Inhibition of NAMPT decreases cell growth and enhances susceptibility to oxidative stress

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Received December 27, 2016; Accepted June 28, 2017

DOI: 10.3892/or.2017.5793

Abstract. Nicotinamide adenine dinucleotide (NAD) is an essential molecule for living organisms and plays a vital role in aging and age-associated diseases. In eukaryotic cells, cellular NAD is mainly generated by the scavenge pathway in which nicotinamide phosphoribosyltransferase (NAMPT) catalyzes the formation of nicotinamide mononucleotide. Inhibition of NAMPT is a therapeutic strategy for cancer treatment. To explore the effects of NAMPT inhibition on cellular processes, cells were treated with 10 nM FK866, an NAMPT inhibitor, resulting in a decrease in the cellular NAD level, a lower growth rate, and enhanced susceptibility to oxidative stress as compared to the untreated cells. Quantitative proteomics revealed that 325 proteins were downregulated in the FK866-treated cells, and were involved in diverse cellular processes including nucleobase-containing compound metabolic process, protein metabolic process, antioxidant and DNA repair processes. Downregulation of 4 selected proteins was confirmed by western blotting and quantitative PCR. Downregulation of antioxidant proteins GRX1 and catalase, and DNA-repair proteins PCNA and PARP1 contributed to the enhanced susceptibility of FK866-treated cells to oxidative stress. FK866 treatment also caused mitochondrial dysfunction through downregulation of mitochondrial ribosomal proteins. Taken together, these results demonstrate that FK866 treatment efficiently decreases the cellular NAD level and induces autonomous changes in proteostasis, leading to cell growth inhibition and increased susceptibility to oxidative stress.

Introduction

NAD is an essential component in life, which plays a vital role in major biological processes (1-3). In cells, the NAD pool is composed of NAD, reduced nicotinamide adenine dinucleotide (NADH), nicotinamide adenine dinucleotide phosphate (NADP) and reduced nicotinamide adenine dinucleotide phosphate (NADPH). NAD and NADP are coenzymes for dehydrogenase (DH)-catalyzed reactions involved in biodegradation and biosynthesis. The NAD/NADH ratio also regulates mitochondrial functions (4-6). NADPH plays an important role in the cell defense system. NAD homeostasis is maintained by NAD synthesis, NAD catabolism, and non-redox NAD-dependent enzymatic reactions. Tryptophan is the precursor for de novo NAD synthesis.

In non-redox NAD-dependent reactions, NAD functions as a substrate for ADP-ribosyltransferases such as poly(ADP-ribose) polymerases (PARPs), mono(ADP-ribosyl)-transferases (ARTs), NAD(+)-dependent deacetylases (sirtuins), tRNA 2’-phosphotransferases, and ADP-ribosyl cyclases. Sirtuins utilize NAD to deacetylate acetyl-lysine residues in proteins (7-10). Sirtuin-catalyzed NAD-dependent deacetylations of nuclear and mitochondrial proteins are the basis for the broad range of cellular functions (11-14). As the substrate for ADP-ribosylation of nuclear proteins, NAD is essential for stress responses (15,16). Recent studies have revealed that NAD levels decline with age in different organisms (17-20). On the other hand, repletion of NAD precursors increases the cellular NAD levels against age-associated diseases and prolongs lifespan extension (21-23). A recent study demonstrated that CD38 levels are increased in aged mice, causing age-related NAD decline and mitochondrial dysfunction (24).

Furthermore, NAD is essential to tumor survival and progression by regulating multiple cellular processes, including metabolism, stress responses, and DNA integrity. Targeting NAD metabolism has evolved to be a new concept in cancer therapy (25,26). The cellular NAD levels are maintained by the NAD scavenge pathway in which the formation of nicotinamide mononucleotide is the rate-limiting step catalyzed by nicotinamide phosphoribosyltransferase (NAMPT). It has been amply documented that NAMPT expression is
highly expressed in tumors, making it a therapeutic target in cancer treatment (26,27). Over the last two decades, extensive studies have identified that FK866 efficiently inhibits NAMPT activity to decrease cellular NAD levels (28,29). Studies show that FK866-induced cell growth inhibition is regulated by p53, mTOR and autophagy pathways (30-34). However, few studies have systematically examined the effects of NAD decrease on proteostasis, cell proliferation and stress responses. Understanding the global impacts of the NAD levels on cellular processes is important for treating cancer and age-associated diseases. In the present study, we treated cells with FK866 and conducted a comprehensive analysis to decipher the effects of the decrease in NAD on proteostasis, cell growth, and cellular responses to oxidative stress.

**Materials and methods**

**Chemicals and reagents.** Phosphate-buffered saline (PBS), penicillin/streptomycin, Dulbecco's modified Eagle's medium (DMEM), normal and dialyzed fetal bovine serum were all purchased from Wisent (Montreal, QC, Canada). SILAC labeling reagents including DMEM medium, isotope encoded L-13C6 lysine-HCl, standard L-lysine-HCl, and standard L-arginine-HCl were purchased from Thermo Fisher (Waltham, MA, USA). Sequencing grade trypsin was purchased from Promega (Fitchburg, WI, USA). Iodoacetamide (IAA) and polybrene were purchased from Sigma (St. Louis, MO, USA). Anti-PARP1 antibody, anti-PCNA antibody and anti-β-actin antibody were purchased from Abmart (Shanghai, China). Anti-p53 antibody, anti-mouse secondary antibody and anti-rabbit secondary antibody were purchased from Cell Signaling Technology (Boston, MA, USA). The Cell Counting Kit-8 (CCK-8) was purchased from Dojindo (Kumamoto, Japan). Hydrogen peroxide was purchased from Aladdin (Shanghai, China). BCA protein assay kit was purchased from Solarbio (Beijing, China). The Total RNA isolation and reverse transcription kit were purchased from Tiangen (Beijing, China).

**Cell culture and SILAC labeling.** Human 293T and A549 cell lines were purchased from the Cell Bank of the Chinese Academy of Sciences (Shanghai, China). The cells were grown in DMEM medium that was supplemented with 10% fetal bovine serum (FBS) and 1% penicillin/streptomycin at 37°C in a humidified incubator with 5% CO2. For SILAC labeling, 293T cells were washed twice with PBS prior to being cultured with the SILAC culture medium, which was prepared by mixing SILAC DMEM medium with 10% D-FBS, 1% penicillin/streptomycin, 146 mg/l isotope labeling L-13C6 lysine-HCl and 84 mg/l standard L-arginine-HCl. The cells were grown for 8 to 10 passages in SILAC medium and the full incorporation of isotope-encoded amino acid into proteins was tested. Cells were cultured for at least 12 h prior to FK866 treatment; cells were treated with FK866 dissolved in DMSO for 12 or 24 h. After treatments, the cells were then washed twice with ice-cold PBS and lysed with RIPA lysis buffer which consisted of 25 mmol/l Tris-HCl pH 7.6, 150 mmol/l NaCl, 0.1% SDS, 1% NP-40, 1% sodium deoxycholate, 1 mmol/l PMSF, and Roche Complete Protease Inhibitor Cocktail for 30 min on ice. Cell lysates were further clarified by centrifugation at 14,000 x g for 20 min at 4°C. The protein concentration in each sample was determined by using a BCA protein assay kit.

**Analysis of NAD and NADH content.** The NAD and NADH contents were determined using a NAD/NADH quantification kit according to the manufacturer's instructions (http://www.sigmaaldrich.com/content/dam/sigma-aldrich/docs/Sigma/Bulletin/1/mak037bul.pdf). Briefly, after cells were washed three times with cold PBS, cell metabolites were extracted using the NADH/NAD extraction buffer for 3 freeze/thawing cycles. After centrifugation, the supernatant was collected. For total NAD (NAD + NADH) detection, the supernatant was mixed with the NAD cycling enzyme to convert NAD to NADH. For NADH detection, the supernatant was heated to 60°C. Then, analytes and NADH standards reacted with a NADH developer reagent. The products were measured at the absorbance wavelength 450 nm (A450) to quantify NAD and NADH concentrations.

NAD and NADH concentrations were also measured by LC-MS analysis. Briefly, the cells were washed with ice-cold PBS and the metabolites were extracted using 80% methanol (~80°C). The metabolites were dried using a speedvac and re-dissolved in 80% methanol and used for LC-MS. The metabolites were separated by a Acquity UPLC BEH amide column (2.1x100 mm, 1.7-μm; Waters, Milford, MA, USA) that was interfaced with the Q-Exactive Mass spectrometer. Metabolites were characterized based on the retention time and the accurate mass measurement with <5 ppm mass accuracy. Trace Finder software was employed to identify the peaks and extract the quantitative information.

**Cell proliferation measurement with CCK-8 kit.** The cells were seeded in 96-well plates at 2,000 cells/well. The cell proliferation rate was measured with the CCK-8 kit according to the manufacturer's protocol (Dojindo Laboratories). Briefly, CCK-8 reagents were added into each well after cells grew for 0, 8, 16, 24, 32, 40, 48, 72, and 96 h, respectively. The absorbance at 450 nm was measured 3 h after CCK-8 addition.

**Susceptibility of FK866-treated and untreated cells to hydrogen peroxide and cisplatin.** Effects of hydrogen peroxide and cisplatin on cell growth in the FK866-treated and untreated cells were analyzed with the CCK-8 kit. Briefly, cells were seeded into 96-well cell culture microplates and incubated with 10 nM FK866 for 6 h prior to H2O2 or cisplatin treatment. Then, untreated and FK866-treated cells were treated with hydrogen peroxide and cisplatin. Susceptibility of FK866-treated and untreated cells to the hydrogen peroxide and cisplatin was measured using the CCK-8 kit. The CCK-8 reagents were added into each well after cells grew for 0, 8, 16, 24, 32, 40, 48, 72, and 96 h, respectively. The absorbance at 450 nm was measured 3 h after CCK-8 addition.

**Identification of differentially expressed proteins by proteomic analysis.** Proteomic analysis was conducted as previously described (35). Briefly, equal amounts of proteins from the FK866-treated and the untreated cells were mixed together and separated by 1D SDS-PAGE. The gel bands were excised,
reduced with 25 mM DTT and alkylated with 55 mM iodoacetamide in 40 mM ammonium bicarbonate at 37 °C overnight. The peptides were extracted twice and analyzed.

The digestion product was analyzed by LC-MS/MS, in which peptides were separated by a 65-min gradient elution at a flow rate of 0.250 µl/min with a Dionex Ultimate integrated nano-HPLC system which was directly interfaced with a Thermo Orbitrap Q-Exactive mass spectrometer. The Q-Exactive mass spectrometer was operated with the data-dependent acquisition method. A full-scan followed by 20 data-dependent MS/MS scans were acquired with normalized collision energy at 33%. Proteomic analysis was conducted in biological triplicates. The SEQuEST searching algorithm in the Proteome Discoverer software (version 1.4.1.14) was used to generate the peak lists from LC-MS/MS analysis, which were searched against the uniprot Human database (release date of November 20, 2015; 20,193 entries). The search criteria were the following: full tryptic specificity was required; one missed cleavage was allowed; carbamidomethylation (C) was set as the fixed modifications; the oxidation (M) was set as the variable modification; precursor ion mass tolerances were set at 10 ppm for all MS acquired in an orbitrap mass analyzer; and the fragment ion mass tolerance was set at 0.02 Da for all MS2 spectra acquired. The peptide false discovery rate was evaluated using the Percolator provided by Proteome Discoverer software. When the q-value was smaller than 1%, the match was considered to be correct. False discovery was evaluated based on peptide spectrum match when searched against the decoy database. Peptides only assigned to a given protein group were considered as unique. Protein quantitation was also conducted with Proteome Discoverer Algorithm (version 1.4). Briefly, the ratio in relative protein expression for each lysine-containing peptide was calculated using the peak area of Lys6 divided by the peak area of Lys0. The protein ratio was then averaged with all peptide ratios for that protein. Quantitative precision was expressed as protein ratio variability.

Western blot analysis. FK866-treated and the untreated cells were harvested and lysed on ice with RIPA lysis buffer. The supernatants were then collected and the protein concentrations were measured with the BCA protein assay kit. Proteins were separated on 1D SDS-PAGE gel and transferred onto a PVDF membrane with electroblotting. The PVDF membrane was blocked with 5% nonfat milk for 1 h at room temperature and incubated with primary antibody overnight at 4°C, washed three times with TBST buffer, and then incubated with anti-mouse or anti-rabbit secondary antibody labeled with HRP. The PVDF membrane was then washed three times with TBST buffer and developed with ECL reagents (Engreen, China). β-actin was used as an internal control and detected with the anti-β-actin antibody.

Quantitative real-time PCR (qPCR). Untreated and FK866-treated cells were cultured. Total RNA was extracted with the Total RNA isolation kit. cDNA was synthesized from 3 µg total RNA using the Reverse Transcription kit. All qPCR experiments were performed with the Roche LightCycler® 480II detection system with SYBR-Green incorporation according to the manufacturer's instructions and β-actin was used as an internal control. The primers were designed and acquired from the Primer Bank (http://pga.mgh.harvard.edu/primerbank/).

Statistical method. Statistical analysis was performed by using GraphPad Prism 5.0 software. Significant differences were determined by the Student’s t-test. p-values of <0.05 were considered to be significant.

Results

FK866 treatment decreases cellular NAD levels and growth rate but increases cell susceptibility to oxidative stress. To examine the effects of FK866 on cellular NAD levels, 293T and A549 cells were treated with FK866 at different concentrations for different periods of time. After FK866 treatment, cells were lysed and cellular metabolites were extracted followed by enzymatic assay or LC-MS analysis. The cellular NAD and NADH levels were determined, showing that the NAD level in the FK866-treated cells was ~5 times lower than that in the 293T cells. Similarly, the level of NADH was found to be 4 times lower in the FK866-treated cells than that in the 293T cells (Fig. 1A and B). Similar results were obtained by LC-MS analysis using the extracted ion currents from mass spectrometric analysis (data not shown). Cell proliferation rates for 293T and FK866-treated cells were determined using
the CCK-8 assay (Fig. 1C) which is a colorimetric assay for analysis of viable cells. The FK866-treated cells grew more slowly than 293T cells. At 96 h, the number of 293T cells was ~5 times more than that of the FK866-treated cells.

To determine the susceptibility of 293T and FK866-treated cells to hydrogen peroxide and cisplatin, cells were pretreated with DMSO or FK866 followed by incubation with different concentrations of hydrogen peroxide or cisplatin for 12 h. The cell viability was determined by the CCK-8 assay. The effects of hydrogen peroxide or cisplatin on cell viability are represented as the percentage of viable cells after a 12 h treatment (Fig. 2). When cells were treated with 200 µM H₂O₂ for 12 h, the percentages of viable cells were 20% and 95% for the FK866-treated and untreated cells, respectively. The percentage of viable cells decreased to 10% when FK866-treated cells were treated with 400 µM H₂O₂ for 12 h, indicating that FK866 treatment made 293T cells extremely sensitive to H₂O₂ treatment. When cells were treated with 50 µM cisplatin for 12 h, the percentages of viable cells were 20 and 50% for FK866-treated and the untreated cells, respectively. Similar results were also obtained for FK866-treated A549 cells, in which FK866 treatment inhibited the growth of A549 cells while increased cell susceptibility to oxidative stress (Fig. 3).

Identification of differentially expressed proteins between 293T and FK866-treated cells. We then carried out proteomic analysis to identify differentially expressed proteins between 293T and FK866-treated cells. Equal amounts of proteins from 293T and FK866-treated cells were pooled and separated by 1D SDS-PAGE. SILAC-based quantitation was used to determine the differentially expressed proteins. The experiments were repeated three times and ~4,200 proteins were identified in each experiment. The false-positive rate was estimated to be <1%. Based on SILAC ratios (>1.5 or <0.67), 384 proteins were found to be differentially expressed between 293T and FK866-treated cells, in which 325 proteins were downregulated and 59 were upregulated (data not shown). In order to understand the biological relevance of the identified proteins, Gene Ontology (GO) was employed to cluster the downregulated proteins according to their biological processes which are summarized via a pie plot using the PANTHER bioinformatics platform (http://www.pantherdb.org/) (Fig. 4). Three hundred and twenty-five proteins were classified into several significant groups of biological processes according to their molecular functions including primary metabolism, amino acid metabolism, DNA repair, and cell cycle regulation.

Verification of differentially expressed proteins by western blotting and qPCR. Among the differentially expressed proteins, protein expression levels of PARP1, p53 and GRX1 were lower in the FK866-treated cells than those in the untreated cells, whereas angiomotin expression was higher. Western blot analysis confirmed that the FK866 treatment
Figure 4. Functional classification of differentially expressed proteins between the untreated and FK866-treated cells with PANTHER (http://www.pantherdb.org).

Figure 5. Confirmation of differentially expressed proteins by western blotting and qPCR. (A-D) Band intensities from western blot images of the selected proteins from the untreated and FK866-treated cells. (E) qPCR analysis of mRNA expression of selected genes from the untreated and FK866-treated cells.
induced downregulation of PARP1, p53, and GRX1, and angiomotin upregulation (data not shown). Band intensities in western blot images were quantified using Image Lab 4.0.1 software (Fig. 5A-D). The changes in the band intensities from western blotting were comparable to the SILAC ratios. Moreover, qPCR analysis was employed to quantify changes in mRNA levels of the selected genes. Results showed that the mRNA expression levels of p53 and PARP1 genes were lower in the FK866-treated cells than those in the untreated cells (Fig. 5E). However, downregulation of mRNA expressions of p53 and PARP1 genes was less significant than the corresponding changes in protein expression.

Discussion

NAD plays an important role in all aspects of biological processes and its cellular levels are regulated by both NAD synthesis and consumption. To characterize the effects of the NAD decrease on cell proliferation, cell responses to oxidative stress, and protein homeostasis, we treated cells with FK866, an NAMPT inhibitor. FK866 treatment led to an 80% decrease in the cellular NAD level. Similarly, the level of NADH was also significantly decreased in the FK866-treated cells as compared to the untreated cells (Fig. 1B). By measuring the proliferation rates of the untreated and FK866-treated cells (Fig. 1C), we observed that FK866-treated cells grew slower than that observed in the untreated cells, indicating that the lower cellular NAD and NADH levels in the FK866-treated cells slowed down cell proliferation. we measured the cell survival rates after treating the untreated and FK866-treated cells with hydrogen peroxide and cisplatin, and found that FK866-treated cells were extremely sensitive to oxidative stress induced by hydrogen peroxide and cisplatin. This indicates that the cellular NAD levels directly affect the responses to oxidative stress.

To identify factors leading to cell growth inhibition and enhanced cell susceptibility to oxidative stress in the FK866-treated cells, we used the SILAC method to quantify proteins differentially expressed between the untreated and FK866-treated cells. We identified approximately 4,000 proteins in three repeated experiments. Among them, 384 proteins were differentially expressed between the untreated and FK866-treated cells, and these were found to participate in a variety of cellular processes, including RNA process, cell metabolism, stress responses, and cell cycle control and DNA repair. Based on GO analysis, 47 downregulated proteins were associated with carbohydrate metabolisms, nucleobase-containing compound metabolic process, and protein metabolism, suggesting that the cellular NAD level adversely affected cellular metabolic process. Quantitative proteomics also showed that proteins associated with mitochondrial mRNA translation were downregulated in the FK866-treated cells, including mitochondrial ribosome subunits, alanine-tRNA ligase, threonine-tRNA ligase and ATP synthase subunits (Fig. 6A), indicating that mitochondrial protein translation was impaired in the FK866-treated cells. These results suggest that downregulation of proteins associated with primary metabolic process and mitochondrial protein synthesis contributes to the FK866-mediated growth inhibition.

Quantitative proteomics also showed that six antioxidant proteins 15 kDa selenoprotein (Sep15), thioredoxin reductase 1 (TXNRD1), selenoprotein H (SELH), catalase (CAT), glutathione peroxidase 1 (GPX1) and thioredoxin-like protein 1 (TXNL1) were downregulated in the FK866-treated cells (Fig. 6B). These proteins are parts of the cellular defense system against oxidative stress: GPX1 catalyzes the reduction of H2O2 and other hydroperoxides and protects cells from reactive oxygen metabolites; and SELH is a novel nucleolar oxidoreductase that protects cells from oxidative damage (36-38). Downregulation of antioxidant proteins in the FK866-treated cells confirmed that the treated cells were more sensitive to oxidative stress. It is worth noting that FK866-mediated PARP1 downregulation was identified by both proteomics and western blotting. PARP1 plays an important role in regulating the cellular NAD level. PARP1 interacts with nuclear NAD synthase to increase the local NAD concentration that regulates PARP1-dependent gene expression, and PARP1 knockdown can increase the cellular NAD level (39,40). Our results revealed that a decrease in NAD induced autonomous PARP1 downregulation. Furthermore, downregulation of PARP1 and other DNA repair proteins, including RAD51, ALKBH1, and HMGB2 evidently led to the increased susceptibility to oxidative stress in the FK866-treated cells.
In conclusion, the present study investigated the effects of the cellular NAD content on cellular responses by quantitative proteomics. The cellular NAD level was decreased by 80% in the FK866-treated cells, which significantly altered the protein homeostasis. Our data indicated that the low cellular NAD level impairs the primary metabolic process and mitochondrial protein translational machinery, leading to the growth inhibition in FK866-treated cells. Downregulation of antioxidant proteins and DNA-repairing proteins contributed to the enhanced susceptibility of the FK866-treated cells to oxidative stress.

Acknowledgements

We thank the Proteomics Facility at Tsinghua University for providing proteomic analysis. This study was supported by Projects of the Natural Science Foundation of Shandong Province (ZR2016CL08, to RH.X.), Medical and Health Technology Development Program of Shandong Province (2016WS0013, to RH.X.), and the National Natural Science Foundation of China (81670064, to C.J.L.).

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