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# COMMUNICATION

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# Total synthesis of incargranine A†

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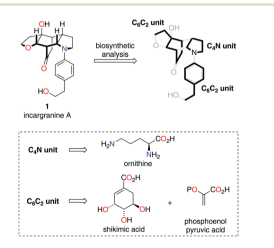
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Synthetic studies into the origins of the alkaloid incargranine A have resulted in the development of a four-step (longest linear sequence) total synthesis. This synthesis has been scaled-up to provide gram-scale quantities of material, which would alternatively require extraction of several metric-tons of dried-whole Chinese Trumpet-Creeper plants (*Incarvillea mairei* var. grandiflora).

In 2009 Zhang and co-workers isolated the alkaloid incargranine A (1) from *Incarvillea mairei* var. *grandiflora*, a Bignonia plant more commonly known as the Chinese Trumpet-Creeper plant (Scheme 1).<sup>1</sup> Incargranine A (1) has not yet succumbed to total synthesis and represents a particularly scarce natural product, constituting just 0.0000002% by weight of the dried whole plant. Therefore, a practical – *i.e.*, efficient and scalable – chemical synthesis of incargranine A (1) might advance a



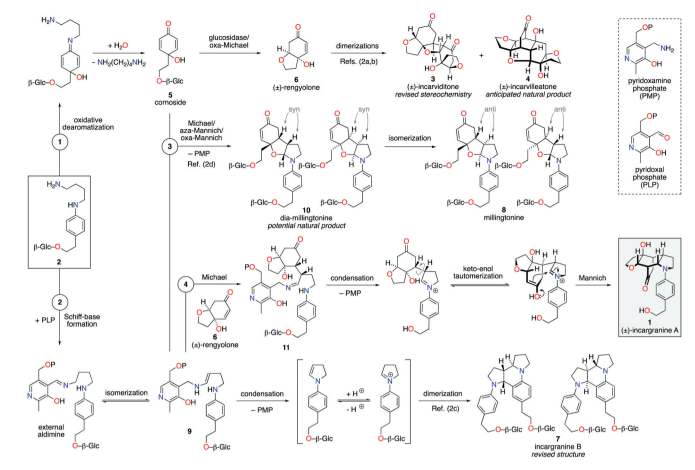
Scheme 1 Structure and biosynthetic analysis of incargranine A.

better understanding of its biological function. The novel framework of incargranine A (1) contains a synthetically daunting bridged-cyclohexane ring, in which all six-carbon atoms are stereogenic. Nevertheless, we were hopeful that if we could gain insight into how nature synthesizes this alkaloid a stepeconomical biomimetic strategy could be developed.

Our biosynthetic analysis, shown in Scheme 1, reveals incargranine A (1) is likely constructed from two shikimatederived C<sub>6</sub>C<sub>2</sub> units linked together by an ornithine-derived C<sub>4</sub>N unit. Our previous biomimetic studies on related phenylethanoid alkaloids provide important clues as to the potential origins of incargranine A (1)<sup>2</sup> We recently proposed that a network of pathways, all originating from a simple biosynthetic precursor, diamine 2, could account for the formation of several structurally distinct phenylethanoid natural products (Scheme 2).<sup>2d</sup> In our proposal, diamine 2 can participate in a pair of divergent oxidative pathways (Scheme 2; pathways 1 and 2). As shown in Scheme 2, pathway 1 terminates in the formation of incarviditone  $(3)^3$  and incarvilleatone  $(4)^4$  via the intermediacy of cornoside  $(5)^5$  and rengyolone (6),<sup>6</sup> whereas pathway 2 results in the production of incargranine B (7).<sup>2*a*-*c*,7</sup> It was proposed that these two divergent pathways could reconverge to give millingtonine (8),8 via a crossed-dimerization of cornoside 5, from pathway 1, and a PLP (pyridoxal phosphate) derived enamine 9, from pathway 2 (Scheme 2; pathway 3).<sup>2d</sup> The chemical feasibility of this re-convergent pathway was demonstrated in our seven-step biomimetic total synthesis of millingtonine (8).<sup>2d</sup> Herein, we propose that an additional reconvergent pathway could give rise to incargranine A (1) (Scheme 2; pathway 4). Thus, a Michael reaction between PLPenamine 9 and rengyolone (6) would give an intermediate imine 11, which would ring-close through a condensation/ Mannich reaction sequence to give incargranine A (1).9 To investigate the feasibility of this second re-convergent pathway, and in the hope of establishing a practical solution to the supply problem associated with incargranine A (1),<sup>1</sup> we decided to pursue the development of a biomimetic synthetic strategy.



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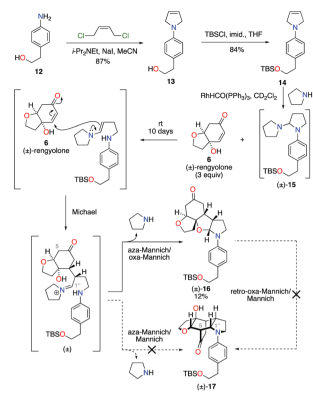


Scheme 2 Proposed network of biosynthetic pathways towards a family of plant-derived phenylethanoid natural products, including incargranine A.

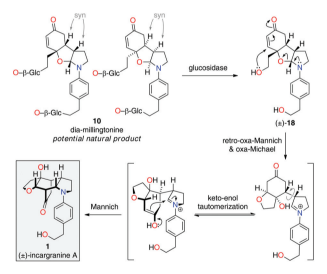
Condensation of 4-aminophenethyl alcohol 12 with (Z)-1,4dichlorobut-2-ene gave N-aryl-2,5-dihydropyrrole 13 in 87% yield (Scheme 3).<sup>10</sup> The primary alcohol functional group was then protected under standard conditions as a tert-butyldimethylsilyl ether, to give alkene 14 in 84% yield. Exposure of alkene 14 to our previously developed RhHCO(PPh<sub>3</sub>)<sub>3</sub> and pyrrolidine reaction conditions gave the expected aminal intermediate 15.<sup>2d,11</sup> Due to the instability of aminal 15, and in the interests of practicality and efficiency, rengyolone (6), which can be readily prepared from tyrosol in 3 steps,<sup>2a</sup> was added directly to this crude reaction mixture. Monitoring the reaction by <sup>1</sup>H NMR spectroscopy revealed it took 10 days at ambient temperature for aminal 15 to be consumed. Purification of the resulting crude reaction mixture by column chromatography resulted in a 12% isolated yield of an unwanted crossed-dimer 16, with no detectable formation of the desired product 17. Hemi-aminal 16 is presumably formed via a domino Michael/aza-Mannich/oxa-Mannich reaction sequence. In contrast, a final Mannich reaction between C5 and C1" would be required for formation of the incargranine A framework 17 (Scheme 3). Although this result demonstrates the viability of a crossed-dimerization between aminal 15 and rengyolone (6), several issues presented themselves with respect to using this strategy to access incargranine A (1). Firstly, rengyolone (6) proved to be relatively unreactive in the crossed-dimerization, taking over a week to give full consumption of starting material 15, while comparable reactions with *para*-quinols were generally complete in 24 h.<sup>2d</sup> Furthermore, the low yield of crossed-dimer 16, even after these prolonged reaction times, was not a promising start to the development of an efficient synthesis. Finally, and most importantly, our attempts to rearrange hemi-aminal 16 to give the incargranine A framework 17, *via* a retro-oxa-Mannich/Mannich reaction sequence, were unsuccessful.<sup>12</sup> This prompted us to reconsider our biosynthetic proposal and synthetic strategy.

Upon further evaluation of the incargranine A (1) framework it became apparent that it might instead be derived from the *syn*-diastereomer of millingtonine, dia-millingtonine (10), which we had previously identified as a potential natural product and direct biosynthetic precursor to millingtonine (8) (Scheme 2; pathway 3).<sup>2d</sup> Specifically, the putative aglycone of dia-millingtonine, diol 18, could undergo a domino retro-oxa-Mannich/oxa-Michael/Mannich reaction sequence to give incargranine A (1) (Scheme 4).<sup>13</sup> If this pathway could be shown to be chemically feasible it would lend further support

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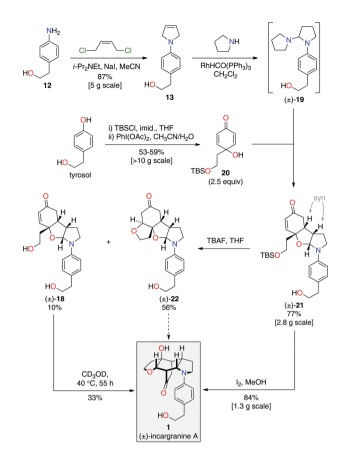
Scheme 3 Failed approach to synthesize incargranine A.



Scheme 4 Revised biosynthetic hypothesis for incargranine A.

to our proposal that dia-millingtonine (1) represents an as-yetundiscovered natural product.<sup>2d</sup>

During the development of this new strategy, it was discovered that protection of the primary alcohol in *N*-aryl-2,5-dihydropyrrole **13** was not necessary for the subsequent alkeneisomerization/hydroamination reaction. Thus, exposure of free alcohol **13** to RhHCO(PPh<sub>3</sub>)<sub>3</sub> and pyrrolidine gave the expected aminal intermediate **19** (Scheme 5).<sup>2d,11</sup> TBS-protected *para*-



Scheme 5 Total synthesis of incargranine A.

quinol **20**, which was prepared in 2 steps from tyrosol,<sup>2*a*</sup> was then added directly to this crude reaction mixture resulting in a kinetically-controlled crossed-dimerization to give *syn*-dimer **21** in 77% yield.<sup>2*d*</sup>

Attention could now turn to the de-protection of crosseddimer 21, a synthetic equivalent of dia-millingtonine (10), and its subsequent conversion to incargranine A (1). Cleavage of the tert-butyldimethylsilyl ether using standard TBAF (tetra-nbutylammonium fluoride) conditions gave the expected diolaglycone 18 in just 10% yield, alongside a cyclized-aglycone 22 in 56% yield (Scheme 5). Remarkably, it was observed that diol-aglycone 18 spontaneously rearranges to give  $(\pm)$ -incargranine A (1) when dissolved in methanol at ambient temperature, albeit very slowly. Ultimately, a 33% isolated yield of (±)-incargranine A (1) was achieved when a  $CD_3OD$  solution of diol-aglycone 18 was warmed to 40 °C for 2 days. The chemical feasibility of our proposed biosynthetic pathway between diamillingtonine (10) and incargranine A (1) had thus been established. All efforts, however, to rearrange the cyclized-aglycone 22 to give incargranine A (1) were unsuccessful, akin to our failure to rearrange hemi-aminal 16 (Scheme 3).<sup>12</sup>

The low yields and lack of selectivity achieved in the final de-protection and rearrangement steps rendered this synthesis unsuitable for scale-up. Alternative deprotection conditions were therefore screened in the hope of favoring production of diol **18**, whilst avoiding formation of the seemingly intractable ring-closed aglycone **22**. Vaino and Szarek have reported iodine in methanol as mild reaction conditions for the cleavage of *tert*-butyldimethylsilyl ethers.<sup>14</sup> Unexpectedly, however, exposure of *syn*-dimer **21** to iodine in methanol did not result in the formation of diol **18**, nor ring-closed aglycone **22**, but instead gave ( $\pm$ )-incargranine A (**1**) directly. Thus, in a single step, 2 new bonds, 2 new rings and 3 new stereogenic centres are formed in an impressive 84% yield. This synthetic sequence was readily scaled-up to provide gram-scale quantities of ( $\pm$ )-incargranine A (**1**), which compares very favorably to the effort required to obtain this material from the natural source; over four metric-tons of dried *Incarvillea mairei* var. *grandiflora* would need to be extracted to isolate one gram of natural incargranine A (**1**).<sup>1</sup>

Zhang and co-workers reported an optical rotation for natural incargranine A (1),  $[\alpha]_{D}^{22} = +2$  (c = 0.175, CHCl<sub>3</sub>).<sup>1</sup> However, given our biosynthetic speculation and the small magnitude of the reported optical rotation value, we consider it likely that natural incargranine A (1) exists as a racemic mixture. Unfortunately, no authentic sample was available to validate this hypothesis.<sup>15</sup> In all other respects, however, the spectroscopic data for our synthetic material matched that reported for natural incargranine A (1).<sup>1,15</sup> We propose that this successful synthesis provides new evidence in support of the proposal that dia-millingtonine (10) is a natural product.<sup>2d,16</sup> In fact, it is possible that incargranine A (1) is only produced from dia-millingtonine (10) during the extraction and isolation process. This would not necessarily mean that incargranine A (1) is an unimportant artifact of human intervention.<sup>17</sup> It is known, for example, that plants can use glycosidic-metabolites as chemical defense systems, wherein damage to the plant brings gycosidase enzymes into contact with the glycosides to release the active aglycones.<sup>18</sup>

#### Conclusions

In just three-linear steps from 4-aminophenethyl alcohol **12** we have selectively formed 2 new C–N bonds, 2 new C–C bonds, 2 new rings, and 6 new contiguous stereogenic centres, in 56% overall yield.<sup>19</sup> Key to the development of this efficient synthetic strategy has been the probing and refinement of a bio-synthetic proposal using chemical synthesis. Ultimately, this has led to new evidence in support of the notion that diamillingtonine (**10**) is an as-yet-undiscovered natural product.<sup>16</sup> Practical quantities of these metabolites are now available for interested parties to study their biological function.

### Conflicts of interest

There are no conflicts to declare.

### Acknowledgements

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## Notes and references

- 1 Y.-Q. Su, Y.-H. Shen, S. Lin, J. Tang, J.-M. Tian, X.-H. Liu and W.-D. Zhang, *Helv. Chim. Acta*, 2009, **92**, 165–170.
- 2 For previous biomimetic studies on related dimeric natural products, see: (a) P. D. Brown, A. C. Willis, M. S. Sherburn and A. L. Lawrence, Org. Lett., 2012, 14, 4537–4539; (b) K. Zhao, G.-J. Cheng, H. Yang, H. Shang, X. Zhang, Y.-D. Wu and Y. Tang, Org. Lett., 2012, 14, 4878–4881; (c) P. D. Brown, A. C. Willis, M. S. Sherburn and A. L. Lawrence, Angew. Chem., 2013, 125, 13515–13517, (Angew. Chem. Int. Ed., 2013, 52, 13273–13275); (d) P. D. Brown and A. L. Lawrence, Angew. Chem. Int. Ed., 2016, 55, 8421–8425).
- 3 Y. Q. Chen, Y. H. Shen, Y. Q. Su, L. Y. Kong and W. D. Zhang, *Chem. Biodiversity*, 2009, **6**, 779–783.
- 4 Y. P. Gao, Y. H. Shen, S. D. Zhang, J. M. Tian, H. W. Zeng,
  J. Ye, H. L. Li, L. Shan and W. D. Zhang, *Org. Lett.*, 2012,
  14, 1954–1957.
- 5 J. S. Rosendal, A. Kjaer and N. B. Juhl, *Acta Chem. Scand.*, 1973, 27, 367–369.
- 6 (a) K. Endo and H. Hikino, Can. J. Chem., 1984, 62, 2011–2014; (b) I. Messana, M. Sperandei, G. Multari, C. Galeffi and G. B. Marini Bettolo, Phytochemistry, 1984, 23, 2617–2619; (c) J. Tian, Q. S. Zhao, H. J. Zhang, Z. W. Lin and H. D. Sun, J. Nat. Prod., 1997, 60, 766–769.
- 7 Y.-H. Shen, Y.-Q. Su, J.-M. Tian, S. Lin, H.-L. Li, J. Tang and W. D. Zhang, *Helv. Chim. Acta*, 2010, **93**, 2393–2396.
- 8 T. Hase, K. Ohtani, R. Kasai, K. Yamasaki and C. Picheansoonthon, *Phytochemistry*, 1996, **41**, 317–321.
- 9 For our earliest biosynthetic proposals, see page 36 of the ESI for ref. 2*c*.
- 10 J. M. Bobbitt, L. H. Amundsen and R. I. Steiner, *J. Org. Chem.*, 1960, **25**, 2230–2231.
- (a) W. A. Herrmann and M. Prinz, Applied Homogeneous Catalysis with Organometallic Compounds, Wiley-VCH, Weinheim, 2nd edn, 2002; (b) T. J. Donohoe, T. J. C. O'Riordan and C. P. Rosa, Angew. Chem., 2009, 121, 1032–1035, (Angew. Chem. Int. Ed., 2009, 48, 1014–1017); (c) A. Vasseur, J. Bruffaerts and I. Marek, Nat. Chem., 2016, 8, 209–219.
- 12 Attempts to rearrange hemi-aminals **16** and **22** failed. Heating solutions in MeOH, EtOH or MeCN returned starting material. Treatment with TFA appeared to give isomerization from the *syn* to the *anti* configuration, with no sign

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of further rearrangement. Treatment with LiOH in refluxing  $MeOH/H_2O$  resulted in slow decomposition.

- 13 Treatment of millingtonine with glucosidase enzymes has been shown to result in a retro-oxa-Mannich/oxa-Michael/ Mannich reaction sequence to give a diastereomer of incargranine A, see ref. 8.
- 14 A. R. Vaino and W. A. Szarek, *Chem. Commun.*, 1996, 20, 2351–2352.
- 15 Professor Zhang very kindly provided pdf files of the processed NMR spectra for natural incargranine A, see the ESI† for direct comparisons.
- 16 For recent examples of natural product anticipation from biomimetic studies, see: (a) D. P. O'Malley, K. Li, M. Maue, A. L. Zografos and P. S. Baran, J. Am. Chem. Soc., 2007, 129, 4762–4775; (b) P. Sharma, B. Lygo, W. Lewis and J. E. Moses, J. Am. Chem. Soc., 2009, 131, 5966–5972; (c) M. Gavagnin, E. Mollo and G. Cimino, Rev. Bras. Farmacogn., 2015, 25,

588–591; (d) A. L. Lawrence, R. M. Adlington, J. E. Baldwin, V. Lee, J. A. Kershaw and A. L. Thompson, Org. Lett., 2010,
12, 1676–1679; (e) V. Sofiyev, J.-P. Lumb, M. Volgraf and D. Trauner, Chem. – Eur. J., 2012, 18, 4999–5005;
(f) S. Strych, G. Journot, R. P. Pemberton, S. C. Wang, D. J. Tantillo and D. Trauner, Angew. Chem., 2015, 127, 5168–5172, (Angew. Chem. Int. Ed., 2015, 54, 5079–5083);
(g) P. Ellerbrock, N. Armanino, M. K. Ilg, R. Webster and D. Trauner, Nat. Chem., 2015, 7, 879–882.

- 17 P. Champy, Artifacts and Natural Substances Formed Spontaneously, in *Biomimetic Organic Synthesis*, ed.
  E. Poupon and B. Nay, Wiley-VCH, Weinheim, 2011, vol. 2, pp. 849–934.
- 18 A. Mithofer and W. Boland, *Annu. Rev. Plant Biol.*, 2012, **63**, 431–450.
- 19 N. J. Green and M. S. Sherburn, *Aust. J. Chem.*, 2013, **66**, 267–283.