

An Alkyne Hydrosilylation-Oxidation Strategy for the Selective Installation of Oxygen Functionality

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Abstract: Alkynes bearing propargylic, homopropargylic, and bishomopropargylic hydroxyl groups are shown to serve as precursors for ketone or α -hydroxy ketone functionality. The approach hinges on the intermediacy of vinylsilanes created through regioselective hydrosilylation catalyzed by the complex [Cp*Ru(MeCN)₃]-PF₆. Several oxidative pathways of linear and cyclic vinylsilanes are studied, and the possibility of diastereoselective epoxidation of cyclic vinylsilanes is demonstrated. The sequences constitute the equivalent of stereoselective aldol, homo-aldol, and bishomo-aldol type processes. The method is applied to a short synthesis of the piperidine alkaloid, spectaline.

Introduction

The controlled introduction of heteroatom functional groups is part and parcel of complex molecule synthesis. Introduction of oxygenation at the ketone or hydroxyl oxidation state from hydrocarbon "feedstock" alkanes, alkenes, and alkynes is a useful way to introduce complex functionality. The use of C-C unsaturation for the installation of oxygenation can reduce the reliance on protecting-group manipulation, thereby improving synthetic efficiency. The selective oxygenation of alkenes, processes such as dihydroxylation, hydration (typically as hydroboration-oxidation), and Wacker-type oxidation, is a successful and pervasive method; the development of alkyne oxygenation strategies, on the other hand, lags far behind, despite the potential for increased flexibility and the potential to access different oxidation states at either alkyne carbon.

In principle, alkynes could serve as the basis for the selective syntheses of 1,2-diols, 1,2-dicarbonyl compounds, or α -hydroxy carbonyl compounds, in addition to the simple ketone or aldehyde hydration products (Scheme 1). This flexibility makes alkynes attractive building blocks, and terminal alkynes have been used to afford carbonyl products available by direct hydration¹ or through the intermediacy of vinylmetal species. In contrast, the selective introduction of oxygenation to internal alkynes is not well developed.^{2b} Hydroboration can be used to provide ketones after oxidation with good control of regiochemistry based on steric differentiation,² and recently the direct hydration of propargylic alcohols has been demonstrated with a gold catalyst.³ In addition, intramolecular diol-alkyne bisScheme 1. Alkyne Building Blocks for Highly Oxygenated Targets



cyclization catalyzed by palladium can be used in the selective synthesis of spiroketals.⁴ Despite these achievements, the use of alkynes as substrates for oxygenation in the context of functionalized molecules remains underutilized. Of additional synthetic benefit, alkyne oxygenation methods may well be orthogonal to known olefin oxidation pathways, allowing multiple oxygen-containing functional groups to be incorporated at a late stage in a controlled manner for facile synthetic design.

Recently, we disclosed preliminary reports of regio- and stereoselective hydrosilylation of internal alkynes, catalyzed by the ruthenium complex $[Cp*Ru(MeCN)_3]PF_6$ (1), providing unique vinylsilanes previously available only by circuitous means (Scheme 2).5-8 Of special interest to this work, propar-

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Scheme 2. Hydrosilylation Directed by Hydroxy Groups^a



 a BDMS = benzyldimethylsilyl; TDMS = 1,1,3,3-tetramethyldisilazane [(HMe₂Si)₂NH)].

gylic,⁷ homopropargylic,⁵ and bishomopropargylic⁵ alcohols all allow selective access to the (*E*)-vinylsilane with distal regioselectivity relative to the hydroxyl group by either intramolecular or intermolecular processes, as appropriate (Scheme 2). The wealth of oxidation chemistry available from vinylsilane intermediates led us to consider the use of alkyne hydrosilylation together with vinylsilane oxidation^{9–12} to introduce functionality distal to a hydroxyl group, which could control regio- and stereochemistry.⁷ In this paper, we provide a full account of these regioselective hydrosilylations and the use of these adducts as an equivalent of an aldol, homo-aldol, and bishomo-aldol process.

Propargylic Alcohols

Propargylic alcohols are surrogates for β -hydroxy ketones provided that a chemo- and regioselective introduction of a carbonyl group at the alkyne carbon β to the alcohol can be performed. Homopropargylic alcohols have been previously utilized for such a purpose via a platinum-catalyzed intramolecular hydrosilylation of homopropargylic alkynes for the selective introduction of a carbonyl group.^{9,11} The asymmetric access to propargylic alcohols, most notably by enantioselective hydrogenation of a ketone¹³ or addition of a zinc acetylide to an aldehyde,¹⁴ makes this approach attractive. Although the direct hydration of a propargylic alcohol is a potentially attractive solution, obtaining regioselectivity has proven difficult,³ and elimination of the hydroxy ketone product under generally acidic hydration conditions is a concern.

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Scheme 3. Hydrosilylation of Propargylic Alcohols



At the onset of our work, the regioselective functionalization of propargylic alcohols by hydrosilylation in a general sense was a challenging problem.¹⁵ We demonstrated efficient intermolecular hydrosilylation with reasonable regio- and stereoselectivity of a variety of propargylic alcohols catalyzed by complex **1**, providing the (*Z*)- β -vinylsilane as the major product.⁷ Recently, a regiocomplementary process has been demonstrated using an extended linker for intramolecular hydrosilylation of 3-octyn-1-ol.¹⁶

We chose to pursue selective incorporation of a vinyl C-Si bond to serve as a versatile ketone precursor. Implicit in the approach is a need for a silane oxidation under conditions mild enough to prevent β -hydroxy elimination and destruction of sensitive functionality. The most obvious silane targets, then, are alkoxy silanes. Although these do function in the hydrosilylation reaction, the selectivity and product stability are compromised with propargylic alcohol substrates (Scheme 3, **9-\beta** and **9-\alpha**). These problems were both solved upon moving to a trialkylsilane reagent (i.e., formation of 10). However, trialkylsilanes are not readily oxidized. Dimethylphenylsilyl is the most studied silyl group for use where robust stability is required,¹⁷ but oxidation of the dimethylphenylsilyl group can be incompatible with sensitive functionality, especially acidsensitive groups. Our search for a balance between stability and ease of oxidation led us to employ benzyldimethylsilane (BDMS-H), an especially attractive silane useful in a number of contexts.^{6,7,18} We have retained a BDMS group through multiple synthetic operations, including those involving organolithium reagents, aqueous acid and base, buffered fluoride necessary for silvl ether deprotection, and strong hydride reductants (i.e., Red-Al). Activation of the BDMS group occurs through loss of toluene and formation of a silvl fluoride or silanol species upon treatment with unbuffered TBAF. Typically addition of 1 equiv of TBAF produces complete conversion in

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5 min at 0 °C in THF or DMF. Subsequent addition of oxidant (aqueous H_2O_2 , urea-hydrogen peroxide adduct, or *m*-CPBA) furnishes the oxidized product. Generally, use of KHCO₃ and H₂O₂ in methanol was preferred.^{18,19} As shown in Scheme 4, the compatibility of the conditions for debenzylation and Tamao-Fleming oxidation with hydrosilylation allows the process to be performed as a one-pot operation. While the proposed intermediates 12 and 13 have not been isolated, their involvement seems probable. It is interesting to note that no dehydration products are observed in this sequence. The involvement of a cyclic silylene 13 wherein such an elimination is geometrically disfavored may account for this behavior. In earlier work using other catalysts to effect intramolecular hydrosilylation of homopropargylic alcohols followed by oxidation to form β -hydroxy ketones, elimination from the initial enol is not possible.9

An additional advantage of this methodology is the ability to further elaborate the vinylsilane 11 prior to unmasking the ketone. As aldol adducts are prone to further reactions such as eliminations, further substitution at the α -carbon becomes difficult.²⁰ Therefore, an important feature of this approach is the ability to introduce additional complexity at the α -carbon. For example, epoxidation occurs with complete diastereoselectivity with *m*-CPBA to provide hydroxy epoxide 15. Others demonstrated the importance of the silvl substituent for good diastereoselectivity.²¹ Thus, as shown in eq 1, path a, this sequence serves as an efficient route to the epoxy alcohols 16 diastereoselectively. On the other hand, treatment of 15 with 1 equiv of TBAF in THF followed by the standard Tamao-Fleming oxidation¹⁰ provides the syn diol **17** (eq 1, path b). The amount of TBAF in the initial stage of the oxidative cleavage is crucial because significant protodesilylation occurs if excess is employed. Purification of the epoxide prior to the Tamao-Fleming oxidation minimized this issue. In this way, the added alkyne unit functions as a hydroxymethyl ketone equivalent with the hydroxy and ketone groups being unmasked simultaneously.



For this aldol surrogate process, the stereochemistry of the β -hydroxy group derives from the acetylide addition. To the extent that can be controlled in terms of both relative and absolute stereochemistry, the utility of this process is much enhanced. We were interested in finding methods to allow both Felkin-Ahn and chelate-controlled alkyne addition and hence





aldol adducts. Diastereoselective addition of acetylide anions to aldehydes has been a challenging problem due to their small size and high reactivity.²² The Felkin-Ahn process has been achieved using a tris-alkoxytitanium acetylide, which presumably acts as a very sterically demanding nucleophile which is coordinatively saturated and thus leaves open no possibility for chelation.²³ Indeed, addition of a titanium acetylide complex converts aldehyde 18 into the anti adduct 19 with a diastereomeric ratio of 9:1 (eq 2). Chelation control, on the other hand, has been achieved with an alkynylzinc species.²⁴ Employing the published conditions, we were able to demonstrate that chelation-controlled addition to the α -alkoxy aldehyde 21 using zinc bromide provides the syn adduct 22 (10:1 dr) (eq 3). In both cases, one-pot hydrosilylation followed by oxidation unmasks the aldol adducts. The final β -hydroxy ketones 20 and 23 are isolated as single diastereomers.



As previously, the intermediate vinyl silane 24 undergoes clean diastereoselective epoxidation (eq 4), and the resultant silvlated epoxide 25 provides the epoxy alcohol 26 as a single diastereomer (path a). Alternatively, oxidative cleavage of the silyl group provides diol 27 (path b). Because the ketone 27

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Scheme 6. Vinylsilane Analysis of γ - and δ -Hydroxy Systems



can undergo diastereoselective reduction, the initial stereogenic center then propagates into three more.



An asymmetric aldol surrogate requires an enantioselective addition of a terminal alkyne to aldehydes or ketones. Following the Carreira protocol¹⁴ with (+)-*N*-methylephedrine, cyclohexanecarboxyaldehyde was converted to the propargyl alcohol **28** in good ee (85% ee) (eq 5). The one-pot hydrosilylation oxidative cleavage then gives the "aldol" product **29** of the same ee. Providing this system via an asymmetric aldol addition of 5-phenyl-2-pentanone would not be trivial considering regioand enantioselectivity issues. Scheme 5 illustrates a similar sequence from isobutyraldehyde **30** with a more functionalized acetylene. An improved enantioselectivity in forming adduct **31** is obtained in this case under identical conditions. Thus, an excellent overall yield of the aldol type product **33** of 94% ee is available in a two-pot operation from isobutyraldehyde. On the other hand, the intermediate vinylsilane **32** may be diastereoselectively oxidized to epoxide **34**. Derivatizing the free alcohol to the MOM ether **35** prior to unmasking the α -hydroxyketone provides the diol product **36** with the two alcohols chemodifferentiated.



Homo- and Bishomopropargylic Alcohol Substrates

We were also interested in developing alkyne nucleophiles as methyl ketone enolate or hydroxymethyl ketone enolate equivalents in homoaldol processes, where traditional aldol processes are not successful. Homoaldol products could be targeted through formation of either the β , γ C–C bond (e.g., a homoenolate-aldehyde coupling) or the α , β bond (e.g., an enolate-epoxide coupling). Several examples of the first, β , γ approach using homoenolate equivalents have been demonstrated, and some recent methods indicate that homoenolates may be important synthetic methods for some target classes.²⁵ Formation of the α , β bond, however, has few effective solutions, because enolate nucleophiles are typically very poor partners with the requisite epoxide electrophiles. Extending the method of the previous section to view terminal alkynes as functional

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Scheme 7. A Methyl Ketone Enolate Equivalent for Homoaldol Processes



equivalents of methyl ketone enolates allows access to a method for the synthesis of the α,β bond because alkynyl anions are good nucleophiles for epoxide-opening reactions.

The homoaldol process requires a γ -silyl homoallylic alcohol. The cyclic siloxane derivative of such compounds is readily constructed through an intramolecular hydrosilylation reaction with the same catalyst **1** as we previously disclosed (Scheme 6, i).⁵

An example of such a process using an alkyne as a methyl ketone enolate equivalent is shown in Scheme 7. Opening the epoxide 37 with an alkynyllithium occurs cleanly to generate highly diastereoselectively the homopropargylic alcohol 38. A regioselective endo-dig hydrosilylation affords cyclic siloxane **39**. At this stage, we encountered some difficulty in performing a silane oxidation. Traditional conditions (aqueous H2O2, KF, KHCO₃, THF/MeOH) provided no turnover even at elevated temperatures (50 °C). Switching the solvent to DMF and heating to 80 °C resulted in decomposition, as did the basic conditions developed by Woerpel.^{26b} However, we found that a little-used protocol utilizing acidic conditions by introduction of acetic anhydride served to furnish the desired hydroxyl ketone, isolated after conversion to the acetate.²⁶ The reasons for the difficulty in obtaining clean oxidation are not obvious for this substrate, although the increased stability of the six-membered ring silacycle relative to acyclic or five-membered substrates, together with the steric interactions from the significant substitution about the silacycle, may play an important role.

In an effort to introduce further molecular complexity to cyclic vinylsilane intermediates such as **39**, we examined epoxidation, which would allow either protodesilylation to synthesize remote *trans*-epoxides with control of stereochemistry or silane oxidation to furnish the hydroxyl ketone functionality. The treatment of silane **39** with *m*-CPBA resulted in clean conversion to a single epoxide product, as judged by ¹H NMR analysis of the crude reaction mixture. However, the epoxide was not stable to silica gel and was isolated in only 31% yield after chromatography. Analysis of ¹H NMR coupling constants or nOe spectra was not instructive, and the relative stereochemistry of epoxide **41** is based solely on conformational speculation and analogy to later examples.

Scheme 8. Synthesis of $\alpha, \gamma\text{-Dihydroxy}$ Ketone by a Cyclic Silyl Epoxide Method



Encouraged by the apparent opportunity for diastereoselective epoxidation of vinyl silacycles, where no hydroxyl directing group is present, we examined alcohol 43, readily available by addition of an alkynylaluminum reagent to benzyl glycidyl ether (Scheme 8). Hydrosilylation produced the vinylsilane, which could be oxidized to the epoxide 45. The use of m-CPBA gave the desired epoxide in reasonable diastereoselectivity (4:1). The stereoselectivity of the nondirected epoxidation of cyclic olefins has not been good; silacycles, which possess significantly more conformational freedom than their carbocycle counterparts, might have been expected to be even worse. Interestingly, the use of dimethyldioxirane²⁷ (DMDO) at low temperatures resulted in a significant increase in selectivity (10:1).²⁸ At this stage, we set about finding a method for vinylsilane oxidation to reveal the dihydroxy ketone. The epoxide substrate 45 was significantly more unstable than the vinylsilane 41 (see Scheme 7), and we initially were plagued by low mass recovery. Sensing that control of the amount of TBAF might be the source of the problem, a solution was found employing a protocol of slow addition (ca. 4 h) of TBAF to a premixed suspension of anhydrous hydrogen peroxide-urea adduct (UHP) in a THF solution of the epoxy-silane. We originally developed this protocol for oxidation of the benzyldimethylsilyl group where protodesilylation competes with oxidation at high fluoride concentration.²⁹ It may well be that creating the pentavalent silicon fluoride is crucial to maintaining an open coordination site for peroxide. Decomposition pathways such as protodesilylation may result from the coordinatively saturated hexavalent silicon difluoride in the presence of significant quantities of TBAF. The protocol has proven useful for diverse reactions including those for which the original rationale does not necessarily apply. Employing those conditions to the problem at hand allowed complete conversion without the extensive decomposition observed when the TBAF is introduced dropwise over just several minutes. The α,γ -dihydroxy ketone was isolated in diastereomerically pure form after protection of the hydroxyl groups as the bis-triethylsilyl ether (46).

The relative stereochemistry of ketone **46** was established through conversion to the acetonide **48** by the sequence summarized in Scheme 9. The acetonide ¹³C NMR shifts establish a 1,3-syn arrangement (¹³C NMR acetal ca. 98.9 ppm, methyl groups $\Delta > 9$ ppm) rather than a 1,3-anti arrangement (¹³C NMR acetal ca. 100.6 ppm, methyl groups $\Delta < 5$ ppm).³⁰

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Scheme 10. Synthesis of α,γ -Dihydroxy Ketone with Selective Methyl Ketal Formation for Diol Differentiation





Because we anticipated that α -face attack would lead to significant torsional strain in the transition state (44 model A, projection down C–Si bond, below), obtention of the product derived from an α -face attack of the oxidant on the vinylsilane 44 was somewhat surprising. The difference between our expectation and experimental observation may be that the longer Si–C bonds remove torsional strain as a significant factor, and the observed product derives from steric screening of the top face (44 model b).



Homopropargylic alcohol **50** behaved similarly, producing ketal **54** by a similar route (Scheme 10). In this instance, treatment of the crude keto-diol **53** with acidic methanol served to form a single ketal product for characterization purposes. Ketalization also has the effect of differentiating the two hydroxyl functionalities for further synthetic operations. The relative stereochemistry of the 1,3-diol **54** is assumed by analogy to the previous example (see Schemes 8 and 9). The stereochemistry at the ketal carbon is assumed to follow from the anomeric effect.

Oxidation problems arose in the context of a more highly substituted system (Scheme 11). A homopropargylic alcohol **56** with a 1,2-syn stereochemical relationship was synthesized by coupling of a propargylic mesylate with isobutyraldehyde using in situ formation of an organozinc compound catalyzed by Pd-(O). Although we employed racemic starting mesylate, it has been shown that enantiomerically pure mesylates do faithfully transmit chirality to the product, allowing access to single enantiomerically pure product alcohols.^{9,31}

Hydrosilylation and epoxidation proceed without incident to afford an epoxide **58**, tentatively assigned the β -epoxide



Figure 1. Conformational analysis of a bishomopropargylic alcohol system.

Scheme 11. Oxidation Problems for a Homopropargylic Alcohol with Additional Steric Hindrance



stereochemistry on the basis of ¹H NMR coupling constants (Scheme 11). The epoxide, however, proved much more stable and resistant to oxidation. Screening of a variety of oxidation conditions did not result in the formation of the desired ketone products. In general, no reaction is observed, and decomposition is seen only at elevated temperatures (80 °C). The vastly different reactivity of **45**, **52**, and **58** toward silane oxidation is curious.

The use of hydroxyl groups to direct selective formation of α -hydroxy ketones from alkyne precursors could prove a valuable means of installing oxygenation functionality with control of diastereoselectivity even at more remote centers. With this in mind, we targeted the bishomopropargylic alcohol 61a (Scheme 12). We performed reduction of ketone 60 as previously reported to allow diastereoselective access to homopropargylic alcohol 61 after alkyne methylation. The original authors reported 15:1 diastereoselectivity in the ketone reduction, directed by coordination to the pendent alkyne, but we observed variable selectivity (ca. 15:1 to 5:1) favoring the syn alcohol 61a and were unable to determine the source of the variation. Pleasantly, a similar hydrosilylation-epoxidation-silane oxidation pathway performed on the mixture of diastereomeric alcohols afforded the analogous 1,4-diol derivitives 63. This time, the product was best isolated as the bis-silvl ether. We do not observe any evidence of the diastereomeric epoxides by crude ¹H NMR analysis of the reaction mixture, a gratifying result given the conformational flexibility of seven-membered rings.

To determine the relative stereochemistry of the newly formed stereocenter, acetonide formation of the mixture of **63a** and **63b** produced a mixture of two inseparable compounds (ca. 5:1) (Scheme 13). The major product **64** exhibited a strong nOe

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Scheme 12. Utilization of Bishomopropargylic Alcohol Substrates



62a (+ minor isomer)

5:1 63a 63b

Scheme 13. Determination of Stereochemistry



correlation between the methyl doublet and the proton α to the ketone, establishing the all-syn stereochemistry of 63a. Alternatively, treatment of the crude product from the oxidation of 62a with acidic methanol produced two compounds in similar amounts: a ketal 65, and another unstable ketal, which decomposed readily on standing. The hydroxylated C3-H of ketal 65 exhibits coupling constants of 6.7, 3.4, and 3.5 Hz in its ¹H NMR spectrum. These peaks are indicative of an axial orientation of the hydroxyl group. The relative stereochemistry of the other groups can be inferred from a 1,3 nOe correlation between the ketal methoxy group and the benzylic C-H, requiring an equatorial arrangement of the aryl group. Finally, the small (3.2 Hz) coupling present for the benzylic peak requires axial arrangement of the neighboring methyl group; this must be the case because syn-1,2 stereochemistry was established in the starting material 81.³²

The observed product derives from β -face attack of the oxidant on the convex face, as depicted in Figure 1. Molecular dynamics calculations indicate that the preferred conformation is the pseudo-boat form as shown (Figure 1).

A Synthesis of (+)-Spectaline Using a Vinylsilane-Based Homoaldol Equivalent

Spectaline is a member of a class of all-cis 2,6-disubstituted, 3-hydroxy piperidine alkaloids that includes carpamic acid, Scheme 14. Retrosynthetic Analysis of (+)-Spectaline



azimic acid, cassine, and prosafrinine. Hydroxylated piperidine alkaloids can possess important biological activity, stemming from their abilities as carbohydrate mimics in the context of enzyme inhibition.³³ These compounds have been the targets of numerous synthetic efforts.³⁴ However, extant syntheses are plagued by protecting group manipulations and functional group interconversions leading to synthetic inefficiency. Reductive amination of an amino-ketone has been shown to be a diastereoselective and efficient strategy for the synthesis of 3-hydroxy piperidine alkaloids, and such a retrosynthetic disconnection in this case furnishes the δ -amino, γ -hydroxyketone 68. The ease of forming such γ -hydroxyketone "homoaldol" products by the method described here interested us in the use of intramolecular hydrosilylation for the selective installation of ketone functionality. This approach to the spectaline alkaloids allows simultaneous late-stage elaboration of ketone and hydroxyl functionality. By installing the amine functionality as an azide group, this overall strategy avoids use of any protecting groups for the synthesis of the piperidine core (Scheme 14). The atom-economical approach employs only dimethylsiloxane and dinitrogen as wasted atoms in the synthesis of the central core and allows a short step count as well.

In the synthetic direction, straightforward processes furnish alkyne 73 (Scheme 15). Starting from the known epoxide 75,

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Scheme 16. Completion of the Synthesis of Spectaline



available by resolution of (\pm) -1-buten-3-ol (**74**) through catalytic asymmetric epoxidation,³⁵ followed by Mitsunobu displacement with $Zn(N_3)_2 \cdot 2pyr^{36}$ gave a volatile azido-expoxide. Although we could isolate the azido-epoxide in low yield, it was far more convenient and efficient to treat the crude azido-epoxide with the alkynyllithium generated from terminal alkyne **73** to furnish the epoxide-opening product **76**.

The azido-alcohol 76 sets the stage for a key regioselective ketone introduction at the distal position of the alkyne. Intramolecular hydrosilylation proceeds well to furnish the product cyclic siloxane in nearly quantitative yield (Scheme 16). The tolerance of the azide functionality in the ruthenium-catalyzed hydrosilylation is demonstrated for the first time with this substrate. The stage is now set for unmasking the ketone. The oxidation of silanes containing azide functionality has been demonstrated for a single substrate,37 so we were encouraged to pursue direct oxidation to the ketone 78. The oxidation was quite slow under typical conditions (H2O2, KF, KHCO3, MeOH/ THF at room temperature provided 38% conversion over 6 d). Unfortunately, raising the temperature or switching to more active oxidation conditions (TBAF, H2O2, DMF) resulted in significant decomposition. We also pursued initial azide reduction prior to silane oxidation. Treatment of the azide 78 with triphenylphosphine or trimethylphosphine allowed clean conversion to an intermediate iminophosphorane. Unfortunately, we had difficulty hydrolyzing the intermediate iminophosphorane in the presence of the rather sensitive cyclic siloxane, and product characterization became troublesome. As a result of these difficulties and the step economy of performing the oxidation in the presence of the azide, we reexamined the Scheme 17. Strategic Approach to Hydroxyketones



Scheme 18. Complementary Behavior Based upon Hydrosilylation Catalyst



oxidation. Again we found that dropwise addition of TBAF over 4 h to a mixture of vinylsilane and oxidant provided the solution. In this case, dropwise addition in THF resulted in a clean reaction, but did not significantly improve the reaction rate. However, switching the solvent to DMF improved the reaction rate and allowed complete conversion without the extensive decomposition observed when all of the TBAF is introduced within 5 min.

It thus remained only to perform the reductive amination and remove the distal ketal protecting group to complete the synthesis. Fortunately, treatment of the azido-ketone with Pd/C under a pressure of H₂ (60 psi) served in one step to reduce the azide, revealing the amine functionality, and to bring about imine formation and reduction to the desired piperidine target (Scheme 16).³⁸ Treatment of the crude mixture with aqueous HCl removed the ketal functionality to allow isolation of the (+)-spectaline target. Of note, our endo-dig hydrosilylation-oxidation strategy allows for a convergent strategy and avoids the need for any traditional protecting groups in the piperidine core and only one protecting group for the side chain. The net result is a synthesis having a longest linear sequence of 9 steps (11 total steps) and proceeding in >18% yield.

Conclusions

Strategically, the availability of the ruthenium-catalyzed regioselective hydrosilylation provides the very useful β -, γ -, and δ -hydroxyketones as illustrated in Scheme 17. The complementary behavior of rhodium, platinum, or palladium-catalyzed hydrosilylation as compared to ruthenium translates into regio-complementary products as summarized in Scheme 18. The masking of the ketone or α -hydroxyketone in the form of the vinylsilane has the additional advantage of serving as a chemically unreactive form. Furthermore, the prospect of introducing substituents other than hydroxyl by electrophilic

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additions to the vinylsilane offer the promise of further broadening of the scope of this strategy.

Here, we present a strategy for the introduction of regio- and stereodefined oxygen functionality from internal alkyne building blocks though vinylsilane intermediates. In addition to discriminating between the two termini of the alkyne, the silyl intermediate serves as a diastereocontrol element, allowing even quite remote stereocontrol of olefin functionalization by enforcing A^{1,3} strain with allylic alcohols or through cyclic intermediates with homo- and bishomopropargylic alcohols. The use of such cyclic vinylsilane intermediates may well provide a means of achieving remote stereocontrol in olefin addition reactions in general.

Experimental Section

(2R,3S)-4-Undecyn-1,2,3-triol 1,2-Isopropylidene Ketal, 19. This procedure was adapted from the method of Shimizu.⁷ A solution of 1-octyne (1.59 mL, 10.8 mmol) in THF (10 mL) under Ar at -78 °C was treated with butyllithium (1.59 mL, 10.8 mmol, 1.48 M in hexanes) dropwise. After 30 min, ClTi(Oi-Pr)3 (2.57 mL, 10.8 mmol) was added, and the flask was warmed to -60 °C. To maintain solubility, additional THF (20 mL) was added, and the solution was stirred for 90 min prior to recooling to -78 °C. (R)-Glyceraldehyde isopropylidene ketal¹ (0.70 g, 5.38 mmol) was then added via syringe in one portion, and the mixture was stirred for 2 h prior to quenching with saturated aqueous NH4Cl (20 mL) at -78 °C. Water (30 mL) was then added, and the reaction was extracted with ether (3 \times 30 mL). The ether extracts were washed with brine (20 mL) and dried over MgSO₄, and the solvents were removed in vacuo. Silica gel chromatography gave 1.19 g (92%) of the desired alcohol as a 9:1 mixture of epimeric diols favoring the trans. ¹H NMR spectra of trans and cis components are very similar to a previously reported addition.8

Data for Major Isomer. ¹H NMR (500 MHz, CDCl₃): δ 4.48 (m, 1H), 4.21 (ddd, J = 6.5, 6.5, 4.0 Hz, 1H), 4.03–4.08 (m, 2H), 2.26 (d, J = 4.4 Hz, 1H), 2.20 (td, J = 7.2, 2.1 Hz, 2H), 1.45–1.51 (m, 2H), 1.46 (s, 3H), 1.37 (s, 3H), 1.24–1.36 (m, 6H), 0.88 (t, J = 7.0 Hz, 3H). ¹³C NMR (125 MHz, CDCl₃): δ 110.0, 87.3, 78.1, 77.0, 65.2, 62.4, 31.3, 28.5, 28.4, 26.3, 25.2, 22.5, 18.7, 14.0. IR (thin film): 3452 (br, OH), 2933, 2234 (w), 1372, 1254, 1216, 1153, 1070, 851 cm⁻¹. [α]²⁶_D +17.8° (c 1.0, CHCl₃). Anal. Calcd for C₁₄H₂₄O₃: C, 69.96; H, 10.07. Found: C, 69.79; H, 10.15.

(2*R*,3*S*)-1,2,3-Trihydroxy-5-undecanone 1,2-Isopropylidene Ketal, 20. The alkyne 19 (120 mg, 0.50 mmol) and BDMS-H (104 μ L, 0.60 mmol) in acetone (1.0 mL) was treated at 0 °C with complex 1 (7.8 mg, 0.015 mmol). The mixture was allowed to warm to room temperature over 30 min. It was then recooled to 0 °C, diluted with THF (1.5 mL), and TBAF (0.60 mL, 0.60 mmol, 1.0 M soln in THF) was added dropwise. After 20 min, MeOH (1.0 mL) was introduced, followed by solid potassium bicarbonate (150 mg, 1.5 mmol) and aqueous H₂O₂ (0.80 mL, 30% soln). The mixture was then stirred for 36 h at room temperature, at which time it was diluted with water (10 mL) and extracted with EtOAc (3 × 10 mL). The organic extracts were washed with brine (10 mL), dried over Na₂SO₄, and concentrated in vacuo. The residue was purified on a silica gel column (eluent: 85: 15:1 pet. ether:EtOAc:MeOH) to give 99 mg (77%) of the desired ketone as a single isomeric compound.

¹H NMR (500 MHz, CDCl₃): δ 4.08 (m, 1H), 3.92–3.95 (m, 3H), 3.27 (br OH, 1H), 2.82 (dd, J = 7.5, 1.8 Hz, 1H), 2.57 (m, 1H), 2.45 (t, J = 7.5 Hz, 2H), 1.57 (m, 2H), 1.39 (s, 3H), 1.34 (s, 3H), 1.26– 1.30 (m, 6H), 0.87 (t, J = 7.0 Hz, 3H). ¹³C NMR (125 MHz, CDCl₃): δ 212.4, 109.4, 77.5, 69.1, 66.9, 45.1, 43.7, 31.5, 28.8, 26.7, 25.1, 23.5, 23.5, 22.4, 14.0. IR (thin film): 3472 (br, OH), 2932 (s), 1713, 1372, 1215, 1066 (s), 851 (w) cm⁻¹. [α]²⁶_D –21.3° (c 1.3, CHCl₃). Anal. Calcd for C₁₄H₂₆O₄: C, 65.09; H, 10.14. Found: C, 64.84; H, 9.92. (*R*)-[(2*S*,3*R*)-3-(Benzyldimethylsilyl)-3-hexyl-oxiranyl]-((*R*)-2,2dimethyl-[1,3]dioxolan-4-yl)-methanol, 25. Following the general procedure for the synthesis of (\pm) -(*R*)-[(2*S*,3*R*)-3-(benzyldimethylsilyl)-3-methyloxiranyl]-phenylmethanol (15), vinylsilane 24 (205 mg, 0.52 mmol) was treated with *m*-CPBA (260 mg, 1.04 mmol) in CH₂Cl₂ (5.0 mL) to give 180 mg (84%) of the desired epoxide as a 9:1 mixture of relative epimers at C2 from the initial alkyne addition. Purification was done on silica gel (eluent: 20:1 to 10:1 pet. ether:acetone).

Data for Major Isomer. ¹H NMR (500 MHz, C₆D₆): δ 7.09–7.13 (m, 2H), 6.94–7.00 (m, 3H), 4.03 (ddd, J = 5.3, 5.3, 1.6 Hz, 1H), 3.92–3.99 (m, 2H), 3.62 (m, 1H), 2.96 (d, J = 6.7 Hz, 1H), 2.31 (d, J = 13.7 Hz, 1H), 2.22 (d, J = 13.7 Hz, 1H), 1.99 (d, J = 4.1 Hz, 1H), 1.90 (m, 1H), 1.35 (3, 3H), 1.22 (s, 3H), 1.08–1.26 (m, 9H), 0.84 (t, J = 7.1 Hz, 3H), 0.09 (s, 3H), 0.08 (s, 3H). ¹³C NMR (100 MHz, C₆D₆): δ 139.5, 128.8, 128.6, 124.8, 109.6, 77.3, 71.6, 67.6, 65.8, 58.7, 37.8, 32.1, 29.7, 26.8, 25.7, 25.3, 24.8, 23.0, 14.3, -3.0, -3.1. IR (thin film): 3452 (br, OH), 2931, 1600 (w), 1251, 1207, 1067, 828, 700 cm⁻¹. [α]²⁶_D +14.3° (*c* 1.0, CH₂Cl₂). This compound is further characterized after conversion to ketone **33**.

(15,25)-1-((*R*)-2,2-Dimethyl-[1,3]dioxolan-4-yl)-1,2-dihydroxynonan-3-one, 27. Following the general procedure for the oxidative desilylation given for 17, employing epoxy silane (90.0 mg, 0.216 mmol), TBAF (0.216 mL, 0.216 mmol, 1.0 M in THF), KHCO₃ (65 mg, 0.64 mmol), and aqueous H_2O_2 (0.3 mL, 30% soln) in THF (0.7 mL) and MeOH (0.7 mL) provided 51.8 mg (85%), of the desired diol, isolated as an inseparable 9:1 mixture of relative epimers at C-2 carried through from the initial alkyne addition. Chromatography eluent: 80: 20:1 to 66:33:1 pet. ether:EtOAc:MeOH.

R_j: 0.37 (67:33:1 pet.ether:EtOAc:methanol). ¹H NMR (500 MHz, CDCl₃): δ 4.41 (d, *J* = 2.4 Hz, 1H), 4.12–4.15 (m, 2H), 4.07 (m, 1H), 3.89 (m, 1H), 3.78 (d, *J* = 3.6 Hz, 1H), 2.48–2.63 (m, 2H), 2.10 (d, *J* = 10.4 Hz, 1H), 1.66 (m, 2H), 1.48 (s, 3H), 1.38 (s, 3H), 1.26–1.35 (m, 6H), 0.89 (t, *J* = 7.0 Hz, 3H). ¹³C NMR (125 MHz, CDCl₃): δ 210.4, 109.4, 76.1, 75.6, 72.4, 66.8, 37.7, 31.5, 28.8, 27.0, 25.0, 23.4, 22.4, 14.0. IR (thin film): 3445 (br, OH), 2933, 1715, 1372, 1216, 1069, 847 cm⁻¹. $[\alpha]^{26}{}_{\rm D}$ +51.4° (*c* 1.0, CHCl₃). Anal. Calcd for C₁₄H₂₆O₃: C, 61.29; H, 9.55. Found: C, 61.42; H, 9.52.

General Procedure for the One-Pot Hydrosilylation and Oxidative Desilylation: (S)-1-Cyclohexyl-1-hydroxy-6-phenyl-hexan-3one, 28. To a solution of (R)-1-cyclohexyl-6-phenyl-2-hexyn-1-ol (21) (50 mg, 0.20 mmol) and BDMS-H (87.9 mg, 0.59 mmol) in CH₂Cl₂ (0.4 mL) was added complex 1 (4.9 mg, 0.01 mmol) at 0 °C. The solution was allowed to warm to room temperature and stirred for 30 min. The solution was cooled to 0 °C, and THF (0.7 mL) and TBAF (0.58 mL, 0.58 mmol, 1.0 M in THF) were added and the solution was stirred for 15 min. Aqueous H₂O₂ (1.9 mL, 10.7 mmol), MeOH (0.7 mL), and KHCO₃ (176 mg, 1.76 mmol) were added to the solution. The solution was then warmed to room temperature and stirred for 18 h. Brine (10 mL) was added, and the mixture was extracted with ether $(2 \times 30 \text{ mL})$. The organic layer was washed with saturated aqueous Na₂S₂O₃ (10 mL) and brine (10 mL) and dried over MgSO₄. Purification on silica gel column (eluent: 8:1 pet. ether:EtOAc) afforded (S)-1cyclohexyl-1-hydroxy-6-phenyl-3-hexanone (46.3 mg, 87%).

R_f: 0.57 (4:1 P.E.:EtOAc). ¹H NMR (300 MHz, CDCl₃): δ 7.15– 7.30 (m, 5H), 3.78 (s, 1H), 2.95 (s, 1H), 2.42–2.64 (m, 5H), 1.61– 1.96 (m, 7H), 0.95–1.43 (m, 7H). ¹³C NMR (75 MHz, CDCl₃): δ 212.4, 141.4, 128.41, 128.36, 126.0, 71.6, 46.1, 42.9, 42.8, 34.9, 28.8, 28.2, 26.4, 26.1, 26.0, 27.9. IR (thin film): 3363 (br), 2915, 2850, 1702 (s), 1446 (w), 1403 (w), 1087 (w), 699 cm⁻¹. Anal. Calcd for C₁₈H₂₆O₂: C, 78.79; H, 9.55. Found: C, 79.00; H, 9.45. [α]²⁵_D +33.9° (*c* 1.0 in CHCl₃). mp 65–67 °C.

(*R*)-Methyl 12-Hydroxy-13-methyl-10-tetradecynoate, 31. This was adapted from the method of Carreira.¹⁴ A flask charged with zinc triflate (360 mg, 1.0 mmol) was placed under vacuum (1 mmHg) and heated to 125 °C for 2 h. (+)-*N*-Methylephedrine (197 mg, 1.10 mmol) was added to the flask, which was placed under an Ar atmosphere.

Toluene (5.0 mL) was added to the flask, and the mixture was stirred for 2 h. Methyl 10-undecynoate (1.18 g, 6.0 mmol) was added to the flask, and, after 15 min, isobutyraldehyde (454 μ L, 5.0 mmol) was added to the flask, which was then heated to 60 °C and stirred for 12 h. The mixture was then quenched by addition of saturated aqueous NH₄Cl (20 mL) and extracted with ether (3 × 15 mL). The extracts were washed with brine (15 mL), dried over MgSO₄, and concentrated under reduced pressure. Purification on silica gel (eluent 80:20:1, then 70:30:1 pet. ether:EtOAc:MeOH) gave the desired alcohol (1.09 g, 81%) as a colorless oil. The enantiomeric excess was determined to be 94%, presumed to be the (*R*) stereochemistry by analogy to Careira's published results. For purposes of ee determination, racemic material was generated in acetonitrile using Zn(OTf)₂.¹⁴

R_f: 0.43 (80:20:1 pet. ether:EtOAc:MeOH). ¹H NMR (300 MHz, CDCl₃): δ 3.66 (s, 1H), 2.30 (t, J = 7.4 Hz, 2H), 2.20 (td, J = 6.8, 1.6 Hz, 2H), 1.77–1.87 (m, 2H), 1.61 (m, 2H), 1.48 (m, 2H), 1.24–1.40 (m, 8H), 0.98 (d, J = 6.1 Hz, 3H), 0.96 (d, J = 6.1 Hz, 3H). ¹³C NMR (75 MHz, CDCl₃): δ 174.3, 86.1, 79.8, 68.1, 51.4, 34.6, 34.0, 29.04, 28.99, 28.8, 28.6, 24.9, 18.6, 18.1, 17.4. [α]²⁶_D +0.94° (*c* 1.3, CHCl₃). Anal. Calcd for C₁₆H₂₈O₃: C, 71.60; H, 10.52. Found: C, 71.60; H, 10.34. Enantiomeric excess determined by HPLC [Chiralcel AS column, eluting with 94:6 heptane/PrOH, 1 mL/min, 225 nm: (*R*)-enantiomer *t*_R 9.32 min, (*S*)-enantiomer *t*_R 10.84 min].

Methyl (*S*)-12-Hydroxy-13-methyl-10-oxo-tetradecanoate, 33. Following the general procedure for one-pot hydrosilylation and oxidative desilylation given for 28, employing alkyne 31 (134 mg, 0.50 mmol), BDMS-H (103 mg, 0.60 mmol), complex 1 (5.0 mg, 0.012 mmol) in CH₂Cl₂ (0.5 mL) at -30 °C, and TBAF (0.60 mL, 0.60 mmol, 1.0 M in THF) and KHCO₃ (150 mg, 1.5 mmol) and aqueous H₂O₂ (0.6 mL, 30% soln) in THF (1.0 mL) and MeOH (1.0 mL) provided the product hydroxyl ketone (114 mg, 80%). Chromatography eluent: 80:20:1 pet. ether:EtOAc:MeOH.

R: 0.31 (80:20:1 pet.ether:EtOAc:methanol). ¹H NMR (500 MHz, CDCl₃): δ 3.83 (m, 1H), 3.69 (s, 3H), 3.02 (br OH, 1H), 2.60 (dd, *J* = 17.3, 2.3 Hz, 1H), 2.51 (dd, *J* = 17.3, 9.8 Hz, 1H), 2.46 (t, *J* = 7.5 Hz, 2H), 2.32 (t, *J* = 7.5 Hz, 2H), 1.57–1.73 (m, 5H), 1.27–1.33 (m, 8H), 0.96 (d, *J* = 6.7 Hz, 3H), 0.93 (d, *J* = 6.8 Hz, 3H). ¹³C NMR (125 MHz, CDCl₃): δ 212.9, 174.3, 72.3, 51.5, 45.9, 43.7, 34.0, 33.0, 29.1, 29.03, 29.02, 24.9, 23.5, 18.3, 17.8. IR (thin film): 3526 (br, OH), 2931 (s), 2857, 1740 (s), 1711, 1438, 1368, 1173 cm⁻¹. [α]²⁶_D –24.7° (*c* 1.0, CHCl₃). Anal. Calcd for C₁₆H₃₀O₄: C, 67.10; H, 10.56. Found: C, 66.94; H, 10.35. HRMS-EI (*m*/ *z*): [M – OH]⁺ calcd for C₁₆H₂₉O₃, 269.2117; found, 269.2127.

General Procedure for Intramolecular Hydrosilylation. (\pm)-6-Benzyloxymethyl-2,2,3-trimethyl-5,6-dihydro-2*H*-[1,2]oxasiline, 44. A round-bottomed flask was charged with 1-benzyloxy-hex-4-yn-2-ol (43) (300 mg, 1.469 mmol) under Ar at room temperature. To the neat alcohol was added 1,1,3,3-tetramethyldisilazane (1.3 mL, 7.344 mmol), and the flask was heated to 50 °C for 2 h. Next, the flask was cooled to ambient temperature and placed under vacuum (ca. 1 mmHg) for 45 min to remove the volatile species. An Ar atmosphere was then reintroduced and the residue taken up in CH₂Cl₂ (3.6 mL). The flask was cooled to 0 °C, and solid [Cp*Ru(MeCN)₃]PF₆ (14.8 mg, 0.029 mmol) was added to the solution. The flask was allowed to warm to room temperature, and, after 2 h, the solution was diluted with ether (15 mL) and filtered through a short plug of florisil, washing with additional ether (20 mL). After concentration under reduced pressure, the cyclic siloxane was obtained as a colorless oil (347 mg, 90%).

R_j: 0.48 (10:1 pet. ether:EtOAc). ¹H NMR (400 MHz, C₆D₆): δ 7.06–7.29 (m, 5H), 6.25 (m, 1H), 4.38 (d, *J* = 12.4, 1H), 4.34 (d, *J* = 12.4 Hz, 1H), 4.14–4.20 (m, 1H), 3.58 (dd, *J* = 9.6, 4.8 Hz, 1H), 3.39 (dd, *J* = 9.6, 6.4 Hz, 1H), 2.19 (m, 2H), 1.62–1.66 (m, 3H), 0.17 (s, 3H), 0.19 (s, 3 H). ¹³C NMR (100 MHz, C₆D₆):

 δ 139.7, 139.3, 135.1, 128.5, 127.9, 127.6, 75.2, 73.4, 71.0, 33.6, 20.5, -1.5, -1.6. IR (thin film): 2855, 1614, 1454, 1251, 1097, 967, 827, 781 cm^{-1}.

(\pm)-(3*S*,5*R*)-6-Benzyloxy-3,5-bis-triethylsilanyloxy-hexan-2one, 46. To a solution of the cyclic siloxane 44 (100 mg, 0.381 mmol) in acetone (1 mL) was added DMDO (19.1 mL, 0.1 M soln in acetone, 1.91 mmol) under Ar at -78 °C via cannula. After being stirred overnight at -78 °C, the reaction was allowed to warm to -20 °C over 2 h. At that temperature, the reaction was quenched with dimethyl sulfide (355 mg, 5.716 mmol). After warming to room temperature and drying over MgSO₄, the mixture was concentrated under reduced pressure, which afforded the crude epoxide 45 (157 mg, 148%) as a colorless oil.

R_f: 0.49 (4:1 pet. ether:EtOAc). ¹H NMR (400 MHz, C₆D₆): δ 7.07–7.28 (m, 5H), 4.33 (s, 2H), 4.21 (m, 1H), 3.38 (dd, *J* = 9.6, 4.8 Hz, 1H), 3.28 (dd, *J* = 9.6, 5.2 Hz, 1H), 2.82 (d, *J* = 3.6 Hz, 1H), 1.96 (ddd, *J* = 15.2, 5.6, 3.6 Hz, 1H), 1.76 (dd, *J* = 15.2, 11.2 Hz 1H), 1.04 (s, 3H), 0.25 (s, 3H), 0.12 (s, 3H).

UHP (179 mg, 1.90 mmol) was added to a solution of the crude epoxide (157 mg, assume 0.381 mmol) in THF (3.8 mL) under Ar at 0 °C. The reaction was allowed to warm to room temperature and treated with TBAF (1.41 mL, 0.81 M soln in THF, 1.143 mmol) dropwise via syringe pump over 4 h. After being stirred for 20 h at room temperature, the reaction was stopped by diluting with EtOAc (15 mL) and aqueous $Na_2S_2O_3$ (10 mL, 0.1 M soln). Extraction with EtOAc (3 × 10 mL) and drying over MgSO₄ afforded a crude oil (189 mg, 208%), which was used directly in the next reaction.

The crude diol (189 mg, assume 0.381 mmol) and imidazole (431 mg, 6.34 mmol) were solved in CH₂Cl₂ (7.9 mL) under N₂ at 0 °C. Et₃SiCl (660 μ L, 3.966 mmol) was then added via syringe at 0 °C. After warming to room temperature, the mixture was stirred for 2 h. The reaction was quenched with H₂O (10 mL) and diluted with diethyl ether (30 mL). The aqueous layer was extracted with Et₂O (3 × 5 mL), and the organic extracts were washed with brine (5 mL) and dried over MgSO₄. The resulting oil was purified on a silica gel column (30:1, then 20:1 pet ether:diethyl ether), which afforded the protected product (138 mg) in 77% yield over three steps.

R: 0.28 (10:1 pet. ether:Et₂O). ¹H NMR (500 MHz, CDCl₃): δ 7.26–7.35 (m, 5H), 4.53 (d, J = 12.1, 1H), 4.49 (d, J = 12.1 Hz, 1H), 4.15 (t, J = 6.5 Hz, 1H), 3.96–4.00 (m, 1H), 3.38–3.43 (m, 2H), 2.14 (s, 3H), 1.76–1.95 (m, 2H), 0.94 (t, J = 7.8, 18H), 0.55–0.62 (m, 12H). ¹³C NMR (125 MHz, CDCl₃): δ 210.9, 138.3, 128.3, 127.6, 127.5, 75.7, 74.3, 73.2, 67.8, 39.9, 24.9, 6.83, 6.81, 6.74, 5.03, 4.87, 4.69. IR (thin film): 2955, 2877, 1717, 1456, 1415, 1352, 1239, 1100, 1006, 740 cm⁻¹. Anal. Calcd for C₂₅H₄₆O₄Si₂: C, 64.32; H, 9.93. Found: C, 64.16; H, 10.08.

(+)-Spectaline, 67. The azido-siloxane 77 (7.1 mg, 0.016 mmol) and solid urea-hydrogen peroxide adduct (6 mg, 0.064 mmol) were taken up in DMF (0.16 mL) and treated at 0 °C with TBAF (63 μ L, 0.063 mmol, 1.0 M in THF) dropwise via a syringe pump over 4 h. The mixture was then allowed to warm to room temperature and was stirred for 10 h. At this time, TLC analysis indicated incomplete consumption of the starting vinylsilane, and so additional urea-hydrogen peroxide adduct (6 mg, 0.064 mmol) was added, followed by additional TBAF (63 µL, 0.063 mmol, 1.0 M in THF) via syringe over 4 h as before. The mixture was then allowed to warm to room temperature and was stirred for 20 h. The mixture was diluted with water (5 mL) and extracted with EtOAc (3 \times 4 mL). The organic layers were dried over Na₂SO₄ and concentrated in vacuo. The residue was purified on a silica gel column (eluent 7:1, then 4:1 pet. ether:EtOAc) to afford 5.2 mg (84%) of the desired ketone 6.129. IR (thin film): 3456 (br, OH), 2923 (s), 2109 (s), 1713, 1463, 1377, 1256, 1060 cm⁻¹. MS-CI (m/z): $[M + Na]^+$ calcd for C₂₂H₄₁N₃O₄Na, 434.3; found, 434.3.

The keto-azide (2.4 mg, 0.0058 mmol) was taken up in EtOH (0.5 mL) and treated with 10% Pd/C (1 mg). The mixture was placed under 60 psi of H_2 and stirred for 24 h. The mixture was filtered through Celite, washing with additional EtOH (2 mL). The filtrated was concentrated in vacuo and taken up in THF (0.5 mL). The solution was treated with aqueous HCl (0.05 mL, 2.0 M soln) and stirred for

24 h at room temperature. The mixture was then quenched by addition of aqueous NaOH (0.5 mL, 1.0 M soln). The mixture was extracted with EtOAc (2×3 mL), dried over Na₂SO₄, and concentrated in vacuo. The residue was purified on a column of basic alumina (eluent 100:1 CHCl₃: EtOH), affording 1.0 mg (53%) of (+)-spectaline.

¹H NMR (500 MHz, CDCl₃): δ 3.54 (m, 1H), 2.75 (qd, J = 6.5, 1.4 Hz, 1H), 2.52 (m, 1H), 2.42 (t, J = 7.5 Hz, 2H), 2.14 (s, 3H), 1.90 (m, 1H), 1.24–1.52 (m, 27H), 1.09 (d, J = 6.6 Hz, 3H). IR (thin film): 2926, 2112, 1251, 1049, 833, 784 cm⁻¹. [α]²⁷_D +5.8 (*c* 0.1, CHCl₃). Lit.¹⁶ [α]_D +8.0 (*c* 0.16, CHCl₃).

(25,35)-2-Azido-9-(2-methyl-[1,3]dioxolan-2-yl)-5-octadecyn-3ol, 76. The terminal alkyne 73 (150 mg, 0.54 mmol) was taken up in THF (2.0 mL) and treated with butyllithium (0.34 mL, 0.54 mmol, 1.6 M in hexanes) at -78 °C and stirred for 30 min. The azido-epoxide 84 (137 mg of 34 wt % solution assuming 100% yield in previous step) was added to THF (2.0 mL) over 4 Å molecular sieves (ca. 0.1 g) and allowed to stand for 10 min before it was added to the reaction flask via cannula. BF₃·OEt₂ (0.067 mL, 0.54 mmol) was then added, and, after 15 min, no epoxide was present by TLC analysis. Saturated aqueous NH₄Cl (3 mL) and water (10 mL) were added, and the mixture was extracted with ether (3 × 15 mL). The organic layers were washed with brine (5 mL) and dried over MgSO₄. Solvent removal in vacuo and purification on a silica gel column (eluent 100:10:1, then 85:15:1 pet. ether:EtOAc:methanol) gave 89 mg (55% from epoxy alcohol) of the alcohol as a colorless oil.

¹H NMR (500 MHz, CDCl₃): δ 3.90–3.97 (m, 4H), 3.64 (m, 1H), 3.58 (dq, J = 5.5 Hz, 1H), 2.43–2.45 (m, 2H), 2.14–2.18 (m, 3H), 1.61–1.64 (m, 2H), 1.48 (m, 2H), 1.24–1.40 (m, 20H), 1.33 (d, J = 6.6 Hz, 3H), 1.31 (s, 3H). ¹³C NMR (125 MHz, CDCl₃): δ 110.2, 83.7, 74.8, 73.2, 64.5, 60.3, 39.2, 29.8, 29.57, 29.56, 29.54, 29.52, 29.48, 29.1, 28.85, 28.84, 24.6, 24.1, 23.7, 18.7, 15.7. IR (thin film): 3458 (br, OH), 2927 (s), 2100 (s), 1377, 1256, 1062 cm⁻¹. [α]²⁷_D +40.4 (*c* 1.0, CH₂Cl₂). HRMS-EI (*m*/*z*): [M – CH₃]⁺ calcd for C₂₁H₃₆N₃O₃, 378.2757; found, 378.2772. Anal. Calcd for C₂₂H₃₉N₃O₃: C, 67.14; H, 9.99; N, 10.68. Found: C, 67.32; H, 10.06; N, 10.58.

(*S*)-6-((*S*)-1-Azido-ethyl)-2,2-dimethyl-3-[3-(2-methyl-[1,3]dioxolan-2-yl)-dodecanyl]-5,6-dihydro-2*H*-[1,2]oxasiline, 77. A mixture of alkyne 76 (50 mg, 0.127 mmol) and 1,1,3,3-tetramethyldisilazane (85 mg, 0.635 mmol) was heated to 70 °C for 20 h. The neat mixture was cooled to room temperature and concentrated in vacuo (ca. 1 Torr) to remove volatile organic species. After 30 min under vacuum, the residue was taken up in CH₂Cl₂ (0.3 mL) and treated with solid 1 (3.2 mg, 0.0064 mmol) at 0 °C. The solution was allowed to warm to room temperature over 2 h and was then filtered through florisil (ca. 3 cm), eluting with ether (5 mL), to afford 56 mg (98%) of spectroscopically homogeneous cyclic vinylsilane without additional purification.

¹H NMR (400 MHz, C₆D₆): δ 6.21 (m, 1H), 3.59 (ddd, J = 11.0, 4.6, 2.6 Hz, 1H), 3.55 (s, 4H), 2.88 (dq, J = 6.7, 4.6 Hz, 1H), 2.22 (m, 1H), 2.08 (t, J = 7.5 Hz, 2H), 1.52–1.78 (m, 6H), 1.34 (s, 3H), 1.30–1.40 (m, 19H), 0.95 (d, J = 6.7 Hz, 3H), 0.26 (s, 3H), 0.22 (s, 3H). ¹³C NMR (125 MHz, C₆D₆): δ 139.9, 138.8, 110.2, 75.3, 64.6, 61.0, 39.8, 35.8, 32.5, 30.4, 30.11, 30.07, 30.0, 29.9, 24.6, 24.1, 15.3, -0.7, -0.8. IR (thin film): 2926, 2112, 1251, 1049, 833, 784 cm⁻¹. [α]²⁷_D –313.4 (*c* 1.0, CH₂Cl₂). HRMS-EI (*m*/*z*): [M – CH₃]⁺ calcd for C₂₃H₄₂N₃O₃Si, 436.3004; found, 436.3006.

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Supporting Information Available: Experimental procedures and characterization data. This material is available free of charge via the Internet at http://pubs.acs.org.

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