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Evolution of a Unified Strategy for Complex Sesterterpenoids: Progress toward Astellatol and Total Synthesis of (–)-Nitidasin

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Abstract

Astellatol and nitidasin belong to a subset of sesterterpenoids that share a sterically encumbered *trans*-hydrindane motif with an isopropyl substituent. In addition, these natural products feature intriguing polycyclic ring systems posing significant challenges for chemical synthesis. Herein, we detail the evolution of our stereoselective strategy for isopropyl *trans*-hydrindane sesterterpenoids. Our endeavors included the synthesis of several building blocks, enabling studies toward all molecules of this terpenoid subclass, and of advanced intermediates of our initial route toward a biomimetic synthesis of astellatol. These findings provided the basis for a second-generation and a third-generation approach toward astellatol that eventually culminated in the enantioselective total synthesis of (–)-nitidasin. In particular, we orchestrated a series of substrate-controlled transformations to install the ten stereogenic centers of the target molecule and forged the carbocyclic backbone in a convergent fashion. Furthermore, we disclose our progress toward the synthesis of astellatol and provide insights into some observed yet unexpected diastereoselectivities by detailed quantum-mechanical calculations.

Keywords

Total Synthesis; Stereoselective Synthesis; Terpenoids; Natural Products; Computational Chemistry

Introduction

Among the large family of terpenoid natural products, sesterterpenoids form a small subclass with ca. 1000 isolated members.^[1] During biosynthesis, the variety of cyclization modes, hydride- and methyl-shifts and oxidation events leads to a plethora of intricate C_{25} -

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Supporting information for this article is given via a link at the end of the document.

architectures ranging from mono- to pentacarbocyclic compounds. One member of the latter rare subset is the fungal metabolite astellatol isolated by Simpson and Sadler in 1989 (Figure 1).^[2] Based on extensive NMR studies, the authors elucidated the relative configuration of the unique carbon framework with a total of ten stereogenic centers, an *exo*-methylene group attached to a four-membered ring and a single oxygenated site.^[3] In 1997, the similarly complex sesterterpenoid nitidasin^[4] was isolated from the Peruvian folk medicine "Hercampuri", which has found an application as a remedy against hepatitis, hypertension and diabetes.^[5] The rare 5-8-6-5 carbon framework is also decorated with ten stereogenic centers, together with its relative stereochemistry being unambiguously proven by single crystal X-ray diffraction. During their isolation program, Kawahara and co-workers found two structurally related metabolites in "Hercampuri". While alborosin^[6] seems to arise by degradation of nitidasin, the biosynthesis of nitiol^[7] did only lead to partial cyclization, not forming the central eight-membered ring.

From a structural perspective, astellatol, nitidasin and alborosin share a sterically encumbered *trans*-hydrindane motif that is substituted with an isopropyl residue *cis* to the angular methyl group. In addition, the five-membered ring has been oxidized to the carbonyl or the alcohol level. This portion has also been found in two inhibitors of mammalian GPI-anchor biosynthesis, YW-3699^[8] and YW-3548.^[9] Retigeranic acid B features the same moiety, although lacking the oxygenation at the five membered ring.^[10] Interestingly, chemists isolated two additional sesterterpenoids, retigeranic acid A^[10] and aleurodiscal,^[11] the isopropyl residue of which adopts the epimeric *trans* relationship to the angular methyl group.

Despite their intriguing architectures and interesting biological profiles, only a few research groups have investigated these natural products.^[12] This might be particularly due to the challenges associated with the synthesis of the polycarbocyclic ring-systems. Only the laboratories of Corey,^[13] Paquette,^[14] Hudlicky^[15] and Wender^[16] succeeded in the total synthesis of retigeranic acid A. Apart from these impressive achievements, Paquette^[17] and Dake^[18] reported the construction of the carbon framework of aleurodiscal and nitiol, respectively, and Tori and co-workers approached the YW-3699 skeleton, without addressing the hydrindane portion.^[19]

A few years ago, we launched a program to develop a general approach to isopropyl *trans*hydrindane sesterterpenoids.^[20] Herein, we provide the full details of our stereoselective strategy that allowed access to advanced intermediates for a biomimetic synthesis of astellatol and resulted in the first total synthesis of (–)-nitidasin.^[21]

Results and Discussion

First-generation retrosynthesis

As outlined in Scheme 1, our program initially focused on the total synthesis of astellatol. We envisioned accessing this natural product *via* a biomimetic cationic cascade from tricycle **1** (*vide infra*). The key intermediate **1**, featuring a strained eleven-membered ring with two trisubstituted double bonds, could be elaborated in a stereospecific manner by different retrosynthetic disconnections that will be detailed in the relevant subsections.

However, all approaches should be based on hydrindane 2 comprising a total of six stereogenic centers and bearing an alkene and a masked ketone as synthetic handles. We envisaged orchestrating a series of diastereoselective reactions to set the stereogenic centers of this key intermediate 2, giving ketone 3 as the logical precursor. While we initially considered employing a Pauson-Khand reaction to construct the five-membered ring of ketone 3, we ultimately decided to use diketone 4 as a starting point for our synthesis, due to the difficulties associated with the installation of the required *trans*-ring junction.^[22] It should be stated that we also aimed at accessing the deoxygenated analogue 5 by diversifying one of the intermediates *en route* to building block 3. Therefore, our route would not only pave the way for the synthesis of astellatol, nitidasin and the YW compounds but it would also provide a key intermediate for studies toward retigeranic acid B and nitiol.

Scheme 2 illustrates the envisaged biomimetic keystep in line with the biosynthesis proposed by Simpson.^[23] We envisioned treating tertiary allylic alcohol **1** with a Lewis or a Brønsted acid to form the stabilized tertiary carbocation **6** (with concomitant deprotection), which could then trigger a ring closure to form tetracycle **7**. Next, a stereospecific 1,2-hydride shift would set the relative configuration at C17 and generate homoallylic carbocation **8** that could engage in a homoallyl-cyclopropylcarbinyl-cyclobutyl rearrangement^[24] *via* intermediate **9** to furnish astellatol.

Domino transformations of this kind are highly challenging to realize in the laboratory in the absence of the biosynthetic machinery, given the number of possible reaction pathways and side products. However, we assumed that the conformational bias associated with the already installed *trans*-hydrindane portion might influence the outcome of the cascade favoring the formation of astellatol. It should be stressed that this biomimetic cascade would rapidly build up structural complexity and provide insight into the biogenesis of astellatol. In addition, potential side reactions could open avenues toward structurally related natural products, *e.g.* nitidasin.

Synthesis of trans-hydrindenone 15

According to our retrosynthetic analysis, our synthetic plan started with the synthesis of diketone **4**, which was readily prepared in both enantiomeric series in 100 g batches (Scheme 3).^[25] Additionally, we elucidated the solid-state structure of this building block that is frequently used for terpenoid synthesis.^[26] We originally intended to conduct a direct *trans*-selective reduction utilizing a bulky CuH species generated *in situ*.^[27] Due to the sensitivity of this reaction and aiming to prepare decagram amounts of material, we decided instead to follow a known, more robust five-step protocol for the installation of the *trans*-ring junction.^[28] Accordingly, chemo- and diastereoselective reduction of the more electrophilic ketone was followed by protection of the resulting alcohol as its *tert*-butyl ether to yield ketone **10**.

Next, reaction with Stiles' reagent $11^{[29]}$ introduced the carboxylic acid moiety as a traceless directing group to allow for a *trans*-selective hydrogenation. The reaction produced intermediate 12, the structure of which was confirmed by X-ray crystallography.

Subsequently, hydrogenation in the presence of Rosenmund catalyst followed by heat and vacuum-promoted decarboxylation provided *trans*-hydrindanone **13** in 59% yield as a 19:1 mixture of diastereomers. An ensuing three-step protocol consisting of ether cleavage, dioxolane formation and Cr-mediated oxidation delivered tricycle **14**. Purification by recrystallization not only removed any traces of the remaining *cis* isomer at this stage, but also provided crystals suitable for single crystal X-ray diffraction. Overall, this route was straightforward, robust and scalable. Thus, 19 g of ketone **14** was prepared in a single batch in 29% yield over eight steps.

With this material in hand, installing the required unsaturation in the five-membered ring was center stage. Initial attempts to achieve this oxidation by exposure of ketone **14** to $IBX^{[30]}$ or *via* synthesis of an α -bromo ketone were not successful. The same remained true for a protocol involving the treatment of a cyclic TIPS enol ether with Pearlman's catalyst in the presence of *t*BuOOH under an O₂ atmosphere.^[31] Contrarily, α -selenylation was straightforward (Scheme 4). Upon oxidation of the resulting selenium species, the desired elimination occurred, but was accompanied by a Bayer-Villiger-oxidation to yield bicycle **15**.^[32] We next evaluated a Saegusa-Ito oxidation,^[33] but attempts to render the reaction catalytic in Pd only gave low conversion to the desired product. Thus, we resorted to the classical conditions treating the intermediate trimethylsilyl enol ether with stoichiometric amounts of Pd(OAc)₂. This protocol provided the desired enone **16** in 82% yield after recycling recovered ketone **14**. The structure of enone **16** was also proven by single crystal X-ray diffraction.

Diastereoselective Michael addition

We next investigated the addition of a suitable three-carbon nucleophile *via* a cuprate addition. To this end, we treated enone **16** with the higher order cuprate derived from isopropylene magnesium bromide and CuCN at -78 °C, obtaining a single diastereomer **17** (Scheme 5).^[34] The selectivity of the addition was initially elucidated by NOESY NMR experiments indicating proximity between one methylene proton and the bridgehead methyl group. This stereochemical outcome was then unequivocally established by single crystal X-ray diffraction. Since we initially assumed that the addition of the cuprate from the β -face of the molecule would be disfavored due to sterical interactions with the bridgehead methyl group and because the selectivity is not obvious from inspecting the crystal structure of enone **16** alone, we investigated this transformation computationally.

We were aware of the difficulties of determining the active species of higher order cuprates in our reaction mixture.^[35] In order to develop a chemically reasonable model for the observed selectivity, we decided to use the less complicated low order cyanocuprate for the calculations. The most stable and reactive [Cu(isopropylene)CN·MgBr]–enone species were calculated to have the (μ_2 -cyano-C,N) ligand connecting the Mg²⁺ and Cu(I) centers (Figure 2). This coordination mode has been observed in X-ray structures of lithium cyanoorganocuprates.^[36] In our model system, magnesium acts as a Lewis acid activating the enone and stabilizing the forming enolate upon isopropylene addition from the cuprate. Other possibilities including the two metal centers being linked by a (μ_2 -Br) ligand were considered. These species were significantly higher in energy and will not be discussed (see

the Supporting Information). Michael additions of organocuprates have been proposed to proceed through an initial conjugate addition of the Cu(I) center to the α,β -unsaturated carbonyl system at low temperature leading to putative Cu(III) intermediates that subsequently transfer the organic nucleophile through reductive elimination upon warming of the reaction mixture.^[37] Similarly to this proposed mechanism, we located diastereomeric reacting complexes (**II**) and transition states (**III**[‡]) for the coordination of the cuprate (**R1**) to bicyclic cyclopentenone (**I**). Approximation from the allegedly less sterically hindered *Re* face, in which the copper atom and the bridgehead methyl group are mutually *anti* (**III**[‡]*anti*), was calculated to be fast ($G^{\ddagger} = +9.3 \text{ kcal mol}^{-1}$ from **II**) and only 0.1 kcal mol⁻¹ more favored than the approach by the opposite face (**III**[‡]*syn*). This was the first indication of steric factors not governing the stereochemical outcome of the reaction.

The structure of the resulting intermediates (IV) resemble Cu(I)- π -complexes more than Cu(III)- α -complexes. However, considering the short Cu–C β distance (2.0 Å), significant pyramidalization of C β carbon (28°) and closeness of the angle between the isopropylene and enone ligands to that of a square planar geometry (92°), as observed in the most stable isomer, the Cu(III) character of these species cannot be ruled out. These intermediates are unstable^[38] and undergo migratory insertion of the isopropylene group into the enone (\mathbf{V}^{\ddagger}) to give the conjugate addition intermediate (VI). This nucleophilic addition of the alkenyl group to the olefin is also fast, but entails the rate-limiting ($G^{\ddagger} = +15.3 \text{ kcal mol}^{-1}$ from **II**) and stereodetermining step of the reaction. At this stage, the alkenyl group transfer from the Si face of the enone $(\mathbf{V}_{syn}^{\ddagger})$ is significantly favored over the approximation from the Re face ($G_{syn-anti}^{\ddagger} = -1.4$ and -2.0 kcal mol⁻¹ at the ω B97X-D and M06-2X levels, respectively, for the lowest energy diastereomeric transition states, see Figure 3), leading to a calculated kinetic ratio of the syn and anti adducts of >97:3 at -78 °C assuming Curtin-Hammett conditions. This calculated stereoselectivity matches the experimental observation, and proves that a larger driving force – favoring the approach of the nucleophile from the more hindered β -face of the enone – overcomes the steric repulsions. The origin of this trend is the different torsional strain arising at the transition states of the syn and anti additions. This well-established effect, proposed initially by Felkin^[39] and applied to a multitude of nucleophilic, radical and electrophilic asymmetric processes, has been recently reviewed for reactions on unsaturated cyclic and bicyclic alkenes.^[40]

As illustrated in Figure 3, the C β -C γ bond in transition state $\mathbf{V}^{\ddagger}_{anti}$ shows a more eclipsed conformation (H β -C β -C γ -C δ dihedral angle = -12°) than in $\mathbf{V}^{\ddagger}_{syn}$ (H β -C β -C γ -C δ dihedral angle = +62°) due to the pyramidalization of C β upon nucleophilic attack and the conformational restrictions imposed by the bicyclic backbone. The C α -C β bond is also less eclipsed in both approximations. Regarding the steric repulsions, the H···H contacts between the incoming alkenyl and the bridgehead methyl groups (2.37 Å) are all larger than twice the van der Waals radius of hydrogen (2.20 Å) in $\mathbf{V}^{\ddagger}_{syn}$, and only contact between the alkenyl group and the axial C δ -H δ bond in the substrate is apparent (2.16 Å). A similar contact between the alkenyl group and the axial C γ -H γ bond in the substrate (2.18 Å) is present in $\mathbf{V}^{\ddagger}_{anti}$. In summary, steric repulsions are a minor contributor to the stereoselectivity of this reaction, which is largely controlled by torsional factors.

The Mg-enolates VI, in which copper forms a π -complex with the transferred isopropylene group, show the opposite and initially expected preference of a *syn* arrangement between the isopropylene and the bridgehead methyl group – both in axial positions of the fivemembered ring – being highly repulsive ($G_{syn-anti} = +4.7 \text{ kcal mol}^{-1}$). In the final products obtained after enolate protonation (VII), the relative stability of the *anti* diastereomer is even higher ($G_{syn-anti} = +6.7 \text{ kcal mol}^{-1}$). This fact will be very relevant in the following step in our synthetic route (*vide infra*). Irrespective of their thermodynamic stability, *syn* conjugate addition products were obtained from **16** under kinetic control, since the alkenyl group transfer is highly exergonic and irreversible ($G = -25 \text{ kcal mol}^{-1}$ from II).

A catalyst-controlled diversification

Since experiments on the use of isopropyl cuprate in the Michael addition gave unsatisfactory results in our hands, we next turned our attention to the seemingly straightforward hydrogenation of the isopropylene moiety and submitted bicycle 17 to standard hydrogenation conditions (Pd/C, H2, MeOH). Interestingly, we noticed a small impurity (less then 5%) by NMR spectroscopy that could be conveniently removed by recrystallization giving pure compound 18 in 91% yield (Scheme 6). Since the structure determination of compound 18 was thwarted by overlaying signals in NOESY NMR experiments, no unambiguous assignment could be established by these means. However, we obtained crystals of ketone 18 suitable for single crystal X-ray diffraction. To our surprise, the structure established that under the hydrogenation conditions epimerization of the former allylic stereogenic center at C4 had occurred. This was attributed to fact that transition metal-catalyzed hydrogenations can lead to double bond isomerization prior to reduction.^[41] This process is often unnoticed in substrates where the isomerized product is subsequently reduced without any consequences, but holds the potential to result in epimerization or racemization. Notably, exposure of alkene 17 to Pd/C in the absence of hydrogen gas resulted in no isomerization to the tetrasubstituted alkene or into conjugation with the ketone functionality.

In order to shed light on this unexpected epimerization, we investigated possible mechanisms by DFT calculations. Although the experimental conditions for hydrogenation are heterogeneous, and the participation of solid Pd/C, or Pd nanoparticles in the epimerization process cannot be ruled out, we only considered homogeneous reactions. Thus, the formation of catalytically relevant Pd(II) species resulting from the oxidative addition of Pd(0) into H–H^[42] (reactant) and MeO–H (solvent) bonds was hypothesized. The most plausible calculated mechanism involves two consecutive Pd(II)-catalyzed transformations (Figures 4 and S2, see the Supporting Information): (1) isomerization of the terminal alkene in the isopropylene group of **VII***syn* into a more stable internal, tetrasubstituted alkene (**XIV**) through an initial barrierless migratory insertion of palladium hydride PdH(MeO)(MeOH)₂ (**IX**[‡]*syn*), followed by a very fast β-elimination of MeOH (**XII**[‡]*syn*, G[‡] = +4.6 kcal mol⁻¹ from **XI***syn*); (2) hydrogenation of **XIV** either with H₂ on Pd/C (not considered), or initial stereoselective migratory insertion of homogenous PdH₂(MeOH)₂ into the terminal alkene (**XVI**[‡]*anti-ipso*), followed by reductive elimination of Pd(0) (**XIX**[‡]) to give ketone **XX***anti*, which is 6.1 kcal mol⁻¹ more stable than its epimer

XX_{*syn*}. The first steps involving alkene isomerization are all calculated to be extremely fast and energetically downhill, except the last β-elimination step that is slightly endergonic. Nevertheless, the overall isomerization process to form the internal olefin is exergonic ($G = -7.0 \text{ kcal mol}^{-1}$ from **XI**_{*syn*}). Migratory hydride insertion of the PdH₂ π-complex **XV** was calculated to occur regio- and stereoselectively at the *ipso* carbon and from the *Re* face of the terminal olefin (**XVI**[‡]_{*anti-ipso*}). Although this process involves a negligible activation barrier, the diastereomeric π-complex and transition state from the *Si* face are both disfavored by 1.6 kcal mol⁻¹. This is due to the incipient repulsive 1,3-diaxial interactions with the bridgehead methyl group. These steric repulsions are even more apparent in the final and rate-limiting reductive elimination step (**XIX**[‡]*anti-ipso*, $G^{\ddagger} = +17.4 \text{ kcal mol}^{-1}$ from **XVII***anti-ipso*) (see the Supporting Information), in which the energy difference between the two epimeric transition states is at a maximum ($G^{\ddagger}_{syn-anti} = +9.1 \text{ kcal}$ mol⁻¹). Overall, this mechanism explains the complete epimerization observed experimentally, and demonstrates the potential for Pd(II) hydrides generated *in situ* from heterogeneous H₂ and Pd/C to promote fast alkene isomerizations.

As a consequence, the choice of the hydrogenation catalyst was revisited and Pd/C was replaced with Adam's catalyst (PtO₂). When conducting the reaction in MeOH, partial reduction of the carbonyl functionality in ketone **19** was observed. This was bypassed when submitting ketone **17** to PtO_2 in EtOH under an atmosphere of hydrogen furnishing ketone **19** in 93% yield (Scheme 7). The obtained NMR spectroscopic data matched the ones of the impurity observed in the hydrogenation with Pd/C and we established the identity of ketone **19** by single crystal X-ray diffraction, indicating that under these reaction conditions the alkene isomerization process is suppressed.

We then turned our attention to the transposition of the oxygen functionality *via* an alkene. Due to the unexpected epimerization, these studies were initially carried out with the aisopropyl ketone **18**. It should be emphasized that the transformations examined on this system provided helpful information for the further synthesis with the desired stereochemistry at C4. Our efforts commenced with the transformation of ketone **18** to tosylhydrazone **22** as a precursor for a Shapiro olefination.^[43] However, the examined reaction conditions (TsNHNH₂, TsOH, NaOAc, TsNHNH₂, TsOH, MgSO₄) did not provide the desired material (Scheme 8). Alternatively, a diastereoselective reduction enabled the synthesis of alcohol **20**, which was subjected to elimination conditions without success. In contrast, formation of the corresponding xanthate **23** followed by Chugaev elimination furnished alkene **21** in low yield.^[44] In parallel, we explored an alternative two-step protocol consisting of enol triflate formation and an ensuing Pd-catalyzed reductive detriflation^[45] leading to the desired alkene **21** in excellent yield.

Such material was then elaborated to ketone **24** by hydrogenation and cleavage of the dioxolane protecting group. We established the structure of building block **24**, which intersects with the routes toward retigeranic acid A developed by Corey^[13] and Hudlicky,^[15] by single crystal X-ray diffraction. The synthesis of ketone **24** was shortened by one step when preparing diene **25**, which again underwent the previously observed epimerization under the hydrogenation conditions.

Next, we were poised to transfer the reductive detriflation protocol to epimeric ketone **19** with the correct C4 stereochemistry. This reaction proceeded in excellent yield providing multigram quantities of alkene **26** (Scheme 9). Subsequently, alkene **26** was subjected to hydrogenation conditions and, in analogy to the hydrogenation of ketone **17**, the use of Pd/C as a catalyst resulted in epimerization giving rise to a 1:2 ratio of diastereomers **27** as indicated by ¹H NMR spectroscopy.

Interestingly, the hydrogenation in the presence of PtO_2 also occurred with partial epimerization in the same ratio. We thus resorted to a non-metal mediated reduction protocol utilizing diimide,^[46] and subsequent acetal cleavage furnished building block **5**, suitable for a synthesis of retigeranic acid B. In addition, regioselective silyl enolether formation followed by ozonolysis could provide an entry for the synthesis of nitiol.^[20]

Installation of the oxygen functionality

We next turned our attention to the introduction of the C5 alcohol. Initially, alkene **26** was converted to epoxide **30** with single crystal X-ray diffraction confirming once more the correct stereochemistry at C4 (Scheme 10). The structure indicates that the trajectory for an epoxide opening *via* a $S_N 2$ displacement is blocked due to geometric constraints. Consequently, treating epoxide **30** with LiEt₃BH up to 140 °C did not result in any conversion.

Next, a screening of alkene hydroboration conditions was carried out. We were aware that the outcome of this reaction would not be favored by electronical factors, but rather by steric effects. We initially employed a slight excess of $BH_3 \cdot SMe_2$ that delivered, after oxidative work up, a 2.7:1 mixture of regioisomeric alcohols **28** and **29** in favor of the desired product **29** (Table 1, Entry 1). Variation of temperature (Entry 2) and equivalents of hydroboration reagent (Entries 3,4) did not result in a significant improvement of the regioselectivity. Since the more bulky boron reagents ThxBH₂, (sia)₂BH and 9-BBN either led to no conversion of the starting material (Entry 5) or acetal cleavage (Entries 6,7), we revisited our findings and noticed that the amount of hydroboration reagent affected the regioselectivity.

Whereas the best result was obtained by quick addition of 0.7 equivalents of BH₃·SMe₂, the ratio dropped to 2.1:1 by employing 1.6 equivalents. This observation was tentatively explained by the *in situ* formation of a chiral hydroboration agent RBH₂. Consequently, alkene **26** was exposed to (+)-IpcBH₂ (**31**), prepared *in situ* from (*S*)-Alpine BoramineTM and BF₃·OEt₂,^[47] which improved the regioisomeric ratio to 3.7:1 (Entry 8). Alcohols **28** and **29** were difficult to separate, but column chromatography enhanced the diastereomeric ratio ranging from 5.5:1 (84% yield) to 12:1 (67% yield), providing 5.20 g of alcohol **29** on the largest scale. Interestingly, the use of (–)-IpcBH₂ (*ent*-**31**) gave an approximately 1:1 mixture of regioisomers **28** and **29** (Entry 9). Therefore, the reaction of alkene **26** with borane **31** constituted a matched case of double diastereoselection. The carbonyl group was then unmasked to yield bicycle **32** and column chromatography allowed for an improvement of the regioisomeric ratio (12:1 to 19:1). Subsequently, alcohol **32** was converted to TBS ether **3**, the structure of which was unambiguously proven by single crystal X-ray diffraction.

Installation of the C9 and C10 stereogenic centers

In the following experiments, we focused on the installation of the C9 and C10 stereogenic centers (astellatol numbering). Based on literature precedence, we were aware that such a setting would be difficult to access. A Tsuji crotylation via a mixed carbonate was ruled out due to the difficulties associated with diastereoselectivity and mixtures of branched and linear products.^[48] In order to evaluate a viable synthetic pathway, these studies were initially carried out with model ketone 13. Firstly, we aimed to establish the required regioselectivity in reactions involving the carbonyl functionality in ketone 13. To this end, we treated ketone 13 with PhMe₃NBr₃ (PTAB) in the presence of HOAc to generate diastereometric α -brown carbonyl compounds 33 that were further elaborated, with enone 34 being isolated as the sole product (Scheme 11). On the other hand, we conducted a kinetic deprotonation with LiHMDS and trapped the enolate with Manders' reagent 36 to give β keto-ester 37.^[49] Furthermore, we investigated the conversion of compound 37 to vinvl allvl ether **38** under Mitsunobu conditions.^[50] However, a brief screen of reaction conditions (*e.g.* ADDP, PBu₃) did not result in any conversion. This was unfortunate as we were particularly interested in this transformation since the latter intermediate 38 could serve as a precursor for a Claisen rearrangement to install the C10 stereogenic center.

Since proof of the desired regioselectivity in transformations of ketone **13** had been obtained, we next focused on alkylation protocols utilizing different electrophiles such as allyl iodide or α -halo acetic acid derivatives. Unfortunately, ketone **13** was largely resistant to the attempts to form product **35** or an *O*-allylated analogue suitable for a Claisen rearrangement.^[49]

Thus, we resorted to a Pd-mediated allylation protocol pioneered by Negishi^[51] and recently applied by Covey^[52] that provided hydrindane **39** on multigram scale (Scheme 12). Subsequently, we examined the reduction of the ketone moiety and found a reagent-controlled reversal of diastereoselectivity. Whereas exposure of the ketone **39** to NaBH₄ delivered alcohol **41** in 79% yield, treatment of intermediate **39** with K-Selectride® exclusively led to α -diastereomer **40**. The structure of alcohol **41** was elucidated based on NOESY experiments and analysis of coupling constants. In order to incorporate a γ -lactone moiety for the diastereoselective installation of the C10 methyl group, we sought to convert alkene **40** into the corresponding aldehyde by ozonolysis. Since the spontaneously formed lactol was quantitatively converted to the acetals by reaction with the protic co-solvents EtOH or MeOH, we submitted alkene **40** to a Lemieux-Johnson oxidation instead.^[53] The desired reaction proceeded smoothly and the obtained crude lactol was oxidized with PCC to yield tricycle **42** in an excellent yield of 90%.

The stage was now set for the installation of the C10 methyl group taking advantage of the conformational bias of lactone **42**. To this end, lactone **42** was sequentially treated with LiHMDS and MeI at -78 °C yielding, as expected, lactone **43** as a single diastereomer (Scheme 13). The stereochemical outcome of this transformation was initially confirmed by analysis of ¹H NMR coupling constants. As shown in Scheme 13, the C10 methine proton was observed as a quartet, indicating a dihedral angle H10–C10–C9–H9 of around 90° according to the Karplus curve. A comparison with the DFT-calculated dihedral angle of 87°

for this structure supported the identity of lactone **43**. In addition, we unambiguously assigned the structure of tricycle **43** by single crystal X-ray diffraction.

The focus was then turned to the installation of a terminal alkene. We initially reduced lactone **43** to the corresponding lactol (DIBAL-H, -78 °C) and exposed the intermediate to a Wittig methylenation. In our hands, this reaction required elevated temperatures (THF, 66 °C) to proceed and provided the desired product only in low yield as a mixture of diastereomers. In contrast, the reaction with ylene **44** at 81 °C yielded ester **45** as a single diastereomer, albeit in moderate yield (Scheme 14).

In view of these results, we decided to pursue another strategy and synthesized the corresponding diol by reaction of lactone 43 with LiAlH₄. In order to avoid an orthogonal protecting group strategy, we exploited the potential of Swern oxidation conditions to convert primary triethylsilyl ethers directly to aldehydes and prepared intermediate 46.^[54] Careful control of the external temperature not exceeding -60 °C and prolonged reaction times (7-8 h) were crucial for a success of the ensuing oxidation since otherwise lower yields and/or the bisoxidized carbonyl compound were observed. Furthermore, the aldehyde was isolated as a single diastereomer, indicating that no epimerization occured under the reaction conditions. Ultimately, a Wittig methylenation provided access to intermediate 47, which served as the branching point for several routes toward the target sesterterpenoids (vide infra). We then set out to adapt the reaction conditions to trans-hydrindane 3 with the correct substitution pattern on the five-membered ring in place. Initially, the TBS protected analogue 3 was converted to ketone 49, which in turn was elaborated to tricycle 51 (Scheme 15). Compared to the model system, the diastereoselective methylation required a slightly higher reaction temperature of -40 °C to achieve full conversion. In analogy to the preparation of the model studies, lactone 51 was carried forward in four steps to alkene 2. Since we found in initial experiments that a chemoselective TES deprotection in bicycle 2 (e.g. CSA, PPTS or HOAc in CH₂Cl₂/MeOH) was accompanied by cleavage of the TBS ether, we decided to employ the more robust, but fluoride labile SEM protecting group in upscale experiments. Thus, protection of alcohol 32 as its SEM ether 48 followed by allylation gave rise to ketone 50, which was converted to alkene 53 via tricycle 52. Overall, the established protocol proved to be excellent, affording building block 53 in 47% yield over nine steps on a reasonable scale.

Toward construction of the macrocycle – the metathesis approach

Having successfully installed the C9 and C10 stereogenic centers, we next turned our attention to the preparation of the precursor for the biomimetic key step. One major challenge concerning this envisaged route was the construction of the highly strained eleven-membered ring in tricycle **1** comprising two (*E*)-configured trisubstituted alkenes (*cf*. Scheme 1).^[55] In order to conserve our precious building block **53**, these studies were conducted using model compound **47**. In our first retrosynthetic proposal, we aimed at closing the macrocycle **54** by a ring-closing metathesis (RCM)^[56] from allylic alcohol **55**, which in turn could arise from alkene **47** by a sequential B-alkyl Suzuki coupling^[57] and an alkenyl lithium addition (Scheme 16).

This approach commenced with a coupling of the *in situ* generated alkyl-boron species of alkene **47** (9-BBN, 40 °C) with alkenyl iodide **56**^[58] and provided intermediate bissilylether **57** that was submitted to the previously established deprotective Swern oxidation conditions (Scheme 17). Upon Wittig olefination, diene **58** was obtained in a good yield of 66% over the three steps. In order to complete the synthesis of the model RCM precursor **55**, the silyl ether within *trans*-hydrindane **58** was cleaved and the resulting secondary alcohol was oxidized with Dess-Martin periodinane (DMP).^[59] The subsequent addition of isopropenyl lithium at -78 °C was surprisingly facile given the steric encumbrance at the carbonyl functionality, and furnished tertiary alcohol **55** as a single diastereomer. The relative configuration at the newly formed stereogenic center could not be assigned unambiguously, but we assumed an equatorial attack due to the shielding effect of the adjacent substituent, blocking an axial attack from the *Si* face.

Next, we embarked on the investigation of the key RCM step.^[60] We initially tested the use of Grubbs second-generation catalyst in refluxing CH_2Cl_2 . These conditions led to the formation of a new product, which was identified as the homocoupled dimer **59** as an undetermined mixture of (*E*)- and (*Z*)-isomers by ¹H NMR and HSQC spectroscopy and ESI mass spectrometry. We assumed that this outcome might be attributed to a high activation barrier for the formation of the strained macrocycle and thus used higher boiling solvents (benzene, DCE, toluene). However, this did not provide any new reaction products. Since the use of the Stewart-Grubbs catalyst,^[61] particularly developed for the synthesis of sterically hindered alkenes by RCM, also resulted only in the formation of dimer **59**, we decided to turn our attention to other ring-forming reactions.

Toward construction of the macrocycle – the B-alkyl-Suzuki approach

In addition to the RCM approach, further strategies were pursued toward closing the elevenmembered macrocycle in tricycle **54** for the envisioned bioinspired cationic cascade. As outlined in Scheme 18, one strategy was based on an intramolecular B-alkyl Suzukicoupling.^[62] The retrosynthetic precursor **60** could arise *via* a Negishi carbalumination and addition of an alkenyl lithium species to a *trans*-hydrindanone prepared from silyl ether **47**.

Our route commenced with a two-step deprotection/oxidation sequence providing ketone **61** in 90% yield (Scheme 19). Next, the alkenyl lithium species derived from alkenyl iodide **62**^[63] was reacted with ketone **61** at -78 °C to afford allylic alcohol **63** as a single diastereomer. The stereochemical outcome of this reaction was verified by NOESY NMR experiments and analysis of the spectroscopic data revealed the preferred conformation of the stereogenic center at C10. Based on the lacking NOE correlation between the C10 methyl group and 9–H, both groups could potentially point into opposite directions. This assumption was further supported by the scalar coupling of the C9 methine (ddd at $\delta = 1.86$ ppm with J = 12.7, 3.7 and 2.3 Hz). While the first two values could be identified to an axial-axial coupling and an axial-equatorial coupling within the six-membered ring (to 8-H), the remaining coupling constant of J = 2.3 Hz accounts for the coupling of 10-H. Thus, this value indicates a dihedral angle of around 67° according to the Karplus curve. Consequently, this observation suggested that a rotation around the C9–C10 bond would be required for an efficient ring closure, constituting an important factor for the activation barrier.

Next, a three-step sequence consisting of oxidative PMB ether cleavage, Swern *oxidation* and homologation with the Ohira-Bestmann reagent (**64**)^[64] cleanly provided access to alkyne **65**. Notably, the oxidation of the intermediate primary alcohol with DMP only proved successful under elevated temperature and a large excess of the reagent. Since a direct Negishi carbalumination of alkyne **65** resulted in an intractable product mixture, we resorted to a Pd-catalyzed silylstannylation.^[65]

It has been shown that alcohol functionalities usually thwart this catalytic process. Thus, we aimed at protecting the tertiary alcohol.^[65b] However, silvlation attempts (TMSOTf, 2,6lutidine, CH₂Cl₂, 40 °C; Et₃SiCl, pyridine, 90 °C; BSTFA, DMF) failed. Consequentially, we assumed that the alcohol might be too burdened in its steric environment to affect the envisaged transformation. Indeed, exposing alkyne 65 to Bu₃SnSiMe₃ under Pd-catalysis gave rise to a single regioisomer 66. Subsequent iododestannylation (I₂, 2,6-di-tert-butyl-4methylpyridine, $-40 \,^{\circ}\text{C}$)^[66] and reaction with Me₂CuLi delivered alkenyl silane 67. In order to access macrocyclization precursor 60, we treated silane 67 with excess NIS (≈ 2.5 equivalents) in MeCN to affect an iododesilylation. Upon isolation of a new product, ¹H NMR spectroscopy revealed that the iododesilylation took place as indicated by a downfield shift of the alkenyl proton from $\delta = 5.20$ ppm to $\delta = 5.89$ ppm. However, the characteristic protons signals for the terminal alkene group disappeared. In addition, proton signals at $\delta =$ 4.53 and 3.61 ppm indicated that heteroatoms were attached to the respective carbon atoms. Thus, we tentatively assigned the new compound as a diastereomeric mixture of tricycle 68. While employing excess NIS in MeCN is usually required for the iododesilylation to secure retention of the double bond geometry,^[67] the use of hexafluoroisopropanol as solvent allows for stoichiometric quantities of NIS.^[68] This protocol resulted in decomposition of alkenyl silane 66 in our hands. We rationalized that a protection of the alcohol functionality might avoid the cyclization, but the examined conditions (KHMDS, Et₃SiCl; Ac₂O, pyridine, 90 °C) were not met with success. Due to material constraints, no further investigations were undertaken at this stage.^[69]

Second-generation approach

In light of our previous endeavors, we sought to establish a more convergent strategy for a potential biomimetic total synthesis of astellatol. Particularly, we opted at a route involving a 5-8-6-5 ring system having the correct relative configuration at C14 already in place and reducing the number of events in the key reaction cascade.

As outlined in Scheme 20, we envisaged diene **69** as key precursor for a biomimetic synthesis of astellatol. Upon treatment with acid, protonation could trigger the proposed homoallyl-cyclopropylcarbinyl-cyclobutyl rearrangement. Closure of the central eight-membered carbocycle should be achieved by a McMurry coupling and it was envisaged that the required diketone could arise from *trans*-hydrindane **53** and triflate **71** using B-alkyl-Suzuki coupling conditions. The latter fragment could in turn be prepared from 3-pentyn-1-ol (**70**) *via* an intramolecular 1,3-dipolar nitrile oxide cycloaddition and subsequent reductive ring opening. In addition, an auxiliary controlled alkylation could render our approach asymmetric.

In the forward direction, Myers alkylation^[70] of pseudo-ephedrinepropionamide (72) with iodide 73 resulted in clean formation of alkyne 74 (Scheme 21).^[71] Subsequently, hydrogenation of alkyne 74 in the presence of Pd/BaSO₄ in pyridine yielded the corresponding *cis* alkene.^[72] Other attempts mostly resulted in over reduction to the respective alkane. Cleavage of the chiral auxiliary with lithium amidotrihydroborate^[70] provided the respective alcohol and characterization of the corresponding Mosher ester^[73] revealed an enantiomeric excess of 98%. Oxidation to the corresponding volatile aldehyde under Swern conditions and transformation to oxime 75 proceeded smoothly and gave an inconsequential 5:2 E/Z mixture of isomers in good yield over the four steps.^[74] Among the various methods for the *in situ* generation of nitrile oxides and ensuing 1,3-dipolar cycloadditions,^[75] treatment of oxime **75** with aqueous NaOCl proved best and gave clean conversion to the 4,5-dihydroisoxazole as an inseparable 2:3 mixture of syn/anti isomers 76 and 77.^[76] Chemoselective hydrogenation to the corresponding β -keto alcohol by reaction with Raney® Ni in the presence of a boron-based buffer^[77] was followed by Swern oxidation to provide ketoenol 78. Notably, exposure of the intermediate alcohol to DMP or TPAP/NMO led to decomposition. Unfortunately, the ensuing conversion of intermediate 78 to triflate **71** by sequential treatment with NaH and PhNTf₂ suffered from a very poor yield.

The low yield of the last step was very surprising to us, since the conversion of commercially available diketone **79** to enol triflate **80**^[78] was high yielding under identical reaction conditions (Scheme 22).

Consequently, we first explored the key step of our proposed strategy with building block **80** and *trans*-hydrindane *ent*-**47**^[79] as model systems. Again, fragment coupling under B-alkyl-Suzuki conditions worked smoothly, delivering product **81** in quantitative yield. Subsequently, desilylation and oxidation yielded ketone **82**. With this material in hand, we were poised to investigate the envisaged McMurry coupling for the synthesis of key intermediate **69**.^[80,81] To this end, substrate **81** was slowly added to a suspension of low-valent titanium, prepared *in situ* from TiCl₃(THF)₃ and Zn–Cu couple in DME.^[82] Interestingly, these reaction conditions did not result in the formation of the desired eight-membered ring in tetracycle **69**, but pentacyclic product **83** was isolated as a single diastereomer. Despite extensive NMR spectroscopic studies, however, we could not unambiguously assign the configuration of the newly formed stereocenters. To the best of our knowledge, this cascade to form the complex 5-5-6-6-5-membered ring system is unprecedented in the literature.

Since reductive carbonyl-carbonyl coupling can also be achieved by means of diol formation, we turned our attention towards SmI_2 -mediated reactions.^[83,84] In agreement with the previous result, the formation of an eight-membered ring was not observed. Instead, the same cyclization mode occurred, affording hemiacetal **84** as a single diastereomer.

Mechanistically, we propose the following reaction pathway for this transformation (Scheme 23). Upon formation of the ketyl radical at the more electrophilic carbonyl in diketone **82**, the resulting intermediate **85** undergoes a Michael addition in a 6-*exo*-trig fashion.^[85] Subsequent one-electron reduction of the resulting acyl radical (not shown) with a second equivalent of SmI₂ forms dimetallic species **87**. Protonation and ketoenol tautomerization –

presumably during aqueous work-up – gives ketone **86** that immediately cyclizes to form hemiacetal **84**. Under the McMurry-coupling conditions, the latter process seems to occur in the reaction mixture and is followed by deoxygenation of the hemiacetal to form an oxonium ion. These steps might be assisted by the Lewis acidic nature and the oxophilicity of the generated titanium species. A final two-electron reduction step would then lead to the corresponding ether **83**. Presumably, the less Lewis acidic properties of SmI₂ combined with the milder reaction conditions account for the fact that in this case no final reductive deoxygenation was observed. A further screening of reaction conditions for a reductive carbonyl-carbonyl coupling proved less fruitful. In light of these findings and the fact that the low yielding enol triflate formation remained a bottleneck in our synthetic roadmap for building block **71**, we abandoned this strategy at this point.^[86]

Third-generation retrosynthesis

Although we had established the B-alkyl Suzuki coupling as a robust and viable C–C-bond formation reaction for our model hydrindane hydrindane **47**, we ultimately decided to pursue other retrosynthetic disconnections. As depicted in Scheme 24, we assumed that we could utilize allylic alcohol **88** to trigger a biomimetic cascade for the synthesis of astellatol. Upon ionization, a *syn* specific 1,2-hydride shift could initiate the homoallyl-cyclopropylcarbinyl-cyclobutyl rearrangement. Notably, the same intermediate **88** could also serve as a synthetic precursor for a total synthesis of nitidasin by sequential oxidations. It was envisaged that the central eight-membered ring of our advanced key intermediate **88** would be constructed by a RCM^[87] and a subsequent chemoselective hydrogenation of the less substituted double bond. Further retrosynthetic simplification would give two building blocks, alkenyl lithium species **89** and ketone **90**, that should be joined by a nucleophilic addition. As we had learned from our previous studies (*vide supra*), such an addition was most likely to be feasible and should result in the correct relative configuration at the tertiary alcohol required for a stereoselective total synthesis of nitidasin.

Synthesis of alkenyl iodide coupling partners

Our studies commenced with the synthesis of the precursor for fragment **89** and related alkenyl iodides. We were aware that the installation of the *cis* relationship of the two adjacent ring substituents would be challenging, therefore we aimed at exploiting substrate control once more and resorted to the chiral pool to implement an asymmetric route (Scheme 25). Accordingly, (–)-citronellene (**91**) was converted in a two-step oxidative protocol to a literature-known aldehyde.^[88] This was followed by a Corey-Fuchs homologation^[89] with quenching of the intermediate lithium species with MeI to furnish enyne **92**. The latter compound was very volatile complicating its isolation and accounted for the poor yield of the four-step sequence. Next, we employed a cyclometallation protocol developed by Negishi^[90] to establish the correct alkene geometry of alkenyl iodide **93**. In this reaction, exposure of enyne **92** to stoichiometric amounts of Cp₂Zr generated *in situ* gave a zirconacycle, which was quenched with excess NIS.^[91,92] The resulting inconsequential mixture of diastereomers **93** was briefly purified before eliminating the primary iodide by treatment with DBU. The so formed diene proved to be highly unstable and was immediately used in a diastereoselective hydroboration (9-BBN, 45 °C) furnishing

alcohol **95** as the major isomer in good yield over the three steps. At this stage, we prepared the corresponding Mosher ester and determined the enantiomeric excess of alcohol **95** to be 60%. This was due to the low optical purity of the commercially available (–)-citronellene (**91**).

In order to explore the envisioned fragment coupling, we next synthesized two building blocks. On the one hand, silylation of alcohol **95** afforded the corresponding silyl ether **94**. On the other hand, sequential Swern oxidation and Wittig olefination delivered diene **96** as a single isomer, indicating that no epimerization occurred at the aldehyde oxidation state. This was assumed to be due to the high 1,3-allylic strain an intermediate enolate would encounter. Overall, the route was straightforward and diene **96** was accessed in 16% yield over nine steps.

Model studies – fragment combination and RCM

Having prepared sufficient amounts of the two alkenyl iodide building blocks **94** and **96**, we were poised to investigate their coupling to the *trans*-hydrindanone **90**. In analogy to our earlier studies and in order to conserve our precious isopropyl substituted fragment **90**, these studies were first carried out with ketone **61**.

Since we assumed that the alkenyl lithium species of iodide **96** might be prone to undergo a 1,3-metallotropic rearrangement, we initially utilized silyl ether **94** in these explorations. To our delight, preparation of the alkenyl lithium compound and subsequent slow addition of ketone **61** resulted in a facile reaction and, after purification, we obtained adduct **97** as a single diastereomer in 48% yield (Scheme 26). This reactivity is particularly worth noting since tetrasubstituted alkenyl lithium species are rarely used in organic synthesis and addition to sterically encumbered ketones such as **61** are unprecedented in the literature.^[93] In order to elaborate the metathesis precursor, we next cleaved the silyl ether and oxidized the primary alcohol under Swern conditions. Interestingly, the formed aldehyde underwent a spontaneous intramolecular cyclization forming lactol **99** as a single diastereomer, the relative configuration of which was not further investigated. While direct olefination of this intermediate **99** proved unviable, this outcome was an indirect proof that the double bond geometry of the alkenyl iodide was retained upon lithiation and under the addition reaction conditions.

Encouraged by this result, we next prepared the alkenyl lithium species derived from iodide **96** at -78 °C. Upon addition of ketone **61**, the fragment coupling was again efficient and provided adduct **98** in 75% yield. It is worth noting that the obtained 10:1 mixture of diastereomers reflects a kinetic resolution of scalemic alkenyl lithium species **89** under the reaction conditions. In addition, we verified the relative configuration of all stereogenic centers and the alkene geometry by single crystal X-ray diffraction. The solid-state structure also revealed a close proximity between the two terminal alkenes, solely requiring a 120° rotation around the C9–C10 bond (astellatol numbering) for the envisaged metathesis. With this in mind, we were excited to explore the key RCM (Scheme 27).

Our efforts commenced by using the Grubbs first-generation catalyst in CH_2Cl_2 , resulting in slow consumption of the major diastereomer **98**, but no formation of the desired product

100. Interestingly, we observed traces of a cyclized product by ¹H NMR spectroscopy and EI mass spectrometry, which we tentatively assigned as tetracycle **102**.^[94] Similar results were obtained when employing the Grubbs second-generation catalyst in CH₂Cl₂ or DCE. When exchanging the solvent for benzene, quick consumption of the starting material **98** occurred and a new product was selectively formed. However, NMR spectroscopic analysis revealed that the tetrasubstituted double bond unexpectedly participated in the metathesis reaction yielding tricycle **101** in 60% yield. Chromatography employing 1% Et₃N in the eluent mixture was required, since otherwise an allylic transposition of the alcohol functionality to a diastereomeric mixture was observed. In analogy to our earlier studies, attempts to increase the sterical demand at the tetrasubstituted alkene in triene **98** by protecting the adjacent alcohol (*e.g.* Et₃SiCI; Et₃SiOTf; TMSOTf; TMSCI; MeI; Me₂SO₄; FSO₃Me) remained unfruitful.

Although our initial attempts to construct cyclooctadiene **100** *via* RCM had been hampered by the formation of tricycle **101**, we were intrigued by the seemingly smooth formation of the minor diastereomer **102**. We rationalized that a relay metathesis strategy (Scheme 28) could prevent initiation of the catalyst at the terminal alkene adjacent to C10 (astellatol numbering), thus possibly giving rise to the desired eight-membered ring.^[95,96]

Modifying the already developed Swern oxidation and Wittig olefination sequence, we were able to access triene **103** from alcohol **95**. Subsequent lithiation and addition to model system **61** furnished the desired metathesis precursor **104** in excellent yield. Again, a kinetic resolution of scalemic alkenyl iodide **103** took place, although in somewhat less of an extent than in the previous explorations. Despite the flexible alkene handle in intermediate **104**, single crystals suitable for X-ray diffraction were obtained. Next, we screened alkene metathesis reaction conditions for the reaction of tetraene **104**. When using the Grubbs first-or second-generation catalyst in benzene, tricycle **101** was formed accompanied with varying amounts of homo-dimer **105** and the previously observed diastereomer **102**. In contrast, the use of DCE as solvent only resulted in observation of the dimer **105**. In order to enhance the catalyst preference to initiate at the least substituted double bond, we next employed catalyst **106**^[97] in benzene, resulting in significantly increased amounts of dimer **105** (62% yield). However, only traces of dimer **105** were observed with this catalyst in DCE.

In view of these results, we decided to block the tetrasubstituted alkene for metathesis by epoxidation, additionally addressing the synthesis of nitidasin. Gratifyingly, exposure of triene **98** to *m*CPBA resulted in a diastereoselective reaction of the tetrasubstituted alkene, furnishing RCM substrate **107** with the correct epoxide geometry for nitidasin (Scheme 29). The structure of this intermediate **107** was again confirmed by single crystal X-ray diffraction and indicated a close proximity of the terminal alkenes necessary for the metathesis. Since we had observed unique reactivity of triene **98** using the Grubbs second-generation catalyst in benzene, we submitted diene **107** to the same reaction conditions. To our delight, the envisaged ring closure occurred smoothly and delivered tetracarbocycle **108** in 98% yield.^[98] Next, we aimed at completing the synthesis of the carbon framework of nitidasin. To this end, alkene **108** was hydrogenated in the presence of Pd/C and the

resulting saturated analogue **109** was fully characterized by NMR spectroscopy in benzene d_6 . In order to compare our model compound with the isolation data of nitidasin, we next employed CDCl₃ as the NMR solvent. Interestingly, this resulted in a rapid conversion of epoxide **109** to *exo*-methylene compound **110** that was attributed to traces of acid present in CDCl₃.

Total synthesis of nitidasin

After successfully establishing a fragment union and the subsequent RCM with model ketone **61**, we were excited to transfer these conditions to a system with the correct substitution pattern at the five-membered ring. To this end, chemoselective cleavage of the triethylsilyl ether in bicycle **53** was followed by oxidation of the resulting secondary alcohol to furnish ketone **90** (Scheme 30). This intermediate was then slowly added to the alkenyl lithium species derived from iodide **96** and *t*BuLi at -78 °C. In analogy to the model system, the reaction proceeded in a facile manner and with complete stereocontrol with respect to the newly formed stereogenic center. The isomers arising from scalemic reagent **89** were separable by column chromatography yielding 7% of the undesired adduct **112**. Interestingly, the desired isomer **111** decomposed upon complete evaporation of the solvents and was thus immediately subjected to the epoxidation conditions. The intermediate was isolated in 71% yield, indicating that again a kinetic resolution of scalemic iodide **96** took place. The ensuing RCM proceeded smoothly to deliver tetracarbocycle **113** in 66% yield for the three steps. At this stage, single crystal X-ray diffraction verified the relative configuration of all eleven stereogenic centers.

In order to complete the total synthesis of nitidasin, we identified the hydrogenation of the cyclooctene and the SEM deprotection as the major tasks. We initially investigated the hydrogenation of alkene **113** under the conditions employed for the model system (0.2 eq. Pd/C, H₂ (1 bar), MeOH). This resulted in comparably slow conversion and the prolonged reaction times caused a mixture of compounds to be formed that we attributed to degradation of product **114**. Thus, we screened further conditions (0.2 eq. Pd/C, H₂ (8, 20 or 30 bar), MeOH or EtOAc; PtO₂, H₂ (1 bar), EtOH; 5 mol-% Rh/Al₂O₃, H₂ (1 bar), EtOH; 10 mol-% Crabtree's catalyst, H₂ (1 bar), CH₂Cl₂), however, without achieving any satisfying result.

Finally, we decided to revisit our initial reaction conditions, but using superstoichiometric amounts of Pd/C in MeOH. This resulted in a fast and relatively clean conversion and delivered the saturated compound **114** in good yield. The structure of this intermediate was again confirmed by single crystal X-ray diffraction. However, the observed sensitivity of tetracycle **114** under the hydrogenation conditions combined with little amounts of material available for testing purposes prompted us to investigate the SEM deprotection at an earlier stage of our synthetic route.

Thus, we screened a variety of conditions on ketone **90** to identify a suitable protocol for this transformation. Among the reagents examined (*e.g.* HF/py; TFA; TASF; MgBr₂/MeNO₂; TBAF; HCl), HF/py proved to be the best (Scheme 31). Interestingly, the reaction did not lead to full conversion and was accompanied by the partial epimerization at C3 (nitidasin

numbering). The isomers were separable and the undesired ketone could be equilibrated by subjection to the reaction conditions together with the recovered starting material, increasing the overall yield. With intermediate **115** in hand, we reasoned that the previously established reaction sequence might also be feasible with the free alcohol functionality. To this end, the fragment combination was carried out with a sacrificial equivalent of the alkenyl lithium species **89** giving a 25:1 mixture of diastereomers **116** originating from scalemic alkenyl iodide **96**.

Subsequent epoxidation and RCM gave tetracarbocycle **117** in an excellent yield of 72% over three steps. The hydrogenation of the double bond under the previously established reaction conditions proceeded smoothly to deliver 'nitidasol', presumably the biosynthetic precursor of nitidasin. Finally, oxidation with TPAP/NMO^[99] furnished the natural product. These conditions had been established with alcohol **117**, in which case DMP had resulted in decomposition. The spectroscopic data of our product matched the isolation data.^[4] However, the use of rigorously acid-free CDCl₃ was required since otherwise, in accordance with our model studies, a conversion to *exo*-methylene compound **118** was observed. This transformation occurred in time frames ranging from minutes to days, depending on the batch of solvent employed. In addition to the correlating spectroscopic data, we verified the structure of our synthetic material by single crystal X-ray diffraction. Furthermore, comparison of the optical rotation of our sample with the authentic value – both being levorotatory – established that we synthesized natural nitidasin.

While this first synthetic route proved viable for a total synthesis of nitidasin, it used a sacrificing equivalent of the alkenyl iodide **96** in the fragment coupling compared to the route with the SEM ether in place. This prompted us to investigate the deprotection in tetracarbocycle **114**. First attempts, using TBAF in DMPU only led to slow decomposition. Interestingly, when using TBAF in THF in the presence of molecular sieves, loss of the TMS moiety was observed rather then complete cleavage of the SEM group. In contrast, the use of excess TASF in HMPA at 75 °C^[100] resulted in clean deprotection along with recovered starting material (Scheme 32). The use of molecular sieves proved crucial for the success of this transformation. The so obtained alcohol **117** was then elaborated to nitidasin as described previously. Overall, 16 mg of the natural product have been prepared *via* the two routes.

Progress toward a biomimetic synthesis of astellatol

As outlined in our third-generation retrosynthetic analysis (*cf.* Scheme 24), we also aimed at accessing allylic alcohol **88** as a precursor for the proposed biomimetic cationic cascade toward the construction of the astellatol ring system. Since our efforts to obtain alkene **88** *via* a cyclooctadiene capitalizing from a RCM proved unsuccessful in model studies (*vide supra*), we changed our strategy and were drawn to the transformation of epoxides into the respective alkenes by the use of a low valent tungsten species generated *in situ* from WCl₆ and *n*BuLi.^[101] In the event, epoxide **114** was cleanly converted to allylic alcohol **88** in excellent yield (Scheme 33).^[102] At room temperature, signal broadening originating from the rotameric nature of cyclooctene **88** thwarted the structure analysis by ¹³C NMR spectroscopy, but performing the studies at 40 °C allowed for clean characterization. In

addition, we resorted to single crystal X-ray diffraction to prove the identity of compound **88**.

We were then in the position to examine the proposed cationic cascade for the synthesis of astellatol. In order to trigger the initial ionization step to transform allylic alcohol 88 into carbocation 119, we investigated the use of Brønsted acids (TsOH; CSA; HCl; HF; TFA; HClO₄) in a variety of solvent systems of different polarities (CDCl₃; CH₂Cl₂; DCE; toluene; CH₃CN; THF/H₂O; 1,4-dioxane; DMSO/H₂O; CH₃NO₂; TFE; MeOH; acetone, pyridine) at temperatures ranging from room temperature up to 110 °C. Unfortunately, the only observation repeatedly made was the formation of several kinds of elimination products that were occasionally accompanied by SEM group removal. The characteristic signals of the *exo*-methylene group of astellatol were never observed by ¹H NMR analysis of the reaction mixtures. Furthermore, the addition of perrhenate species such as Re₂O₇, which are known to catalyze the transposition of allylic alcohols,^[103] could not avoid undesired elimination reactions. The same held true when Lewis acids (e.g. BF₃·OEt₂, Ti(OiPr)₄, TiCl₄, Cp₂ZrCl₂, AlCl₃) were employed instead. Presumably, the required initial 1,2hydride shift to form intermediate 8 exhibits an energetic barrier too high to compete with elimination processes. Accordingly, future work will be directed to the synthesis of the corresponding homoallylic alcohol of cation 8.

Conclusions

In summary, we have presented the evolution of our campaign for the total synthesis of isopropyl *trans*-hydrindane sesterterpenoids. Starting from diketone 4, we have developed a robust and scalable synthetic route to several versatile building blocks that was based on a series of substrate-controlled and diastereoselective transformations. Since some of the selectivities were surprising at first glance, we investigated their origin by DFT calculations and established reasonable reaction pathways. The initial efforts of our unified approach aimed at a biomimetic synthesis of astellatol. However, attempts to close a strained elevenmembered ring remained unsuccessful and prompted us to pursue alternative routes by preparing a 5-8-6-5 tetracarbocyclic backbone in a convergent fashion. In our first strategy to close the central eight-membered ring by a reductive coupling of two carbonyl functionalities, we identified the intriguing yet undesired pentacyclic products 83 and 84, which from a cascade that sets four stereogenic centers in a single step. Our second approach involved the addition of a complex alkenyl lithium species to a transhydrindanone, a chemo- and stereoselective epoxidation and an olefin-metathesis to construct a 5-6-8-5 tetracyclic carbon framework. While our attemots to access astellatol based on this intermediate remained unfruitful, we successfully synthesized (-)-nitidasin and established its absolute configuration. Our stereoselective route proceeded in a longest linear sequence of 34 steps and delivered the natural product in 1.9% overall yield. A secondgeneration synthesis of key ketone 90 and studies toward the synthesis of astellatol and the YW compounds are currently under investigation in our laboratories and will be reported in due course.

Experimental Section

Experimental, computational and crystallographic details as well as compound characterization data and copies of ¹H and ¹³C NMR spectra are available in the Supporting Information for this article on the WWW under http://dx.doi.org/10.1002/chem. 2015XXXXXX.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgements

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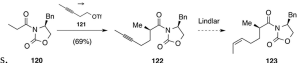
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epimerization of the selectively installed stereogenic center took place in the pursued route toward

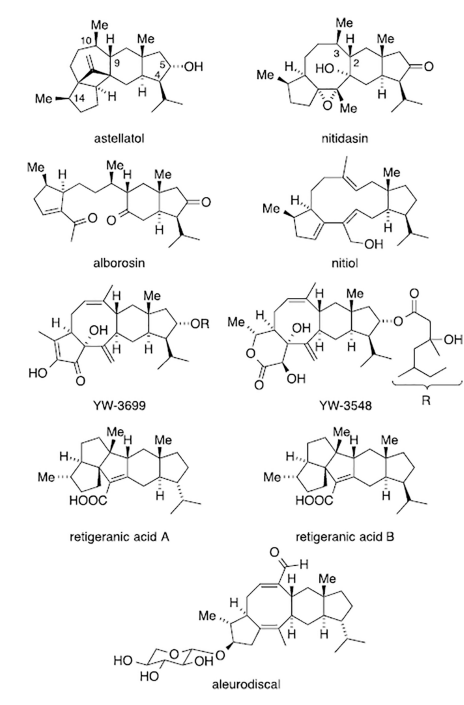


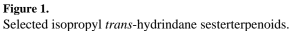
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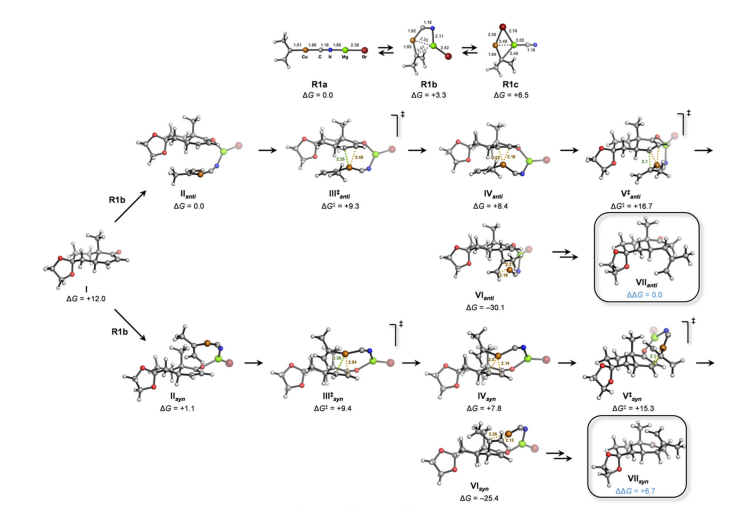


Figure 2.

Mechanism of the conjugate addition of organocuprate [Cu(isopropylene)CN·MgBr] to bicyclic cyclopentenone **16** calculated at the SMD_{THF}/ ω B97X-D/6-311+G(2d,p) +SDD(Cu,Mg,Br)// ω B97X-D/6-31G(d)+SDD(Cu,Mg,Br) level. Free energies are given in kcal mol⁻¹ and distances in angstroms. Different structures were considered for reactant (**R1**). Explicit solvation-desolvation of the metal atoms was not considered. Therefore, both the geometies of **R1** and the initial complexation energies should be considered only qualitatively.

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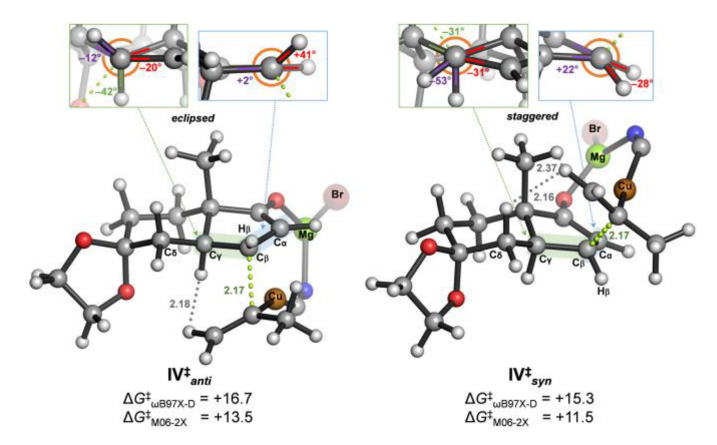
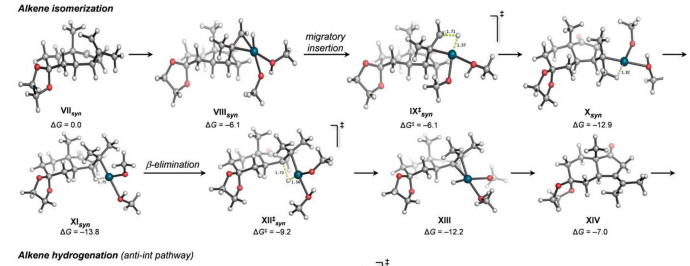


Figure 3.

Lowest energy transition state structures for the rate-limiting and stereodetermining addition step for the addition of isopropylene cyanocuprate to bicyclic cyclopentenone **16**, calculated at the SMD_{THF}/DFT/6-311+G(2d,p)+SDD(Cu,Mg,Br)// ω B97X-D/6-31G(d) +SDD(Cu,Mg,Br) [DFT= ω B97X-D and M06-2X] levels. Free energies are given in kcal mol⁻¹ and distances in angstroms.



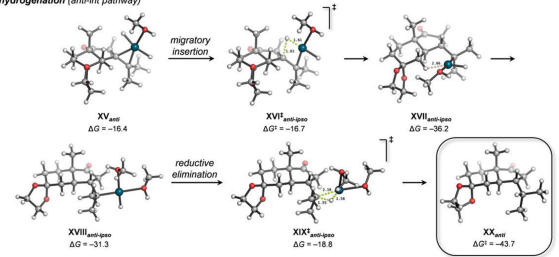
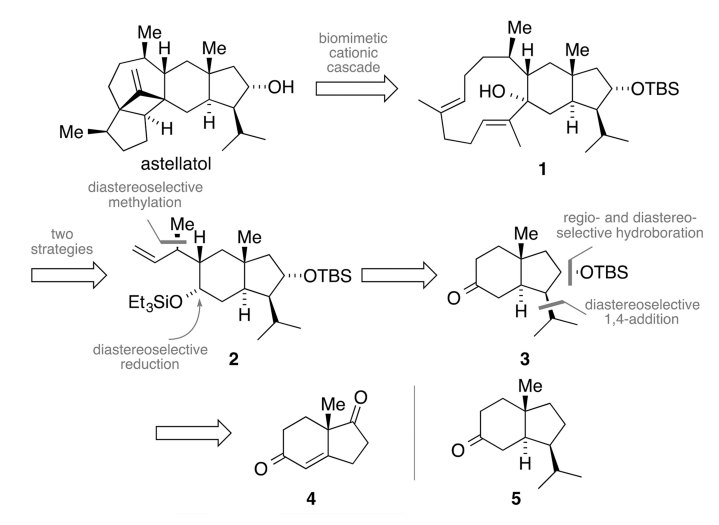


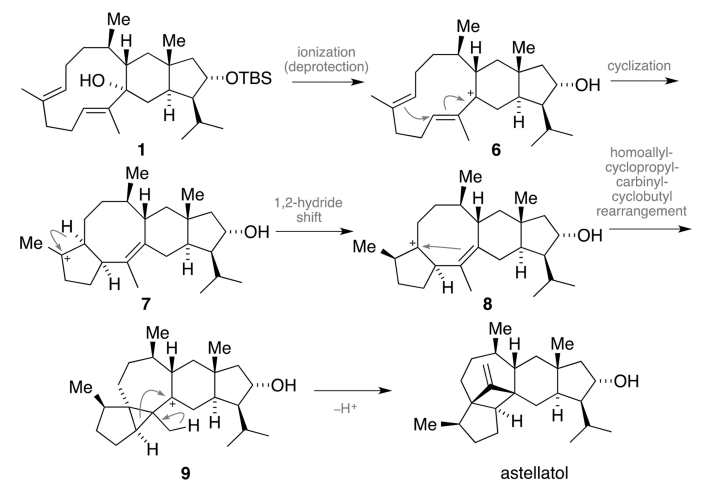
Figure 4.

Minimum energy pathway for the epimerizing hydrogenation of ketone **17** involving an initial alkene isomerization, calculated at the SMD_{MeOH}/ ω B97X-D/6-311+G(2d,p) +SDD(Pd)// ω B97X-D/6-31G(d)+SDD(Pd) level. Free energies are given in kcal mol⁻¹ and distances in angstroms.



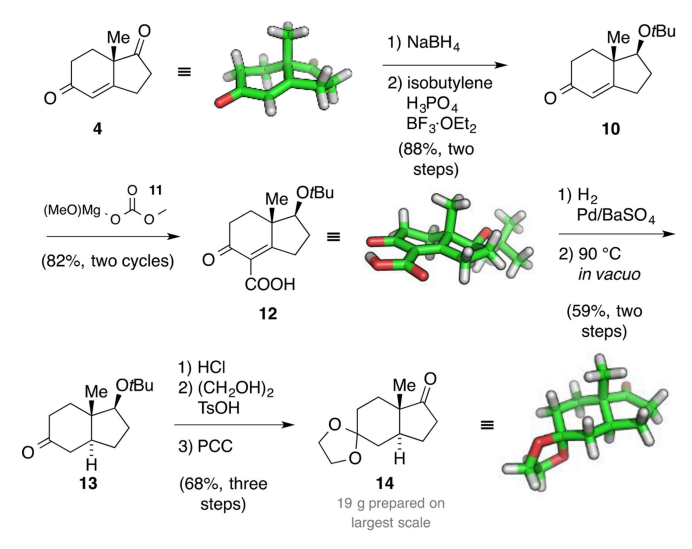
Scheme 1. Retrosynthesis of astellatol based on a biomimetic cascade.

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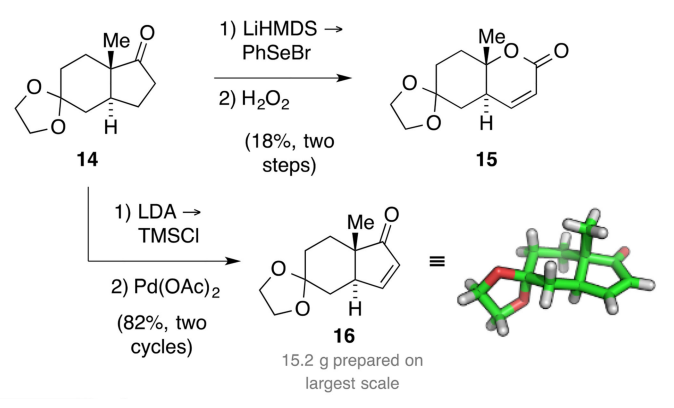


Scheme 2. Proposed biomimetic cationic cascade toward astellatol.

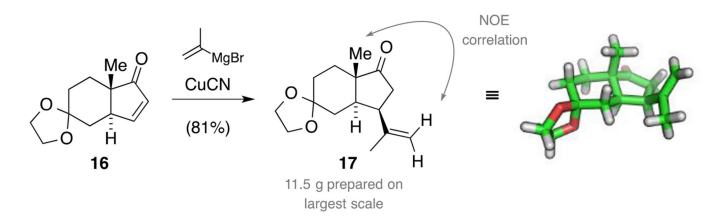
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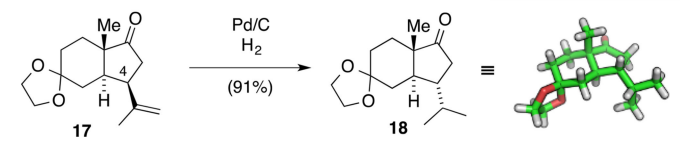
Scheme 3. Synthesis of *trans*-hydrindanone 14.



Scheme 4. Synthesis of enone 16 and lactone 15.

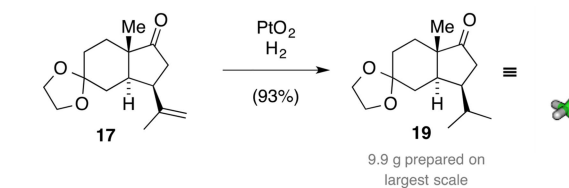


Scheme 5. Synthesis of ketone 17 by a diastereoselective 1,4-addition.



Scheme 6. An unexpected epimerization to form ketone 18.

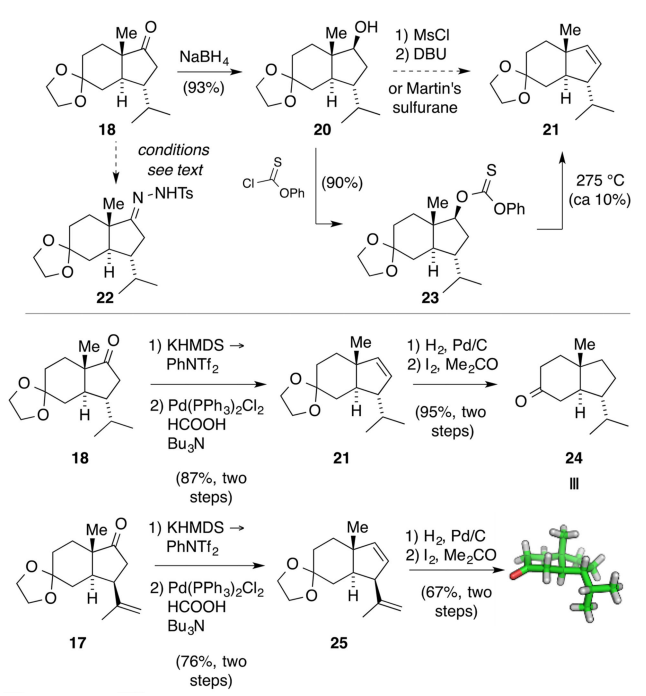
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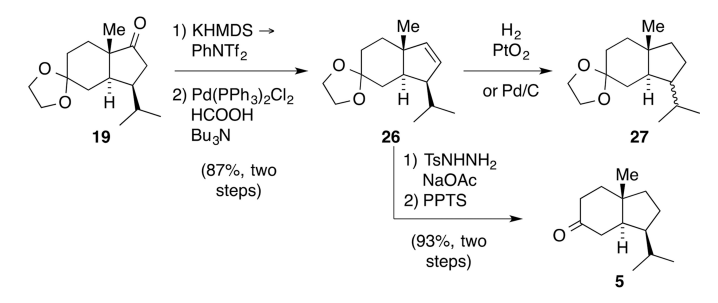
Scheme 7. Synthesis of ketone 19.

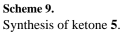
Chemistry. Author manuscript; available in PMC 2016 September 21.

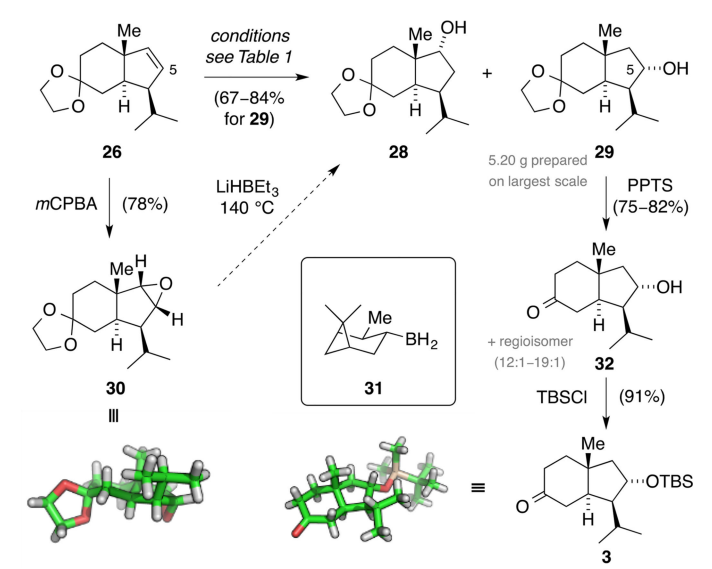
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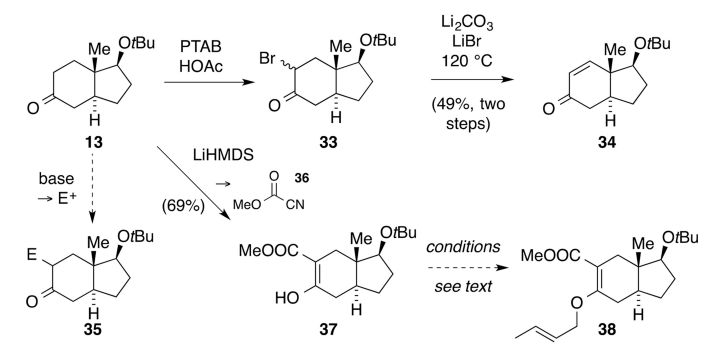
Scheme 8. Initial attempts to prepare alkene 21 (top) and synthesis of ketone 24 (bottom).





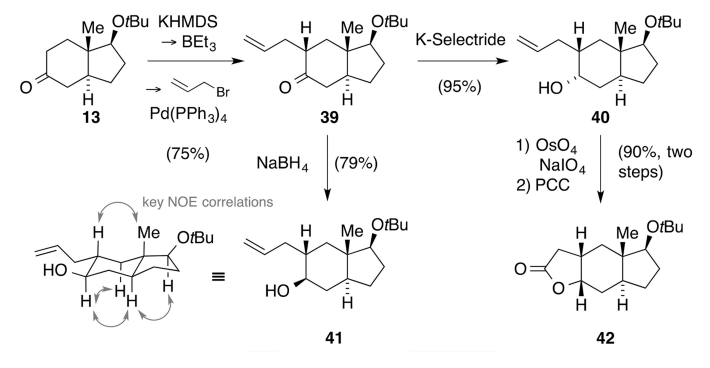


Scheme 10. Synthesis of ketone 3 and epoxide 30.

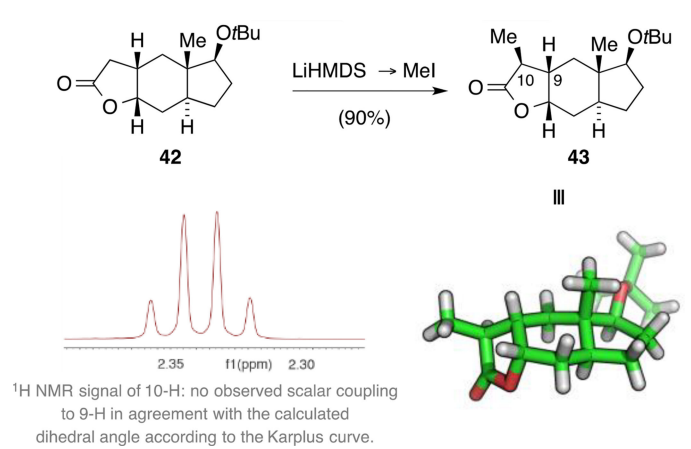


Scheme 11.

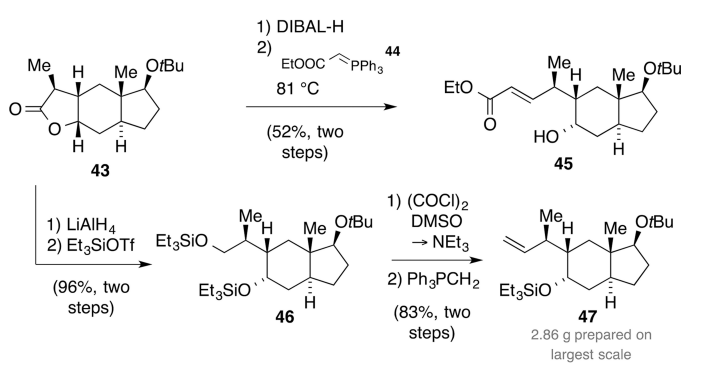
Proof of regioselective reactivity of ketone 13: synthesis of enone 34 and attempts toward Claisen rearrangement precursor 38.



Scheme 12. Preparation of lactone 42.

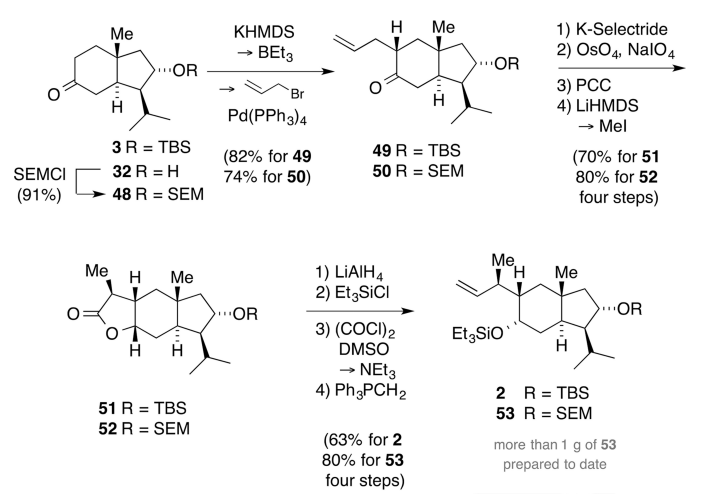


Scheme 13. Diastereoselective synthesis of lactone 43.

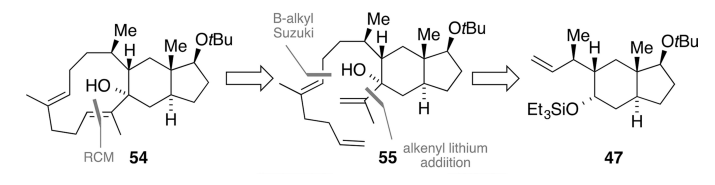


Scheme 14. Synthesis of ester 45 and of alkene 47.

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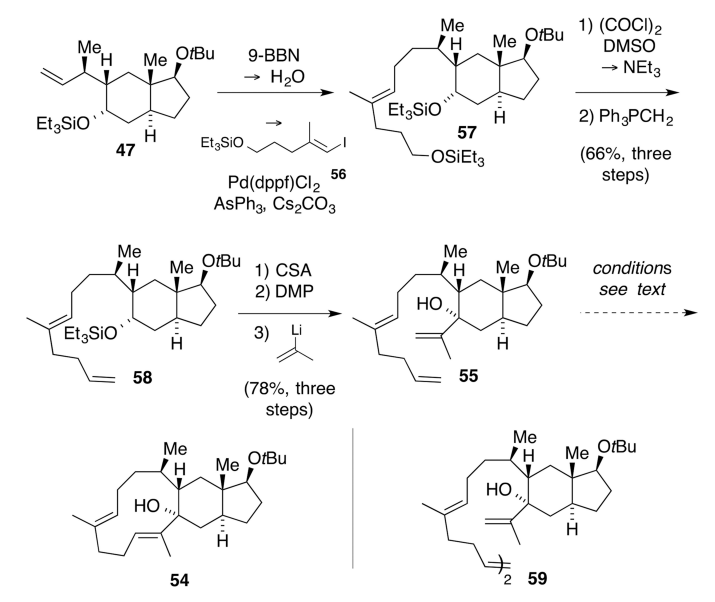


Scheme 15. Synthesis of alkenes 2 and 53.

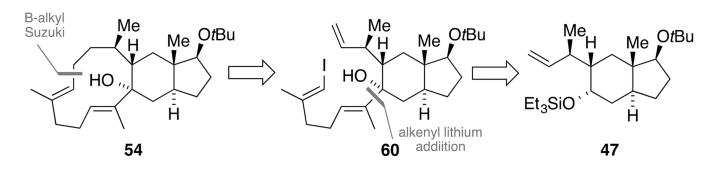


Scheme 16. Retrosynthesis of macrocycle 54 involving a RCM.

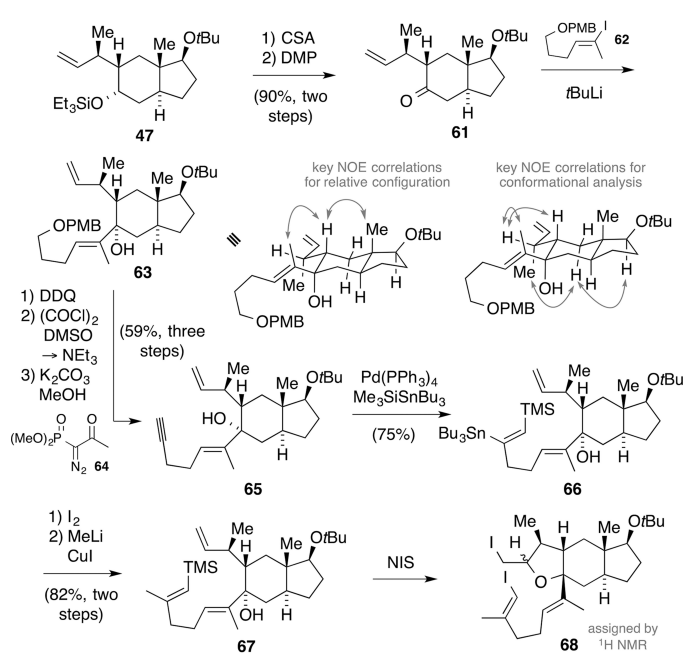
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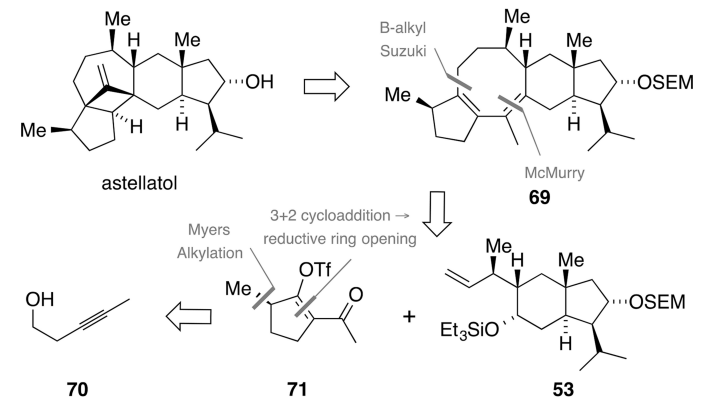
Scheme 17. Synthesis of triene 55 and observed dimer 59.



Scheme 18. Retrosynthesis of macrocycle 54 with a B-alkyl Suzuki coupling.

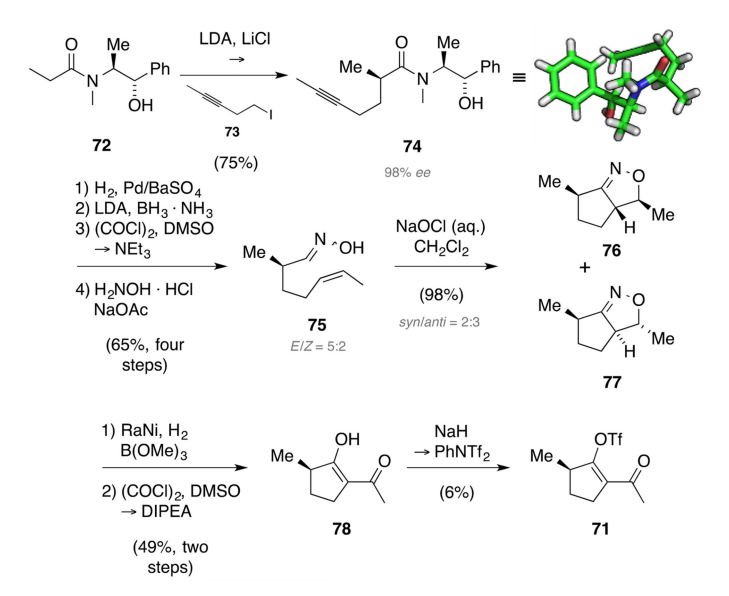


Scheme 19. Synthesis of alkenyl silane 67 and observed iodoetherification.



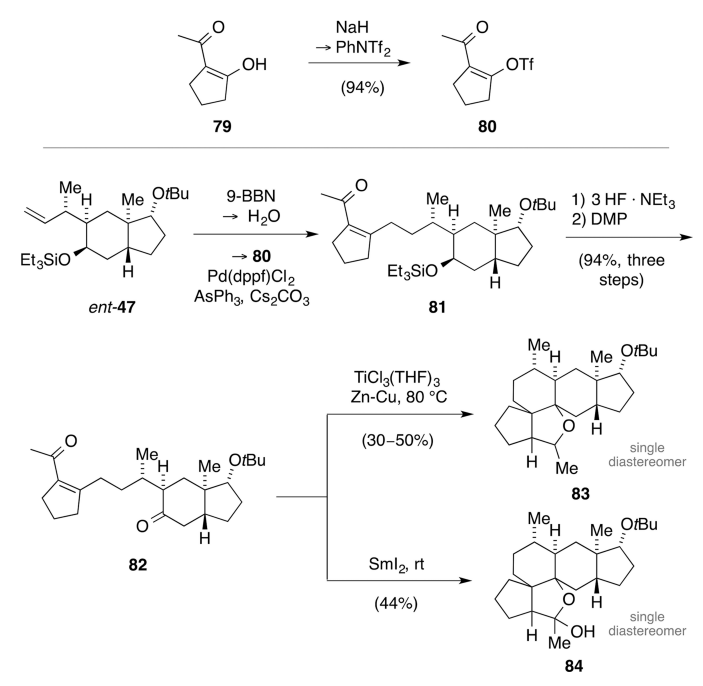
Scheme 20.

Second generation retrosynthetic analysis of astellatol: diene **69** as precursor for the biomimetic cationic cascade.



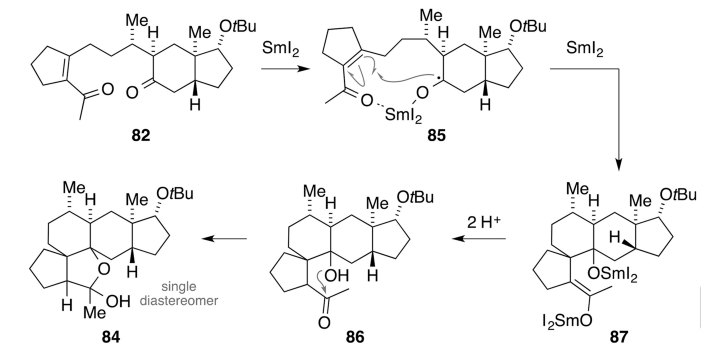
Scheme 21.

Enantioselective synthesis of enol triflate 71.



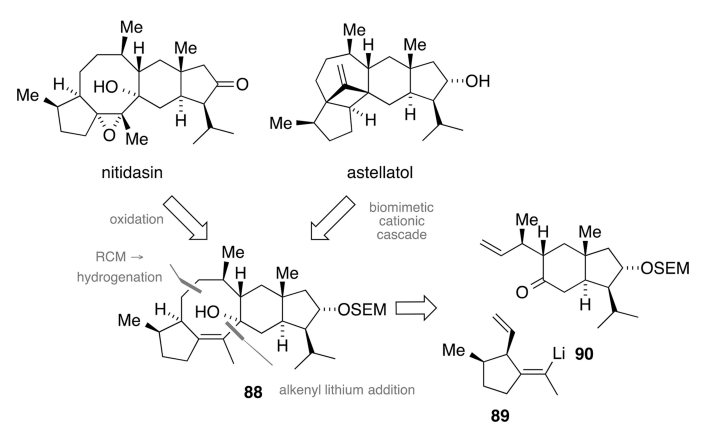
Scheme 22.

Synthesis of enol triflate **80** (top) and formation of ether **83** and hemiacetal **84** under reductive carbonyl coupling conditions (bottom).



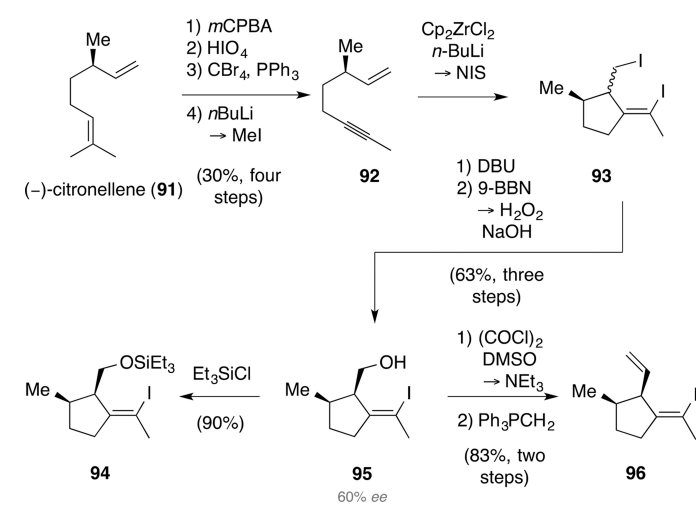
Scheme 23.

Proposed mechanism for the formation of pentacycle **84** under reductive carbonyl coupling conditions.

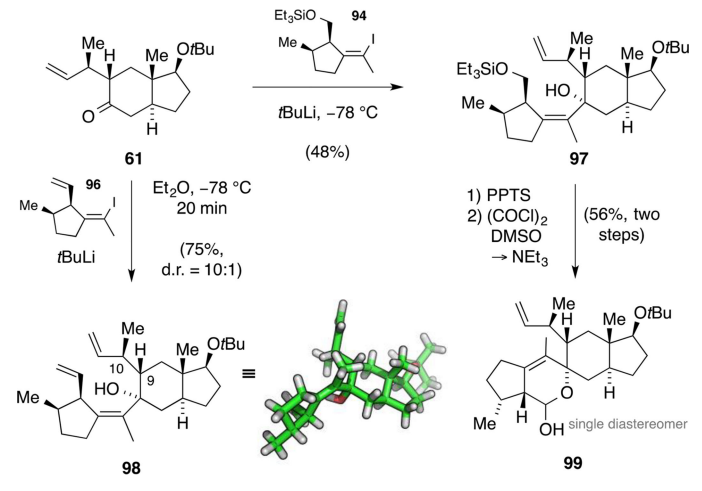


Scheme 24.

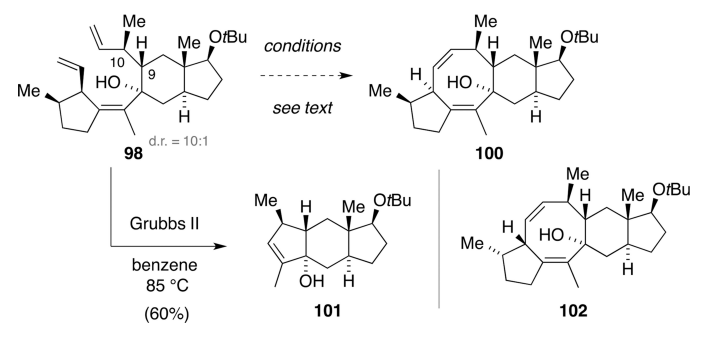
Third-generation retrosynthesis: convergent access to tetracycle **88** as precursor for astellatol and nitidasin.

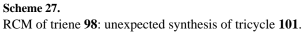


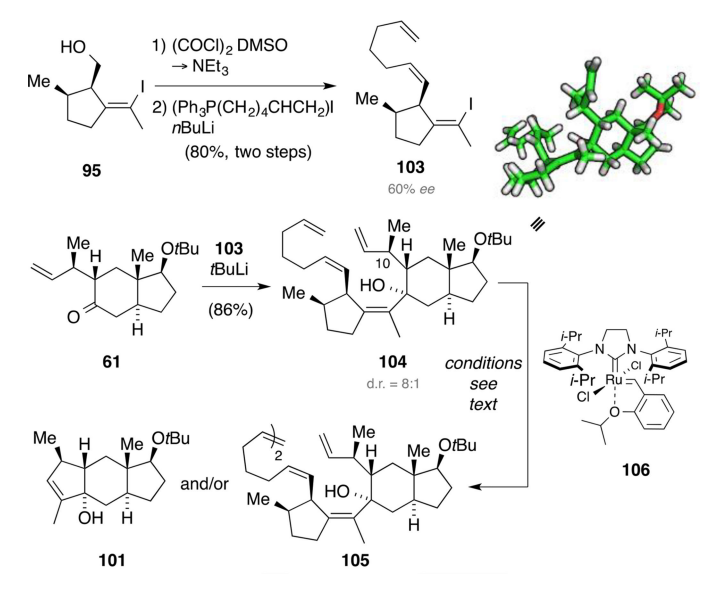
Scheme 25. Diastereoselective synthesis of alkenyl iodides 94, 95 and 95.



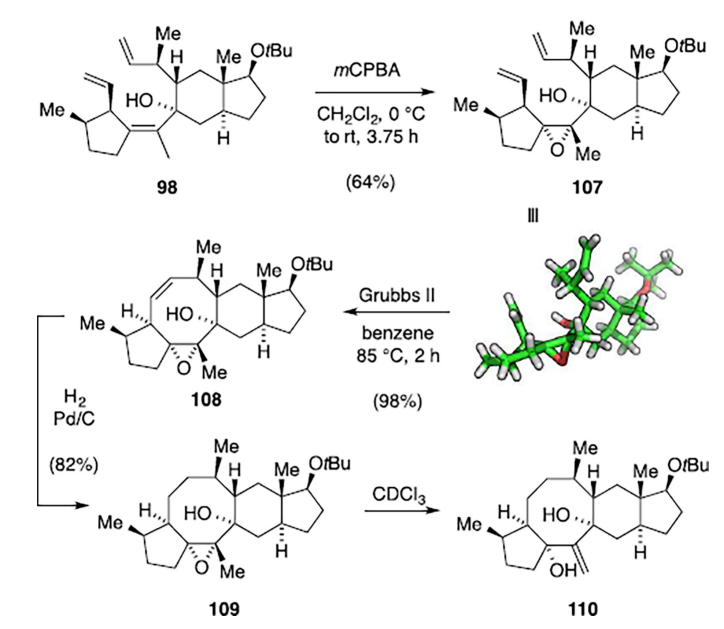
Scheme 26. Model fragment coupling: synthesis of triene 98 and lactol 99.

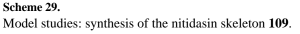


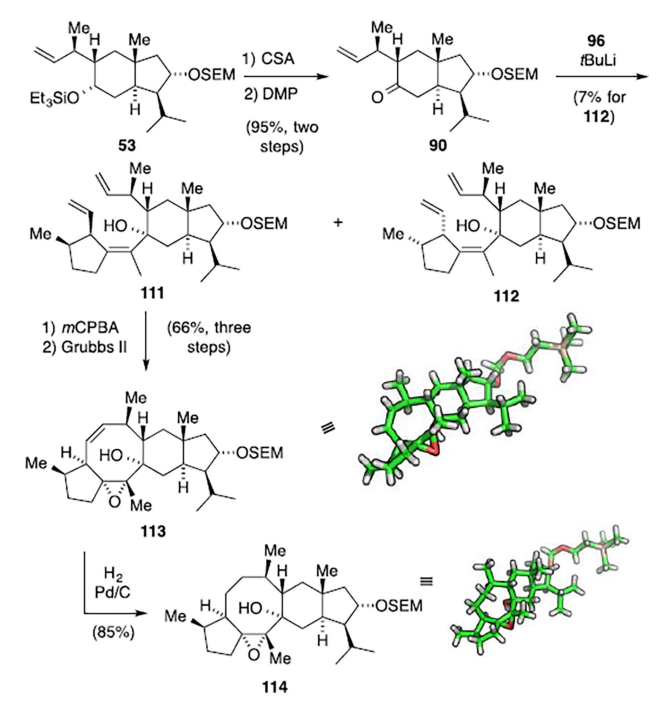




Scheme 28. Relay RCM of tetraene 104.

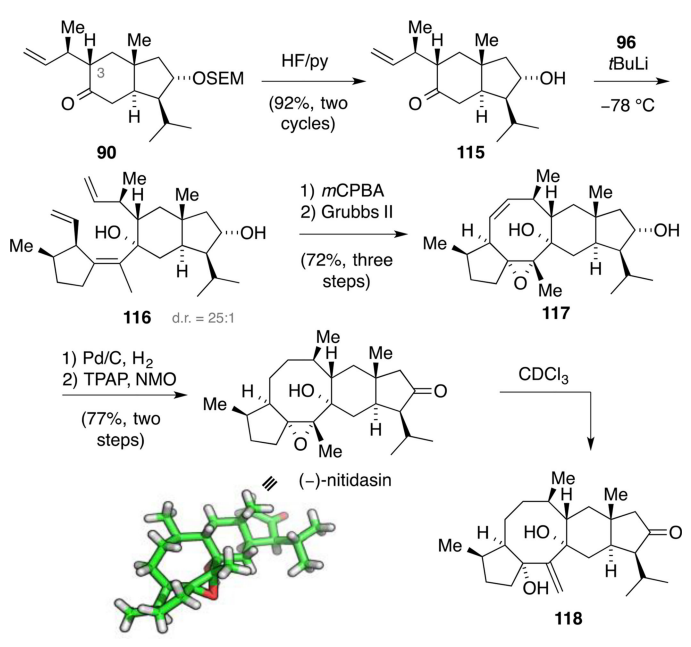




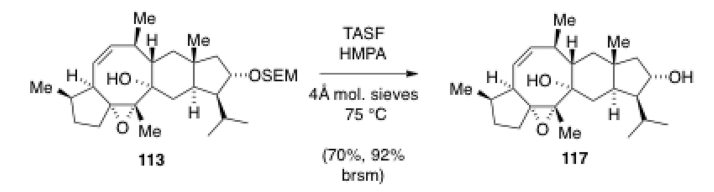


Scheme 30. Construction of the nitidasin backbone 114.

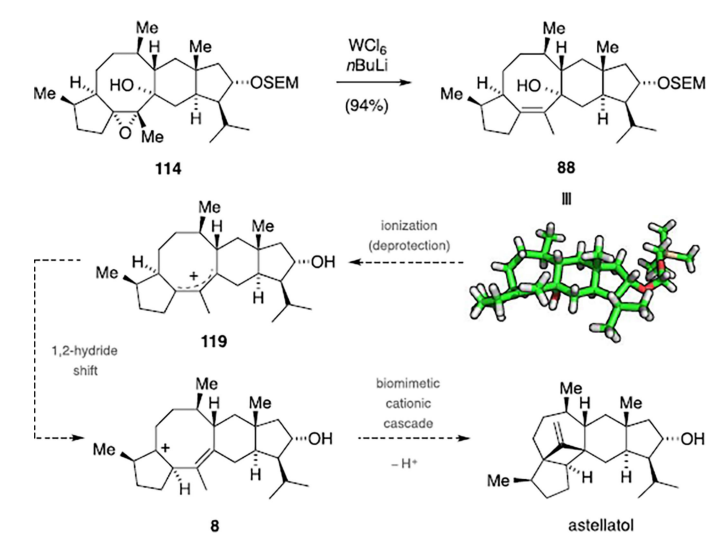
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Scheme 31. Total synthesis of (–)-nitidasin.



Scheme 32. Late stage SEM ether cleavage using substrate 113.



Scheme 33.

Deoxygenation of epoxide **114** and proposed pathway for the investigated biomimetic cationic cascade toward astellatol.

Table 1

Hydroboration of alkene 26: variation of the reaction conditions.^[a]

Entry	borane (equiv.)	T [°C]	t [h]	29:28 ^[b]
1	BH ₃ ·SMe ₂ (1.3)	0 to rt	0.5	2.7:1
2	BH ₃ ·SMe ₂ (1.3)	rt	0.5	2.6:1
3	BH ₃ ·SMe ₂ (1.6)	0 to rt	0.5	2.1:1
4	BH ₃ ·SMe ₂ (0.7)	0 to rt	0.5	2.9:1
5	9-BBN (5.0)	0 to 66	15	_[c]
6[d]	ThxBH ₂ (1.6)	0 to rt	1	_[e]
$\gamma[d]$	(sia) ₂ BH (2.5)	0 to rt	1	_[e]
₈ [d]	(+)-IpcBH ₂ (1.5)	0 to rt	0.5	3.7:1[f]
9[d]	(–)-IpcBH ₂ (1.5)	0 to rt	0.5	1:1

[a] All reactions were carried out on 10 mg scale in THF (c 0.03M) unless otherwise stated. All reactions were followed by oxidative, basic workup (NaOH, H2O2).

 $^{[b]}$ As determined by ¹H NMR spectroscopy of the crude reaction mixture.

[c] No conversion was observed.

[d] The hydroboration agent was prepared immediately prior to use.

[*e*] The reaction condition resulted in acetal cleavage

[f] The ratio was enhanced to 5.5:1 (84% yield) to 12:1 (67% yield) by flash column chromatography. rt = room temperature.