COMMUNICATION

Fluorometabolite biosynthesis: Isotopically labelled glycerol incorporations into the antibiotic nucleocidin in *Streptomyces calvus*.

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5 *Receipt/Acceptance Data* [DO NOT ALTER/DELETE THIS TEXT] *Publication data* [DO NOT ALTER/DELETE THIS TEXT] DOI: 10.1039/b000000x [DO NOT ALTER/DELETE THIS TEXT]

Deuterium and carbon-13 labelled glycerols have been fed to *Streptomyces calvus* fermentations and isotope incorporation into

- 10 the fluorine containing antibiotic nucleocidin have been evaluated by ¹⁹F-NMR. A single deuterium atom was incorporated from $[{}^{2}H_{5}]$ - and (R)- $[{}^{2}H_{2}]$ - glycerol into C-5' of the antibiotic, suggesting that an oxidation occurs at this carbon 30 after ribose ring assembly from glycerol (pentose phosphate
- 15 pathway), during nucleocidin biosynthesis.





Scheme 1. Fluorinase reaction of AdoMet 3 to give FDA 4 and then the relationship between FDA and FAc 1 and 4-FT 2. Incorporation of (*R*)-[²H₂]-20 glycerol determined that both fluoromethyl hydrogens at C5' of FDA 4 are retained in the fluorometabolites.⁹ Nucleocidin 7 is a metabolite of *S. calvus*.

Fluorine containing natural products are exceedingly rare in nature. The toxin fluoroacetate (FAc) **1** is the most widely distributed natural **65** monitored, even at low production levels, directly from extracts by

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product of this class where it has been identified in many plants² and
25 a few bacteria.^{3,4} The mode of fluoroacetate biosynthesis in plants has not been established, however its biosynthesis in bacteria is known.⁵ Streptomyces cattleya is a bacterium that co-produces FAc 1 and 4-fluorothreonine (4-FT) 2.³ Studies carried out with *S. cattleya* determined that the fluorination enzyme (fluorinase) catalyses a
30 nucleophilic substitution reaction between fluoride ion and *S*-

- adenosyl-L-methionine (Ado-Met) **3** to generate 5° -fluorodeoxyadenosine (FDA) **4**.⁶ This is the only biosynthetic fluorination enzyme known. It has subsequently been identified by gene mining in several other bacteria.⁷ FDA **4** is then processed to
- 35 FAc 1 and 4-FT 2. A combination of isotopic labelling studies, including isotopically labelled glycerol 5, and an understanding of the individual steps in the biosynthesis of 1 and 2 have determined that the C5' fluoromethyl and C4' carbon of FDA 4 are converted to FAc 1 and C3 and C4 of 4-FT 2, the circled regions in Scheme 1.⁸ It has

40 also been established that both hydrogens of the fluoromethyl group of FDA 4 are retained in the conversion to the fluoromethyl groups of 1 and $2.^9$

The antibiotic nucleocidin 7 produced by the actinomycete soil bacterium, *Streptomyces calvus* is another example of the rare

- 45 fluorometabolites.¹⁰ Its biosynthesis is unknown; however, the presence of the fluorine atom at the C4' position of the ribose ring¹¹ in 7 is inconsistent with the involvement of the fluorinase found in *S. cattleya*. Full genome sequencing of the nucleocidin producer *S. calvus* ATCC 13382 has not identified a fluorinase gene,¹² thus *S.*
- **50** *calvus* may employ a very different process for enzymatic C-F bond formation relative to bacterial FAc 1 production. Ever since its isolation, from *S. calvus* in 1956,¹⁰ nucleocidin 7 production has proved problematic, with consistently low titres and
- unpredictable production. Publically deposited strains appear to have 55 lost the ability to produce nucleocidin.¹²⁻¹⁴ Some progress has been made to understand the inconsistent production, and it has been observed that a mutation of the *bldA* gene, encoding a Leu-tRNA, is apparent in the publically available strains. Correction of this mutation was recently shown¹² to re-establish production in *S. calvus*.
- **60** In another study¹³ mutation of the *rpoB* gene (rifamycin resistance) led to an apparent increase in nucleocidin production. In this study we use an in-house strain of *S. calvus* T-3018 held by Pfizer. This strain does not have a *bldA* mutation, although production is still fickle with very low titres. Nucleocidin production can however be
- ¹⁹F{¹H}-NMR. As a first step towards elucidating the biosynthetic pathway to fluorometabolite 7 we now report the incorporation of various isotopically labelled glycerols **5** a-d.
- For this study we have supplemented fermentation cultures of *S*. **70** *calvus* with glycerols $[2-^{13}C]$ **5a**, $[^{2}H_{5}]$ **5b**, $(R)-[^{2}H_{1}]$ **5c** and $(S)-[^{2}H_{1}]$ **5d**. Despite the low titres, we envisaged that the incorporation of deuterium or carbon-13 isotopes close to the fluorine

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atom of the antibiotic could be reported visually by ${}^{19}F{}^{1}H$ -NMR due to heavy atom induced chemical shifts of the fluorine resonance, 110 value is intermediate between these extremes and consistent with 75 and/or ¹³C-¹⁹F-spin-spin coupling in the case of carbon-13

incorporation. A first experiment explored pulse feeding of glycerol $[2-^{13}C]$ - 5a to a final concentration of 8mM, in a shake flask fermentation of S.

- 80 described.¹³ Nucleocidin is excreted from the bacterial cells and thus the cells were spun down in a centrifuge and the supernatant was extracted into butanol. The extract was concentrated and was then analysed by ${}^{19}F{}^{1}H$ -NMR. The resultant spectrum is shown
- 85 accompanying satellites due to ${}^{1}J_{CF}$ (232Hz) coupling, consistent with a population (~25%) of nucleocidin 7 molecules enriched with carbon-13 at C4'. The satellite signals are off-centre relative to the natural abundance ¹⁹F-NMR signal due to a heavy atom α shift.⁸ This incorporation is entirely consistent with pentose125
- 90 phosphate pathway involvement in ribose biosynthesis where carbons C3-C5 of D-ribose derive from the exogenously added glycerol, feeding into the C₃ metabolite pool via sn-glycerol-3phosphate 6.15



95 Figure 1. ¹⁹F{¹H}NMR of nucleocidin 7 (-119.8ppm) after glycerol 5 feeding experiments to S. calvus fermentations. An unknown metabolite (-118.1ppm) is also present in some spectra. (a) [2-¹³C]-Glycerol 5a; (b) $[^{2}H_{5}]$ -glycerol **5b**; (c) (*R*)- $[^{2}H_{2}]$ -glycerol **5c**; (d) (*S*)- $[^{2}H_{2}]$ -glycerol **5d**. A pictorial summary of the isotopic labelling outcome is given in each case.

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Addition of perdeuterated glycerol 5b (final conc 8 mM) to a S. calvus fermentation led to the incorporation of deuterium (~5%) into nucleocidin 7 as determined by an upfield shift in the resultant {¹H}¹⁹F-NMR spectrum as shown in Figure 1b. The

105 magnitude of this upfield shift at 0.22ppm is consistent with a single vicinal deuterium incorporated β - to the fluorine atom.155 There is a F-C-C-D angular dependence on the magnitude of the shift ranging from 0.15ppm (0°) to 0.35ppm (180°) as determined

in classic studies by Lambert and Greifenstein.¹⁶ The observed a single vicinal C-D (γ -shift) bond approximately at 90° to the C-F bond. This experiment does not however distinguish the location of the incorporation between C3' or C5' of the ribose ring of 7. To achieve such a distinction, comparative calvus. Cultures were worked up after 20 days as previously 115 incorporation experiments were explored with the enantiomeric glycerols (R)-[²H₂]- 5c and (S)-[²H₂]- 5d. It is already established that the pro-R arm of glycerol delivers C5' of ribose through the pentose phosphate pathway, and that the pro-S arm delivers C3' of ribose. Glycerols 5c and 5d were prepared synthetically with in Figure 1a. It is clear that the fluorine signal for nucleocidin has 120 enantiomeric purities > 95% e.e., as previously described.⁹ The resultant ¹⁹F{¹H}-NMR spectra from each of these glycerol feeding experiments are shown in Figure 1c and Figure 1d. It is clear that only glycerol (R)-[²H₂]- **5c** results in the incorporation $(\sim 8\%)$ of deuterium into nucleocidin 7, and again only a single deuterium atom survives from the double labelled precursor. It follows from the known stereochemistry of glycerol processing through the pentose phosphate pathway, that this deuterium is located at C5^{1,15,17}

In this experiment a second fluorometabolite was observed in the 130 ¹⁹F{¹H}-NMR spectrum with a signal at -118.1ppm. This metabolite is detected periodically in nucleocidin extracts although its occurrence and level is unpredictable in our experience. Such a fluoro metabolite was also observed in the recent study where nucleocidin production was elicited by 135 reversing the *bldA* mutation.¹² Its structure is unknown although given the similar level of deuterium incorporation from glycerol 5c, it appears to be metabolically related to nucleocidin either as a biosynthetic precursor, or as a metabolite.

For isomer (S)- $[{}^{2}H_{2}]$ - 5d the absence of any deuterium at C3' in 7 140 is consistent with sn-glycerol-3-phosphate 6 being processed to ribulose-5-phosphate via xylulose-5-phosphate, and the action of D-ribulose-5-phosphate 3-epimerase, with exchange of the surviving deuterium with bulk solvent.¹⁸ The predicted labelling pattern of ribose-5-phosphate from glycerol is illuystrated in Scheme 145²



Scheme 2. The pentose phosphate pathway showing the fate of glycerol through to ribose-5-phosphate. Glycerol 5e has a virtual labelling pattern 150 which is a composite of the experimentally administered glycerols 5a-d, and also highlights the stereochemical fate of the (pro-R) hydroxymethyl carbon (■). This pathway predicts incorporation of a C₃ glycerol unit (bold C-C bonds) into C-3-C-5 of ribose-5-phosphate, with the retention of two deuteriums at C-5 and the loss of deuteriums at C-3 and C-4.

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The outcome where both **5b** and **5c** glycerols contribute only one deuterium to C5' of nucleocidin can be contrasted with the biosynthesis of FAc, **1** and 4-FT, **2** in *S. cattleya* discussed above**225**

and shown in Scheme 1. In those cases two deuterium atoms were incorporated into the fluorometabolites, and by implication into the C5' ribose carbon of FDA 4.^{9a} For nucleocidin only one deuterium is incorporated.

In summary, this study provides the first biosynthetic data on 165 nucleocidin 7 assembly from isotope labelling studies. It is shown that glycerol is incorporated into the ribose ring moiety of nucleocidin, consistent with expectation *via* the pentose

- phosphate pathway. However, the presence of a single deuterium only at the C5' position of nucleocidin 7 after feeding
 170 experiments with glycerols 5b and 5c suggests that a hydrogen is lost from this carbon after ribose ring assembly but prior to, or concominant with, fluorine introduction. This observation places
- a constraint on working hypotheses addressing nucleocidin biosynthesis.
 - Acknowledgement: We are grateful to Dr Alessandra Eustáquio for helping to establish this research project.

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