

Novel synthetic route to the C-nucleoside, 2-deoxy benzamide riboside

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ABSTRACT

2-Deoxy-C-nucleosides are a subcategory of C-nucleosides that has not been explored extensively, largely because the synthesis is less facile. Flexible synthetic procedures giving access to 2-deoxy-C-nucleosides are therefore of interest. To exemplify the versatility and highlight the limitations of a synthetic route recently developed to that effect, the first synthesis of 2-deoxy benzamide riboside is reported. Biological properties of this novel C-nucleoside are also discussed.

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C-Nucleosides are an important class of hydrolytically and enzymatically stable compounds and have been found to be important probes in the pursuit of novel antiviral and antiproliferative drugs.¹ There are several known ribose-based C-nucleosides that are potent antiproliferatives. Famous examples are formycin, tiazofurin and benzamide riboside (Fig. 1).² Carboxamide derivatives of C-nucleosides, such as, benzamide riboside and tiazofurin, are based on nicotinamide riboside (NR), one of three NAD precursor vitamins.³ As prodrugs, these compounds must gain entry to target cells, be phosphorylated by NR kinase 1 or NR kinase 2^{3a} and then be adenylated by NMN adenylyltransferase (NMNAT)⁴ to a derivative of NAD.⁵ Benzamide riboside (BR) has been shown, as its benzamide adenine dinucleotide (BAD) metabolite, to exhibit significant cytotoxicity against a variety of human tumour cell lines.⁶ The mechanism of cytotoxicity of BAD and TAD depends on inhibition of inosine monophosphate dehydrogenase (IMPDH),⁵ thus resulting in the depletion of the intracellular GTP pool.⁷

Additionally, the discovery that select nucleosides with L-configuration have biological activity has led to a breakthrough in antiviral chemotherapy. The antiviral mechanism has shown that such L-nucleosides were phosphorylated by cellular kinases, and selectively interacted with viral polymerases, while remaining relatively innocuous to normal cellular processes.⁸ These results highlight the need for new synthetic strategies generating both enantiomers of nucleoside analogues equally well, while retaining

the possibility for synthetic desymmetrisation, if either or both analogues show interesting antiproliferative properties.⁸

The syntheses of C-nucleosides are, in general, quite difficult, low yielding and do not allow access to large series of derivatives for SAR studies, in particular with regards to the aromatic moiety.⁹ Strategies have remained linear, resulting in a limited number of options with regards to the possible structural and functional combinations that can potentially enhance pharmacodynamic/pharmacokinetic properties of the novel compounds.

Our goal is to synthesise analogues of BR to probe each step in the process of NR to NAD biotransformation: stability of the glycosidic linkage to hydrolysis/phosphorolysis^{3c} import, phosphorylation by NR kinases,^{3a,d} adenylation,⁴ and inhibition of subsequent target enzymes. Definition of NR analogue specificity may allow probes to be developed for cancer or for dissection of the metabolic effects of NR.^{3f,h} Analogues with variation on the aromatic moiety, such as tiazofurin and its selenium equivalent, selenofurin have yielded precursors to potent IMPDH inhibitors. Similarly, the 2' position of tiazofurin was shown to be tolerant to manipulation, with various compounds retaining cytotoxicity, for example, the 2'-amido-tiazofurin analogue.¹⁰ Similarly the epoxide functionality across the 2'-3' positions displayed the most pronounced increase in cytotoxic activity against certain lymphoma cell lines, being almost 30 times more potent than the parent compound tiazofurin.¹¹ While BR and tiazofurin share similar biological profiles and potencies, similar analogues have not been reported for BR. We therefore proposed that BR-analogues incorporating different carbohydrate substitution could also offer novel biological opportunities. As a

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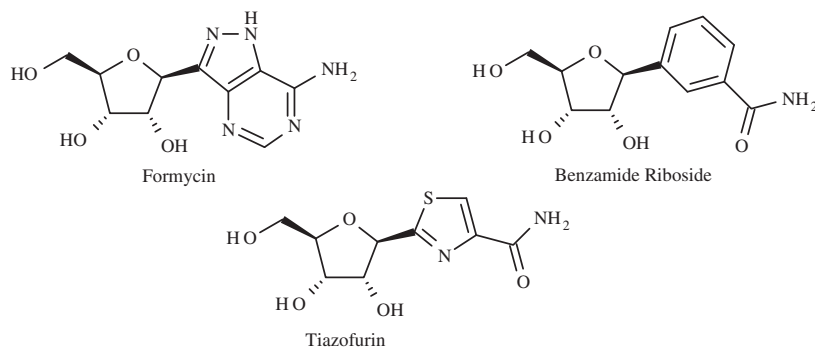


Figure 1. Examples of potent antiproliferatives. Formycin benzamide riboside and tiazofurin.

result, 2- and 4-deoxypyranoside analogues of BR have been generated.¹² We also proposed to test whether biological properties might be retained in the 2-deoxyriboside series of C-nucleosides.

To access this type of novel 2-deoxy C-nucleoside furanosides, we have devised a synthetic methodology involving key *exo*-alkene intermediates.¹³ Herein, we report the synthesis of the *meta*-vinyl substituted phenyl *exo*-alkene intermediate which provided the necessary chemical flexibility to access the *meta*-carboxamide substituent essential to BR analogues. The *meta*-vinyl group was chosen as a direct result of the electron withdrawing nature of the preferred amide or nitrile aromatic functionality, which had proven incompatible to the TMSNTf₂ mediated cyclization conditions necessary to access the *exo*-alkene furanoside intermediates.¹³

The procedure began with the synthesis of acetal **3** from the reaction between easily accessible intermediates allylsilane **1** and chloroacetal **2** in 59% yield.¹³ Subsequently, using optimized conditions, acetal **3** was intramolecularly cyclised to provide a 1.3:1 (*cis*:*trans*) mixture of furanoside **4** using 2 equiv of TMSNTf₂ in 54% combined yield.¹⁴ With this intermediate mixture in hand, the double oxidation of the unsaturated functionalities was performed using optimized conditions for ozonolysis,¹³ to achieve the *cis* and *trans* isomers of keto-aldehyde product **5** in 87% combined yields. This diastereomeric mixture of **5** was easily separated on silica with the desired *cis*-isomer isolated in 50% yield. Unfortunately, attempts failed to convert the *trans*-isomer to the *cis*-isomer by epimerisation. Density Functional Theory (DFT) calculations showed that the *trans*-isomer was 16 kcal/mol lower in energy relative to the *cis*-isomer,¹⁵ and this was therefore the preferred configuration for the compound, with too high an energy difference to easily achieve efficient epimerisation from *trans*- to *cis*-configuration without decomposition.

Difficulties in accessing the essential amide functionality directly from aldehyde **5** led us to proceed through the carboxylic acid derivative **6**. An oxidising system of sodium chlorite, hydrogen peroxide and sodium dihydrogen phosphate was successfully implemented to obtain **6** in high yields.¹⁸ At this point some conflicting issues arose: the amide formation was inevitably going to involve an amine, which had the potential to epimerise the stereocentre adjacent to the ketone, as observed in previous studies.¹³ The most straightforward route from a carboxylic acid to an amide is via an acid chloride and subsequent reaction with the required amine. The latter was successfully carried out using aqueous ammonia. Gratifyingly, this reaction did not cause noticeable epimerisation of the vulnerable stereocentre and quantitatively provided compound **7**.¹⁷

Deprotection of the TBDPS moiety could not be carried out using standard protocols, either by using a fluoride source or other basic conditions without causing the epimerisation.¹⁶ Indeed, attempts to use TBAF gave a mixture of products in low yields, with complete scrambling of the stereocentre. However, when using *p*-toluene

sulfonic acid, the silyl protecting group was cleaved at 60 °C in DMF, albeit in 56% yield. The final reduction using sodium triacetoxymethylborohydride proceeded smoothly, giving the final compound, 2-deoxy-benzamide riboside **cis-8**, in 95% yields, with the relative stereochemistry confirmed by NOE experiments, (Scheme 1).^{13,19}

To the best of our knowledge, this compound has not yet been synthesised, and thus never been tested for targeted biological and enzymatic activity. It was anticipated that the removal of the 2'-hydroxyl group on the ribose would affect the ability of the compound to hydrogen bond with polar residues of the binding pockets in NRK,^{3d,g} NMNAT²⁰ and IMPDH.²¹ Therefore to retain activity, it is vital that the resulting NAD analogue structure not only fit into the IMPDH pocket to achieve inhibition, but that its metabolic precursors be recognised and metabolised effectively by the enzymes that convert it into an NAD analogue.^{7,22}

The target molecule **cis-8** was therefore tested for activity against a number of cancerous cell lines. The cell lines were: RKO—Human colon carcinoma, HT29 - Human colorectal adenocarcinoma, HCT116—Human colorectal carcinoma, H460 - Human large lung cell carcinoma, MCF7—Human breast adenocarcinoma (pleural effusion metastatic site), T47D Human breast ductal carcinoma, PC3—Human prostate adenocarcinoma (bone metastatic site) and DU145—Human prostate carcinoma (brain metastatic site). Some of these cell lines have been shown to be sensitive to tiazofurin and benzamide riboside at concentrations that range from 50 μM (benzamide riboside in MCF7) to 500 μM (cell line PC-3).²³ Despite incubation at concentrations up to 1 mM, no cell line displayed a significant level of inhibition upon incubation with **cis-8** compared to the blank sample of pure DMSO.

In work preceding the discovery of the NRK-dependent NAD biosynthetic pathway³ and the evidence that tiazofurin was an excellent substrate for NRK,^{3d} tiazofurin and benzamide riboside were proposed to enter cells via nucleoside transporters. While a nicotinamide riboside specific transporter has been identified in yeast, the human homolog still remains undiscovered. To test the hypothesis that **cis-8** fails as a cytotoxic agent because of a deficit in the biotransformation to NAD rather than or in addition to limited transport, we incubated the compound at 0.5 mM with 1 mM ATP and sufficient NR kinase 2 for complete conversion of 0.5 mM NR to NMN in 30 min. Reactions monitored for 2 h by HPLC and LC–MS indicated that the compound is totally resistant to conversion to the corresponding monophosphate. Earlier a 2-deoxy derivative of tiazofurin²⁴ was synthesized and shown to be lacking in cytotoxic activity. Our data strongly suggest that 2-deoxy substitutions are not tolerated by NR kinases,^{3a,d} thereby severely compromising their pharmacological activities.

The present work provides the direction which the chemistry of C-nicotinamide riboside modified on the sugar moiety must take and that is attempting to by-pass the NRK step in accessing in situ NAD analogues, putative inhibitors of IMPDH and other

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