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Original Article Evaluation of 9-cis retinoic acid and mitotane as antitumoral agents in an adrenocortical xenograft model

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Abstract: The available drug treatment options for adrenocortical carcinoma (ACC) are limited. In our previous studies, the in vitro activity of 9-cis retinoic acid (9-cisRA) on adrenocortical NCI-H295R cells was shown along with its antitumoral effects in a small pilot xenograft study. Our aim was to dissect the antitumoral effects of 9-cisRA on ACC in a large-scale xenograft study involving mitotane, 9-cisRA and their combination. 43 male SCID mice inoculated with NCI-H295R cells were treated in four groups (i. control, ii. 9-cisRA, iii. mitotane, iv. 9-cisRA + mitotane) for 28 days. Tumor size follow-up, histological and immunohistochemical (Ki-67) analysis, tissue gene expression microarray, quantitative real-time-PCR for the validation of microarray results and to detect circulating microRNAs were performed. Protein expression was studied by proteomics and Western-blot validation. Only mitotane alone and the combination of 9-cisRA and mitotane resulted in significant tumor size reduction. The Ki-67 index was significantly reduced in both 9-cisRA and 9-cisRA+mitotane groups. Only modest changes at the mRNA level were found: the 9-cisRA-induced overexpression of apolipoprotein A4 and down-regulation of phosphodiesterase 4A was validated. The expression of circulating hsa-miR-483-5p was significantly reduced in the combined treatment group. The SET protein was validated as being significantly down-regulated in the combined mitotane+9-cisRA group. 9-cisRA might be a helpful additive agent in the treatment of ACC in combination with mitotane. Circulating hsa-miR-483-5p could be utilized for monitoring the treatment efficacy in ACC patients, and the treatment-induced reduction in protein SET expression might raise its relevance in ACC biology.

Keywords: Adrenocortical cancer, mitotane, 9-cis retinoic acid, xenograft, SET protein, circulating microRNA

Introduction

Adrenocortical carcinoma (ACC) is a rare tumor (incidence is 0.5-2 cases/million people/year) with poor prognosis in its advanced stages [1]. The primary treatment is surgical resection. Even with the most widely used EDPM (etoposide, doxorubicin, cisplatin, mitotane) chemotherapeutic regimen, the median overall survival was only 14.8 months in the recent FIRM-ACT trial [2, 3]. Mitotane (o,p-DDD) is the only adrenal specific drug that is registered for the treatment of ACC [4]. Despite its numerous side effects and largely unknown mechanism of action this agent is used in the clinical practice for more than 50 years [5]. Intensive efforts are therefore going on for finding novel, effective drugs with fewer side effects.

In our previous *in silico* study of adrenocortical tumors genomics data, we have found that retinoic acid signaling via retinoid X receptor is an important pathogenic pathway in ACC [6]. Based on these *in silico* data, we have performed *in vitro* studies on the NCI-H295R ACC cell line and found that 9-cis retinoic acid

(9-cisRA) was able to diminish hormone synthesis and cell viability in a concentration- and time-dependent manner, and it induced significant gene expression changes [7]. In a pilot *in vivo* xenograft study in nude mice, 9-cisRA reduced tumor growth, as well [7].

In the present study, we aimed to further dissect the antitumoral effect of 9-cisRA on ACC in a large xenograft study involving both mitotane and 9-cisRA and their combination. We have applied multimodal molecular biological, genomic and proteomic approaches to decipher the molecular way of action of the agents tested.

Materials and methods

Cell culture

The human adrenocortical NCI-H295R cell line was obtained from the American Type Culture Collection (Manassas, VA, USA). Cells were cultured in Dulbecco's modified Eagle's medium/ Nutrient Mixture F-12 Ham (DMEM: F12) supplemented with 3.61*10⁻⁸ M selenium (Sigma-Aldrich, St. Louis, MO, USA), 1.92*10⁻⁵ M linoleic acid, 0.001 Minsulin, 7.81*10⁻⁸ M transferrin (Sigma-Aldrich), 1.899*10⁻⁵ M bovine serum albumin and adjusted to a final concentration of 1% HEPES, 1% Penicillin/Streptomycin (Sigma-Aldrich), 2.5% Nu-Serum (BD Biosciences, San Jose, CA, USA) and 2.5% L-glutamine (Sigma-Aldrich) at 37°C in a humidified 5% CO₂ atmosphere. The medium was changed two or three times a week and subcultured once or twice a week.

Xenograft model

44 male BALB/c SCID (severe combined immunodeficiency) mice aged 6-8 weeks with average weight between 21-23 g were injected subcutaneously with NCI-H295R cell suspension (10^7 cells/200 µl PBS). Treatment was started when the solid tumor reached 3 mm mean diameter (37 days after the H295R inoculation). The animals have received the treatment by oral gavage for 28 days. Four groups were established: 1. control: 200 µl corn oil/day; 2. mitotane: 200 mg/kg/day/200 µl corn oil; 3. 9-cis retinoic acid: 5 mg/kg/day/200 µl corn oil; and 4. mitotane 200 mg/kg/day/100 µl corn oil + 9-cis retinoic acid 5 mg/kg/day/100 µl corn oil. The tumors were measured twice a week with a caliper by the same investigator. The volumes of the tumors were calculated by the following formula: $(a*a*b*\pi)/6$ [8]. On day 29 after starting the treatments, the animals were killed by cervical dislocation in ether anes-thesia. Only one animal died before day 29 due to technical problems.

Tumors were removed and their weights were measured. One half of the tumor was fixed in formalin for histological and immunohistochemical examination, the other half was frozen in liquid nitrogen and stored at -80°C until use. Lungs, heart, kidneys, spleen and liver were also removed for histological analysis. Whole blood was collected, then plasma was isolated and stored at -80°C until use. All animal experiments were conducted according to the ethical standards of the animal Health Care and Control Institute, Csongrád County, Hungary, permit No. XVI/02037-2/2008.

Histological and immunohistochemical analysis

Four µm sections of formalin-fixed paraffinembedded tissues were dewaxed with xylene and ethanol and processed either for hematoxylin-eosin (HE) staining or Ki-67 immunostaining. Ki-67 immunostaining (antibody: Cat. No. M7240, DakoCytomation, Glostrup, Denmark; dilution 1:400) was conducted using a Leica BOND-MAX automated immunostaining system (Leica Biosystems, Wetzlar, Germany). Slides were scanned by Pannoramic Flash II scanner, and digital slides were analyzed by Pannoramic viewer software (3DHistech, Budapest, Hungary). Different regions of the viable xenograft tumor were annotated and Ki-67 positive and negative cells were individually marked on these regions of interests (ROIs). Proliferation index was given in percentages of positive cells. 4-8 slides of each group with good quality were selected for analysis, and 3-5 ROIs/slides were counted with 4498-6273 cells in average. Ki-67 expression was scored by three independent pathologists in a blinded fashion.

RNA isolation from tumor tissue

Total RNA was isolated from the frozen tumors with Qiagen miRNeasy Mini Kit according to the manufacturer's protocol (Qiagen, Hilden, Germany). RNA concentration was measured with NanoDrop 2000 spectrophotometer. (Thermo Fisher Scientific, Waltham, Mam USA). RNA integrity was determined by Agilent 2100 Bioanalyzer System (Agilent Technologies, Santa Clara, CA, USA). Samples with an RNA integrity number (RIN) above 8.0 were used for further analysis. RNA was stored at -80°C until use.

Messenger RNA (mRNA) expression profiling

Gene expression profiling was performed on 16 samples (4-4 samples from each group) using a single-color array method by 4x44K Agilent Whole Genome Microarray slides (Agilent Technologies).

Total RNA (200 ng) was labeled and amplified using the low Input Quick Amp Labeling Kit according to the instructions of the manufacturer. Labeled RNA was purified and hybridized to Agilent Human Gene Expression Microarray 4x44K array slides, according to the manufacturer's protocol. After washing, array scanning and feature extraction was performed with default scenario by Agilent DNA Microarray Scanner. Fluorescence intensities of spots were quantified, background subtracted, and dye normalized by Feature Extraction software, version 11.0.1 (Agilent Technologies). The raw data were analyzed with GeneSpring 12.6 software (Agilent Technologies).

Reverse transcription quantitative polymerase chain reaction (RT-qPCR)

Seven genes were selected for validation by real-time RT-qPCR with Taqman gene expression assays (Thermo Fisher Scientific): *MYC* (Hs00153408_m1), *APOA4* (Hs00166636_m1), *CXCR3* (Hs01847760_s1), *BAALC* (Hs00-227249_m1), *PDE4A* (Hs00183479_m1), *PR-DM1* (Hs00153357_m1) and *TGFBI* (Hs009-32747_m1). Based on the selection criteria of Cheng et al. [9] and our previous study [10], *ZNF625* (Hs00377010_m1) was chosen as reference gene.

Total RNA (10 ng) was reverse transcribed using High-Capacity RNA-to-cDNA Kit (Thermo Fisher Scientific). Quantitative RT-PCR was performed by TaqMan Fast Universal PCR Master Mix (2×) (Thermo Fisher Scientific) on a 7500 Fast Real-Time PCR System (Thermo Fisher Scientific) according to the manufacturer's protocol. Samples were run in triplicate. For the evaluation of the data we used the ddCT method [11] using Microsoft Excel 2010 (Microsoft Corporation).

RNA isolation from plasma

Total RNA was isolated from 100 µl plasma samples with Qiagen miRNeasy Serum/Plasma Kit according to the attached manufacturer's protocol (Qiagen) amended with addition of a spike-in control miRNA *cel-miR*-39 (2594091) (Qiagen). RNA concentration was measured with NanoDrop 2000 spectrophotometer (Thermo Fisher Scientific). RNA was stored at -80°C until use.

RT-qPCR for circulating microRNA measurement

Based on our previous studies and literature data, four microRNAs were selected for validation by RT-qPCR with Taqman miRNA assays (Thermo Fisher Scientific): *hsa-miR-181b* (00-1098), *hsa-miR-184* (000485), *hsa-miR-210* (000512) and *hsa-miR-483-5p* (002338). *Celmir-39* (000200) was used as reference gene [12].

Total RNA (1 μ I) was reverse transcribed using specific TaqMan miRNA assays and TaqMan MicroRNA Reverse Transcription Kit (Thermo Fisher Scientific) on Proflex Base PCR System (Thermo Fisher Scientific). Quantitative RT-PCR was performed by TaqMan Fast Universal PCR Master Mix (2×) and TaqMan miRNA assays on a 7500 Fast Real-Time PCR System. PCR reactions were run in triplicate. Negative control reactions did not include cDNA templates.

RT-qPCR for tissue microRNA measurement

Considering the results from the circulating microRNA measurements, tissue *hsa-miR-483-5p* (002338) was studied by RT-qPCR with Taqman miRNA assays according to the instructions of the manufacturer. Four reference genes were used: *U6* (001973), *RNU6B* (001093), *RNU44* (001094) and *RNU48* (001006).

Proteomics study

Equal amount of protein lysates (20-20 μ g of each sample, 3 samples from each group) were separated by SDS-PAGE on a 10% minigel and stained with colloidal Coomassie Brillant Blue. The gel lanes were cut to 10 pieces and their protein content were in-gel digested with trypsin (Promega, Fitchburg, WI, USA) for 4 hours at 37°C after reduction with dithiotreitol and alkylation with iodoacetamide. The digestion reaction was stopped by acidification with formic acid, and the tryptic peptides were extracted from the gel and dried.

Samples were redissolved in 30 µl of 0.1% formic acid and subjected to LC-MSMS (Liquid Chromatography-Mass Spectrometry and Liquid Chromatography-Tandem Mass Spectrometry b, on a Waters nanoAcquity ULC online coupled to an Orbitrap Elite Mass spectrometer) analysis. 5 µl sample was loaded onto a Symmetry C18 (5 µm, 180 µm ×20 mm) trap column (Waters, 186003514) in 3% of solvent B and analyzed on a BEH300 C18 (1.7 µm, 075 µm ×250 mm) nanoAcquity UPLC column (Waters 186003815) using a gradient elution (3-40% of solvent B during 37 min, solvent A: 0.1% formic acid in water, solvent B: was 0.1% formic acid in acetonitrile:DMSO (95:5)). The flow rate was 300 nl/min. The LTQ-Orbitrap Elite mass spectrometer operated in data dependent mode, each survey scan (full MS measured in the Orbitrap R=60000, m/z range: 380-1600) was followed with ion-trap collision induced dissociation (CID) scans of the 10 most intense peaks. Dynamic exclusion was used for 30 seconds and single charged ions were not selected for fragmentation.

Mass spectrometry raw data were converted to MSMS peak list files using PAVA script [13]. Peak list files corresponding to the same sample (10 to each) were merged and searched against the human, mouse species specified UniProtKB random concat (06.11.2014) protein database, containing 136244 human and 74540 mouse proteins and their randomized sequences. The search was done with the following parameters: fully specific tryptic peptides with maximum 2 missed cleavages were allowed. All cysteines were considered as carbamidomethylated and some variable modification has been allowed: oxidized methionine, acetylation of protein N-termini and pyroglutamic acid modification at peptide N-terminal glutamine. Parent mass tolerance was set to 10 ppm, while fragment mass tolerance was 0.6 Da. All database search were completed on our in-house ProteinProspector (ver. 5.14.1) (Baker, P.R. and Clauser, K.R. http://prospector. ucsf.edu accessed: July 22. 2015) search engine.

Spectral count was used for semi-quantitation. The number of spectra identified a protein were normalized by the number of all identified spectra in the sample. These relative spectral counts were compared in between the different samples.

Western-blot analysis

Based on our proteomic results and literature data, the SET (SET nuclear proto-oncogene) protein was chosen for Western-blot analysis on 3 samples from each group of the xenograft study and on altogether 6 human adrenocortical samples (2 normal adrenal cortices, 2 ACAs (adrenocortical adenoma) and 2 ACCs (adrenocortical cancer)). Normal human adrenal cortices were obtained from patients operated for hypernephroma [14]. The study on human samples was approved by the Ethical Committee of the Hungarian Health Council and informed consent was obtained from all patients involved.

Tumor tissues were pulverized in liquid nitrogen and lysed in 1 ml of Lysis Buffer (0.02 M Tris, 0.15 M NaCl, 0.002 M EDTA, 0.5% Triton X-100, 0.5% Protease Inhibitor Cocktail (Sigma Aldrich), 0.01 M NaF, 0.002 M Na, VO, on ice for 30 minutes. Next, samples were centrifuged at 13000 rpm for 15 min. Supernatants were kept and protein concentrations were measured as described before by Bradford [15]. 20 µg of total protein per sample was mixed with loading buffer containing *B*-mercaptoethanol and was incubated at 99°C for 5 min. Denatured samples were loaded onto a 10% polyacrylamide gel and were run for 30 min at 200 V on a Mini Protean vertical electrophoresis equipment (Bio-Rad, Hercules, CA). Proteins were transferred onto a PVDF membrane (Milipore, Billerca, MA) by blotting overnight at 75 mA. Ponceaustaining was applied to see the loading and blotting efficiency. Membranes were blocked with 5 w/v% non-fat dry milk (Bio-Rad) in TBS (0.15 M NaCl, 0.02 M Tris-HCl pH=7.5) for 1 hour followed by incubation with either anti-SET antibody (ab1183, Abcam, Cambridge, MA, dilution 1:1000) or anti-β-actin antibody (A22-28, Sigma Aldrich) for 16 h at 4°C. Membranes were washed 5 times with TBST (TBS+0.05% Tween-20) then incubated with secondary, HRP-conjugated anti-rabbit (P0448, DakoCytomation) or anti-mouse (P0447, DakoCvtomation) antibodies for 1 hour. SuperSignal West Pico Chemiluminescent Substrate Kit (Pierce/



Figure 1. A. Normalized tumor volumes; B. Normalized tumor volumes at the end of the treatments on day 28; C. Representative images of the Ki-67 immunhistochemical analysis from the four treatment groups; D. Numerical values of the Ki-67 immunhistochemical analysis. (Average results by three independent investigators).

Thermo Fisher Scientific) and Kodak Image Station 4000 MM Digital Imaging System were used to visualize the signals. β -actin served as loading control. Band density was measured by ImageJ software (National Institutes of Health, Bethesda, MD, USA).

Statistical analysis

For the statistical analysis of tumor growth, Mann-Whitney U test was applied (SPSS Statistics 20, IBM). The analysis of Ki-67 index was performed by One-way ANOVA followed by Tukey's post Hoc test or Kruskal-Wallis ANOVA & Median Test (STATISTICA 7.0) depending on the results of the Shapiro-Wilks normality test. P<0.05 was considered to be significant.

The results from the microarray data were analyzed by Genespring 12.6 software (Agilent

Technologies). A filter on expression at the 20th percentile of raw signal values, then a 2-fold change filter was used, and One-way ANOVA was performed, followed by Tukey's post Hoc test without Benjamini-Hochberg false discovery rate calculation. RT-qPCR and proteomics data were analyzed by One-way ANOVA followed by Tukey's post Hoc test (STATISTICA 7.0).

Western-blot was evaluated by Kruskal-Wallis ANOVA & Median Test (STATISTICA 7.0).

Results

Analysis of xenograft tumor growth

The normalized tumor volume (relative to the first measured volume) was smaller in all treated groups than in controls during the whole experiment. At the end of our experiment, the



Figure 2. Results of RT-qPCR validation of APOA4 and PDEA4 (n=6).

average values of normalized tumor volumes were 22.48-fold, 9.3-fold, 12.3-fold and 8.22fold higher in the control, mitotane treated, 9-cisRA-treated, and 9-cisRA+mitotane groups relative to the starting tumor volume, respectively. The reduction in tumor size has been significant in the mitotane only and 9-cisRA+ mitotane groups (**Figure 1A** and **1B**).

Histology, Ki-67 scoring

Tumors in the 9-cis RA receiving groups appeared to be more differentiated compared to the control and mitotane groups by hematoxylin and eosin (H&E) staining. The Ki-67 proliferation index scored blindly by three independent pathologists was significantly lower in the 9-cisRA (32.64 ± 3.07%) group in comparison to the control (54.23 ± 3.71%) and mitotane groups (46.35% ± 2.2%) (P=0.00018 vs. control; P=0.00022 vs. mitotane). The lowest Ki-67 index was noted in the combined 9-cisRA+ mitotane group (25.6 ± 4.09%) that was significantly smaller compared to all other groups (P=0.00018 vs. control; P=0.00018 vs. mitotane; P=0.01684 vs. 9-cisRA) (Figure 1C and **1D**). Among the other tissues, only moderate fatty lesions were observed in the liver, probably due to the corn oil vehicle. No signs of treatment toxicity have been found.

Microarray analysis

4 samples from each group (altogether 16 samples) have been subjected to microarray. By analyzing the raw data, we have found 483 significant gene expression changes after the statistical analysis, but only by omitting the Benjamini-Hochberg false discovery rate (FDR). By using FDR, only two significantly differentially expressed transcripts emerged whose biological relevance is unknown. Microarray data are accessible at Gene Expression Omnibus (GEO; www.ncbi.nlm.nih.gov/geo, accession number: GSE73417).

RT-qPCR validation

Seven genes have been selected for validation based on the microarray data chosen from genes with the highest positive and negative fold changes, taking into account literature data, as well. However, only two genes could be validated to be significantly differentially expressed (**Figure 2**). The well-known 9-cis retinoic acid target gene *APOA4* (apolipoprotein A4) turned out to be significantly overexpressed in the combined treated group relative to the control group (P=0.0045), whereas *PDE4A* (phosphodiesterase 4A) appeared to be significantly underexpressed in the combined treatment group (P=0.0024).

Proteomics analysis and Western-blot

47 significant protein changes have been found between the groups (**Table 1**). By considering literature data regarding the tumor biological relevance of these proteins, the SET protein [16-20] was chosen for validation by Westernblotting.

SET protein expression was smaller in all treated groups relative to control, but only reached the level of significance in the combined treated group (**Figure 3A** and **3C**). To look at the

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UniProt ID	Gene Name	Protein Name
P30050	RPL12	60S ribosomal protein L12
Q13765	NACA	Nascent polypeptide-associated complex subunit alpha
B1AK87	CAPZB	Capping protein (Actin filament) muscle Z-line, beta
Q01105	SET	Phosphatase 2A inhibitor I2PP2A
Q86SZ7	PSME2	PSME2 protein
P14314	PRKCSH	Protein kinase C substrate 60.1 kDa protein heavy chain
Q5U8W9	HRMT1L2	Protein arginine methyltransferase 1 isoform 4
Q6DC98	LMNB1	LMNB1 protein
M0QZL1	BLVRB	Flavin reductase (NADPH)
P61158	ACTR3	Actin-related protein 3
Q9UBQ7	GRHPR	Glyoxylate reductase/hydroxypyruvate reductase
075347	TBCA	Tubulin-specific chaperone A
Q6NVY0	CACYBP	Calcyclin binding protein
K7ES63	TUBB6	Tubulin beta-6 chain
P42126	ECI1	Enoyl-CoA delta isomerase 1
Q5T1M5	FKBP15	FK506-binding protein 15
075477	ERLIN1	Erlin-1
H7BZ09	ANP32A	Acidic leucine-rich nuclear phosphoprotein 32 family member A
E9PK45	FTH1	Ferritin heavy chain
P62714	PPP2CB	Serine/threonine-protein phosphatase 2A catalytic subunit beta
P31040	SDHA	Succinate dehydrogenase
H7C333	GBAS	Protein NipSnap homolog 2
Q15404	RSU1	Ras suppressor protein 1
Q92882	OSTF1	Osteoclast-stimulating factor 1
D6RC06	HINT1	Histidine triad nucleotide-binding protein 1
P62888	RPL30	60S ribosomal protein L30
Q969G3	SMARCE1	SWI/SNF-related matrix-associated actin-dependent regulator of chromatin subfamily E member 1
P61326	MAGOH	Protein mago nashi homolog
Q16539	MAPK14	Mitogen-activated protein kinase 14
P10599	TXN	Thioredoxin
Q9Y333	LSM2	U6 snRNA-associated Sm-like protein LSm2
P26885	FKBP2	Peptidyl-prolyl cis-trans isomerase FKBP2
Q9UJV8	PURG	Purine-rich element-binding protein gamma
P04080	CSTB	Cystatin-B
E5RJH5	EN03	Beta-enolase
P28062	PSMB8	Proteasome subunit beta type-8
P67775	PPP2CA	Serine/threonine-protein phosphatase 2A catalytic subunit alpha isoform
Q53G26	DNAJA3	DnaJ (Hsp40) homolog, subfamily A, member 3 variant
Q9UBI6	GNG12	Guanine nucleotide-binding protein G(I)/G(S)/G(O) subunit gamma-12
Q8NHP8	PLBD2	Putative phospholipase B-like 2
D6R9R5	SAR1B	GTP-binding protein SAR1b
Q00169	PITPNA	Phosphatidylinositol transfer protein alpha isoform
P63173	RPL38	60S ribosomal protein L38
Q5T6V5	C9orf64	UPF0553 protein C9orf64
A4D2P2	RAC1	Ras-related C3 botulinum toxin substrate 1

 Table 1. Treatment-induced significant protein changes (47 proteins) based on the results of proteomic analysis (P<0.05)</th>

potential relevance of SET in human adrenocortical tumors, we have analyzed some human tissues in a preliminary study, and found that SET protein is undetectable in normal and ACA tissues, whereas it is expressed in ACC (**Figure 3B**).



microRNA expression

From the four selected circulating miRNAs, *hsa-miR-483-5p* has been significantly underexpressed in the combined group relative to control (P=0.028) (**Figure 4**). The expression of the other three tested circulating microRNAs was unchanged. However, we have not found significant changes in the expression of tissue *hsa-miR-483-5p*. From the four reference genes tested for tissue microRNAs, *RNU44* and *RNU6B* turned out to be the best for normalizing the results.

Discussion

Mitotane is the only available adrenal cortex specific agent in the therapy of adrenocortical cancer. Despite being used for more than 50 years, its precise mechanism of action is largely unknown [5]. The degeneration of mitochondria, free radicals, inhibition of steroid biosynthetic enzymes, lipid-mediated endoplasmic reticulum stress have all been suggested to take part in the action of mitotane [5, 21]. The narrow therapeutic range and frequent side effects of mitotane constitute serious problems in clinical practice. Based on our previous *in vitro* and pilot xenograft study [7], 9-cisRA appeared to be a promising agent in ACC treatment. To further dissect its actions and to compare it to mitotane, we have performed a largescale xenograft study involving 9-cisRA, mitotane and their combination. We have observed that these agents act together in a synergistic manner on tumor growth.

Contrary to the several human *in vivo* studies applying mitotane, there are only few studies on mitotane-treated xenograft models to date [22-27]. Since there was no uniformly defined dose of mitotane given to mice, we have chosen the average amount used in published studies. The results of these studies on the effectiveness of mitotane in animal models are also contradictory. Lindhe et al. described that mitotane has antitumoral effect given simultaneously with the tumor cells inoculation [24], whereas Doghman et al. did not find any longlasting effect in a similar study [25]. Here, we have observed the antitumoral activity of mitotane.



Figure 4. The expressional difference of circulating hsa-miR-483-5p (n=6).

Retinoids, the natural and synthetic derivates of vitamin A play crucial roles in cell differentiation, growth, and death. The anti-proliferative and antitumoral effects [28] of 9-cisRA acting via the retinoid X receptor are well documented in many different types of tumors *in vitro* and *in vivo* e.g. gastric cell line [29], head and neck squamous cell line [30], neuroblastoma [31] and breast cancer [32].

The dose of 9-cis retinoic acid (5 mg/kg) was much less than the documented toxic dose [30] and it was comparable to other *in vivo* doses used in the literature [31, 33-35].

We have observed antitumoral effects of both 9-cisRA and mitotane when used alone, but significant reduction in tumor volume could only be achieved by groups receiving mitotane alone and the combination of mitotane and 9-cisRA. The Ki-67 index was the lowest in the group receiving the combined treatment. Despite mitotane was more effective than 9-cisRA as a single agent for reducing tumor size, the Ki-67 index was lower in the 9-cisRA-treated group than in the mitotane only group (**Figure 1**). The background for this phenomenon is unclear.

To decipher the molecular mechanisms behind the observed effects on tumor growth and proliferation, we have performed molecular analysis at mRNA and protein levels. In our previous *in vitro* studies on the adrenocortical cell line NCI-H295R, we have observed that 9-cisRA affected the expression of more than 2000 genes [7] whereas the effect of mitotane on gene expression was more limited [10]. In contrast to these in vitro data and the observed changes in tumor size and Ki-67 levels, we could observe only modest changes in gene expression in the present study. By omitting the Benjamini-Hochberg False Discovery Rate, 483 genes were found to be significantly genes, and from the 7 selected genes only 2 (APOA4, PDEA4) was validated to be significantly differentially expressed.

APOA4, a member of apolipoprotein family is a known 9-cisRA target gene that has important roles in fat and glucose metabolism [36], in antioxidant [37], and anti-atherogenic processes [38], but it has been suggested also as a predictive factor in inflammatory bowel disease [39, 40], and as a novel serum biomarker in ovarian, cervical cancer, and acute lymphocytic leukemia [41-44]. There are no data, however, on the potential relevance of *APOA4* in adreno-cortical tumors.

PDE4A, as a member of PDE (phosphodiesterase) family, act as crucial cAMP level regulator [45], and it is involved in the basic cell pathways and also takes part in cancer progression. Its inhibition has been associated with many antitumoral actions e.g. regression of brain tumors [46], reduced proliferation and angiogenesis in lung cancer [47], reduced motility and invasion in colon [48] and breast cancer cell line [49], decreased growth of malignant melanoma [50]. In line with these observations, we have also observed reduced expression of *PDE4A* in the combined treatment group, thus *PDE4A* might also be involved in ACC pathogenesis or progression.

We hypothesize that after the long treatment period of 28 days, mRNA expression does not really reflects the altered tumor behavior. We have therefore turned to the protein level, where we have found 47 significantly differentially expressed proteins by proteomics analysis. We have chosen the SET protein from the significantly differentially expressed proteins for validation by Western-blot. As expected, SET expression was down-regulated by the treatments, showing the lowest expression in the combined treatment group.

Protein SET is an inhibitor of tumor suppressor PP2A (protein phosphatase 2A) that has widespread actions in the cellular functions implying cell cycle [18], apoptosis [51] and cell migration [19]. Due its target PP2A, it also influences ß-catenin [20], c-Myc [17] and Akt [16] pathways, and via inhibition of nm23-H1 (NME/ NM23 nucleoside diphosphate kinase 1) it also promotes metastatic potential [52], and thus it is associated with tumor progression. SET overexpression is described in many tumors e.g. Wilm's tumor [53], acute lymphoblastic leukaemia [54], chronic lymphocytic leukemia and non-Hodgkin lymphoma [55], lung [56], colon [57], pancreatic [58], prostate [59] and ovarian cancer [60], as well. To the best of our knowledge, our study is the first to raise the potential relevance of SET in ACC biology.

We have performed a preliminary evaluation of SET protein expression in human adrenocortical tumor samples, as well. SET was weakly expressed in ACC, but absent from benign adenomas and normal adrenocortical samples. This finding that certainly awaits validation on a larger cohort appears to support our xenograft findings where SET expression was the highest in the untreated xenograft, and suppressed in the treated samples parallel to tumor shrinkage.

In addition to our mechanistic efforts aimed at deciphering the molecular way of action of 9-cisRA and mitotane in our xenograft model, we have also studied some circulating microR-NAs as potential markers for treatment efficacy monitoring based on our previous studies in humans. Circulating hsa-miR-483-5p appears to be the best circulating microRNA marker of adrenocortical malignancy [61-63]. From the four analyzed circulating microRNAs, only hsamiR-483-5p expression was modulated by the treatments, and its expression was significantly suppressed by the combined 9-cisRA + mitotane treatment. Circulating hsa-miR-483-5p thus appears to be a marker of treatment efficacy, and this could be relevant in the clinical setting, as well. Regarding the tissue expression of hsa-miR-483-5p, however, no significant differences have been observed. We can conclude that the circulating and tissue *hsamiR-483-5p* alter independently. There are data that microRNAs can be changed parallel, contrary, or independently between the circulation and tissues [64, 65], but the molecular mechanisms underlying these discrepancies and the regulation of adrenocortical *hsa-miR-483-5p* expression are largely unknown, yet.

In conclusion, 9-cisRA might represent an alternative additive treatment option in ACC that seems to be most efficient when combined with mitotane. The tumor size, Ki-67, SET protein, and circulating hsa-miR-483-5p were all lowest in the group receiving the combined treatment that suggest synergistic action of 9-cisRA and mitotane in our xenograft model. The molecular background of the observed synergistic mitotane-9-cisRA action is, however, unclear. We have found only modest gene expression changes, and although the number of affected proteins is higher, their list is not extensive, either. The decrease of SET protein expression by the treatments might be noteworthy, as paralleled by its expression in human adrenocortical tumors. The potential applicability of circulating hsa-miR-483-5p for monitoring ACC treatment efficacy might be relevant in the clinical setting, as there is no reliable bloodbased tumor marker of ACC at present. Circulating microRNA might be exploited for treatment monitoring in other diseases, as well.

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Disclosure of conflict of interest

None.

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