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Asymmetric synthesis of multiple quaternary stereocentre-containing cyclopentyls by oxazolidinone-promoted Nazarov cyclizations†

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Carbometalation of oxazolidinone (Ox)-substituted ynamides is used to generate highly substituted Ox-divinyl (and aryl vinyl) ketones for use in Nazarov cyclizations. The Ox-group serves as a remarkably effective chiral activating group, enabling the torquoselective Nazarov cyclization of these sterically congested substrates to be performed under mild conditions. It also serves as a charge-stabilizing group in the intermediate oxyallyl cation, suppressing undesired [1,2]-sigmatropic shifts of neighboring substituents and facilitating the regio- and stereoselective incorporation of nucleophiles to yield cyclopentanoids containing up to three contiguous all-carbon quaternary (4°) stereocentres.

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Introduction

The enantioselective synthesis of quaternary (4°) stereocentres is a major challenge in organic synthesis, hindering access to sp³-rich scaffolds in drug discovery and natural products synthesis.^{1,2} Particularly problematic is the enantioselective formation of multiple 4° -stereocentres, which requires control over both relative and absolute stereochemistry.

The Nazarov cyclization offers inherent control over relative stereochemistry through conservation of orbital symmetry and constitutes an attractive route to multistereocentre-containing cyclopentanoids.³ However, the potential of the Nazarov cyclization for 4°-stereocentre formation has not yet been fully realized due to two significant challenges: (i) stereoselective access to highly substituted divinyl (and aryl vinyl) ketone substrates⁴ and (ii) torquoselective⁵ ring closure. In a landmark study, Tius and co-workers⁶ reported chiral Brønsted acid-catalyzed Nazarov cyclizations of divinyl ketones 1 (Scheme 1a) leading to cyclopentenols 3 containing two new vicinal 4°-stereocentres ($R^{1-3} \neq H$) with high enantioselectivities (often er > 97:3). Careful design of the divinyl ketone 1 with dual-activating electron donor (OCHPh₂) and acceptor

(CO₂R) elements was key to attaining efficient cyclization.⁶ Electrofugal release of Ph₂HC⁺ from the intermediate oxyallyl cation 2 further promoted the cyclization and suppressed competing Wagner–Meerwein rearrangements ([1,2]-sigmatropic shifts of R¹⁻³ within 2). Herein, we report that highly substituted aryl vinyl and divinyl ketones 5 can be readily accessed through carbometalations of oxazolidinone (Ox)-substituted ynamides 4 (Scheme 1b).⁷ The Ox-group proves to be remarkably effective as a single chiral activating group for the Nazarov cyclizations of these highly substituted and sterically congested substrates 5, giving *exo*-methylene cyclopentanones 7 under remarkably mild conditions, with excellent and predictable enantiocontrol. Furthermore, since no electrofugal release is required for

Dual activating (polarizing) groups: Ph₂HCO and CO₂R

b) This work:

Ox

$$R^5$$
 R^4
 R^3
 R^2
 R^1
 R^5
 R^4
 R^3
 R^2
 R^1
 R^3
 R^2
 R^3
 R^2
 R^4
 R^3
 R^2
 R^4
 R^3
 R^2
 R^4
 R^3
 R^2
 R^4
 R^4
 R^3
 R^2
 R^4
 R^4
 R^3
 R^2
 R^4
 $R^$

Scheme 1 Nazarov substrate activation modes for the enantiose-lective synthesis of 4° -stereocentres.

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substrate activation or suppression of Wagner–Meerwein rearrangements, the oxyallyl cation $\bf 6$ can be exploited in nucleophilic trapping 7,8 to afford multistereocentre-centre-containing products $\bf 8$ with up to three all-carbon $\bf 4^\circ$ -stereocentres. The rapid assembly of such levels of complexity from a prochiral starting material highlights the powerful activating and stereocontrolling influence of the $\bf Ox$ group. Using theoretical calculations, we show that the exceptional activating properties of $\bf Ox$ originate from a combination of covalent and non-covalent transition-state stabilizing effects.

Results and discussion

Two different carbometalation strategies were developed to give access to Ox-containing divinyl and aryl vinyl ketones 5 ($R^2 = alkyl/aryl$, Table 1). Firstly, Cu-catalyzed

carbomagnesiation of Ox-ynamides 4 with Grignard reagents gave 9 (M = MgBr); alternatively, Rh-catalyzed carbozincation of 4 with ZnEt₂ gave 9 (M = ZnEt).¹⁰ Addition of iodine to organometallics 9 (M = MgBr or ZnEt) gave the key building block alkenyliodides 10a (68%) and 10b (79%). Carbonylative Stille coupling (Method A) of 10a and 10b with tributyl(cyclohexen-1-yl)stannane afforded divinyl ketones 5a/ \mathbf{a}' and $\mathbf{5b/b'}$, respectively, each as a 5 : 1 mixture of E/Z-isomers about the Ox-substituted double bond (entries 1-4).11 Despite this partial isomerization, the major isomers, 5a and 5b, were isolated in 55% and 52% yield, respectively. All other divinyl and aryl vinyl ketones 5 shown in Table 1 were accessed by reaction of 9 (M = MgBr) with the corresponding aldehyde followed by Dess-Martin periodinane oxidation of the crude alcohols (Method B) giving 5c-i in yields of 31-91% (entries 5-12).

Table 1 Synthesis of Nazarov substrates 5 and their cyclization to 4° -stereocentre-containing cyclopentanoids 7

^a Nazarov substrates 5 formed from 4 using Method A (A) or Method B (B), as indicated. ^b Isolated as a minor isomer using Method A. ^c Cyclized with BF₃·THF or TfOH in CH_2Cl_2 at various temperatures (ranging from -78 °C to 40 °C) depending on acid and substrate; see text and ESI for details. ^d Diastereomeric ratio (dr) refers to stereochemistry at C1 relative to **Ox** (determined by ¹H NMR). Some products 7 were isolated as a mixture of C2-epimers, indicated by a wavy bond (see ESI for ratio), these give a single enantiomer upon **Ox** removal (eqn (1), ref. 7). ^e Isolated yield of C1-(S) isomer, an additional 24% was isolated as a 3:1 (R):(S)-C1 mix.

Nazarov cyclizations of divinyl and aryl vinyl ketones 5a-j were performed using either BF₃·THF or TfOH as catalyst in CH₂Cl₂, giving cyclopentanoids 7a-i (5f and 5j did not cyclize) containing one new 4°-stereocentre (Table 1). Broadly speaking, these Nazarov cyclizations performed very well, particularly where the "inner" substituent (R²) in 5 was Me, Et or Ph (BF₃·THF or TfOH). Use of TfOH as catalyst allowed the Nazarov cyclization to be conducted at temperatures as low as −78 °C, but generally the reactions were performed at 0 °C to rt or in refluxing CH2Cl2 (40 °C) using either TfOH or BF3-·THF.‡ The torquoselectivities were very high (dr > 20:1 for C1 relative to Ox), with the sole exception of 7d (dr = 2:1 (S):(R)-C1, entry 6). X-ray crystal structure and density functional theory (DFT) studies have shown that Ox auxiliaries of this configuration consistently favor anticlockwise conrotation leading to R¹-β stereochemistry (see below);^{7b} we have therefore assigned this stereochemistry to each product in Table 1. Most likely, the cyclization of 5d, which required heating to 40 °C due to the sterically encumbering isopropyl group ($R^2 = iPr$), gave lower selectivity due to partial Z/Eisomerization of the oxazolidinyl-alkene prior to cyclization, rather than because of poor stereoinduction by the auxiliary (see also below).

The presence of two aliphatic substituents on the tetrasubstituted alkene terminus, as in $\mathbf{5e}$, led to slower cyclization, but the stereoinduction remained high (entry 7). Diaryl-substituted alkene $\mathbf{5f}$ underwent undesired side reactions to give multiple minor products along with return of starting material (entry 8). In a number of cases, the presence of epimers at C2 (the carbon bearing \mathbf{Ox}) was apparent, but both epimers lead to the same product once the auxiliary is removed by reductive cleavage (see below). Cyclizations of electron-rich aryl vinyl ketones were successful (entries 9–11), even for the very hindered substrate $\mathbf{5h}$ where $\mathbf{R}^2 = i\mathbf{Pr}$. For the less activated aryl vinyl ketone $\mathbf{5j}$, alkene isomerization to form $\mathbf{\beta}$, $\mathbf{\gamma}$ -unsaturated ketone $\mathbf{11}$ became the dominant pathway and no Nazarov cyclization was observed.

As has been demonstrated in our previous study utilizing a diverse array of less substituted Nazarov products 7 ($R^2 = R^3 = H$), the oxazolidinone can be removed by reductive-cleavage using lithium naphthalenide (LiNph).^{7c} Two examples are given as part of this work (eqn (1)): reductive cleavage of the **Ox** group in **7c** and **7i** gave **12** (79%) and **13** (55%), respectively, both in high enantiomeric purity (er > 98 : 2).

Also, as per our previous work, additional stereochemical complexity can be built up by nucleophilic trapping of the intermediate oxyallyl cations $6.^{7c}$ Accordingly, the highly substituted divinyl ketone 5c was converted into the indoletrapped product 14 (75%) (eqn (2)). Notably, this tandem

sequence generates four new contiguous stereocentres, including two 4° -centres, with excellent control over both relative and absolute stereochemistry: only a single isomer was observed.

Having achieved stereoselective Nazarov cyclizations leading to products with adjacent 3° and 4°-stereocentres, we next addressed the formation of vicinal 4°-stereocentres. To prepare the fully substituted Nazarov substrate 17 we developed a convergent carbometalation approach starting from two alkynes: ynamide 4a and 3-hexyne (Scheme 2a). Cucatalyzed addition of MeMgBr to 4a, followed by in situ formylation with ethylformate, afforded 15 (52%) stereoselectively. Carboalumination of 3-hexyne to give 16,11 followed by 1,2-addition of 16 to 15 and oxidation with DMP, gave divinyl ketone 17 (71%). The C2-C3 double bond retained its Z stereochemistry while the C5-C6 double bond was formed as a 3:1 E:Z mixture. Separation of these isomers proved challenging; however, a pure sample of (2Z,5E)-17 was isolated in 24% yield (from 15). We also prepared the fully substituted ketone 20 (Scheme 2b) bearing a tethered nucleophile (electron-rich aryl group). Access to 20 commenced with formation of vinyl bromide 19 from bromoalcohol 18.13 Lithiation of 19, followed by addition to a solution of 15 and AlMe₃ (Lewis acid) and DMP oxidation of

Scheme 2 Syntheses of fully substituted Ox-divinvl ketones 17 and 20.

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Scheme 3 Nazarov cyclization of 17 in CH₂Cl₂.

the crude carbinol (not shown) afforded **20** (44%) as a single alkene-stereoisomer.¶

Nazarov cyclization of (2Z,5E)-17 with MeSO₃H (CH₂Cl₂, -78 °C) gave 21 as a complex mixture of C2,3-diastereomers, keto/enol-tautomers and E/Z-isomers (Scheme 3). Warming the mixture to ambient temperature resulted in a double Wagner-Meerwein shift of the C3-ethyl and C2-phenyl substituents in the reversibly formed oxyallyl cation 22 to give (3R)-23 and (3S)-23 in a 2:5 ratio (Scheme 3).14 The stereochemistry of these products was confirmed by X-ray crystallography of (3S)-23.‡ We believe that the origin of this epimeric mixture is partial double-bond isomerization of (2Z,5E)-17 to (2E,5E)-17 under the acidic conditions prior to Nazarov cyclization. While this isomerization was undesired, the rapid (<2 h) cyclisation of both isomers of 17 at -78 °C demonstrates the remarkable ability of the Ox group to activate the Nazarov reaction. Upon further experimentation with reaction conditions (acids and solvents) to avoid double-bond isomerization of (2Z,5E)-17 to (2E,5E)-17, we found that treatment of (2Z,5E)-17 with MeSO₃H in 1,4-dioxane with mild heating gave cyclopentanone 24 stereoselectively in 52% isolated yield (eqn (3)). The stereochemistry of (E)- and (Z)-24 were confirmed by X-ray crystallography and 2D NMR, respectively.‡ Replacing CH₂Cl₂ with 1,4-dioxane as solvent appears to exert different effects on the rates of the various competing reactions involved in the formation of 21, 23 and 24 (Scheme 3 and eqn (3)). Solvation of MeSO₃H by 1,4-dioxane likely reduces the rates of all of these reactions, however, its strongest effects appear to be the suppression of C2-C3 double-bond isomerization in 17 and Wagner-Meerwein rearrangement in 22, leading to the observed stereo- and chemoselective formation of 24.15 Cyclization of 20 (eqn (4)) under these conditions was also successful, yielding the intramolecularly trapped product 25 as the only product discernable by ¹H-NMR (53% isolated yield). Conversion of 20 to 25 forms two new rings and three contiguous 4°-stereocentres, underscoring the effectiveness of the Ox-controlled Nazarov reaction for synthesis of structurally complex, 4°-stereocentre-containing scaffolds. The asymmetric formation of three contiguous 4°-stereocentres entirely from prochiral carbons is a rare transformation; a Diels-Alder reaction reported by Nicolaou et al. is the only other example known to us.16

These **Ox**-promoted Nazarov cyclizations are remarkably facile, allowing efficient generation of sterically congested products at temperatures as low as -78 °C. This points to a powerful activating influence of the **Ox** auxiliary. In order to determine the origins of this activation, we performed DFT calculations (Fig. 1).‡ Calculations with M06-2X show that in the absence of an oxazolidinone, the activation energies (ΔG^{\ddagger}) for Nazarov cyclizations of **26–28** leading to zero, one, or

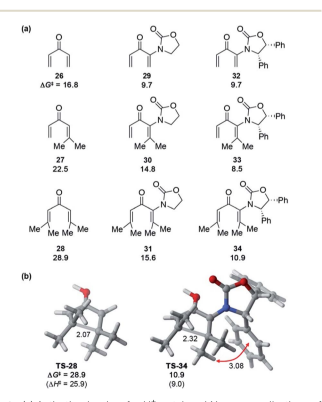


Fig. 1 (a) Activation barriers for H⁺-catalyzed Nazarov cyclizations of model divinyl ketones 26–34 and (b) transition states for cyclizations of 28 and 34, calculated with M06-2X/6-311+G(d,p)//M06-2X/6-31G(d) in implicit (SMD) dichloromethane. Distances in Å, ΔH^{\ddagger} and ΔG^{\ddagger} in kcal mol⁻¹.

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two 4°-centres are 16.8, 22.5, and 28.9 kcal mol⁻¹, respectively. Each new 4°-stereocentre raises the barrier by 6 kcal mol⁻¹.‡ An achiral oxazolidinone devoid of Ph substituents ($\mathbf{OxH_2}$, see $\mathbf{29}$ –31) lowers the cyclization barrier by 7–13 kcal mol⁻¹ ($\Delta G^{\ddagger} = 9.7$ –15.6 kcal mol⁻¹) relative to the oxazolidinone-free substrates, while the diphenyl-oxazolidinone (\mathbf{Ox} , see 32–34) provides further activation still, leading to cyclization barriers of only 8.5–10.9 kcal mol⁻¹. These very low barriers are consistent with the facile ring closures observed for 5, 17, and 20.

The transition states (TSs) for OxH2- and Ox-promoted cyclizations benefit from several stabilizing effects. Firstly, the nitrogen lone pair affords resonance stabilization of the incipient oxyallyl cation. Secondly, the oxazolidinone-containing TSs feature a longer forming C-C bond than the corresponding oxazolidinone-free TSs, leading to reduced steric repulsion between the Me groups about the forming C-C bond (see Fig. 1b). A third activating influence of Ox is evident from a comparison of the cyclizations of 33 and 34 (containing Ox) with those of 30 and 31 (containing OxH₂). The two Oxsubstituted TSs have ΔG^{\ddagger} values about 6 kcal mol⁻¹ lower than those of the corresponding OxH2 derivatives. The additional activation by Ox can be traced to a CH- π interaction in the TS between the "inner" substituent on C2 (R2, rotating downwards) and the nearby Ph substituent on Ox (see red arrow in Fig. 1). Together, these three TS-stabilizing influences of Ox make it an exceptionally powerful activating group, capable of reducing the barrier for vicinal 4°-centre formation by almost 18 kcal mol⁻¹ (28 vs. 34). Indeed, computations predict that when the R¹ substituent is an aryl group, like in many of our substrates (5, 17, and 20) (with R^2 = alkyl) the barrier for cyclization is even lower still.‡

Conclusions

To conclude, carbometalation of Ox-ynamides affords direct access to highly substituted Ox-divinyl and -aryl vinyl ketones, which undergo exceptionally facile Nazarov cyclizations leading to 4°-stereocentre-containing cyclopentanoids. In addition to the powerful activating and stereodirecting influence of Ox in the Nazarov cyclization, the Ox auxiliary helps suppress undesired Wagner-Meerwein rearrangements in the intermediate oxyallyl cations, and facilitates nucleophilic trapping of these intermediates enabling rapid assembly of multiple stereocentres (including vicinal 4°-stereocentres) with excellent stereochemical control. Theoretical studies allowed us to discover the electronic origin of the strong activating effect of the Ox, which is traced to a combination of covalent (lone pair donation to the incipient oxyallyl cation and reduced steric crowding about the newly forming bond) and non-covalent (CH- π interaction) effects which are generally applicable across most of the divinyl (or aryl vinyl) ketones reported here.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

‡ See the ESI.†

 \S The basis of the isomerization of the C5-alkenyl unit has not yet been fully discerned. Treatment of **16** with I₂ gave only the expected *E*-iodoalkene, whereas 1,2-addition of **16** to **15** gives the corresponding earbinol (not shown) as \sim 3:1 mixture of the *E*- and *Z*-isomer.

 \P In the absence of AlMe₃ the reaction affords mostly an acyl migration product involving ring opening of the oxazolidinone:

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