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# COMMUNICATION

## Synthesis and activity of the archazolid western hemisphere†

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A convergent and scalable synthesis of the archazolid western hemisphere has been completed. The V-ATPase inhibitory activity of this compound along with a previously prepared eastern domain was then tested using a convenient *Arabidopsis*-based V-ATPase assay.

The vacuolar-type H\*-ATPase (V-ATPase) is among the most widely distributed ATP-driven ion pumps in nature. It is expressed in all eukaryotic cells, where it participates in the acidification of intracellular organelles that are essential for many cellular processes.¹ The elucidation of the physiological role of the V-ATPase has revealed the important function these proteins play in a wide array of pathological processes including osteoporosis,² renal acidosis,³ and cancer.⁴ Thus, the V-ATPase is regarded as a promising pharmacological target and selective inhibitors of this enzyme represent important leads toward a better understanding of these diseases and the development of effective drugs for their therapy.

In 2003, as part of a program directed toward the identification of new anti-cancer agents, Höfle and coworkers reported the isolation of two novel cytotoxic compounds from the culture broth of strains of the myxobacteria Archangium gephyra.<sup>5</sup> Archazolids A (1) and B (2) showed high activity against a number of human and mammalian cancer cell lines with IC50 values ranging from 0.1 to 1 ng ml<sup>-1</sup> (Fig. 1). An incubation of PtK<sub>2</sub> potoroo cells with archazolid A (5 ng ml<sup>-1</sup>) led to the formation of vacuoles in the endoplasmic reticulum, a phenomenon that is typical for inhibitors of V-ATPases. Later studies of PtK<sub>2</sub> kidney cells incubated with archazolid A showed that these cells lost intact acidic lysosomes, also suggesting that the archazolids interfere with vacuolar-type ATPase.<sup>6</sup> This was confirmed by measuring the inhibitory activity of archazolids A and B on purified V-ATPase from M. sexta. Both archazolids A and B were shown to inhibit the activity of the purified V-ATPase holoenzyme halfmaximally at a concentration of 20 nM (IC<sub>50</sub> ca. 0.8 nmol per mg

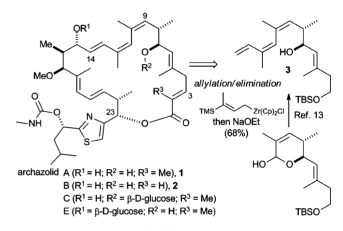


Fig. 1 Archazolid natural products.

enzyme), in the same range as the well-established plecomacrolidic V-ATPase inhibitors bafilomycin<sup>7</sup> and concanamycin.<sup>8</sup> Previous studies had demonstrated that the archazolids share, at least partially, the same binding site within subunit c of the  $V_0$  domain as bafilomycin and concanamycin.<sup>6</sup> Very recent point mutation experiments combined with the use of a labeled semi-synthetic archazolid derivative, however, indicate that archazolid V-ATPase binding is strikingly different than the plecomacrolides.<sup>9</sup>

Stereochemical assignment of archazolids A and B was first reported in 2006,  $^{10}$  and was later confirmed by total synthesis.  $^{12}$  Since their initial discovery, the family has been extended to include archazolids C–F.  $^{11}$  Each of the archazolids share a 30-carbon linear polyketide backbone that has been incorporated into a highly functionalized 24-membered macrolactone. Embedded within the core is a  $C_9$ – $C_{14}$  Z,Z,E-conjugated triene unique to the archazolids. Recently our group reported a direct approach to this subunit utilizing a tandem lactol TMS-allylation/Peterson elimination and a completion of the  $C_3$ – $C_{13}$  fragment (3).  $^{13}$ 

Elaboration of 3 into archazolids A and B was envisioned to occur by first olefination at  $C_2$ – $C_3$  followed by macrocycle formation by ring-closing metathesis (RCM) (Fig. 2). Herein we report a concise synthesis of the remaining  $C_{14}$ – $C_{23}$  and sidechain fragment (4) common to each of the archazolids required to complete our synthesis. Key disconnections include Horner–Emmons (HWE) olefination at  $\Delta^{18,19}$  and side-chain installation by addition of a suitable organometallic reagent derived from thiazole 5. Additionally, a tractable *Arabidopsis*-based V-ATPase assay has been developed that was then used to test the V-ATPase

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<sup>†</sup> Electronic supplementary information (ESI) available: Complete experimental details including spectroscopic and analytical data for compounds 4, 11, 12, 16, 17, 18, 19, and 20, Mosher ester data for compound 4, and V-ATPase assay dose response curves and methods for compounds 3, 20, and 21. See DOI: 10.1039/c1ob06446k

Fig. 2 Archazolid retrosynthesis.

inhibitory activity of both the "eastern" and "western" archazolid hemispheres.

Our synthesis began from known Weinreb amide 7,14 prepared on a ten-gram scale by a titanium-mediated Evans' aldol reaction with acrolein (8) (Scheme 1). 15 Conversion to the ketophosphonate 9 then allowed for a Horner–Emmons coupling<sup>16</sup> with (R)-Roche ester derived aldehyde 10.17 A comparison of deprotonation methods revealed that Ba(OH)2.8H2O18 proceeded with the highest trans-selectivity and has delivered multigram quantities of 11. Reduction of the ketone with NaBH<sub>4</sub> gave the corresponding alcohol with > 10:1 diastereoselectivity as determined by NMR analysis.<sup>19</sup> After some experimentation, it was found that methylation of the newly generated hydroxyl-group could be performed on gram scale by deprotonation with a freshly prepared precooled solution of LiHMDS followed by alkylation with methyl triflate (MeOTf) at -78 °C. Allowing the reaction mixture to warm resulted in a significant amount of 1,3-silyl migration<sup>20</sup> and a mixture of products. Selective removal of the primary TBS-group<sup>21</sup> in 12 and oxidation with Dess-Martin periodinane<sup>22</sup> then completed a synthesis of aldehyde 6.

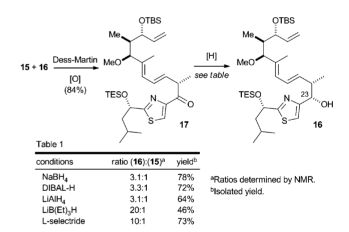
Scheme 1 Gram-scale synthesis of aldehyde 6.

The final fragment was prepared from known ketothiazole 13,<sup>23</sup> available in one step from commercially available 2,4-dibromothiazole (Scheme 2). Enantioselective reduction<sup>24</sup> and

Scheme 2 Thiazole fragment coupling.

TES-protection gave bromide 14. Treatment of 14 with *tert*-butyllithium resulted in a rapid lithium–halogen exchange<sup>25</sup> giving the corresponding highly-colored organolithium that added smoothly to 6 producing an inseparable mixture of 15 and 16 with modest Felkin-control (*vide infra*).

Oxidation of this mixture with Dess–Martin periodinane gave 17 which then allowed for installation of the desired  $C_{23}$ -O stereocenter by selective hydride addition (Scheme 3). It was found that L-Selectride gave the best balance of yield to selectivity, affording 16 in 73% yield as a 10:1 (NMR) mixture of diastereomers (Table 1). It



**Scheme 3** Installation of the  $C_{23}$ -O stereocenter.

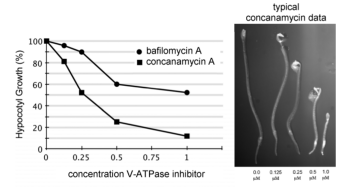
Completion of the fragment was then achieved in four steps from **16** by acylation of the free hydroxyl followed by selective TES-deprotection affording alcohol **18** (Scheme 4). Installation of the carbamate and removal of the acetate with DIBAL-H delivered **4** in 62% overall yield from **16**.

Scheme 4 Completion of the archazolid western hemisphere.

It was thought that the dense functionalization and various rigidifying elements within 4 might render compounds of this type V-ATPase inhibitors themselves.<sup>28</sup> To test this, the V-ATPase inhibitory activity of compounds **20**, **21**, and the previously prepared C<sub>3</sub>–C<sub>13</sub> fragment (3) was measured using a convenient *Arabidopsis*-based V-ATPase assay (Scheme 5).

Scheme 5 Synthetic western hemisphere derivatives.

Acidification of the plant vacuole by the V-ATPase facilitates cell expansion by generating turgor pressure through solute accumulation.<sup>29</sup> Selective inhibition by bafilomycin A, in fact, was a key component in identifying the first plant V-ATPase<sup>29</sup> and it is known that the selective inhibition of concanamycin A inhibits *Arabidopsis* cell elongation in a dose dependent manner.<sup>30</sup> The first V-ATPase mutant identified in plants, *det3*, exhibits a reduction in subunit *c* and in V-ATPase activity, resulting in the loss of the etiolated (dark) growth habit in seedlings.<sup>31</sup> A key component in the etiolated habit is stem elongation driven by V-ATPase mediated cell expansion. Cell growth in *Arabidopsis* seedlings was assayed by measuring etiolated seedling stem length in the presence of varying amounts of inhibitors (Fig. 3).<sup>32</sup>



Treatment <sup>A</sup>	IC <sub>50</sub> (μM)
concanamycin A	0.25
bafilomycin A	1.0
compound 20	>100
compound 21	>100
compound 3	>100

<sup>A</sup>Seeds were germinated and grown on media containing assayed compounds.

Fig. 3 Arabidopsis hypocotyl-growth V-ATPase assay.

While the plants proved highly sensitive to the known V-ATPase inhibitors bafilomycin A and concanamycin A, none of the synthetic archazolid derivatives displayed any significant V-ATPase inhibitory activity in this assay.<sup>33</sup> This data would

suggest that, like the plecomacrolides,<sup>34</sup> macrolactone formation is essential for archazolid V-ATPase binding and inhibition.

#### Conclusions

A convergent synthesis of the archazolid "western hemisphere" has been completed that is particularly well-suited for the large-scale preparation of advanced intermediates and access to a series of side-chain analogues. The V-ATPase inhibitory activity of this material was then evaluated using a tractable *Arabidopsis*-based V-ATPase assay. Assay results indicate that the macrocyclic structure of the archazolids is critical to their V-ATPase inhibitory activity. Efforts are ongoing to complete the synthesis of the natural product and analogues for future SAR studies using our plant-based assay method.

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