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Nicotinamide Nucleotide Transhydrogenase as a novel treatment
target in adrenocortical carcinoma

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Abstract

Adrenocortical Carcinoma (ACC) is an aggressive malignancy with poor response to chemotherapy. Here we evaluated a potential new treatment target for ACC, focusing on the mitochondrial NADPH generator Nicotinamide Nucleotide Transhydrogenase (NNT). NNT has a central role within mitochondrial antioxidant pathways, protecting cells from oxidative stress. Inactivating human NNT mutations result in congenital adrenal insufficiency. We hypothesized NNT silencing in ACC cells will induce toxic levels of oxidative stress. To explore this, we transiently knocked down NNT in NCI-H295R ACC cells. As predicted, this manipulation increased intracellular levels of oxidative stress; this resulted in a pronounced suppression of cell proliferation and higher apoptotic rates, as well as sensitization of cells to chemically-induced oxidative stress. Steroidogenesis was paradoxically stimulated by NNT loss, as demonstrated by mass spectrometry-based steroid profiling. Next, we generated a stable NNT knockdown model in the same cell line to investigate the longer-lasting effects of NNT silencing. After long-term culture, cells adapted metabolically to chronic NNT knockdown, restoring their redox balance and resilience to oxidative stress, although their proliferation remained suppressed. This was associated with higher rates of oxygen consumption. The molecular pathways underpinning these responses were explored in detail by RNA sequencing and non-targeted metabolome analysis, revealing major alterations in nucleotide synthesis, protein folding and polyamine metabolism. Our study provides the first pre-clinical evidence of the therapeutic merit of antioxidant targeting in ACC as well as illuminating the long-term adaptive response of cells to oxidative stress.
Introduction

Adrenocortical carcinoma (ACC) is a rare but aggressive malignancy. The majority of patients present with, or eventually develop, metastatic disease, which shows limited or no responsiveness to cytotoxic chemotherapy (1, 2). A recent randomized trial revealed a median survival of <15 months for patients with disseminated disease receiving combination chemotherapy (3). Glucocorticoid or androgen excess often constitutes an additional clinical burden on ACC patients, undermining their quality of life (1). Unfortunately, the obvious need for more effective medical treatment options in ACC patients remains unmet, despite the remarkable progress in our understanding of the molecular biology of ACC in the last two decades (1).

Recent genetic studies have provided new insights into adrenal pathophysiology, revealing that inactivating mutations in the gene encoding the antioxidant enzyme Nicotinamide Nucleotide Transhydrogenase (NNT) underlie a rare, hereditary form of primary adrenal insufficiency (4). Affected patients present in early childhood with failure to thrive, hypotension and hypoglycemia, due to inability of the adrenal glands to produce sufficient cortisol (4). Intriguingly, despite the key role of NNT in preserving cellular redox balance and its ubiquitous expression, the adrenal glands are the only affected organ in most patients; this observation suggests a selective sensitivity of the adrenal glands to NNT loss (4, 5). Supportive of this, NNT-deficient mice harbor adrenal glands with disorganized cortical architecture and high apoptotic rates in their adrenal zona fasciculata, the location of glucocorticoid synthesis, but no other abnormality (4).

NNT is a dimeric proton pump that resides in the inner mitochondrial membrane of eukaryotic cells and uses the transmembrane proton gradient to catalyze the transfer of reducing equivalents from NADH to NADP⁺, according to the reaction
NADH + NADP$^+$ + H$^+$\text{Intermembrane} ↔ NAD$^+$ + NADPH + H$^+$\text{matrix} \ (6, 7)

NADPH is an essential donor of reducing power to the two main mitochondrial antioxidant pathways, the glutathione and the thioredoxin pathways, which protect the mitochondria from the deleterious effects of oxidative stress with their capacity to detoxify reactive oxygen species (e.g. hydrogen peroxide, H$_2$O$_2$). Reactive oxygen species (ROS), the molecular mediators of oxidative stress, are continuously produced within the mitochondria by electron leakage along the respiratory chain complexes and the tricarboxylic acid cycle (TCA); in adrenocortical mitochondria, steroidogenesis represents an important additional source of ROS (8-11). Excessive levels of oxidative stress lead to irreversible DNA, protein and lipid damage, which can culminate in apoptotic cell death (Fig. 1A) (12).

Drawing on these data, which indicate a distinct metabolic vulnerability of the adrenal cortex to oxidative stress, we explored the value of antioxidant targeting as a novel therapeutic approach in ACC, focusing on NNT as a putative treatment target. Interrogating a publicly available whole genome gene expression database (13), we observed that NNT is up-regulated in ACCs in comparison to benign adrenocortical adenomas and healthy adrenals (Fig. 1B). Therefore, we hypothesized NNT silencing in ACC cells will impair their antioxidant capacity and lead to progressive accumulation of ROS, inducing unsustainable oxidative toxicity within the mitochondria and eventually triggering cellular apoptosis. We also postulated that steroidogenesis will be suppressed as a result of NADPH depletion and/or increased oxidative stress.
Materials and Methods

Cell culture protocol and cell line validation

NCI-H295R (RRID:CVCL_0458) ACC cells (passage 10-25) were cultured under standard conditions using DMEM/Ham’s F-12 medium (Gibco, Thermo Fisher) supplemented with 2.5% Nu serum (Corning), 1% penicillin-streptomycin (Gibco, Thermo Fisher) and 1% ITS+ universal cell culture premix (Corning). Cell line identity was confirmed through Short Tandem Repeat (STR) genetic analysis performed by the DNA Diagnostics Company (London, UK) followed by comparison to genetic profiles provided by the American Tissue Culture Collection (ATCC) (http://www.lgcstandards-atcc.org) (Suppl. Table 1).

Small-interfering RNA (siRNA) transfection

Transient NNT gene silencing was achieved through transfection of NCI-H295R cells with small interfering RNA (siRNA), using Viromer® Blue (Lipocalyx) molecules as transfection vehicles. Three alternative siRNAs targeting different areas of the NNT gene were tried (Life Technologies, HSS118900, HSS118901 and HSS118902), and the one exhibiting the most consistent efficiency in knocking down NNT (HSS118902) was selected for subsequent experiments. HSS118901 was used to corroborate results in proliferation and apoptosis assays, whose results are open to confounding by off-target effects. A scrambled, non-sense siRNA (Silencer Select 1 negative control, Life Technologies) was used as negative control (SCR siRNA). Viromer-siRNA transfection was performed according to the manufacturer’s instructions in 6-well plates (300,000 cells/well) and 96-well plates (6-8,000 cells/well).

Short-hairpin RNA (shRNA) transfection

Lentiviral vectors were obtained from Dharmacon (UK) in a p.GIPZ backbone and contained
5 shRNAs specific for human NNT (RHS4430-98851990; RHS4430-98913600; RHS4430-98524425; RHS4430-101033169; RHS4430-101025114) under the control of the CMV promoter, as well as the puromycin resistance and green fluorescence protein (GFP) genes. Vectors expressing non-sense, scrambled shRNA (SCR shRNA) were used as negative controls. HEK293T cells (packaging cells, RRID:CVCL_0063) were transfected with the shRNA particles by lipofectamine transfection (Thermo Fisher), according to the manufacturer’s instructions. Cell media containing the viral particles was collected 48-72 h post-transfection and used to transduce NCI-H295R cells. Four days after transfection, GFP-positive cells were selected in 4μg/ml puromycin. Transduction efficiency was determined by fluorescence microscopy and Western Blotting for NNT expression.

**Gene expression**

Gene expression for NNT and steroidogenic enzymes was evaluated at a transcriptional level by quantitative real-time polymerase chain reaction (qRT-PCR). RNA extraction was performed using the RNeasy Mini kit (Qiagen) following the manufacturer’s instructions. Reverse transcription to generate complementary DNA (cDNA) was carried out using the TetrocDNA Synthesis Kit (Bioline), following the manufacturer’s instructions (500 – 2,000 ng of RNA per reaction used). cDNA concentration was determined by use of a fluorescent DNA dye (Quant-iTTM PicoGreen® dsDNA reagent, Thermo Fisher), comparing sample fluorescence to the fluorescence exhibited by a dilution series of samples of known concentrations (Wallac Victor 1420 multilabel counter). Gene expression was then quantified by qRT-PCR, using the Taqman Gene Expression System (Thermo Fisher). Reactions were run in a 7500 ABI qRT-PCR analyzer [50°C incubation for 2 minutes, 95°C for 10 minutes, followed by 40 cycles of 95°C for 15 seconds (denaturation) then 60°C for 1 minute (annealing-extension)]. All reactions were normalized
against the housekeeping gene RPLPO (large ribosomal protein). Data are expressed as ΔCt values

\[ \Delta C_t = (C_t \text{ of the target gene}) - (C_t \text{ of the housekeeping gene}) \] or fold-change to control cells (2^\DeltaΔCt), where ΔΔCt = ΔCt (NNT knockdown cells) – ΔCt (control cells).

**Protein expression**

Protein lysate generation was performed by applying RIPA buffer (Sigma-Aldrich) with protease inhibitor cocktail (Sigma-Aldrich) to adherent cells grown in 6-well plates and subsequent collection by scraping. Total protein concentration was estimated colorimetrically using the BCA Protein Assay Kit (Thermo Fisher) as per the manufacturer’s instructions, measuring absorbance at 560 nm (Wallac Victor 1420 multilabel counter). NNT protein expression level was assessed by Western Blotting. Samples were run in 10% SDS-PAGE (polyacrylamide) Gels (Thermo Fisher) and transferred to a nitrocellulose membrane using the iBLOT™ Dry Transfer System (Thermo Fisher). Membranes were subsequently probed with anti-NNT antibody produced in rabbit (HPA004829, Sigma-Aldrich, RRID:AB_1079495) at a 1:500 dilution and secondary anti-rabbit antibody (sc-2030, Santa-Cruz, USA, RRID:AB_631747) at a 1:2,000 dilution. β-actin was used as control protein (primary antibody A5441, RRID:AB_476744 from Sigma-Aldrich and secondary anti-mouse antibody from Santa-Cruz (sc-2005, RRID:AB_631736), dilutions 1:10,000 and 1:20,000, respectively).

**Reduced to oxidized glutathione ratio (GSH/GSSG)**

Total cell glutathione (GSH + GSSG) and oxidized glutathione (GSSG) were measured by luminescence in cells growing in opaque-walled 96-well plates, using the GSH/GSSG-Glo Assay (Promega) according to the manufacturer’s instructions. The resulting luminescent signal was measured in Wallac Victor 1420 multi-label counter, using triplicate samples per treatment group and subtracting blank measurements to produce net results. GSH/GSSG ratios were calculated
directly from Net Relative Luminescence Units (RLU) measurements using the equation
\[
\text{GSH/GSSG ratio} = \frac{\text{Net total glutathione RLU} - \text{Net GSSG RLU}}{\text{Net GSSG RLU}/2}.
\]

**Metabolic Flux Analysis (Seahorse XF)**

Metabolic Flux analysis in a Seahorse XF 24 Analyzer was used to assess the effect of NNT knockdown on mitochondrial bioenergetics, applying the Seahorse XF Cell Mito Stress kit. Cells were plated in Seahorse XF microplates the day before the experiment at a density of 100,000 cells/well. Changes in oxygen concentration provide the oxygen consumption rate (OCR), which is a measure of mitochondrial respiration. Changes in proton concentration (or pH) provide the extracellular acidification rate (ECAR), reflective of the rate of glycolysis. Measurements were taken at baseline and after successive application of compounds interfering with oxidative phosphorylation: oligomycin (complex V inhibitor, 2 μM), Carbonyl cyanide-p-trifluoromethoxyphenylhydrazone (FCCP, mitochondrial uncoupler, 1 μM) and Antimycin A + Rotenone (Complex I and III inhibition, 1 μM). Results were normalized to protein concentration, measured by the BCA Protein Assay Kit (Thermo Fisher).

**Cell proliferation and apoptosis**

Cell proliferation was assessed in 96-well plates (loading concentration 6-8,000 cells/well), using the CyQuant Proliferation Assay Kit (Thermo Fisher) and following the manufacturer’s instructions. Cell DNA fluorescence was measured at the end of the time course, i.e. 166 hours post siRNA transfection and/or 96 hours after treatment. The beginning of treatment was used as the baseline time point (t=0) for each proliferation series; for siRNA knockdown experiments, 72 hours post transfection was taken as the baseline time point. Proliferation rates were provided by the following ratio: \([(\text{end cell number} - \text{baseline cell number})]/\text{baseline cell number}\).
Cellular apoptosis was assessed using the Caspase-Glo 3/7 Assay kit (Promega), a luminescence-based assay measuring Caspase 3 and 7 activity in cell lysates, and following the manufacturer’s instructions. Luminescent signals were quantified using the Wallac Victor 1420 multilabel counter. At the end of the assay, media and reagents were removed from all wells and stored at -80°C. The next day, relative quantification of cell number was performed by use of the CyQuant® Proliferation Assay Kit, as described above. Luminescence values obtained in the caspase assay were normalized to the fluorescence results of the proliferation assay.

Paraquat and auranofin were purchased from Sigma-Aldrich (UK). Buthionine sulfoximine (BSO) was purchased from Cayman Chemical (USA).

**In vitro steroid profiling by liquid chromatography-tandem mass spectrometry (LC-MS/MS)**

Steroid synthesis by NCI-H295R cells was assessed by comprehensive multi-steroid profiling employing liquid chromatography-tandem mass spectrometry (LC-MS/MS), as described previously (14, 15). Steroid extraction and analysis by LC-MS/MS are discussed in Supplementary Methods.

**RNA sequencing**

RNA was prepared in triplicate from NCI-H295R KD siRNA, SCR siRNA (72 hours post-transfection), KD shRNA and SCR shRNA cells using the RNeasy Mini kit (Qiagen). Libraries were generated using the TruSeq Stranded mRNA Library Prep Kit (Illumina, USA). 4 nM library (containing the 16 pooled libraries) was sequenced on a NextSeq500 System (Illumina). Pathway analysis on sequencing data was completed using GAGE V2.22 package from Bioconductor release 3.2 and referencing the KEGG pathways. Differentially expressed genes were considered significant applying a false discovery rate of <5% (q<0.05). Differentially regulated pathways
were called at a p value of <0.01. A detailed description of the methodology for RNA sequencing and pathway analysis can be found in Supplementary Methods.

In addition, RNA sequencing data from recently published work on three different mouse strains (Nnt inactivating mutation, C57BL/6J (RRID:MGI:3702942); wild-type, C57BL/6NHSd (RRID:MGI:2161078); and transgenic Nnt overexpressor, C57BL/6J BAC (16)) were re-analyzed employing the same pathway analysis as for the human cell-based model; detailed information on this dataset can be found in Supplementary Methods.

Metabolome analysis

Cell and media samples were prepared for non-targeted metabolome analysis through quenching cell metabolism with a mix of acetonitrile, methanol and water (Sigma-Aldrich). The process of sample generation and analysis is described in more detail in the Supplementary Methods.

Statistical analysis

Statistical analysis was performed using GraphPad Prism 7 Software (RRID:SCR_002798). Data are represented as mean ± SEM values, unless otherwise stated. Comparisons were made using Student’s paired t-test for normally distributed data or Wilcoxon’s signed-rank test for data not following a Gaussian distribution. Multiple comparisons (BSO and auranofin treatments) were performed by one-way ANOVA followed by post-hoc multiple comparison testing. Statistical methods for the RNA sequencing and untargeted metabolome analysis are detailed in Supplementary Methods.
Results

Transient and stable NNT knockdown

Transient NNT silencing by siRNA knockdown was employed to explore the acute effects of NNT loss on ACC cells. NNT siRNA transfection in NCI-H295R cells yielded efficient gene silencing for at least 166 hours post-transfection with two different siRNAs (Suppl. Fig. 1A-B). All subsequent experiments were performed with the siRNA that gave the best knockdown results on real-time PCR and Western Blotting (referred to here as KD siRNA). The second siRNA (KD siRNA2) was used to corroborate the results of proliferation and apoptosis assays, whose results are most likely to be distorted by off-targets effects.

Stable NNT silencing by shRNA knockdown was used to delineate the long-term effects of NNT loss on ACC cells. Stable NNT knockdown in NCI-H295R cells was achieved by lentiviral transfection with shRNA-expressing plasmids and selection with puromycin, and resulted in permanent NNT silencing (Suppl. Fig. 1C-D).

NNT siRNA knockdown increases cellular oxidative stress

Given the central role of NNT within the mitochondrial ROS scavenging network, we hypothesized that NNT knockdown will increase oxidative stress in NCI-H295R cells. To test this, we measured the intracellular ratio of reduced to oxidized glutathione (GSH/GSSG), an established marker of oxidative stress; a decrease in the GSH/GSSG ratio indicates that the proportion of oxidized intracellular glutathione is increased as a result of higher intracellular ROS levels. Indeed, we observed a statistically significant (p<0.05) decrease in the GSH/GSSG ratio in NNT KD siRNA-transfected cells 96 hours post-transfection (Fig. 2A).

NNT siRNA knockdown suppresses cell proliferation and induces apoptotic cell death

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Cell proliferation rates were assessed over the time window from 72 to 166 hours post-transfection, a period with consistent NNT knockdown confirmed at protein level (Suppl. Fig. 1). NNT knockdown by KD siRNA transfection led to a marked decrease in cellular proliferation rates (Fig. 2B). These results were corroborated by use of a second siRNA against NNT, which completely obliterated cell proliferation (Suppl. Fig. 2).

To establish whether the increased oxidative stress observed with NNT KD leads to higher rates of apoptosis - as predicted by ROS physiology- we measured intracellular caspase 3 and 7 activity 120 hours post-transfection. We also quantified relative cell numbers by DNA fluorescence at the same time-point to standardize results to cell number. NNT KD siRNA cells exhibited significantly higher caspase 3/7 activity than SCR siRNA cells (p<0.05), confirming our hypothesis that NNT knockdown triggers cell death by apoptosis (Fig. 2C). The effect was even more marked with the alternative siRNA against NNT (Suppl. Fig. 2).

NNT siRNA knockdown sensitizes cells to oxidative stress

Next, we evaluated changes in mitochondrial respiration by direct measurement of the cellular oxygen consumption rate (OCR), using Extracellular Flux analysis. Despite the location of NNT in the inner mitochondrial membrane, we observed no statistically significant difference between NNT KD siRNA- and SCR siRNA-transfected cells, either at baseline or in response to mitochondrial respiration disruptors (Fig. 2D). Baseline extracellular acidification rate (ECAR), representative of the glycolytic rate, was also similar between the two groups (Fig. 2E).

Considering the integral role of NNT in mitochondrial antioxidant defense and the detrimental impact of NNT inhibition on redox balance, we further hypothesized that NNT loss will render NCI-H295R cells more sensitive to chemically induced oxidative stress. To assess this assumption, we treated NCI-H295R cells with a sub-toxic dose of paraquat, a pesticide which
induces oxidative stress in vitro generating superoxide. Treatment with 10 μM of paraquat for 96 hours led to a statistically significant decrease in cell proliferation in cells transfected with KD siRNA, but not in their counterparts that had been transfected with SCR siRNA (Fig. 2F).

**Redox adaptation develops with stable NNT knockdown**

To explore the long-term metabolic consequences of NNT silencing in NCI-H295R cells, we employed a different model, involving stable transfection with shRNA against NNT. With long-term culture under persistent NNT silencing (4-12 weeks post-transfection), cells managed to restore their redox balance to the levels of their SCR shRNA-transfected counterparts (Fig. 3A).

**ACC proliferation remains suppressed with stable NNT knockdown**

The distinct metabolic consequences of NNT silencing in the stable knockdown setting, in comparison to acute knockdown by NNT siRNA, translated into an attenuated response with respect to cellular proliferation and viability. Proliferation rates remained significantly lower in KD shRNA-transfected cells compared to the SCR shRNA-transfected controls; however, this was less pronounced than the decrease in proliferation we observed with siRNA-mediated knockdown (Fig. 3B). Apoptotic rates did not differ between SCR shRNA and KD shRNA cells (Fig. 3C), in keeping with the restoration of redox homeostasis we had ascertained based on the reduced/oxidized glutathione ratio.

Interestingly, NNT KD shRNA cells consumed more oxygen than SCR shRNA cells at baseline (Fig. 3D). This finding potentially reflects higher energy needs in NNT deficient cells. The same trend was observed in ECAR, a surrogate marker of glycolysis, but without reaching statistical significance (Fig. 3E). Finally, stable NNT knockdown did not enhance cell sensitivity to oxidative stress induced by paraquat. (Fig. 3F).
Transient, but not stable, NNT knockdown paradoxically stimulates steroidogenesis

The effects of NNT silencing on steroidogenesis were evaluated by comprehensive multi-steroid profiling in cell media by LC-MS/MS, as well as gene expression analysis by qRT-PCR. We postulated that NNT silencing will disrupt steroidogenesis, either depriving mitochondrial steroidogenic monoxygenases [cholesterol side-chain cleaving enzyme (CYP11A1), 11β-hydroxylase (CYP11B1), aldosterone synthase (CYP11B2)] of their essential electron donor NADPH, or due to oxidative stress-induced down-regulation of key steroidogenic enzymes. Surprisingly, NNT KD siRNA-transfected cells actually produced significantly more glucocorticoids (cortisol) and androgens (androstenedione) than controls (Fig. 4A-B). Individual enzyme activities were determined as product-to-substrate ratios for three key steroidogenic enzymes, 11β-hydroxylase (CYP11B1), 21-hydroxylase (CYP21A2), and CYP17A1 17/20-lyase activity; all three displayed higher activity in NNT KD siRNA-transfected cells, in keeping with a paradoxical generalized stimulation of steroidogenesis by acute NNT loss (Fig. 4C-E).

By contrast, in the shRNA-transfected cells with chronic NNT silencing we observed no significant impact on steroidogenesis, with similar rates of cortisol or androstenedione synthesis between KD shRNA and SCR shRNA cells (Fig. 4A-E).

We also explored the gene expression alterations underpinning the enhanced steroid production of cells transfected with KD siRNA, comparing the expression of core steroidogenic genes (StAR, CYP11A1, CYP21A2, CYP17A1, 3βHSD2) between KD siRNA and SCR siRNA cells by qRT-PCR. There was a statistically significant increase in the expression of cytochrome P450 (CYP) type 2 steroidogenic enzymes located in the ER CYP21A2 (p<0.05), CYP17A1 (p<0.05), as well as the ER dehydrogenase HSD3B2 (p<0.01) in NNT KD siRNA cells (Suppl.
Table 2). CYP11B1 and CYP11B2 expression levels were too low to be quantified by qRT-PCR in our cells.

Analysis of gene expression by RNA sequencing in an extended panel of 14 steroidogenic genes indicated a significant up-regulation of CYP21A2 (q<0.05) and CYP17A1 (q<0.01), as well as the ACTH receptor MC2R (q<0.01) in the transient NNT knockdown model; no significant changes were observed in the stable knockdown model (Fig. 4F).

NCI-H295R cells are sensitive to glutathione depletion and thioredoxin reductase inhibition

Given the effects of NNT inhibition on NCI-H295R cell proliferation, we went on to evaluate the sensitivity of ACC cells to isolated inhibition of each of the two pillars of mitochondrial antioxidant defense: the glutathione pathway and the thioredoxin pathway.

We used buthionine sulfoximine (BSO), a potent inhibitor of the glutathione-producing enzyme c-glutamylcysteine ligase, to deplete intracellular glutathione. We observed a decline in cell proliferation with a BSO dose of ≥100 μM after 96 hours of treatment (Fig. 5A).

Pharmacological manipulation of the alternative mitochondrial antioxidant pathway, the thioredoxin pathway, was achieved by auranofin, a gold complex agent with a well-established capacity to inhibit thioredoxin reductase. NCI-H295R treatment with doses of ≥1 μM was associated with major cytotoxicity (Fig. 5B).

Finally, dual inhibition of the glutathione and thioredoxin pathways by co-administration of low-dose BSO (50 μM) and auranofin (0.2 μM) resulted in marked cytotoxicity, suggesting that a potent synergistic effect can be achieved by dual pathway targeting (Fig. 5C).

Whole transcriptome and metabolome analyses reveal extensive metabolic perturbations with transient NNT knockdown, as well as changes in protein processing and polyamine metabolism
with stable NNT knockdown

In order to uncover the molecular mechanisms that underpin the effects we observed in the two models and the discrepancies between them, we applied whole transcriptome analysis in RNA extracted from four groups of NCI-H295R cells (NNT KD siRNA vs SCR siRNA, NNT KD shRNA vs SCR shRNA). 842 genes were differentially regulated between NNT KD siRNA and SCR siRNA cells; 247 genes were differentially regulated between NNT KD shRNA and SCR shRNA cells (q<0.05). Of note, only 17 of the genes regulated differentially between KD and SCR cells were identical in the two models, NNT KD siRNA and shRNA (Suppl. Table 3).

Differentially regulated pathways are visualized in Fig. 6A+B (p<0.01) and tabulated in Suppl. Table 4. In NNT KD siRNA cells, significant changes were observed in crucial pathways affecting cellular proliferation and viability (p53 pathway, MAPK pathway, checkpoint kinases). Interestingly, in KD shRNA cells one of the borderline significantly altered pathways (p<0.01, q=0.11) controlled protein processing in the ER, with up-regulation of genes encoding heat shock proteins (predominantly in the HSP40 family), chaperone proteins that facilitate correct protein folding and transfer of misfolded proteins to proteasomes for degradation (Suppl. Table 4) (17, 18). Other significantly up-regulated pathways with stable NNT KD included ribosomal genes (p<0.01, q<0.05) and pyrimidine metabolism (p<0.01, q=0.11), including an up-regulation of RNA II polymerases. Taken together, these findings hint at a higher protein turnover which may allow cells to swiftly replace proteins that have sustained irreversible oxidative damage.

Of note, recent rodent-based work by Meimaridou et al. performed RNA sequencing on mouse adrenal glands derived from three different mouse strains: a strain which carries an inactivating mutation of Nnt (C57BL/6J), a strain with wild-type Nnt expression (C57BL/6N), and transgenic mice overexpressing Nnt on the background of the NNT-deficient mouse strain (rescue...
model; C57BL/6J$^{\text{BAC}}$) (16). We have extended this work here, carrying out additional, pathway analysis on this data. Detailed information on significantly dysregulated pathways in C57BL/6J vs C57BL/6N mice and C57BL/6J vs C57BL/6J$^{\text{BAC}}$ mice are presented in Suppl. Table 5. Significant dysregulation of the major cell signaling pathway of mitogen activated signaling kinases (MAPK) was one of the salient molecular changes in both comparisons; the same pathway was also dysregulated with transient NNT knockdown in NCI-H295R cells. Oxidative phosphorylation was up-regulated in both comparisons, but changes were much more pronounced in the C57BL/6J vs C57BL/6N comparison and likely to reflect a strain, rather than gene, effect.

RNA sequencing in NCI-H295R cells was complemented by whole metabolome analysis performed separately in cells and corresponding cell culture supernatant. In the siRNA knockdown model, NNT silencing was associated with a significant metabolic perturbation when compared to the SCR siRNA cells. An increase in the presence of 44 oxidized fatty acids supports a shift to a more oxidized intracellular microenvironment (Fig. 6C and Suppl. Tables 6 and 7). This was not observed in the stable NNT knockdown model. Pathway enrichment analysis demonstrated that six important metabolic pathways were enriched (q<0.05): tricarboxylic acid (TCA) cycle, arginine and proline metabolism, pyrimidine metabolism, nicotinate and nicotinamide metabolism and glutathione metabolism. Additional to this, we observed statistically significant changes (q<0.01) for 16 acyl carnitines, 25 fatty acids, 15 acyl amino acids, 20 purine and pyrimidine metabolites and 5 metabolites present in the nicotinate and nicotinamide metabolic pathway. Taken together, these indicate a perturbation in mitochondrial fatty acid beta-oxidation (as shown by changes in fatty acids, acyl carnitines and TCA metabolites), changes in nucleotide synthesis and a potential overload of acetyl units.

In the stable NNT shRNA knockdown model, pathway enrichment analysis highlighted
changes in purine metabolism ($q<0.05$) and NNT knockdown cells exhibited a significant ($p<0.05$) rise in several purine and pyrimidine metabolites (Fig. 6D and Suppl. Tables 6 and 7), mirroring the results of the transcriptome analysis. Polyamine (spermine, spermidine) metabolism was also significantly modified in both models: KD siRNA cells displayed a dramatic increase in polyamine catabolism (accumulation of diacetyl-spermine, diacetyl-spermidine, spermine dialdehyde) leading to depletion of spermine and spermidine, a response that has been associated with arrest of cell growth (Suppl. Tables 6 and 7) (19). Conversely, KD shRNA cells exhibited a significant rise in spermine, a polyamine that can act as a ROS scavenger (20, 21).
Discussion

With this work, we have explored the immediate and longer-term impact of NNT silencing on ACC cells with respect to redox balance, mitochondrial bioenergetics, cell proliferation and viability, and steroidogenesis, using two distinct in vitro knockdown models in the human adrenocortical carcinoma cell line NCI-H295R. Our aim was to establish whether NNT inhibition can have therapeutically beneficial effects with respect to control of tumor growth and steroid excess. We hypothesized that NNT inhibition would compromise the ability of adrenocortical mitochondria to deal with oxidative stress, leading to progressive accumulation of ROS. ROS excess has multiple toxic sequelae, and can directly impair cell viability triggering apoptosis (8, 12). Importantly, the adrenal-specific clinical phenotype in humans and the reported increased rate of adrenocortical cell apoptosis in otherwise healthy NNT mutant mice suggest that this manipulation may selectively target ACC cells, sparing other organs. This susceptibility of adrenocortical cells to mitochondrial antioxidant pathway disruption can be explained by the fact that enzymes involved in the rapid adrenal steroid response to stress represent a major additional source of ROS in the adrenals, increasing their dependence on efficient ROS scavenging (10, 11).

In keeping with our hypothesis, we found that in the acute setting (siRNA-mediated transient knockdown), NNT loss increased intracellular oxidative stress. Redox balance perturbations in response to NNT loss have been previously demonstrated in a limited number of cell lines in vitro, as well as in lymphocytes derived from NNT mutant patients ex vivo (4, 22-25). These findings are in line with the biological role of NNT as a major mitochondrial generator of NADPH, the essential provider of reducing equivalents to the two main antioxidant pathways (26).

Importantly, in our study we observed that NNT silencing led to an immediate and marked inhibition of cell proliferation accompanied by increased apoptotic rates. This anti-tumor effect
was even more pronounced when using a second anti-NNT siRNA (KD siRNA2). This apparent difference in degree (but not in direction) of cell response raised the possibility of additional, off-target effects triggered by KD siRNA2; on interrogation of the NCBI Basic Local Assignment Search tool, however, neither of the two siRNAs share substantial homology with any genes that would be expected to impact cell proliferation and viability. The association between excessive oxidative stress and mitochondrial apoptosis has been well established in the literature (8, 12), but data on the effects of NNT loss on cellular proliferation and viability are limited. Transient NNT silencing was previously shown to increase rates of apoptosis in PC12 (rat pheochromocytoma) cells (22); stable NNT knockdown in human melanoma cells was associated with reduced viability and high apoptotic rates in vitro, as well as slower growth of melanoma xenografts in mice (27). Meimaridou et al reported high levels of apoptosis in the zona fasciculata of the adrenal cortex from NNT mutant mice, as well as NCI-H295R cells stably transfected with shRNA against NNT in vitro (4). Although ROS have typically been associated with a stimulation of cellular proliferation, a number of in vitro models have demonstrated the opposite effect (suppression of cell division), in a complex relationship that may depend on the magnitude of ROS excess and/or tissue type (28, 29). NNT inhibition may also interfere with cellular proliferation in a ROS-independent way, curtailing the amount of NADPH available to fuel the pressing anabolic needs of malignant cells. In keeping with the major impact on cellular viability and proliferation, we observed far-reaching metabolic effects of NNT knockdown implicating several areas of cell metabolism, including mitochondrial fatty acid oxidation, polyamine metabolism and nucleotide synthesis. The enhanced cellular sensitivity to oxidative stress in the aftermath of NNT silencing (paraquat treatment) is translationally important, as oxidative stress is induced by a number of classic chemotherapy agents, contributing to their cytotoxic effect (8, 30). NNT inhibition could
represent a feasible strategy to sensitize ACC to such drugs

The longer-term effects of NNT loss on ACC cells, as delineated in the stable knockdown model, were disparate from the ones encountered in the acute setting. Importantly, with long-term culture under constant NNT silencing, NCI-H295R ACC cells managed to restore their redox balance. This compensation abrogated the pro-apoptotic early impact of NNT loss. Interestingly, a persistent proliferative handicap was demonstrated, though this was less marked than the one observed in the acute setting. This may be attributable to the limited supply of NADPH in the absence of NNT. Extracellular flux analysis revealed higher rates of oxygen consumption in KD shRNA cells, a response that may reflect higher energy needs or be driven by the spare NADH which fails to be converted to NADPH in NNT-deficient cells. Previous studies on the effect of NNT silencing on oxygen consumption have shown mixed results, which may be cell type-dependent (22, 25, 31).

Redox adaptation to oxidative stress has been previously described in tumor models in vitro; this process is driven by the strong selective pressure applied by oxidative toxicity and promoted by the genomic instability which characterizes the oxidized intracellular microenvironment (32). We obtained insights into how this adaptation was facilitated in our model by comprehensive transcriptome and metabolome analysis. In NNT KD shRNA cells, we observed an up-regulation of genes that are involved in protein folding in the ER, as well as in the identification and degradation of damaged proteins. Purine and pyrimidine metabolism was activated in these cells, and ribosomal genes were up-regulated. Taken together, these findings hint at increased protein turnover, involving degradation of damaged protein and acceleration of new protein synthesis. This may represent a key compensatory mechanism against oxidative stress, achieving the timely removal and replacement of irrevocably damaged (oxidized) proteins. The
observed increase in oxygen consumption could provide additional energy to fuel this process. Of note, we recently described up-regulation of chaperone proteins in the adrenals of NNT-deficient mice (16). The additional pathway analysis we performed on the same RNA sequencing data from that rodent work displayed otherwise limited overlap with our in vitro model, likely reflecting the expected biological differences between a healthy mouse adrenal and a malignant human adrenal cell line.

Interestingly, polyamine metabolism exhibited dramatic shifts in opposite directions in the two models. Polyamines (spermine, spermidine) are versatile cationic molecules involved in a number of cell processes, including ROS scavenging and cell proliferation (19, 33). High endogenous polyamine levels have been found in a number of cancer types (34). Acute NNT loss was accompanied by a rapid accumulation of acetylated catabolic products of polyamines, leading to depletion of spermine and spermidine. Polyamine catabolism can both be triggered by oxidative stress and generate hydrogen peroxide (H$_2$O$_2$), creating a vicious cycle that propagates ROS accumulation (33, 35). Indeed, stimulated polyamine catabolism has been associated with growth arrest and cell death in various in vitro models (19, 36). Conversely, in the chronic setting, stable NNT knockdown cells demonstrated increased spermine concentrations and no evidence of accelerated polyamine catabolism. This response is likely to represent a major facilitator of the successful redox adaptation in this model. Our findings underscore the importance of polyamine homeostasis in adrenocortical carcinoma cells.

Within the same framework, we also explored alternative antioxidant targets focusing on the glutathione and thioredoxin pathways. Pertinently, human mutations in thioredoxin reductase 2 have also been shown to result in isolated glucocorticoid deficiency (37). We used BSO to inhibit glutathione synthesis. BSO has shown anti-proliferative effects against a number of cell lines in
vitro (38-43). We observed a significant suppression of cell growth with doses of ≥100 μM, i.e. at doses that are clinically attainable in plasma with no serious toxicity (38). Auranofin, a gold complex-based agent able to inhibit thioredoxin reductase, also suppressed cell proliferation at doses ≥1μM and was associated with marked cytotoxicity at doses of ≥2 μM. Auranofin has also displayed anti-tumor activity against a number of cell lines in vitro and is currently being investigated in clinical trials against leukemia (44-47). Applying combined treatment with low doses of both agents, we observed a dramatic cytotoxic impact, suggesting that dual antioxidant targeting can achieve potent synergistic results.

The observed effects of NNT silencing on NCI-H295R steroidogenesis were surprising. In the acute setting, i.e. NNT siRNA knockdown, we observed a generalized stimulation of steroidogenesis, leading to increased glucocorticoid and adrenal androgen output by the cells. This was corroborated by a significant up-regulation of a number of steroidogenic enzymes. This response is contrary to what one might have anticipated considering mitochondrial NADPH is an essential cofactor to the steroidogenic cytochrome P450 enzymes CYP11A1, CYP11B1 and CYP11B2. Elucidating the mechanisms that drive this transient effect will require additional studies. The few studies exploring the relationship between ROS and steroidogenesis (mostly on testicular Leydig cell tumor cells) have reported a down-regulation of steroidogenic enzymes with oxidative stress (14, 48-50). Human patients with inactivating NNT mutations (4) and a murine Nnt deletion model (16) has been shown to have disrupted steroidogenesis; the data from our in vitro NNT knockdown models suggest that NNT loss is not limiting for adrenal steroidogenesis. Interestingly, Zhao et al. demonstrated a biphasic relationship between ROS and steroidogenesis, indicating that the direction of the effect is dose-dependent (51).

Taken together, we show that NNT silencing can induce cytotoxicity and impede cell
growth in adrenocortical carcinoma cells, as well as sensitize them to chemically-induced oxidative stress. Moreover, we have demonstrated how the plasticity of ACC cells can lead to the development of a compensatory molecular response with time and described how changes in polyamine metabolism and ER protein processing are involved in this process (Fig. 7). A limitation of our work is that it is based on a single cell line; however, NCI-H295R remains the only established, well characterized steroidogenic human ACC cell line. These results merit further exploration with *in vivo* studies to corroborate the effectiveness of mitochondrial anti-oxidant pathway targeting and explore its durability, alone or in combination with other pro-oxidant agents. The unique features of adrenocortical cells, with their high-volume ROS generation due to steroidogenesis, make ACC a most amenable target to this approach.
References


**Fig. 1:** A, Interaction between NNT and the mitochondrial antioxidant pathways. ETC: electron transfer chain; ATP: adenosine triphosphate; GSR: glutathione reductase; GSSG: oxidized glutathione; GSH: reduced glutathione; GPX1: Glutathione peroxidase 1; TXNRD2: thioredoxin reductase 2; TXN: oxidized thioredoxin; TXN-SH: reduced thioredoxin; GPX3: peroxiredoxin 3; O2\(^{-}\): superoxide; H\(_2\)O\(_2\): hydrogen peroxide; SOD2: Superoxide dismutase 2. B, NNT expression in patients with ACC (n=33), compared to patients with adrenocortical adenomas (n=22) and healthy adrenals (n=10). Gene expression was quantile-normalized and log-transformed as described (13); bars represent median and interquartile range (IQR). Gene expression across groups was compared applying the Kruskal-Wallis test, followed by post-hoc Bonferroni test. **p<0.01.

**Fig. 2:** Effects of transient (siRNA-mediated) NNT silencing on NCI-H295R cell redox balance, respiration, proliferation and viability. Bars represent mean ± SEM values, unless otherwise stated. A, GSH/GSSG ratio in NCI-H295R cells transfected with KD siRNA (96 hours post-transfection), normalized to the corresponding ratio of SCR siRNA-transfected cells. Significant suppression of the GSH/GSSG ratio in KD siRNA cells suggests higher intracellular oxidative stress. Bars represent median ± IQR values. *p<0.05; n=8 independent experiments. B, Proliferation rates observed in siRNA-transfected NCI-H295R cells, 72-166 hours post-transfection. ***p<0.001; n=14. C, Caspase 3/7 activity ratio in KD siRNA cells to SCR siRNA-transfected cells, after standardization to cell numbers (120 hours post-transfection). *p<0.05; n=8. D, Seahorse XF24 analysis of cellular oxygen consumption rate (OCR) at baseline and after successive application of three mitochondrial respiration inhibitors (166 hours post-transfection). Results were standardized to protein concentration. Bars represent median ± IQR values. FCCP: Carbonyl cyanide-p-trifluoromethoxyphenylhydrazone. A-mycin/Rot: Antimycin A plus Rotenone. p>0.05; n=4. E, Extracellular acidification rate (ECAR), surrogate marker of anaerobic glycolysis, standardized for protein concentration. p>0.05; n=4. F, Proliferation under low-dose chemically induced oxidative stress (paraquat 10 μM) in KD siRNA and SCR siRNA-transfected cells, normalized to corresponding cell proliferation without paraquat treatment. *p<0.05; n=6.

**Fig. 3:** Effects of stable (shRNA-mediated) NNT silencing on NCI-H295R cell redox balance, respiration, proliferation and viability. Bars represent mean ± SEM values, unless otherwise stated. A, GSH/GSSG ratio in NCI-H295R cells transfected with KD shRNA, normalized to the corresponding ratio cells transfected with SCR shRNA. Bars represent median ± IQR values. p>0.05, n=10. B, Proliferation rates over a 96-hour period. *p<0.05; n=13. C, Caspase 3/7 activity ratio in KD shRNA cells to SCR shRNA-transfected cells, after standardization to cell...
numbers. p>0.05; n=4. D. Seahorse XF24 analysis of cellular oxygen consumption rate (OCR) at baseline and after successive application of three mitochondrial respiration inhibitors. Results were standardized to protein concentration. Bars represent median ± IQR values. FCCP: Carbonyl cyanide-p-trifluoromethoxyphenylhydrazone. A-mycin/Rot: Antimycin A plus Rotenone. *p<0.05; n=7. E. Extracellular acidification rate (ECAR) standardized for protein concentration. p>0.05; n=7. F. Proliferation under chemically induced oxidative stress (paraquat) in KD shRNA and SCR shRNA cells, normalized to corresponding cell proliferation without paraquat treatment. p>0.05; n=13.

Fig. 4: Effects of NNT silencing on NCI-H295R steroidogenesis, delineated by LC-MS/MS steroid profiling in serum-free cell media and RNA sequencing. Cortisol production (A) and androstenedione production (B) over a 48-hour period in NCI-H295R cells transfected with siRNA or shRNA. A significant stimulation of cortisol and androstenedione synthesis was observed 72-120 hours post-transfection with KD siRNA. **p<0.01, *p<0.05; n≥5. C-E, specific enzyme activity derived from product to substrate ratios for 11β-hydroxylase (CYP11B1) (C), 21-hydroxylase (CYP21A2) (D) and 17,20-lyase (CYP17A1) (E) in siRNA and shRNA-transfected cells. **p<0.01, ***p<0.001; n≥5. F, Heat-map representation of steroidogenic gene expression changes induced by transient and stable NNT knockdown, as revealed by RNA sequencing. Scale represents log-fold changes in NNT knockdown cells compared to their respective (siRNA or shRNA) scrambled controls. *q<0.05, **q<0.01; n=3.

Fig. 5: Effect of glutathione and thioredoxin pathway inhibition on NCI-H295R cell proliferation. A, 96-hour treatment with incremental doses of BSO (0-200 μM), inhibitor of glutathione synthesis. Control cells were treated with vehicle only. *p<0.05, n=9. B, 96-hour treatment with incremental doses of auranofin (0-5 μM), a thioredoxin reductase inhibitor. Control cells were treated with vehicle only. Negative proliferation rates indicate net decrease in cell number after 96 hours of treatment. **p<0.01, ****p<0.0001; n=9. C, Combined glutathione and thioredoxin inhibition by use of low-dose BSO (50 μM) and auranofin (0.2 μM). *p<0.05; n=3.

Fig. 6: Whole transcriptome sequencing and non-targeted metabolome analysis of KD siRNA, SCR siRNA, KD shRNA and SCR shRNA-transfected NCI-H295R cells. A+B, Whole transcriptome sequencing in siRNA and shRNA-transfected NCI-H295R cells. A, Significantly altered molecular pathways (p<0.01, q<0.05) between KD siRNA and SCR siRNA cells and number of associated genes that are up/down-regulated in KD siRNA cells. B, Significantly altered molecular pathways (p<0.01) between KD shRNA and SCR shRNA cells and number of associated
genes that are up/down-regulated in KD shRNA cells. C+D, Non-targeted metabolome analysis in siRNA and shRNA-transfected NCI-H295R cells. C, Significantly up- and down-regulated intracellular metabolites (q<0.01) in KD siRNA-transfected cells, as compared to SCR siRNA-transfected cells. D, Significantly up- and down-regulated intracellular metabolites (p<0.05) in KD shRNA-transfected cells, as compared to SCR shRNA-transfected cells (no metabolites with q<0.05 in this comparison).

Fig. 7: Response of NCI-H295R cells to NNT silencing in the acute (transient knockdown) and chronic (stable knockdown) setting, with proposed redox adaptation mechanisms. Acute NNT knockdown induces oxidative stress as predicted by NNT’s role as a major NADPH generator; enhanced steroidogenesis and polyamine catabolism further accentuate ROS accumulation, triggering apoptosis and a sharp decline in cell proliferation. With time (stable knockdown), cells manage to adapt removing damaged proteins and enhancing spermine synthesis as an alternative, NADPH-independent ROS scavenger. This restores redox homeostasis and abrogates the original pro-apoptotic effect, but cellular proliferation remains suppressed. Horizontal arrows represent paucity of change.