1 **Title:**

2 Minimal lactazole scaffold for *in vitro* production of pseudo-natural thiopeptides

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26 Abstract

27 Lactazole A is a cryptic thiopeptide from Streptomyces lactacystinaeus, encoded by a compact 9.8 kb 28 biosynthetic gene cluster. Here, we established a platform for *in vitro* biosynthesis of lactazole A, 29 referred to as the FIT-Laz system, via a combination of the flexible in vitro translation (FIT) system 30 with recombinantly produced lactazole biosynthetic enzymes. Systematic dissection of lactazole 31 biosynthesis revealed remarkable substrate tolerance of the biosynthetic enzymes, and led to the 32 development of the "minimal lactazole scaffold", a construct requiring only 6 post-translational 33 modifications for macrocyclization. Efficient assembly of such minimal thiopeptides with FIT-Laz 34 enabled access to diverse lactazole analogs with 10 consecutive mutations, 14- to 62-membered 35 macrocycles, and up to 18 amino acid-long tail regions. Moreover, utilizing genetic code 36 reprogramming, we demonstrated synthesis of pseudo-natural lactazoles containing 4 non-37 proteinogenic amino acids. This work opens possibilities in exploring novel sequence space of pseudo-38 natural thiopeptides.

40 Introduction

41 Thiopeptides are natural products defined by a six-membered nitrogenous heterocycle, usually 42 pyridine, grafted within the backbone of a peptidic macrocycle.¹ Multiple azole rings, dehydroamino 43 acids, and other optional nonproteinogenic elements further contribute to the resulting structural 44 complexity characteristic of thiopeptides. More than a hundred thiopeptides isolated to date are defined by strong antibiotic activity against Gram-positive bacteria, including methicillin resistant 45 Staphyrococcus aureus (MRSA).¹⁻³ For instance, thiostrepton has been used as a topical antibiotic in 46 47 veterinary medicine, and LFF571, a synthetic derivative of naturally occurring GE2270A, underwent clinical trials as a treatment against Clostridium difficile infections.⁴ 48

A decade ago, thiopeptides were shown to be of ribosomal origin.^{5–8} During biosynthesis, a structural 49 50 gene encoding a thiopeptide precursor is transcribed and translated, and the resulting peptide undergoes 51 post-translational modifications (PTMs) introduced by cognate enzymes colocalized with the structural 52 gene in a biosynthetic gene cluster (BGC). Commonly, these enzymes utilize the N-terminal leader 53 peptide (LP) region of the precursor as a recognition sequence, and act on the core peptide (CP) to 54 introduce PTMs such as azole and dehydroalanine (Dha). Eventually, a pyridine synthase catalyzes formation of a six-membered heterocycle in the CP and eliminates the LP, yielding a macrocyclic 55 56 thiopeptide. Thus, thiopeptides represent a group of ribosomally synthesized and post-translationally 57 modified peptide (RiPP) natural products.9

RiPP biosynthetic logic is highly conducive to bioengineering.^{10,11} Simple nucleotide substitutions in the structural gene yield novel compounds, provided that these mutations are tolerated by the biosynthetic machinery. For BGCs encoding promiscuous enzymes, e.g. lanthipeptides and cyanobactins, this strategy can be applied to construct combinatorial libraries of natural product analogs. Recent studies demonstrated that such libraries can be screened to improve or completely reprogram antibacterial activities of the underlying RiPPs.^{12–19}

In contrast, thiopeptide bioengineering proved to be significantly more challenging. Single-point 64 mutagenesis studies²⁰⁻²⁴ and a few complementary reports (e.g., BGC minimization,²⁵ and an 65 incorporation of a single non-proteinogenic amino acid (**npAA**) suitable for bioconjugation)²⁶ represent 66 67 the bulk of the work on this topic. The challenges in thiopeptide bioengineering can be attributed to a 68 highly cooperative yet only partially understood biosynthesis process. For many thiopeptides, the roles of individual biosynthetic enzymes are only beginning to be elucidated.²⁷⁻³¹ Chemoenzymatic and 69 semisynthetic approaches^{32–38} may circumvent the limitations imposed by biosynthetic machinery, but 70 71 due to the structural complexity of thiopeptides, these strategies present a number of challenges of their 72 own.

We previously reported isolation and characterization of lactazole A, a cryptic thiopeptide from *Streptomyces lactacystinaeus* (**Fig. 1a**).³⁹ It is biosynthesized from a compact 9.8 kb *laz* BGC encoding just five enzymes essential for the macrocycle formation (**Fig. 1b**). Lactazole A has a low Cys/Ser/Thr content, a 32-membered macrocycle, and bears an unmodified amino acid in position 2 (Trp2), all of which are unusual features among thiopeptides (**Fig. 1c**).⁴⁰ Moreover, lactazole A shows no antibacterial

activity, and its primary biological function remains unknown. Recent bioinformatic studies indicated

- that the lactazole-like thiopeptides, characterized by an unmodified amino acid in position 2, comprise
- 80 close to half of all predicted thiopeptides (251 out of 508 annotated BGCs), and yet the prototypical *laz*
- 81 BGC remains the only characterized member of this family to date.⁴¹ Overall, lactazole-like thiopeptides
- 82 remain a rather enigmatic family of natural products, as close to nothing is known about their function,
- 83 structural diversity, and biosynthesis.
- 84 Intrigued by the uniqueness of *laz* BGC, we set out to reconstitute *in vitro* biosynthesis of lactazole A. 85 We sought to establish rapid and reliable access to lactazole A and its analogs in order to evaluate the 86 applicability of *laz* BGC for bioengineering, and to pave the way for future characterization of enzymes 87 and BGCs from the lactazole family. To this end, we report construction of the FIT-Laz system, a combination of flexible *in vitro* translation (FIT)⁴² with PTM enzymes from *laz* BGC, as a platform for 88 89 facile in vitro synthesis of lactazole-like thiopeptides (Fig. 1d). Taking advantage of the FIT-Laz system, 90 we explored the substrate plasticity of laz BGC, and found that in vitro lactazole biosynthesis is 91 remarkably tolerant to mutation, insertion and deletion of multiple amino acids, including npAAs. A 92 systematic dissection of the pathway led to the identification of the "minimal lactazole scaffold", a CP 93 with only 5 amino acids indispensable for the macrocyclization process. Our work opens a possibility 94 to tap into an unexplored sequence space of pseudo-natural thiopeptides, and use them as molecular
- 95 scaffolds in drug lead discovery against protein targets of choice.

96 **Results**

97 In vitro reconstitution of lactazole biosynthesis

We began with recombinant production of Laz enzymes in *Escherichia coli* BL21(DE3). The five enzymes (LazB, LazC, LazD, LazE, and LazF) were expressed and purified as soluble His-tagged proteins (**Fig. S1**). The FIT system was used to establish access to the precursor peptide (LazA;⁴³ **Fig. 2a**). Linear DNA template encoding LazA was assembled by PCR and incubated with the *in vitro* reconstituted translation machinery from *E. coli* supplemented with T7 RNA polymerase. This scheme for precursor peptide production parallels the previously established FIT-PatD and FIT-GS systems, used for the synthesis of azoline-containing peptides⁴⁴ and goadsporin analogs⁴⁵, respectively.

105 With all components in hand, we turned to reconstitution of lactazole biosynthesis. Maturation of 106 goadsporin, a distantly related linear azole-containing RiPP, is initiated with the formation of azoles, while Dha installation is dependent on it,46 and biosynthesis of thiomuracin also follows a similar 107 108 modification order.⁴⁷ Based on these results, we hypothesized that azole formation is the starting point 109 in lactazole biosynthesis, and therefore attempted to reconstitute the activity of LazDEF (LazD, LazE and LazF) first. LazDE is a split YcaO cyclodehydratase^{40,48} characteristic of thiopeptide BGCs: LazD 110 is predicted to bear a RiPPs recognition element, necessary for LP binding, ⁴⁹ and LazE contains an 111 ATP-binding domain,⁵⁰ utilized for ATP-dependent cyclodehydration of Cys/Ser residues in the CP (Fig. 112 113 1d). LazF is an unusual bifunctional protein that features a fusion between an FMN-dependent dehydrogenase which oxidizes azolines installed by LazDE to azoles,⁵¹⁻⁵³ and a glutamate elimination 114 115 domain, tentatively participating in the formation of Dha (see below). After LazA precursor peptide

expressed with the FIT system (**Fig. 2b**) was incubated with LazDE, the mixture was treated with iodoacetamide (**IAA**) and analyzed by LC-MS. The resulting broad-range extracted ion chromatogram (^{br}EIC; see S.I. 2.10 and Fig. S2–S5 for detailed description of ^{br}EIC) indicated that the LazDE reaction yielded a mixture of two, three, four and five dehydrations (**Fig. 2c**). Either LazD or LazE alone had no activity (**Fig. S6a and b**). In contrast, incubating LazA with LazDEF afforded a single product containing 4 azoles (**Fig. 2d**). No alkylation occurred on this peptide by IAA suggesting that all Cys residues were cyclized, and MS/MS analysis of this product supported the native azole pattern, *i.e.* three

- 123 thiazoles in positions 5, 7, 13 and one oxazole in position 11 (Fig. S7).
- 124 Next, we attempted to reconstitute the Dha-forming activity (Fig. 1d). LazBF is a split dehydratase, widely conserved in thiopeptide BGCs (Fig. 1b).54,55 These proteins are homologous to class I 125 lanthipeptide dehydratases, which utilize Glu-tRNA^{Glu} to glutamylate Ser or Thr residues in the CP of 126 127 a substrate (using the glutamylation domain),⁴⁶ and then catalyze elimination of the glutamate to yield Dha (using the elimination domain). In laz BGC, LazB is annotated as a glutamylation domain, and the 128 129 N-terminal part of LazF is an elimination domain. Even though we assumed that azole formation 130 precedes Dha synthesis, we first attempted to test LazB activity on the unmodified LazA. Surprisingly, LazB glutamylated LazA once when incubated in the presence of synthetic tRNA^{Glu} originating from 131 S. lactacystinaeus and glutamyl-tRNA synthetase (GluRS) from S. lividans (Fig. 2e), while reactions 132 lacking any one component led to no modification (Fig. S6f-h). These results indicated that like 133 homologous enzymes,^{56,57} LazB utilizes Glu-charged tRNA^{Glu} and catalyzes glutamylation of LazA. 134 Because *E. coli* tRNA^{Glu} and GluRS present in the translation mixture were not accepted, LazB appears 135 to be specific for the Streptomyces tRNA^{Glu} and GluRS. The complete dehydratase activity was 136 137 reconstituted with the addition of LazF to the mixture, in which case the reaction yielded a singly 138 dehydrated product (Fig. 2f). Extending the reaction time led to sluggish second and third dehydrations 139 (Fig. S6i). These results indicated that LazBF can catalyze formation of some but not all Dha in LazA 140 independent of azole formation.

We next studied whether LazBF forms remaining Dha in an azole-dependent fashion. To this end, we incubated the LazDEF-treated LazA, bearing 4 azoles, with LazBF, tRNA^{Glu} and GluRS. This reaction led to a major product 90 Da lighter than the 4-azole LazA, consistent with the formation of 5 Dha, suggesting that all available Ser in the CP were dehydrated (**Fig. 2g**). Coincubation of LazA with LazBDEF, tRNA^{Glu} and GluRS resulted in the formation of a complex mixture with the same major product (**Fig. S6j**).

Finally, we tested reconstitution of the entire biosynthetic pathway by adding LazC to the reaction. The putative pyridine synthase LazC is weakly homologous to TclM and TbtD, two well-studied enzymes catalyzing analogous reactions.^{38,58} Both enzymes are believed to initiate a [4+2]cycloaddition reaction leading to the formation of a macrocyclic product bearing dihydropyridine, which is further aromatized by eliminating LP as a C-terminal amide (**LP-NH**₂) to give rise to a pyridine ring. Accordingly, incubation of the aforementioned LazA bearing 4 azole/5 Dha with LazC afforded LP-NH₂ accompanied by a thiopeptide 18 Da lighter than expected, indicating that Ser4, unmodified in 154 lactazole A, was dehydrated (Dha4-lactazole A) (Fig. 2h and i). MS/MS analysis of this product 155 confirmed its structure (Fig. S11). Lactazole A was a minor product under these reaction conditions. 156 Changing the order of the enzyme addition (LazDEF followed by LazBC, tRNA^{Glu} and GluRS) 157 suppressed the formation of the overdehydrated product, but still, a mixture of thiopeptides formed (Fig. 158 2j). In contrast to the stepwise reactions, coincubation of LazA with the full enzyme set (LazBCDEF, tRNA^{Glu} and GluRS) resulted in the formation of lactazole A and LP-NH₂ with only a trace amount of 159 160 Dha4-lactazole A (Fig. 2k). LC-MS analysis of the *in vitro* synthesized lactazole A showed that its 161 molecular weight and HPLC retention time were identical to the authentic in vivo synthetized standard 162 (Fig. S9). Additionally, both samples had matching, annotatable CID MS/MS spectra (Fig. S8 and S10),

163 indicating that the one-pot reaction yielded the authentic thiopeptide.

In summary, here we demonstrated that the translation product of *lazA* accessed with the FIT system can be treated with the full set of Laz enzymes to yield lactazole A. We refer to this series of transformations as the FIT-Laz system.

167 Analysis of substrate tolerance of Laz enzymes

168 To understand the overall substrate plasticity of *laz* BGC, we next investigated whether the FIT-Laz 169 system can produce lactazole analogs. We commenced with Ala-scanning mutagenesis and prepared 14 170 single-point Ala mutants in the CP region of LazA. The precursor peptides were expressed and modified 171 with the FIT-Laz system, and the reaction outcomes were analyzed by LC-MS as above (Fig. 3a). Only 172 4 Ala mutants, S1A, S10A, S11A, and S12A, abolished formation of thiopeptides, whereas other 173 constructs led to the formation of corresponding lactazole analogs and LP-NH₂. During maturation, 174 Ser1 and Ser12 are converted to Dha and are then utilized by LazC for pyridine 175 formation/macrocyclization. Moreover, pyridine synthases are known to recognize the modification pattern around the 4π component,³⁸ which is consistent with the abrogation of biosynthesis in S10A and 176 177 S11A mutants. On the other hand, C13A mutant was converted to a thiopeptide without modifications 178 in the tail region, suggesting that the Dha10-oxazole11-Dha12 moiety is the minimal recognition motif 179 around the 4π component for the LazC-catalyzed macrocyclization. Significant accumulation of linear 180 side-products and partially processed peptides for W2A and G3A mutants indicates that these amino acids are also important for smooth lactazole biosynthesis. Ala mutants in positions 4–8, 15 and 16, 181 182 including those disrupting azole and Dha installation, were tolerated, albeit in some cases a mixture of

- 183 thiopeptides formed (Fig. S12 and S13).
- 184 Intrigued by these results, we examined whether non-essential modifications inside the macrocycle 185 (Ser4–Cys7) can be removed altogether. Indeed, a tetra-Ala mutant, LazA S4-C7A, was converted to a
- 186 thiopeptide containing just 2 azoles and 1 Dha upon treatment with the full enzyme set (**Fig. 3b**). A
- 187 pentamutant LazA S4-C7A, C13A also afforded a thiopeptide, but at a much lower overall efficiency,
- 188 as a number of partially processed linear peptides accumulated after overnight treatment (**Fig. S14a**).
- 189 Based on these results, we concluded that the five residues undergoing PTM in LazA S4-C7A (Ser1,
- 190 Ser10, Ser11, Ser12 and Cys13) are essential for efficient maturation. We termed the resulting
- 191 thiopeptide as the minimal lactazole scaffold, and the corresponding precursor peptide as the minimal

lactazole precursor (LazA^{min}). Because *in vitro* biosynthesis of the minimal lactazole proceeded as
 efficiently as the wild type, we decided to investigate enzymatic processing of LazA^{min} and its potential
 for bioengineering applications in more detail.

195 In the next series of experiments, we examined the tolerance of Laz enzymes to the presence of charged amino acids in the CP, and performed Lys- and Glu-scanning of LazA^{min} CP. Charged amino 196 acids are rarely found in CPs of thiopeptides,³⁹⁻⁴¹ and RiPP enzymes from other classes are also known 197 198 to disfavor charged amino acids in general, especially negatively charged Asp and Glu close to the 199 modification site. We prepared 11 single-point Lys mutants and 11 single-point Glu mutants in the nonessential positions of LazA^{min} and analyzed their processing as above (Fig. S15). The results of Lys-200 201 scanning revealed that a positively charged amino acid is well tolerated in 9 out of 11 positions, whereas W2K and A14K mutants suffered from inefficient processing. Glu was less accepted than Lys overall. 202 203 In addition to inefficient processing of W2E and A14E, mutants Q8E, A9E and P16E also resulted in 204 little to no thiopeptide formation. These data suggest that in addition to the five previously identified amino acids, Trp2 and Ala14 also play an important role in LazA^{min} maturation. 205

206Next, we sought to establish the minimal and maximum macrocycle sizes accessible with FIT-Laz. 207 All known thiopeptides range between 26- (thiocillin, thiostrepton and nosiheptide) and 35-membered macrocycles (berninamycin).⁵⁹ Additionally, previously reported bioengineering of thiocillin BGC led 208 209 to 23-membered artificial variants.⁶⁰ Here, we prepared amino acid insertion and deletion variants of LazA^{min}, and, as before, expressed and modified them with the FIT-Laz system. The results of LC-MS 210 211 analysis are summarized in Fig. 3c. Deletion of up to 3 amino acids between Ser1 and Ser10 was well 212 tolerated, and led to the efficient formation of 29- to 23-membered thiopeptides. The 4-6 amino acid 213 deletion mutants were also competent substrates and yielded 20- to 14-membered macrocycles, but at 214 relatively low overall efficiencies, as a number of partially processed linear peptides accumulated. 215 Formation of an 11-membered thiopeptide (deletion of 7 residues) was not observed. Thus, it appears 216 that 14-membered thiopeptides are the smallest accessible with Laz enzymes, 9 atoms smaller than the previously smallest thiocillin variants.⁶⁰ In contrast, no upper limit on the ring size could be placed. All 217 tested substrates were accepted by the enzymes: the largest synthesized product bore a 62-membered 218 219 macrocycle, which corresponds to a 10 amino acid insertion. The overall processing efficiency decreased linearly with increasing the cycle size; where LazA^{min} itself was efficiently converted to a 220 221 macrocycle and LP-NH₂, substrates with multiple amino acid insertions had substantial accumulation 222 of linear intermediates and side-products. A recent study demonstrated that TbtD, a LazC homologue 223 from thiomuracin biosynthesis, could perform an *inter*molecular [4+2]-cycloaddition, whereas TclM, 224 an enzyme from thiocillin BGC, could not,⁶¹ suggesting that LazC might function similarly to TbtD. 225 While certainly not intermolecular, the enzyme catalyzed formation of remarkably large macrocycles. Sequence extension outside of the macrocycle was also easily achievable, as 3 LazA^{min} variants with 226 227 the C-terminal tail extensions of up to 15 amino acids were efficiently converted to thiopeptides (Fig. 228 3d and S14b-d).

229 Encouraged by these results, we examined whether FIT-Laz can accommodate sequence

randomization inside the macrocycle. We prepared 10 LazA^{min} variants containing 10 consecutively 230 231 randomized amino acids each, which corresponds to the simultaneous insertion of 3 amino acids and 232 mutation of residues 3–9 in LazA^{min} CP (see S.I. 2.7 for sequence choices). Expression and modification 233 of these peptides by FIT-Laz and the subsequent LC-MS analysis (Fig. 4a) revealed that 9 out of 10 substrates produced thiopeptides as efficiently as LazA^{min}. One substrate (10aa-sub-4) led to the 234 formation of multiple thiopeptides owing to three Ser and Thr residues in the inserted region undergoing 235 236 differential dehydration (Fig. S16), and another variant (10aa-sub-10) had major accumulation of 237 partially processed linear peptides, albeit with detectable formation of the thiopeptide.

- Finally, we combined sequence randomization inside the macrocycle with the C-terminal extension, and constructed a LazA^{min} mutant with a 34 amino acid-long CP. Despite its size, this substrate efficiently generated a 3.7 kDa thiopeptide when treated with Laz enzymes (**Fig. 4b, S17**), highlighting the scaffolding ability of key residues in LazA^{min}.
- Taken together, these data indicate an unprecedented flexibility of *laz* BGC. Many individual enzymes and entire RiPP pathways are similarly promiscuous, but thiopeptide biosynthesis is usually sensitive to much more modest perturbations. These results point to potential applications of *laz* BGC in bioengineering.

246 Synthesis of hybrid thiopeptides with FIT-Laz

247 One advantage of the FIT system is its amenability to genetic code reprogramming. Incorporation 248 of multiple npAAs can be achieved by adding appropriate orthogonal tRNAs precharged with npAAs of choice by the use of flexizymes⁴² to the translation mixture lacking certain proteinogenic amino acids 249 250 and cognate aminoacyl-tRNA synthetases. The FIT system was previously used to synthesize peptides 251 containing a variety of npAAs, including D-, β -, N-methylated-, and α . α -disubstituted-amino acids as well as hydroxyacids.⁶² Recently, a combination of genetic code reprogramming in the FIT system with 252a promiscuous RiPP enzyme also enabled synthesis of peptides containing exotic azoline residues.⁶³ 253 Such npAAs are often found in peptidic natural products, both in RiPPs⁹ and in non-ribosomally 254 synthesized peptides (NRPs).⁶⁴ We reasoned that if LazA precursors containing ribosomally installed 255 256 npAAs are accepted by Laz enzymes, various "hybrid" thiopeptides may be accessible with the FIT-257 Laz system.

- 258We began by testing the ability of FIT-Laz to produce N-methylated thiopeptides, and prepared 12 259 lazA^{min} mutants bearing a single Met codon (AUG) in the CP. The Met codon was reassigned to either *N*-methylglycine (^{Me}Gly) or *N*-methylalanine (^{Me}Ala) by expressing these genes from a Met-depleted 260 translation mixture in the presence of precharged ^{Me}Gly-tRNA_{CAU} or ^{Me}Ala-tRNA_{CAU} (see S.I. 2.8 for 261 details). Treatment of these translation products with the full enzyme set (^{Me}Gly- and ^{Me}Ala-scanning 262 263 mutagenesis) and the subsequent LC-MS analysis revealed that, similarly to the results of Lys/Glu-264 scanning, either of the tested N-methylated amino acid was easily accepted in 9 positions, while 265 mutations at Trp2, Cys13 and Ala14 were detrimental, affording little to no mature thiopeptide (Fig. 266 S18).
- 267 Next, we tested whether more diverse npAAs can be incorporated into the thiopeptide scaffold

following the same logic. For this study, we focused on $lazA^{min}$ bearing the AUG codon in position 5, and analogous to the experiments above, prepared LazA^{min} variants containing D-Ala, D-Ser, cycloleucine (cLeu), pentafluorophenylalanine (Phe(F₅)), 5-hydroxy-tryptophan (Trp(5-OH)), lactic acid (^{HO}Ala), β-Met, and β-homoleucine (β-hLeu). All of these substrates were smoothly converted to the corresponding thiopeptides by the action of Laz enzymes, affording hybrid thiopeptides containing a variety of npAAs (**Fig. 5a**).

274 Finally, we studied whether multiple different npAAs can be simultaneously incorporated into the 275 minimal lactazole scaffold to generate highly artificial "pseudo-natural" macrocycles. Due to the presence of a 38-residue LP in LazA^{min}, the codon boxes available for reprogramming are limited (see 276 277 S.I. 2.8 for details). After some experimentation, we opted to reprogram four codons (AAG, CAU, UGG and UUU), and reassigned them as MeGly, cLeu, Phe(F5) and MeAla, respectively. To this end, a DNA 278 279 template encoding lazA^{min} with 4 codons of interest (Fig. 5b and 5c) was incubated in a 280 Lys/His/Phe/Trp-depleted translation reaction with ^{Me}Gly-tRNA_{CUU}, ^{Me}Ala-tRNA_{AAA}, cLeu-tRNA_{GUG}, and Phe(F₅)-tRNA_{CCA} to yield a LazA precursor peptide bearing 4 npAAs. Treating this substrate with 281 282 LazBDEF/tRNA^{Glu}/GluRS afforded a fully processed linear precursor bearing 2 azoles and 3 Dha (Fig. S19), while the reaction utilizing the full enzyme set led to the formation of the predicted thiopeptide 283 284 accompanied by LP-NH₂ (Fig. 5d). The identities of this macrocycle and its linear precursor were 285 confirmed by CID MS/MS analysis (Fig. S19 and S20). From these experiments, we conclude that the 286 FIT-Laz system offers facile access to previously inaccessible hybrid thiopeptides, including novel 287 heavily modified pseudo-natural architectures. These results additionally underscore the promiscuity of 288 Laz enzymes, as all tested substrates containing disruptive amino acids outside of the canonical 289 Ramachandran space were efficiently converted to mature thiopeptides.

290 Discussion

291 In this study, we completed *in vitro* reconstitution of *laz* BGC, which is responsible for biosynthesis 292 of lactazole A, a cryptic thiopeptide from S. lactacystinaeus. This is the first in vitro reconstitution of 293 an entire thiopeptide BGC, and the second reconstitution of biosynthetic enzymes involved in the 294 formation of a primary thiopeptide macrocycle.⁶⁵ The FIT-Laz system established in this study enabled rapid access to numerous LazA variants; the entire workflow from PCR assembly of lazA DNA 295 296 templates to LC-MS analysis of reaction outcomes fits within two working days. An added benefit of 297 working with an in vitro reconstituted BGC is the ability to decouple self-immunity, export and 298 proteolytic stability issues, so often complicating in vivo studies, from direct assaying of enzymatic 299 activities. Conversely, in vitro experiments provide no insight into in vivo fates and metabolism of the 300 underlying natural product, and thus, should be interpreted accordingly.

The results presented here indicate that *all* Laz enzymes tolerate substantial disruptions in the structure of the precursor peptide (**Fig. 6a**), which stands in contrast to other thiopeptide BGCs characterized to date. Out of 102 structurally diverse precursor peptides tested in this work, 92 yielded lactazole-like thiopeptides, 73 of which were accessed with efficiencies comparable to wild type lactazole A (Table S8). How the enzymes manage to properly modify such a diverse set of substrates

306 remains to be demonstrated. For now, it is apparent that Laz enzymes are highly cooperative. For 307 instance, addition of LazF to LazDE orchestrates efficient cyclodehydrations (Fig. 2c vs. Fig. 2d), LazC 308 prevents overdehydration by LazBF (Fig. 2h vs. Fig. 2j), and unaminoacylated S. lactacystinaeus 309 tRNA^{Glu} somehow affects azole formation mediated by LazDEF in the presence of LazB (Fig. S6e vs. 310 Fig. 2d and S6c-d). Combined with the fact that some Dha can form independent from azole 311 installation, it is likely that the biosynthetic mechanism is more elaborate than the "azoles form first, Dha second" paradigm observed during thiomuracin biosynthesis⁴⁷ and frequently assumed for other 312 313 RiPPs. Investigations into the nature of this cooperativity are a subject of our ongoing studies.

314 Broad substrate scope of Laz enzymes enabled development of the minimal lactazole scaffold (Fig. 315 6a and b). This thiopeptide requires only 6 PTM events (formation of 2 azoles, 3 Dha and a pyridine 316 heterocycle) for macrocyclization, and is biosynthetically the simplest known thiopeptide to date. The 317 5 amino acids undergoing these modifications - Ser1, Ser10, Ser11, Ser12 and Cys13 - are 318 indispensable for efficient macrocycle assembly, and further experiments demonstrated that the residues 319 adjacent to the modification sites, Trp2 and Ala14, are also important for efficient biosynthesis. 320 Remaining positions (3-9, 15, 16) accept a variety of amino acids, including disruptive npAAs. Modification of minimal lactazole precursor, LazA^{min}, in the FIT-Laz system is robust, and tolerates 321 322 massive sequence variations. Specifically, the macrocycle can be contracted or expanded to synthesize 323 14- to 62-membered thiopeptides (2 to 18 unmodified amino acids inside the macrocycle; Fig. 6c and 324 6d), and the variants with up to 18 amino acid-long tails are accessible as well (Fig. 6e). Most 325 importantly, LazA^{min} can accommodate mutations of consecutive amino acids, as demonstrated by the synthesis of thiopeptides with 10 randomized amino acids inside the macrocycle (Fig. 6e). 326

327 This flexibility of the lactazole biosynthetic machinery, combined with the minimal size of laz BGC, 328 which contains only the genes essential for macrocyclization, suggest that the minimal thiopeptide 329 scaffold may be an excellent candidate for bioengineering. Because continuous randomized epitopes can be displayed inside a thiopeptide backbone, we envision that combinatorial libraries based on this 330 scaffold can be generated and screened akin to the recent reports on lanthipeptide bioengineering.¹²⁻¹⁶ 331 332 In those cases, lanthipeptide libraries were prepared with the use of promiscuous lanthipeptide synthases, 333 and could be screened against a protein target of interest with the use of phage/yeast display or with the reverse two-hybrid system. These studies resulted in the discovery of lanthipeptide inhibitors of HIV 334 budding process,¹⁵ urokinase plasminogen activator¹⁶ and $\alpha_{v}\beta_{3}$ integrin binders.¹³ Similarly, we 335 anticipate that the integration of the FIT-Laz system with powerful in vitro screening techniques such 336 337 as mRNA display⁶⁶ will provide access to artificial thiopeptides with desirable pharmacological profiles for drug discovery purposes. 338

339 Synthesis of thiopeptide hybrids with other RiPP and NRP classes represents another bioengineering 340 avenue explored in this work. Combinatorial biosynthesis is a concept from NRP and PKS fields, where 341 enzymes from different BGCs are combined to act on a single substrate to generate novel natural 342 products.^{67,68} This concept has recently been applied to RiPPs either via simultaneous use of enzymes 343 from near-cognate BGCs⁶⁹ or by devising chimeric LPs,⁷⁰ demonstrating that multiple promiscuous

344enzymes can act together to produce nonnatural hybrid RiPPs. In vitro genetic code reprogramming, 345 easily achievable with FIT-Laz, offers an alternative route to similar hybrids, many of which are 346 inaccessible by existing methods. We demonstrated that thiopeptide-NRP hybrids (macrocycles 347 containing hydroxyacids, D-, β -, N-methylated-, and α , α -disubstituted-amino acids), thiopeptide-RiPP hybrids (N-methylation and D-amino acids are found in borosins,⁷¹ lanthipeptides,⁵⁴ proteusins,⁷² 348 phallotoxins⁷³ and many other RiPPs families⁹), and thiopeptides with "anthropogenic" amino acids not 349 350 found in nature (Phe(F_5) and cLeu) can be routinely accessed with FIT-Laz. Such noncanonical hybrid 351 architectures can further expand the range of available molecular complexity for biotechnology and 352 drug discovery. Overall, we believe that the established FIT-Laz system opens exciting new 353 opportunities for thiopeptide engineering and characterization of natural thiopeptide diversity.

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361 and S.A.

362 **References**

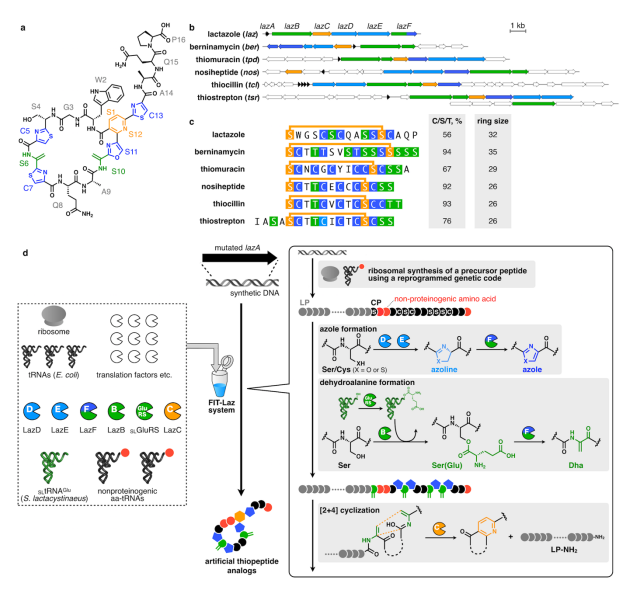
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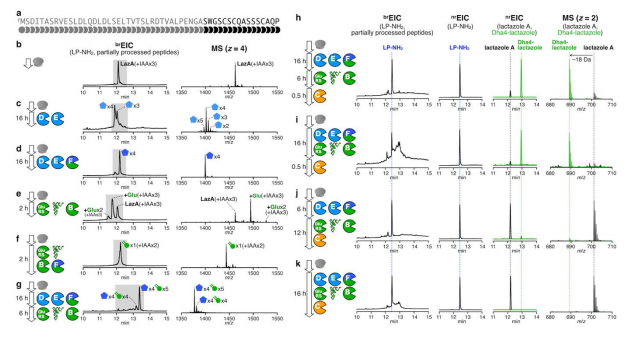
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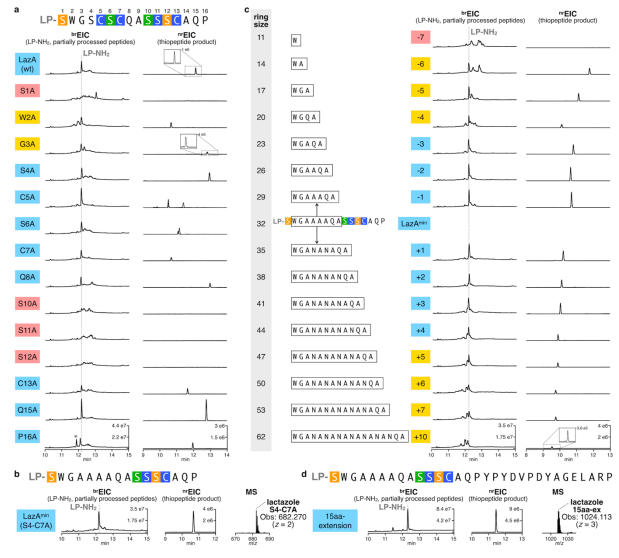
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Figure 1. Lactazole A and its biosynthesis with the FIT-Laz system. (a) Chemical structure of lactazole 526 A. (b) Comparison of *laz* BGC to other prototypical thiopeptide BGCs. Homologs of *laz* genes are 527 color-coded. Genes encoding enzymes responsible for the installation of azolines, azoles, 528 529 dehydroalanine, and pyridine are shown in light blue, blue, green, and orange, respectively. Precursor 530 peptide structural genes are shown in black, and ancillary genes absent from *laz* BGC are in white. (c) 531 Comparison of primary sequences for thiopeptides from panel (b), with the same PTM color coding. The comparison reveals an unusual macrocycle size, low C/S/T content and the absence of azole 532 533 modification in position 2 as unique features of lactazole. (d) Summary of the FIT-Laz system and the 534 roles of individual enzymes during lactazole biosynthesis. In FIT-Laz, synthetic DNA templates 535 encoding LazA or its mutants are in vitro transcribed and translated to generate precursor peptides, 536 which undergo a cascade of PTMs introduced by lactazole biosynthetic enzymes to yield lactazole A or its artificial analogs. 537



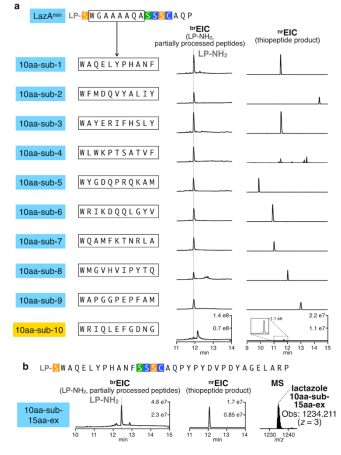
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540 Figure 2. Reconstitution of in vitro lactazole A biosynthesis. (a) Primary amino acid sequence of LazA precursor peptide. (b) – (g) Reconstitution of azole and Dha formation in FIT-Laz. LazA precursor 541 542 peptide produced with the FIT system was treated with a combination of Laz enzymes as indicated in each panel and the reaction outcomes were analyzed by LC-MS. Displayed are ^{br}EIC LC-MS 543 544 chromatograms and composite mass spectra integrated over a time period shaded in the corresponding 545 chromatograms. See S.I 2.6 and 2.10 for details on reaction conditions and the explanation of ^{br}EIC chromatograms. (h) – (k) Reconstitution of lactazole A biosynthesis in FIT-Laz. Displayed are LC-MS 546 chromatograms (left to right: ^{br}EIC; ^{nr}EIC at *m/z* 1026.77 for LP-NH₂ generated during the final 547 macrocyclization step; overlaid ^{nr}EICs at m/z 701.20 for lactazole A shown in black, and at m/z 692.20 548 549 for Dha4-lactazole in green) and overlaid mass spectra for the lactazole A and Dha4-lactazole. These 550 results demonstrate that the order of enzyme addition is critical to the success of lactazole A biosynthesis. 551



552

553 Figure 3. Substrate scope of the FIT-Laz system. (a) Ala scanning of the LazA CP. Single-point Ala mutants of LazA were treated with the full enzyme set and the outcomes were analyzed by LC-MS. 554 Displayed are LC-MS chromatograms (^{br}EIC chromatograms on the left showing partially processed 555 linear peptides and LP-NH₂ after enzymatic treatment, and ^{nr}EIC chromatograms on the right for 556 557 expected thiopeptides generated at m/z 0.10 tolerance window). For mutants highlighted in light blue 558 biosynthesis proceeded efficiently; yellow highlighting indicates inefficient thiopeptide formation 559 accompanied by the accumulation of linear intermediates and side-products; red - mutants that failed 560 to yield a detectable thiopeptide. Peaks denoted with an asterisk (*) indicate translation side-products. 561 Mutants C5A and S6A gave 4 and 2 thiopeptides, respectively, annotations of which can be found in 562 Fig. S12 and S13. Y-axes are scaled between samples for each chromatogram type. (b) LC-MS chromatograms as in (a) for the enzymatic processing of LazA^{min} on the left with a zoomed-in mass 563 564 spectrum of the produced thiopeptide on the right. (c) LC-MS chromatograms as in (a) for ring expansion and contraction study of LazA^{min}. (d) LC-MS chromatograms and mass spectrum as in (b) 565 566 for a LazA^{min} variant containing a 15-amino acid extension in the tail region. Collectively, these data point to the remarkable substrate tolerance of Laz enzymes. 567



568

Figure 4. Synthesis of lactazole-like thiopeptides with randomized sequences. (a) LazA^{min} variants 569 containing 10 consecutively randomized amino acids were first treated with LazDEF, and then with 570 LazBC, tRNA^{Glu} and GluRS (see S.I. 2.6 for details) and the outcomes were analyzed by LC-MS. 571 572 Displayed are LC-MS chromatograms (^{br}EIC on the left showing partially processed linear peptides and 573 LP-NH₂ after enzymatic treatment, and ^{nr}EIC chromatograms on the right for expected thiopeptides 574 generated at m/z 0.10 tolerance window). Mutants highlighted in light blue indicate efficient thiopeptide assembly; in yellow – inefficient thiopeptide formation accompanied by the accumulation of linear 575 576 intermediates and side-products. One construct, 10aa-sub4, resulted in 8 different thiopeptides, partial 577 annotation of which can be found in Fig. S16. Efficient *in vitro* biosynthesis observed in 9 out of 10 578 cases underscores the substrate plasticity of FIT-Laz. (b) LC-MS chromatograms as in (a) for the enzymatic processing of a 34 amino acid-long LazA^{min} variant on the left with a zoomed-in mass 579 spectrum of the produced thiopeptide on the right. 580

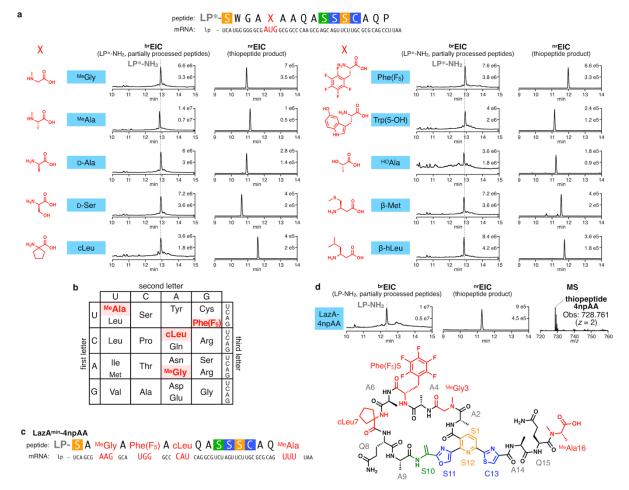
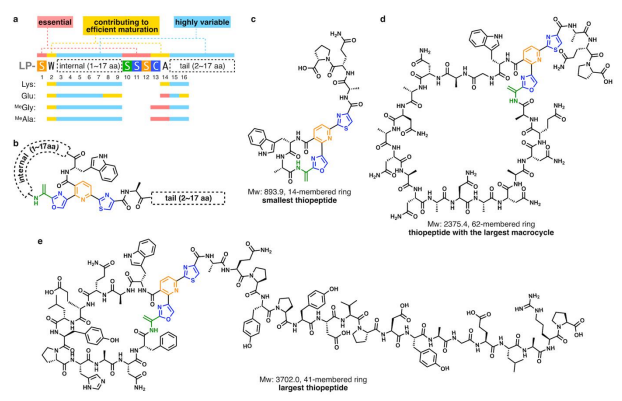


Figure 5. Synthesis of hybrid thiopeptides by genetic code reprogramming with the FIT-Laz system. 583 (a) Incorporation of a single npAA in a permissible position 5 of LazA^{min} using the Met AUG codon. 584585 LazA^{min} mutants accessed with *in vitro* genetic code reprogramming were treated with the full enzyme 586 set and the reaction outcomes were analyzed by LC-MS. Displayed are LC-MS chromatograms (^{br}EIC 587 chromatograms on the left showing partially processed linear peptides and LP*-NH₂ after enzymatic treatment, and "EIC chromatograms on the right for expected thiopeptides generated at m/z 0.10 588 tolerance window). LP* stands for LazA LP sequence where formyl-Met is replaced with N-589 590 biotinylated-Phe (see S.I. 2.8 for details). (b) Reprogrammed genetic code utilized for the synthesis of 591 a pseudo-natural lactazole containing 4 npAAs, and (c) its mRNA sequence. (d) LC-MS chromatograms as in (a) for the enzymatic processing of the LazA^{min} variant from (c), and the chemical structure of the 592 resulting thiopeptide. Taken together, these data suggest that diverse hybrid pseudo-natural thiopeptides 593 594 are accessible with the FIT-Laz system.



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Figure 6. The summary of the work. (a) Primary sequence representation of the minimal lactazole scaffold with the outcomes of the Lys, Glu ^{Me}Gly and ^{Me}Ala scanning experiments mapped to the resulting consensus sequence. (b) Chemical structure representation of the minimal lactazole scaffold. (c) – (e) Structural diversity of thiopeptides accessible with the FIT-Laz system. Displayed are chemical structures of the smallest artificial lactazole (c), thiopeptide with the largest macrocycle (d) and the largest construct (e) synthesized in this work.