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The Biosynthetic Gene Cluster of Pyrazomycin—A C-Nucleoside Antibiotic with a Rare Pyrazole Moiety — Source link

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1	The biosynthetic gene cluster of the C-nucleoside antibiotic		
2	pyrazomycin with a rare pyrazole moiety		
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15			
16	• Abstract		
17	Pyrazomycin is a rare C-nucleoside antibiotic with a naturally occurring pyrazole ring, whose		
18	biosynthetic origin has remained obscure for decades. In this study, we report the identification of		
19	the gene cluster responsible for pyrazomycin biosynthesis in Streptomyces candidus NRRL 3601,		
20	revealing that StrR-family regulator PyrR is the cluster-situated transcriptional activator governin		
21	pyrazomycin biosynthesis. Furthermore, our results from in vivo reconstitution and stable-isotop		
22	feeding experiments support that PyrN is a new nitrogen-nitrogen bond forming enzyme linking t		
23	ϵ -NH ₂ nitrogen of L-N ⁶ -OH-lysine and α -NH ₂ nitrogen of L-glutamate. This study lays t		
24	foundation for further genetic and biochemical characterization of pyrazomycin pathway enzyme		
25	constructing the characteristic pyrazole ring.		
26			
27	• Main text		
28	Nitrogen-nitrogen (N-N) containing natural products are group of specialized metabolites with		
29	diverse structures and a variety of biological activities. ^[1] These molecules have been isolated from		

30 different sources, including bacteria, fungi and plants. Despite extensive studies on the genetic and 31 biochemical basis of natural product biosynthesis over the past three decades, the biochemical routes 32 leading to enzymatic N-N bond formation are only starting to be revealed.^[2–14] We recently reported 33 a heme-dependent piperazate synthase responsible for N-N cyclization in piperazic acid, a building 34 block for many nonribosomal peptide (NRP) or NRP-polyketide hybrid molecules (Fig. 1).^[2] The biosynthetic route to piperazate begins with the N-hydroxylation of L-ornithine, giving a 35 hydroxylamine precursor, which is reminiscent of valanimycin biosynthesis, which also begins with 36 37 a hydroxylamine precursor.^[9] By contrast, N-N bond formation in other N-N bond containing 38 natural products including compounds cremeomycin, fosfazinomycin and kinamycin starts instead with the generation of nitrous acid, [5,6,8,10] and the biosynthesis of the N-nitroso streptozocin 39 40 originates with oxidation of the guanidine of L-arginine.^[3,4] However, whether other pathways to N-41 N bond containing molecules might begin from hydroxylamine precursors was unknown. Recent studies into hydrazone unit formation in the dipeptide s56-p1 resulted in the discovery of a pathway 42 that links the hydroxylamine L-N⁶-OH-Lys to hydrazino acetic acid formation.^[7] In this pathway, 43 44 Spb40, a fusion protein with cupin and aminoacyl-tRNA synthetase (aaRS)-like domains, appears 45 from expression studies in E. coli to couple glycine and L-N⁶-OH-Lys by forming a N-N bond between the α -NH₂ of glycine and the ϵ -N atom of L-N⁶-OH-Lys, the latter of which is in turn 46 47 generated by Spb38-catalyzed N⁶-hydroxylation of L-Lys. An Spb40-like enzyme was also reported in the gene cluster for triacsins, but the details of this reaction, and whether this type of enzymology 48 49 might play a role in pathways to other N-N bond containing structures, remains obscure.^[14]

50 Despite progress toward understanding the enzymology of N-N bond formation, biosynthetic 51 routes to aromatic structures containing N-N bonds is largely unexplored. Construction of an N-N 52 bond embedded in an aromatic ring might require a very different biosynthetic logic from formation 53 of a hydrazine or other N-N linkage. Among N-N bond-containing structures, a key example is 54 pyrazomycin (PZN, also known as pyrazofurin), a C-nucleoside with a rare pyrazole moiety.^[15] As 55 a nucleoside analog, PZN is a potent inhibitor of orotidine 5'-monophosphate decarboxylase and 56 possesses broad-spectrum antiviral and antitumor activities.^[16] Early biosynthetic studies on PZN 57 and its structural analog formycin used isotope-labeled precursors to establish that the C-3 to C-6 58 of the pyrazole ring in PZN derive from C-4 to C-1 of glutamate or α -ketoglutarate. Similarly, the 59 C-9, C-4, C-5, C-6 of the pyrazolopyrimidine ring in formycin derive from these precursors (Fig.

60 1).^[17] However, the direct precursors of the two pyrazole nitrogen atoms and the biosynthetic
61 machinery driving N-N bond formation have remained obscure.

62 To answer these long-standing questions, we sequenced the genome of the PZN producing 63 strain Streptomyces candidus NRRL 3601, and identified a ~28 kb gene cluster as a candidate for 64 the PZN biosynthetic gene cluster, which we name here as the pyr cluster (Fig. 2a, Table S1). This 65 gene cluster carries a four-gene sub-cluster (pyrOPQS) that has been linked to the biosynthesis of coformycin, whose production seems to be associated with many nucleoside family natural 66 67 products.^[17–19] Moreover, the *pvr* cluster also contains *pvrE*, which shows sequence homology to genes encoding β -ribofuranosylaminobenzene 5'-phosphate (β -RFAP) synthase. In the biosynthesis 68 of methanopterin, β-RFAP synthase catalyzes the condensation of p-aminobenzoic acid with 69 70 phosphoribosylpyrophosphate through a C-glycosidic linkage.^[20] A similar reaction for C-71 glycosidic bond formation seems to be also required during PZN biosynthesis. Furthermore, another interesting feature about the pyr cluster is the presence of spb38 and spb40 homolog genes pyrM 72 73 and *pvrN*, indicating that this cluster encodes a potential hydrazine-producing pathway analogous 74 to that of the s56-p1 pathway. which is also consistent with the N-N bond containing structure of PZN.^[7] We note that the pvr cluster share many homologous genes with the recently reported 75 formycin biosynthetic gene cluster (for cluster).^[19] Considering the great structural similarity 76 77 between pyrazomycin and formycin, it is not unexpected that they utilize similar biosynthetic genes. Taken together, our *in silico* analysis suggests that the *pyr* cluster might be responsible for PZN 78 79 assembly in S. candidus NRRL 3601.

80 Although strain S. candidus NRRL 3601 was known as a PZN producer, we failed to detect any PZN production with various media we tested, including the one used in early studies (Fig. 2b-81 2d).^[15] Introduction of a vector carrying the whole putative pyr cluster into the common 82 83 Streptomyces host S. albus J1074 also did not produce any detectable PZN. To explore why no PZN 84 is produced, we isolated total RNA from S. candidus NRRL 3601 and performed transcriptional 85 analysis of the pyr cluster by RT-PCR. We found that most of the pyr genes were not actively 86 transcribed under the culture condition we used (Fig. 2d). We thus turned to regulatory engineering 87 of the PZN pathway. Analysis of the pvr genes revealed the presence of a single putative 88 transcriptional regulator *pyrR*, which shows sequence homology to StrR-family regulators. As the 89 prototype of this family, the streptomycin biosynthesis regulator StrR is the transcriptional activator

controlling the expression of streptomycin biosynthetic genes by interacting with multiple promoter 90 regions within the gene cluster.^[21,22] Members of this family also include NovG and Bbr, which are 91 located in the biosynthetic gene clusters of novobiocin and balhimacin, respectively.^[23,24] This 92 93 bioinformatic analysis suggested that *pvrR* likely encodes a pathway-specific activator of the PZN 94 biosynthetic gene cluster. To interrogate whether *pvrR* could activate this biosynthetic pathway, we introduced an additional copy of pyrR into S. candidus NRRL 3601 under the control of a 95 96 constitutive promoter to release the transcriptional control from higher-level regulatory mechanisms. 97 Subsequent metabolic profiling of the resulting strain S. candidus pyrR-OE by LC-MS together with 98 NMR analysis of isolated PZN demonstrated that PZN production was successfully recovered, with 99 a PZN yield of ~10 mg/L (Fig. 2b and 2c, Fig. S1). Furthermore, RT-PCR analysis of selected pyr 100 genes revealed their active transcription in S. candidus pyrR-OE, including the spb40 homolog pyrN, 101 which is located in a putative operon consisting of genes *pyrKLMN* (Fig. 2d). Altogether, these 102 results demonstrated that PyrR is a transcriptional activator governing PZN biosynthesis, and by 103 extension, the pyr cluster is responsible for PZN biosynthesis.

104 We next interrogated the biosynthetic origin of N-1 and N-2 in the pyrazole ring of PZN, which 105 remain obscure despite the previous isotope feeding experiments and our in silico analysis of the 106 pyr cluster. The presence of spb38 and spb40 homologs pyrM and pyrN in the pyr cluster indicate 107 that a N-N bond forming mechanism analogous to that of the s56-p1 pathway might operate in PZN 108 biosynthesis.^[7] We envisaged two possible routes that could afford the hydrazine moiety in the 109 pyrazole ring (Fig. 3a). In one scenario, the Spb40 homolog PyrN could catalyze N-N bond formation between the N-6 nitrogen of L-N6-OH-Lys with another amine-containing substrate to 110 111 generate a molecule as a hydrazine carrier, which would be followed by the transfer of the hydrazine 112 mojety to a glutamate or α -KG derivative by downstream pathway enzymes. In this scenario, PyrN 113 could be functionally equivalent to Spb40 and produce the product Lys-Gly for the subsequent 114 hydrazine transfer. A second scenario involves the direct installation of hydrazine moiety on 115 glutamate or its derivative, through linking the α -NH₂ of glutamate (or its derivative) to N-6 nitrogen of L-N⁶-OH-Lys. The resulting product could then be processed by other *pyr* enzymes to furnish the 116 117 pyrazole ring. To distinguish between these two possible routes, we first separately fed L^{-15} N-Gly 118 or L-15N-Glu (at concentrations of 3 mM) into the culture of S. candidus pyrR-OE and analyzed the isotope incorporation of PZN by LC-MS. We found that the nitrogen from the α -NH₂ of glutamate 119

efficiently incorporated into PZN, with the relative intensity of +1 Da peak increasing from 11% to 46% when compared to unlabeled PZN (**Fig. S2**). Furthermore, the +2 Da peak increased to 16 %. By contrast, only minor incorporation of the nitrogen from ¹⁵N-Gly was detected. Although it is likely that the observed incorporation pattern is due to or partially attributed to scrambling of the labels by primary metabolism, this result favors the second scenario, where the N-1, and by extension, C-3 to C-6 atoms, in the pyrazole moiety might derive from L-Glu as an intact unit (**Fig. 3a**).

127 To gain further insights into the product from the PyrN-catalyzed reaction, we adopted an *in* vivo reconstitution method by introducing pyrN into the E. coli strain expressing nbtG. NbtG, a 128 129 homolog of PyrM and Spb38, is a well-characterized lysine N^6 -hydroxylase and thus could provide 130 endogenous L-OH- N^6 -Lys in vivo. LC-MS analysis of culture supernatant from the E. coli strain 131 carrying both *nbtG* and *pyrN* revealed the presence of a molecule with a MS signal at m/z 292, which 132 is absent from that of E. coli strain expressing either *nbtG* or *pyrN* (Fig. 3b). This MS signal (m/z) 133 292) is consistent with the $[M+H]^+$ ion of a Lys-Glu conjugate product (1) (Fig. 3a). No MS signal 134 (m/z 220) of Lys-Gly was detected, supporting that Gly is not a substrate of PyrN, in contrast to 135 Spb40. To track the origin of compound 1, we fed the strain E. coli/nbtG+pyrN with different combinations of ¹³C/¹⁵N-labeled and unlabeled amino acid precursors, including L-¹³C₆-Lys, L-¹⁵N-136 Glu, L-¹⁵N-Gly (Fig. S3). Compared with cultures supplemented with unlabeled amino acids, the 137 compound 1 produced in the presence of L- $^{13}C_6$ -Lys showed the appearance of a strong +6 Da MS 138 139 signal (m/z 298.1), supporting the incorporation of an intact lysine carbon skeleton. While 1 produced with the addition of ¹⁵*N*-Gly showed a similar isotope pattern to the one fed with unlabeled 140 141 amino acids, addition of ¹⁵N-Glu resulted in significant enrichment of +1 Da signal, with an increase 142 in the relative intensity from 12 % to 45 %. Fmoc-chloride treatment of the E. coli culture supernatants supplemented with ¹³C₆-Lys, followed by LC-HR-MS analysis, resulted in the 143 detection of a strain-specific metabolite from E. coli/nbtG+pyrN with MS signals at m/z 514.2164 144 145 and 520.2364, which are consistent with the $[M+H]^+$ ion of Fmoc-Lys-Glu and Fmoc- $({}^{13}C_6)$ Lys-Glu 146 conjugate, respectively (Fig. 4a and Fig. S4). Our attempt to isolate this molecule for NMR analysis 147 was hampered by its low yield and instability. However, the fragmentation pattern of 1 from LC-148 HR-MS/MS analysis after Fmoc-Cl derivatization, together with the sequence homoloy between 149 PyrN and Spd40, supports that 1 most likely arises from linking the α -NH₂ of glutamate with the ϵ - 150 NH₂ of lysine (Fig. 4b and 4c).^[7] Altogether, the above results suggested that PyrM and PyrN
151 together mediate the conjugation L-Glu and L-Lys though a N-N linkage.

152 As in Spb40, the residues potentially involved in ATP-binding and metal-chelating are also conserved in the aaRS-like and cupin domains of PyrN, respectively (Fig. S5 and S6). Consistent 153 154 with this in silico analysis, inductively coupled plasma mass spectrometry (ICP-MS) analysis of the 155 purified His-tagged PyrN demonstrated that it contains two equivalents of zinc, supporting the 156 presence of a single, conserved zinc coordination site in each domain (Fig. S7). Even though we 157 were able to obtain soluble recombinant PyrN, our preliminary screening of reaction conditions for 158 the in vitro assay of PyrN did not produce any detectable product so far, which could be attributed 159 to yet-unknown factors necessary for this reaction. Nonetheless, we expect that the PyrN reaction 160 mechanism is likely to be similar to that has been suggested for Spb40, where there is a similar lack 161 of *in vitro* work reported in the literature.^[7] Based on the results from our *in vivo* reconstitution in E. coli and previous isotope tracking experiments on Spb40 (Fig. S8), the C-terminal aaRS-like 162 domain of PyrN is likely responsible for glutamate carboxylate activation, to afford glutamyl-163 164 adenylate or further to give glutamyl-tRNA, which could then undergo ligation to the hydroxyl 165 group of L-OH-N⁶-Lys to form an ester intermediate, and followed by rearrangement to generate 1, 166 which is likely mediated by the N-terminal cupin domain. Formation of an ester intermediate by the conjugation of an amino acid to a hydroxylamine, using aminoacyl-tRNA as a carrier, is not 167 unprecedented. In the biosynthesis of azoxy-containing valanimycin, the servl-tRNA synthetase 168 169 VlmL produces L-seryl-tRNA, followed by VlmA-mediated seryl transfer to isobutylhydroxylamine 170 to form O-seryl-isobutylhydroxylamine, which then undergoes subsequent transformation(s) for N-N bond formation.^[9] Following the formation of 1, the next step in the PZN pathway is likely 171 172 catalyzed by PyrL, whose encoding gene pyrL appears to be located in the same operon with pyrM 173 and pyrN, indicating their functional relevance (Fig. 2a). PyrL displays sequence homology to 174 saccharopine dehydrogenase. In lysine metabolism, saccharopine dehydrogenase mediates 175 reversible conversion of saccharopine to L-2-aminoadipate-6-semialdehyde (AASA) and L-Glu (Fig. **S9**).^[25] PyrL might thus catalyze an analogous reaction, converting 1 to AASA and 2-176 177 hydrazinoglutaric acid (2), the latter of which could then be processed by downstream pathway enzymes for pyrazole ring assembly (Fig. 4d). 178

179

In conclusion, we have identified the gene cluster responsible for pyrazomycin biosynthesis in

180	<i>S</i> . <i>c</i>	candidus NRRL 3601, and demonstrated that PyrR is a new member of the StrR family	
181	transcriptional activator controlling PZN biosynthesis. Furthermore, our current data strongly		
182	supports that PyrN is a new N-N bond forming enzyme, and the formation of the lysine-glutamate		
183	conjugate through a N-N linkage could be the starting point for the PZN biosynthetic pathway. This		
184	study paves the way for further biochemical characterization of PyrN and other PZN pathway		
185	enz	ymes involved in the assembly of the unique pyrazole ring.	
186			
187	Exp	perimental Section	
188	Full experimental details are available in the Supporting Information		
189			
190	Acl	knowledgements	
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195			
196	Keywords: biosynthesis • N-N bond • pyrazole • C-nucleoside • pathway activator		
197			
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236 Figure 1. Examples of natural products containing a nitrogen-nitrogen linkage.

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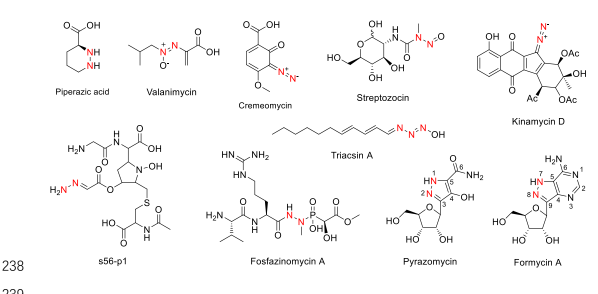
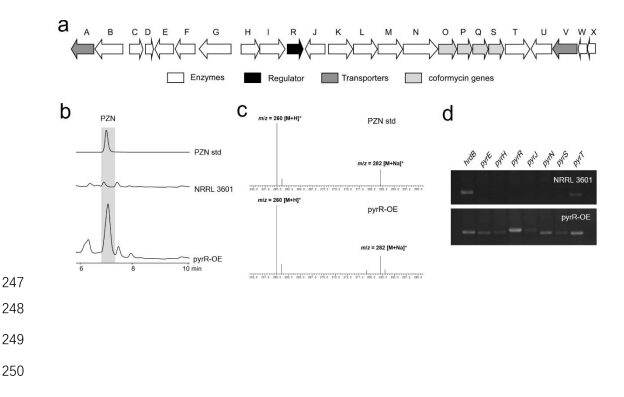
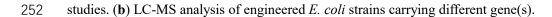
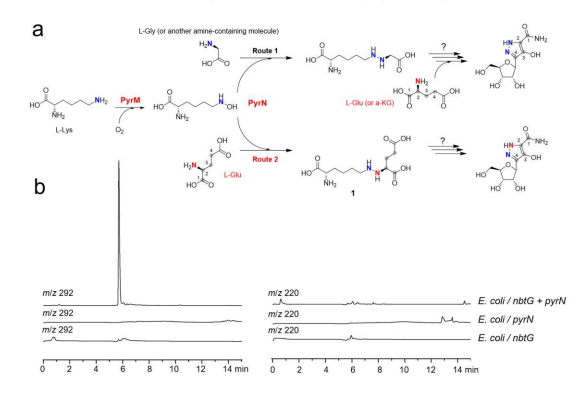


Figure 2. Identification of the gene cluster and cluster-situated activator for pyrazomycin biosynthesis. (a) The biosynthetic gene cluster of pyrazomycin. (b) HPLC analysis of the culture supernatants from strain *S. candidus* NRRL 3601 and its pyrR-overexpression mutant pyrR-OE. (c) MS analysis of the pyrazomycin standard compound and the new product produced by strain pyrR-OE. (d) Transcriptional analysis of selected genes from *pyr* gene cluster in strain NRRL 3601 and pyr-OE by RT-PCR.



251 Figure 3. (a) Possible biosynthetic routes to the pyrazole ring in pyrazomycin based on previous





- Figure 4. The *in vivo* reconstitution of the PyrN-catalyzed reaction in *E. coli* and proposed
- 256 biosynthetic origin of the pyrazole ring in pyrazomycin. (a) HPLC analysis of culture supernatants
- 257 from the *E. coli* strains containing different gene combinations after Fmoc-Cl derivatization. Note:
- Lys458 is a residue potentially involved in ATP-binding in PyrN (see below in the main text). (b)
- and (c) LC-HR-MS/MS analysis of the strain-specific metabolites from E. coli / nbtG + pyrN
- supplemented with $L^{-13}C_6$ -Lys under negative and positive modes, respectively. Note: the red
- traces are from the MS/MS analysis of unlabeled molecules, and the black traces are from ${}^{13}C_{6}$ -
- 262 labeld molecules. (d) Proposed biosynthetic origin of the pyrazole moiety in pyrazomycin based
- 263 on the results from this study and previous stable-isotope precursor feeding experiments.

