SHORT REPORT

Kynurenine Pathway Metabolism is Involved in the Maintenance of the Intracellular NAD⁺ Concentration in Human Primary Astrocytes

Ross Grant¹,², Susan Nguyen¹ and Gilles Guillemin¹,³
¹School of Medical Sciences, University of New South Wales, Sydney NSW Australia. ²Australiasian Research Institute, Sydney Adventist Hospital, Sydney NSW Australia. ³St. Vincent's Centre for Applied Medical Research, St. Vincent's Hospital, Sydney NSW Australia. Corresponding author email: r.grant@unsw.edu.au

Abstract: Efficient synthesis of NAD⁺ is critical to maintaining cell viability in all organs of the body. However, little is known of the pathway(s) by which cells of the central nervous system produce NAD⁺. The aim of this study was to investigate the relationship, between tryptophan degradation via the kynurenine pathway (KP) and de novo NAD⁺ synthesis in human astrocytes, a major cell type within the brain. In this study we observed that inhibition of single enzymes of the KP resulted in significant decreases in NAD⁺ levels in astroglial cells after a 24 hr period. We also observed that astrocytes cultured in media deficient in tryptophan, nicotinic acid and nicotinamide resulted in a 50% decrease in NAD⁺ levels after 24 hrs. This decrease in NAD⁺ was partially restored by supplementation of the culture media with either tryptophan or kynurenine, or nicotinic acid or with supply of the salvage pathway precursor nicotinamide.

Keywords: NAD, Tryptophan, Astrocyte, nicotinic acid, nicotinamide

International Journal of Tryptophan Research 2010:3 151–156

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**Introduction**

Nicotinamide adenine dinucleotide (NAD\(^+\)), as the parent molecule for the pyridine family of nucleotides, (NADH, NADP and NADPH) is an important coenzyme enabling electron transfer in the oxidative production of ATP and for hydride ion transfer in many enzyme reactions. Depleted mitochondrial NAD\(^+\) impairs respiration and ATP synthesis resulting in energy crisis and cell death.\(^1\) More recently NAD\(^+\) has also been identified as a primary substrate for other important enzymes including poly (ADP-ribose) polymerase (PARP) and the sirtuin family of de-acetylase enzymes. PARP is a nuclear enzyme, activated by DNA strand breaks that are involved in DNA repair and maintenance of genomic integrity. PARP uses up NAD\(^+\) to make ADP ribose polymers. An increase in DNA damage (often due to oxidative stress) can rapidly deplete the cell of NAD\(^+\) resulting in reduced ATP production and cell death.\(^2,3\) NAD\(^+\) is also a substrate for a relatively new class of NAD\(^+\) dependent de-acetylase enzymes called the sirtuins (SIRT1-7). Sirtuin enzymes consume NAD\(^+\) in the process of removal of acetyl groups from histone and non-histone moieties. Sirtuin targets consist of a variety of transcription factors and proteins involved in important processes, such as stress resistance and cell survival.\(^4\) Enhancing sirtuin activity has been shown to promote longevity in a range of organisms.\(^5\) As NAD\(^+\) plays a central role in these cellular systems, synthesis of NAD\(^+\) is essential to ensuring the health of the cell and associated organ systems.

*De novo* synthesis of NAD\(^+\) from tryptophan through the kynurenine pathway (KP) has been studied in the periphery and established as a primary source of NAD\(^+\), in the kidney and liver.\(^6\) We have previously reported that in the rat central nervous system (CNS), NAD\(^+\) concentrations can be regenerated in astrocytes using nicotinic acid, nicotinamide or quinolinic acid\(^7\) and that some downstream KP metabolites can affect NAD\(^+\) levels in human astrocytes.\(^8\) However the relative importance of KP and/or salvage pathway (Fig. 1) metabolism to NAD\(^+\) synthesis in primary human brain cells is yet to be clarified. We have earlier reported that human astrocytes are deficient in one of the enzymes of the tryptophan to NAD\(^+\) metabolic cascade.\(^9\) It is therefore not clear whether tryptophan catabolism is involved in NAD\(^+\) synthesis in these cells. Clarification of this question is important as NAD\(^+\) depletion is becoming increasingly recognised as a cause of cell death in CNS inflammatory and degenerative disorders.\(^10\)

The aim of this study was therefore to investigate the relationship, between kynurenine pathway (KP) metabolism and *de novo* NAD\(^+\) synthesis in astrocytes, the most numerous cell type of the CNS. Results from this study, showing the dependence of CNS cells on KP metabolism for NAD\(^+\) synthesis, implies that the KP could represent a significant clinical and therapeutic target.

![Figure 1. Schematic of NAD\(^+\) Biosynthesis; de novo and Salvage Pathways. (a) Indoleamine 2,3 dioxygenase 1 (1&2), (b) kynurenine formylase, (c) kynurenine 3-hydroxylase, (d) kynureninase, (e) 3-hydroxyanthranilic acid oxidase, (f) quinolinic acid phosphoribosyltransferase, (g) nicotinic acid phosphoribosyltransferase, (h) nicotinic acid mononucleotide adenylyltransferase (i) Glutamine dependent NAD\(^+\) synthetase, (j) Poly (ADP ribose) polymerase (PARP), (k) nicotinamide phosphoribosyltransferase OR pre-B-cell enhancing factor, (l) nicotinic acid mononucleotide adenylyltransferase. (α) This reaction proceeds non-enzymatically. Abbreviations: NIC, nicotinic acid; NAMN, nicotinic acid mononucleotide; NAAD, nicotinic acid adenine dinucleotide; NAD, nicotinamide adenine dinucleotide; NAM, nicotinamide; NMN, nicotinamide mononucleotide.](image-url)
Materials and Methods

Reagents and chemicals
All cell culture media and supplements were purchased from Invitrogen (Australia) unless otherwise stated. All reagents and chemicals used in experiments were purchased from Sigma Aldrich Chemical Co. (Australia) unless otherwise stated.

Cell cultures
Human primary astrocytes were grown in uncoated flasks (Falcon) and maintained in complete media (cRPMI), which contains RPMI 1640 media supplemented with 10% foetal bovine serum (FBS), 1% 2 mM glutamine (Sigma Aldrich, Australia) and 1% penicillin/streptomycin (Sigma Aldrich, Australia).

As part of its normal commercial formulation cRPMI contains 24 µM tryptophan and 8 µM nicotinamide. The cell medium was changed twice a week, and all cell cultures were kept incubated at 37 °C in 5% CO₂.

A day prior to experimental treatments, cultures were trypsinised (Trypsin 0.25%) and seeded at desired cell density, into 24 well plates (Falcon).

NAD⁺ precursor depleted RPMI
Depleted RPMI (dRPMI) was prepared using a standard mix formula for RPMI that included all nutrients except tryptophan (TRYP) and nicotinamide (NAM). Note that standard RPMI does not contain nicotinic acid (NIC) or kynurenine (KYN).

The dRPMI medium containing 10% FBS, 1% 2 mM glutamine (Sigma Aldrich, Australia) and 1% penicillin/streptomycin (Sigma Aldrich, Australia) was supplemented as required with the addition of substrates TRYP (25 µM), KYN (25 µM) NAM (10 µM) or NIC (10 µM). The concentrations chosen for TRYP and NAM supplementation correspond to that found in the commercially prepared RPMI (cRPMI). As KYN is not present in RPMI media this upstream metabolite was added at a concentration comparable to TRYP, which has been shown to increase in the culture medium of stimulated astroglial cells. NIC (the acid form of vitamin B3) is also not normally present in RPMI media and was added at the same concentration as its amide form, NAM to allow direct comparison. All supplements were prepared as sterile solutions in purified (>10 mΩ) water.

NAD⁺ precursor supplementation
Astrocytes were seeded into a 24-well culture plate at a density of 1 × 10⁶ cells/ml in and left to equilibrate in 5% CO₂ at 37 °C in complete RPMI (cRPMI) for 24 hr. The culture medium was then aspirated and each well was washed twice with warm phosphate buffered saline (PBS) before addition of 1 ml of fresh dRPMI (containing no TRYP, NAM or NIC). Selected NAD⁺ substrates TRYP (25 µM), KYN (25 µM) NAM (10 µM) or NIC (10 µM) were then added to appropriate cultures for 24 hrs before analysis.

Kp enzyme inhibition
Human primary astrocytes were plated into 24-well plates at a density of 1 × 10⁵ cells/ml in cRPMI and left to equilibrate for 24 hours before treatment. Cultured medium was aspirated from each well, washed twice with warm phosphate buffered saline (PBS), before addition of 1 ml of fresh cRPMI followed by addition of specific inhibitors (Table 1) for 24 hrs.

NAD⁺ analysis
Cultured medium was aspirated from each well. The cells were washed twice with PBS, before being homogenised by sonication in 500 µl of PBS (containing 12 mM nicotinamide as a PARP inhibitor). Quantification of NAD⁺ (H) in the cell was measured using the Thiazolyl blue microcycling assay adapted to a 96-well format.

Total protein assay
Cultured medium was removed from each well and the cells were washed twice with PBS, before

Table 1. Inhibitors used in the experiments.

| Inhibitor | Concentration (mM) | Action/notes |
|-----------|--------------------|--------------|
| 1-methyl-Tryptophan (1-MT) | 1 | Competitive IDO inhibitor |
| 3-ethoxy-β-carboline (3-EBC) | 0.2 | Non-competitive IDO inhibitor |
| Phthalic Acid (PHTH) | 0.04 | 'Dead-end' inhibitor of QPRT |

Notes: Previous studies have shown maximal IDO1 inhibition using 1 mM 1-MT. We have previously shown that 0.2 mM 3-EBC results in a maximal decrease in IDO1 activity in astroglial cells without measurable toxicity. Dose response analysis showed a maximal effect using PHTH at 0.04 mM (unpublished data).
being homogenised by sonication in 500 µl of PBS (containing 12 mM nicotinamide as a PARP inhibitor). The total protein per sample was measured using the commercially available Bradford assay method and reagents (BIORAD, Sydney, Australia), adapted to a 96-well microtitre plate format, using a 595 nm filter in a Multiskan MS microplate reader.

**Statistical analysis**
For all culture samples, significant differences between treatment groups, at $P < 0.05$ level of significance, were determined using One-way ANOVA and Dunnett's multiple comparison test. All results are represented as mean ± standard error mean.

**Results**

**Involvement of KP enzymes in de novo NAD$^+$ synthesis**
To determine whether tryptophan catabolism via the KP played a role in NAD$^+$ synthesis, we used small molecule inhibitors that reduced the activity of critical enzymes at the beginning and end of the KP.

Inhibition of the initial rate limiting enzyme indoleamine 2,3 dioxygenase (IDO1) with either the competitive inhibitor, 1-methyl tryptophan (1-MT) or the non-competitive inhibitor, 3-ethoxy-β-carboline (3EBC) significantly decreased intracellular NAD$^+$ by 14% and 26% respectively after 24 hrs. In a similar fashion, inhibition of the downstream KP enzyme, quinolinic acid phosphoribosyl transferase (QPRT) with the dead-end inhibitor phthalic acid (PHTH) resulted in a significant 29% decrease in intracellular NAD$^+$ levels after 24 hrs (Fig. 2).

**Importance of de novo and salvage pathway precursors to NAD$^+$ synthesis**
To help verify that the results obtained for enzyme inhibition were specific to KP metabolism, we tested whether the removal of KP or salvage pathway precursors (i.e. tryptophan, nicotinic acid and nicotinamide) from the incubation medium affected cellular NAD$^+$ levels.

We found that in the absence of all of these precursors a significant decrease of 50% in intracellular NAD$^+$ levels after 24 hrs was observed (Fig. 3).

Subsequent supplementation of the incubation medium with the upstream KP precursor tryptophan alone, increased NAD$^+$ levels by 75% within 24 hrs compared to control. Addition of nicotinic acid to the culture medium also increased NAD$^+$ levels significantly by 54% after 24 hrs (Fig. 3). Though an apparent increase in NAD$^+$ was observed with kynurenine and nicotinamide supplementation, these did not reach statistical significance (Fig. 3).

**Discussion**
We have previously shown that KP metabolism is linked to the maintenance of NAD$^+$ levels in murine primary astrocytes and human and murine cell lines. However, following the observation that primary human astrocytes do not express the KP enzyme, kynurenine 3-hydroxylase, we were unsure whether these cells could efficiently use tryptophan for NAD$^+$ synthesis. Using the dual approach of enzyme inhibition and NAD$^+$ substrate depletion, we investigated the role of the KP and salvage pathway metabolism in the maintenance of NAD$^+$ concentrations in primary human astrocytes.

In the present study, we observed that intracellular NAD$^+$ levels decreased significantly following inhibition of either the initial or downstream KP enzymes IDO1 or QPRT, respectively (Fig. 2). These results are consistent with our previous findings showing a clear relationship between IDO activity and NAD$^+$
substrates in the incubation medium (Fig. 3). Absence of either KP metabolites or salvage pathway applied in equimolar concentrations (Fig. 3) is indicative of preferred metabolism from this source. This is again consistent with our previous report in murine primary astrocytes showing NIC was also more efficient at promoting NAD synthesis from KP clearly results in NAD+ production, significant synthetic capacity is also possible via nicotinic acid or nicotinamide and the salvage pathway (Fig. 3).

With growing interest in IDO1 inhibition as a therapeutic target coupled with the fact that maintaining NAD+ levels is an essential part of organ health characterisation of NAD+ metabolism in other cells of the CNS such as neurons and microglia are needed. Further research is also required to confirm whether NAD+ levels can be effectively maintained when IDO1 or other KP enzymes are inhibited by concurrent addition of NIC or salvage pathway precursors such NAM.

Acknowledgements
This study was supported by a research grant from the Australasian Research Institute.

Disclosures
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