

Total Synthesis of (+)-Isomigrastatin**

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Isomigrastatin (**1**), recently isolated^[1] from *S. platensis*, has been a target for total synthesis in our laboratory. It is structurally related to migrastatin (**2**).^[1,2] The ability of migrastatins to retard cell migration might conceivably be exploitable in inhibiting the progression of clinically manageable tumors to less containable metastatic states. Earlier, we had described a successful inaugural total synthesis of **2**^[3a,b] and how that effort had paved the way for a program in diverted total synthesis (DTS).^[4b,c] DTS in turn enabled the discovery of migrastatin-inspired structures which are considerably more potent than **2** in suppressing colonization of tumors in vivo.^[4]

Our interest in isomigrastatin was further enhanced by its chemical vulnerability. Thus, Shen and co-workers demonstrated^[5a,b] that the ester (lactonic) linkage in **1** is rapidly hydrolyzed, thereby leading to a family of previously known biologically active systems, that is, the dorrigocins (Figure 1, see bold black and gray arrows). Moreover, as very elegantly shown, compound **1** undergoes extremely ready solvolytic and thermally induced rearrangement to migrastatin itself (Figure 1, see thin arrows). Indeed, **2** may actually not be a primary metabolite. Rather, it might well arise through a biosynthesis pathway leading to isomigrastatin (**1**), which, in turn, is transformed to **2** under isolation conditions.

Presumably, the ready transformation of **1** to **2** reflects the fact that the two *E*-configured double bonds in the macrolactone are better accommodated in the context of a 14-membered (cf. **2**) rather than a 12-membered (cf. **1**) macrolide. The pursuit of the total synthesis of compounds at the margins of viability can be conducive to the discovery of new

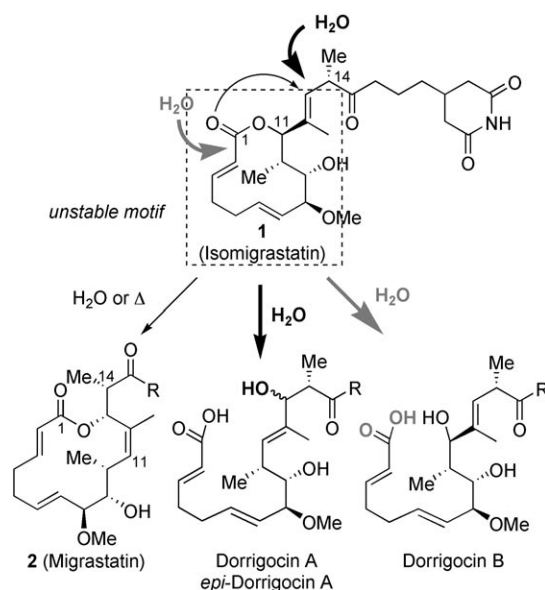


Figure 1. Isomigrastatin rearrangements (see text for details).

chemistry.^[6] Despite the deceptively simple allylic relationship between **1** and **2**, the present target presents a much greater challenge and the route required to reach **1** had very little homology with that used for **2**.

As the lability of **1** probably arises from the strain inherent in its *trans-trans* dienolide core, a responsive total synthesis strategy (Figure 2) would seek to delay implementation of the two *E*-configured double bonds until the latest possible stages of the effort. Another area of challenge in reaching **1** by total chemical synthesis is in the C11–C14 region. Of course, we hoped to take advantage of lessons learned from our earlier total synthesis of **2**. However, the *sp*³-configurational information in migrastatin (**2**), from C11 outward, is vested solely

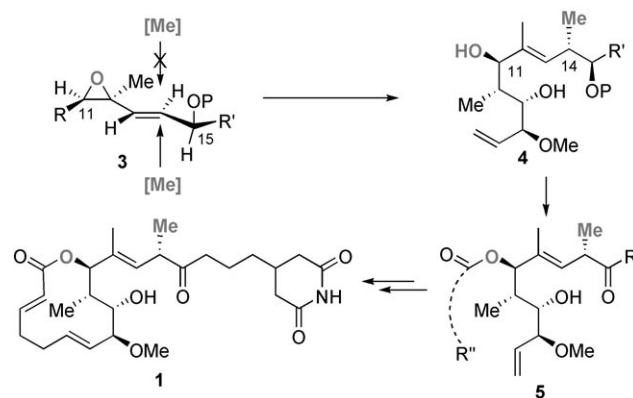


Figure 2. Synthetic strategy to **1** (P: protecting group).

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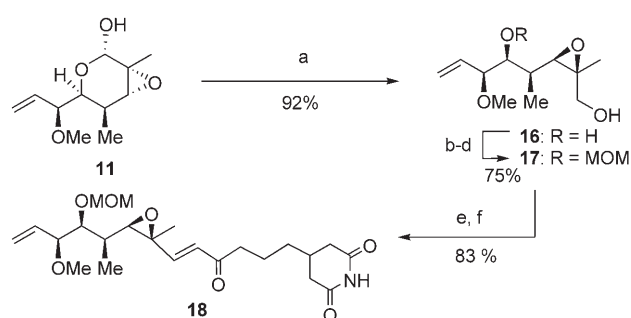
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in the vicinal C13–C14 relationship. Accordingly, the migrastatin experience^[3] does not inform as to the means for management of the relatively remote C11–C14 stereoconnectivity in **1** flanking the C12–C13 *E*-configured double bond. In Figure 2 is adumbrated the logic we hoped to employ in organizing the stereochemistry and functionality of the isomigrastatin, beyond C11. In the key stereo-defining step (see structure **3**), chirality information from a C11–C12 oxirane and possibly from C15 would hopefully govern the stereochemical sense of nucleophilic attack at C14, thereby establishing the C11–C14 stereoconnectivity. With its stereo-guidance role (if any) accomplished, the C15 oxygen atom would eventually be converted to the ketone level (see structure **5**) in the context of paving the way for the ring-closure phase of the synthesis to generate the dienolide en route to **1**.

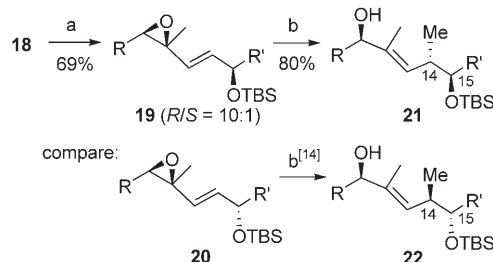
The initial steps of the venture, drawn from our migrastatin synthesis^[3a] occurred smoothly (Scheme 1). LACDAC reaction of **6** and **7** led to **8** and thence to **9**. Following Luche reduction^[7] and aqueous Ferrier rearrangement,^[8] as shown, **10** was in hand. Epoxidation of **10** afforded the α -oxirane **11** in 43 % yield (after recrystallization) along with approximately 5 % of the β -oxiranyl isomer (not shown).^[9] In a route featuring maximum convergence, the known Wittig reagent **13**^[10] served to alkylidenate the known aldehyde **12**.^[11] Hydrogenation of **14** afforded the appropriately functionalized Wittig reagent **15**.

In principle, coupling of **11** with **15** would be ultimately convergent in reaching the required series. In practice, this union could not be effected in our hands,^[12] thus obliging us to proceed in a more circumspect manner. Reduction of the lactol arrangement in **11** was accomplished (Scheme 2) with lithium borohydride, leading to alcohol **16**. This compound could be converted into its C9 MOM ether, **17**. Oxidation followed by coupling with phosphorane **15** afforded the required aldehyde **18**.

Precedent from the pioneering work of Marshall et al.^[13] could have been taken to suggest S_N2' attack (Scheme 3) by a first-order cyanocuprate at C14 would occur *anti* to the epoxide, regardless of the configuration of the proximal C15 OTBS group. Enone **18** was reduced with sodium borohydride to give a 1:1.2 mixture of epimeric allylic alcohols. This step was followed by protection, as shown, to afford the

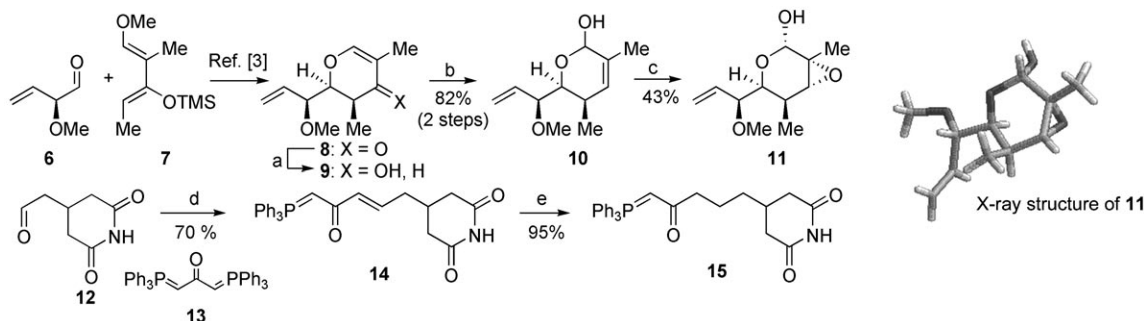


Scheme 2. Fragment coupling. Reagents and conditions: a) LiBH_4 , THF/ H_2O ; b) Ac_2O , pyridine; c) MOMCl, $i\text{Pr}_2\text{NEt}$; d) K_2CO_3 , MeOH; e) $(\text{COCl})_2/\text{DMSO}$, -78°C , then $i\text{Pr}_2\text{NEt}$; f) **15**, DMSO/ CHCl_3 , room temperature, 18 h. MOM: methoxymethyl.



Scheme 3. Matched–mismatched S_N2' epoxide opening. Reagents and conditions: a) 1.5 equiv (*S*)-Me-CBS, 1.1 equiv $\text{BH}_3\text{-Me}_2\text{S}$, then TBSCl, DMAP, Et_3N ; b) MeCuCNLI , Et_2O , $-20^\circ\text{C} \rightarrow \text{RT}$ (see Ref. [14]). (*S*)-Me-CBS: (*S*)-methyl oxazaborolidine; TBS: *tert*-butyldimethylsilyl; DMAP: 4-dimethylaminopyridine.

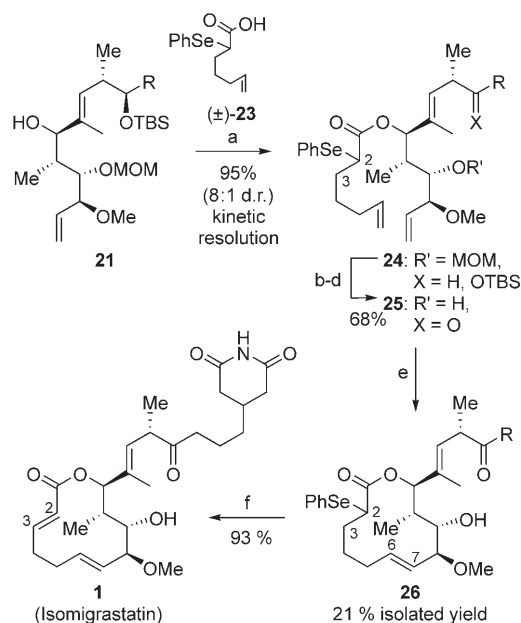
corresponding mixture of TBS ethers **19** and **20** (Scheme 3). In the event, both components of the mixture^[15] underwent clean S_N2' displacement with lithium cyanomethylcuprate with high selectivity for attack *anti* to the C15 OTBS group as implied in Figure 2, conformer **3**. Thus, unlike the case of Marshall et al., the configuration at C15 is apparently a very important stereochemical marker in determining the sense of the S_N2' substitution.^[13] This connectivity provided a strong incentive to reach a particular (in this case *R*) C15 epimer with stereocontrol. As indicated above, metal borohydride



Scheme 1. Fragment synthesis. Reagents and conditions: a) NaBH_4 , $\text{CeCl}_3\text{-H}_2\text{O}$, MeOH, $-20 \rightarrow 0^\circ\text{C}$, 2 h; b) CSA, THF/ H_2O 9:1, reflux, 3 h; c) mCPBA, K_2CO_3 , CH_2Cl_2 , 0°C , 24 h; d) DMSO, room temperature; e) H_2 , Pd/C, THF/MeOH, 16 h. TMS: trimethylsilyl; CSA: camphorsulfonic acid; mCPBA: *m*-chloroperbenzoic acid; DMSO: dimethyl sulfoxide.

reduction drawing upon resident stereochemical biases had led to a mixture of C15 epimers. Fortunately for our purpose, the powerful reagent-controlled technology of Corey et al.^[15] provided the required solution. In our case, use of the (*S*)-Me-CBS Corey catalyst^[15b] in the reduction of **18** afforded **19** with good *R* selectivity (approximately 10:1).

Following addition of lithium cyanomethylcuprate to the enriched mixture, purification was possible, providing **21** in 80% yield. After extensive exploration of macroannulation strategies, it was determined that successful olefin metathesis macrocyclization hinged on the absence of a double bond at C2–C3.^[16] Acylation of the C11 alcohol **21** with racemic selenoacid **23**^[17] proceeded with substantial kinetic resolution, providing **24** as an approximately 8:1 mixture of inseparable C2 epimers (Scheme 4). After removal of the TBS protecting



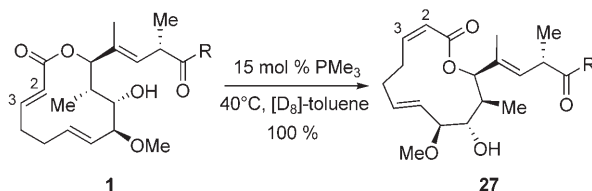
Scheme 4. Synthesis of (+)-isomigrastatin (**1**). Reagents and conditions: a) (±)-**23** (excess), EDCI, DMAP, (3 equiv each) CH₂Cl₂, 0 °C; b) pyridine–HF (1.1:1 mol/mol), 40 °C; c) (COCl)₂, DMSO, –78 °C, then *i*Pr₂NEt; d) Me₂BBr, *i*Pr₂NEt, –78 °C, then THF/NaHCO₃(aq); e) 20 mol% Grubbs' second-generation catalyst, toluene, 110 °C, 2 min; f) mCPBA, –78 °C, then *i*Pr₂NEt, –78 °C → RT. EDCI: 1-ethyl-3-(3'-dimethylaminopropyl)carbodiimide.

group, oxidation at C15, and MOM deprotection, ring-closing metathesis of **25** afforded the desired *E*-configured cyclized product **26** in 21% isolated yield, along with 36% of the C6–C7 *Z* isomer^[18] (not shown). At this stage, the C2 epimers of **26** could be separated. Gratifyingly, oxidative deselenation of either C2 epimer afforded isomigrastatin (**1**) with very high selectivity for the *E* geometry at C2–C3.^[19] ¹H and ¹³C NMR spectra of synthetic **1** were identical with those obtained from a naturally derived sample of isomigrastatin.^[20]

Moreover, synthetic and natural **1** exhibited identical behavior on TLC and provided the same product distribution under the hydrolytic conditions reported by Shen and co-workers,^[5a,b] as monitored by LC/MS analysis. Finally, the optical rotation of synthetic **1** ([α]_D = +178, CHCl₃, *c* = 0.18)

agreed, within error, with that of the natural sample ([α]_D = +170, CHCl₃, *c* = 0.18), confirming the absolute configuration as drawn.

Action of trimethylphosphine^[21] (Scheme 5) on synthetic **1** resulted in its complete conversion into **27**, showing rigorously that, as expected, the *Z*-configured 2,3 double-



Scheme 5. Isomerization of **1** to the 2,3-*cis* isomer.

bond linkage is more stable than the *E* variant.^[22] This stability order raises the question as to what factors are responsible for selective *E*-olefin formation (in **26** → **1**). We would argue that the preference for a *pro-E* over a *pro-Z syn* selenoxide transition state must be governed by minimization of vicinal nonbonded (van der Waals) interactions. Apparently, the transannular steric strain factors which eventually make the *E* isomer less stable than the *Z* isomer are not felt at this early point along the reaction coordinate.^[23]

In summary, the inaugural total synthesis of the very elusive isomigrastatin, while still a work in progress from a process perspective (see the low yield of **26**), illustrates important principles in the control of sp³-level stereochemistry and provides several pleasing examples of convergence. The properties of the broad family of migrastatins continue to stimulate research in our laboratory.

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- [14] Compound **20** was subjected to the cuprate S_N2' reaction as a mixture with **19**, from which it was virtually inseparable. However, the corresponding products **21** and **22** were easily separated and isolated with a combined yield of 75%.
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