence and no metastases, resulting in 8 out of 8 animals without detectable tumours or metastases.

Table I summarises the effect of adjuvant treatment after resection of orthotopic PDAC in mice.

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<th>Local recurrence</th>
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<td>Resected animals, untreated</td>
<td>8/8</td>
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<tr>
<td>Adjuvant treatment with MRON</td>
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<td>Adjuvant treatment with gemcitabine</td>
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<td>Adjuvant treatment with gemcitabine and MRON</td>
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established tumours as a conservative treatment. Fig. 5 shows the distribution of pancreatic tumour weight for conservative treatment of established tumours. The weight of untreated tumours ranges closely around a mean of 217 mg. When treated with a daily dose of 1 mg/kg BW of MRON, the mean tumour weight was 236 mg with a spread of individual weights from 380 mg to 123 mg. Daily treatment with 3 mg/kg BW of MRON reduced the mean tumour weight significantly (p=0.016) to 118 mg. Furthermore, the individual tumour weight was rather homogenous, arguing for sensitivity of preformed tumours towards this dosage of MRON. When applied at a 3 day interval at a dosage of 3 mg/kg BW, MRON reduced the mean tumour weight to 152 mg, representing a 30% reduction; however, this was not statistically significant (p=0.098). Higher individual dosages of MRON at 6 mg/kg BW did not improve the reduction of tumour weight, while increasing side-effects (data not shown).

Table I summarises the effect of adjuvant treatment after tumour resection using MRON, gemcitabine or a combination of both. Animals, which were resected but left untreated, showed 100% local recurrence, and 7 out of 8 animals had metastasis formation. Adjuvant treatment with MRON at a dosage of 3 mg/kg BW at 3 day intervals resulted in 2 animals showing no local recurrence and 7 out of 8 animals with metastases. Adjuvant treatment with gemcitabine resulted in local recurrence in 3 animals and liver metastasis in 1 animal. Adjuvant treatment with a combination of MRON and gemcitabine showed no local recurrence and no metastases, resulting in 8 out of 8 animals without detectable tumours or metastases.

**Discussion**

In vivo application of antisense oligonucleotides in animal models has always suffered from the critique that the antisense-effect was not properly documented (21). We have shown before that p53 antisense oligonucleotides reduced the level of p53 expression in vitro (22) and had an inhibitory effect on tumour development, when applied in vivo (13). The study presented here shows that the effectivity of oligonucleotides was strongly affected by their terminal modification. Depending on the length and lipophilic properties of terminal modifications, the apoptotic effectiveness is strongly enhanced in vitro. The most effective terminal modification proved to be a 5’3’-D,L-α-tocopherol residue, in all examined parameters in vitro. A totally sequence-independent random control oligonucleotide with 5’3’-D,L-α-tocopherol modification was the most effective agent in vitro. MRON was further tested in vivo for its effect on established orthotopic pancreatic tumours and in a clinically adapted resection model for adjuvant therapy.

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Gemcitabine is well known to effectively reduce pancreatic tumour weight in mice (20). However, during therapy with gemcitabine alone, 3 animals out of 8 showed local recurrence and liver metastasis. Combinational adjuvant therapy with MRON and gemcitabine not only inhibited local recurrence in all animals treated, but also inhibited formation metastases, thus combination of MRON and gemcitabine proved to be more effective than gemcitabine as monotherapy.

Investigating the molecular mechanism of MRON activity, we analysed NF-κB activity. It has been demonstrated that PDAC shows enhanced constitutive activity of NF-κB/RelA (17, 23). Constitutive activity appears to play an essential role in resistance of PDAC against gemcitabine (24) and metastasis (25). In PancTu-1 cells, displaying a moderate constitutive NF-κB activity, MRON further activated NF-κB activity in a dose-dependent manner, low doses induced a slight, short increase in DNA binding, while higher doses of MRON induced a stronger signal that lasted longer and, thus, might induce a different set of target genes (26) than a short burst of activity. This is supported by the observation that low doses of MRON have no effect on DNA damage but slightly induce proliferation, while high doses of MRON induce a strong inhibition of DNA synthesis and subsequently DNA fragmentation. One possible explanation for the observed combination effects might be the induction of cyclins (27) accelerating the cell cycle, thus raising the susceptibility of cells to anti-metabolites.

In humans, gemcitabine has been established as a new standard treatment for patients with advanced pancreatic cancer (28). Besides a significant improval in clinical benefit response, median survival is only modestly prolonged to approximately 6 months (29). Lately, a multi-centre study (ESPAC I) has shown a significant survival effect of adjuvant treatment with 5'-fluorouracil after pancreatic resection (30). Currently, an ongoing trial (ESPAC II) evaluates the potential of gemcitabine to reduce local recurrence and systemic relapse in the same adjuvant setting. Our studies, employing the clinically adapted model of resection of orthotopic pancreatic tumours (20) clearly support the absolute need for adjuvant therapy. In this study investigating a new oligonucleotide-based substance for treatment of pancreatic adenocarcinoma we found a definite synergistic activity between MRON and gemcitabine, even if MRON alone was by far not as efficient in mice as gemcitabine. As to the molecular mechanism of MRON activity, uptake experiments in tumour-bearing mice have shown a selectivity of MRON for tumour cells and interstitial cells. In normal mouse pancreas, MRON is clearly enriched in interstitial fibroblast-like cells (data not shown). In tumour-bearing mice, MRON is enriched in interstitial fibroblast-like cells surrounding acini and in the tumour cell mass. Furthermore, Cy3-labelled MRON accumulates in the cytoplasm and nuclei of tumour cells. Investigation of the liver revealed that uptake into hepatocytes is by far not as efficient as into interstitial cells. This finding argues for differential uptake mechanisms for MRON (or modified oligonucleotides in general, as we have shown previously). One mechanism might be passive uptake via pinocytosis, i.e. an unaided traversal across the plasma membrane, which would account for diffuse labelling as observed in hepatocytes. But the striking selectivity of labelling in interstitial cells and tumour cells versus parenchymal cells of the pancreas and liver argues for a receptor-based activity and cellular entry, which cannot be explained by phagocytosis as MRON is soluble, not particulate. As orthotopic tumours of PancTu-1 cells in mice do not induce a profound desmoplastic reaction, they do not build up an encapsulated environment restricting flow of Cy3-MRON. In vivo uptake experiments showed biological availability and stability. Cellular staining was pronounced after 3 h of treatment and started to fade after 24 h of systemic application; thus, MRON is effectively delivered to orthotopical tumours.

Recent research on combinational therapy of pancreatic tumours employing chemotherapeutical drugs with agents directly addressing molecular targets clearly indicates the way forward. It is most unlikely that a single agent can address the variety of escape mechanisms that tumour cells have developed, but combined activation of different cellular signalling pathways and responses appears to be a more effective method of anti-tumour therapy (31). MRON appears to target signalling pathways which are likely to activate tumour cells, which are then hit more efficiently by a second strike with the metabolic blockade introduced by gemcitabine. This appears to have a profound effect on aggressively growing activated residual tumour cells after resection. Obviously further investigation is needed regarding molecular mechanisms and optimal combinations of MRON with other therapeutic agents, but a relative selectivity towards certain cell types is a promising aspect of modified oligonucleotide-based anti-tumour agents beyond antisenes strategies.

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