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Introduction

a-Diimines (1,4-diaza-1,3-dienes) are widely used as ligands for organometallic complexes,^{1,2} perhaps most notably in Brookhart-type catalysts for α -olefin polymerization.³⁻⁵ The α diimine ligand scaffold has proven to be remarkably versatile due to its facile stereoelectronic tunability.⁶–⁹ For example, modifications to the backbone and N-aryl substituents of α diimines can exert control over the molecular weight and microstructure of α -olefin polymers by attenuation of chain walking processes.^{1,10} As ligands, they are often redox noninnocent, which allows for richer and more complex redox processes.¹¹–¹⁴ a-Diimines are also precursors to valuable chiral 1,2-diamines through asymmetric hydrogenation,^{15,16} or to Nheterocyclic carbene (NHC) ligands through cyclization with paraformaldehyde and subsequent deprotonation.¹⁷–²⁰

Typically, a-diimines are synthesized through condensation of a-diketones and amines. Despite the apparent simplicity of this approach, accessing unsymmetrical α -diimines through stepwise condensations is synthetically challenging due to poor chemoselectivity.²¹ Furthermore, since imine formation is reversible, attempts at sequential selective condensations can result in complex mixtures (for example, see Fig. 1A and S95–

α -Diimine synthesis via titanium-mediated multicomponent diimination of alkynes with C-nitrosos†

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 α -Diimines are commonly used as supporting ligands for a variety of transition metal-catalyzed processes, most notably in α -olefin polymerization. They are also precursors to valuable synthetic targets, such as chiral 1,2-diamines. Their synthesis is usually performed through acid-catalyzed condensation of amines with α -diketones. Despite the simplicity of this approach, accessing unsymmetrical α -diimines is challenging. Herein, we report the Ti-mediated intermolecular diimination of alkynes to afford a variety of symmetrical and unsymmetrical α -diimines through the reaction of diazatitanacyclohexadiene intermediates with C-nitrosos. These diazatitanacycles can be readily accessed in situ via the multicomponent coupling of Ti=NR imidos with alkynes and nitriles. The formation of α -diimines is achieved through formal $[4 + 2]$ -cycloaddition of the C-nitroso to the Ti and γ -carbon of the diazatitanacyclohexadiene followed by two subsequent cycloreversion steps to eliminate nitrile and afford the α -diimine and a Ti oxo. **EDGE ARTICLE**
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S98†). Imines are also notoriously difficult to isolate because they are prone to hydrolysis, making methods that generate product mixtures impractical. Highlighting this problem, although amine condensation with glyoxal to afford α -diimines
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unsymmetric NHCs synthesized via condensation with glyoxal are rare.19,22,23

There are comparatively few examples of unsymmetrical α diimines, and these are mostly limited to modification of N-aryl substituents, usually involving very sterically encumbered groups,3,9,24 rather than substituents on the backbone (Fig. 1B), 11,25 although there are several examples from aldimine cross coupling.26,27 Selective trapping of unsymmetrical diimines via cyclization has also recently been reported.²⁸ Given this methodology gap, developing a route to unsymmetrical α -diimines from simple feedstocks would provide a useful tool for the development of more diverse ligand scaffolds and pharmaceutically relevant building blocks.^{3,15,16} Our group has reported several examples of Ti-catalyzed oxidative functionalizations of alkynes for the synthesis of multisubstituted N-heterocycles that overcome limitations of classical condensation reactions.²⁹–³² We envisioned that a complementary synthetic route to unsymmetrical a-diimines could be achieved through Ti-mediated oxidative diimination of alkynes. Examples of alkyne diamination or diimination are scarce³³⁻³⁶ despite many reports of alkene diamination.³⁷–⁴² To the best of our knowledge, there is only a single example of a multicomponent intermolecular alkyne diamination⁴³ and no examples of alkyne diimination. Nevertheless, there are several elegant examples of alkyne difunctionalizations⁴⁴ using Ti including alkyne carboamination⁴⁵⁻⁴⁷ and iminoamination⁴⁸ that provide motivation for further exploring Ti-catalyzed or -mediated diamination/diimination. Chemical Science

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Previously, we reported that diazatitanacyclohexadienes (prepared from the multicomponent coupling of Ti imidos, alkynes, and nitriles) could undergo oxidation-induced N–N coupling to yield pyrazoles (Fig. 2, top).³² In the interest of further expanding the utility of this unique intermediate, we have begun examining its reactivity with various group transfer reagents. Herein, we report the intermolecular diimination of alkynes by Ti imidos and C-nitrosos to afford unsymmetrical adiimines (Fig. 2, bottom). This diimination reaction proceeds through a cascading sequence of formal cycloaddition and retrocycloaddition reactions from this key diazatitanacyclohexadiene intermediate. This approach is a useful strategy for synthesizing unsymmetrical α -diimines, which are challenging to access through traditional condensation reactions.

Fig. 2 In situ generated diazatitanacyclohexadiene intermediates. (top) Previous work on oxidative N–N coupling to pyrazoles; (bottom) intermolecular diimination of alkynes (this work).

Results and discussion

Addition of C-nitrosos to diazatitanacyclohexadienes

Addition of PhNO (2a) to diazatitanacycle 1 resulted in rapid, near-quantitative formation of α -diimine 3a (81% by ¹H NMR)
with the concentrant formation of n tolunitrile (Fig. 3). It is with the concomitant formation of p -tolunitrile (Fig. 3). It is proposed that this occurs through $[4 + 2]$ -cycloaddition of PhNO to the Ti and γ -carbon of the ligand backbone, followed by [4 + 2]-retrocycloaddition to eliminate p-tolunitrile, and a second cycloreversion to afford 3a and a Ti=O species (vide infra).

The scope of the C-nitroso reactants examined is shown in Table 1. Conveniently, nitrosoarenes can be readily prepared from the corresponding aniline via Oxone® oxidation.⁴⁹ In most cases, the α -diimine products were obtained in very good isolated yields (70% to 85%).

Use of sterically demanding nitrosos 2b and 2c resulted in good yields of 3b and 3c (77% and 76%), which are commonly used as bulky α -diimine substituents.³ This method tolerates both electron-poor (3d–3f) and electron-rich (3g–3i) parasubstituted (3d-3i) nitrosoarenes, as well as *ortho-substituted* $(3j-3l)$ nitrosoarenes $(3l)$ is formed in a 6 : 1 ratio with its enamine tautomer). The reaction also proceeds cleanly with aliphatic nitrosos: 2-methyl-2-nitrosopropane 2n gives a mixture of tertbutyl substituted α -diimine 3n and its enamine tautomer in a 3.75 : 1 ratio (71%). Similarly, 1-nitrosoadamantane 2o afforded a-diimine 3o and its enamine tautomer in a 3.13 : 1 ratio (70%). While the yield of the reaction was mostly consistent irrespective of nitrosoarene substituent, there were a couple of exceptions. For the reaction with electron-rich 2i, the formation of α -diimine 3i (49% yield by ¹H NMR) was accompanied by the formation of unidentified side products that produced isolation formation of unidentified side products that precluded isolation. Also, the reaction with 2-nitrosopyridine 2m yielded only a small amount of 3m by ${}^{1}H$ NMR (20%) relative to the amount of ptolunitrile byproduct (85%).

In situ multicomponent diimine synthesis

Diazatitanacycles such as 1 can be synthesized via the multicomponent coupling of Ti imidos, alkynes, and nitriles.³² With this strategy, one-pot alkyne diiminations were carried out (Fig. 4 and Table 2).

Fig. 3 Reaction of a C-nitroso with 1 yields an α -diimine 3a via a sequence of cycloaddition/retrocycloaddition steps.

^a Conditions: 1 (0.2 mmol), 2a-o (0.2 mmol, 1 equiv.), 3 mL C₆H₆, N₂ atmosphere (glovebox). Isolated yields. $\frac{b}{b}$ Mixture of imine/enamine tautomers. See ESI for enamine characterization. c^c 1H NMR yield vs. 1,3,5-trimethoxybenzene internal standard = 0.2 M.

In an initial experiment, reactions with 1 equiv. MeCN led to moderate yields of 5b (40%) from reaction of 4b and PhNO (Fig. 4A). MeCN was chosen over p-tolCN (the formal nitrile component from the reactions in Table 1) because of the easier removal of MeCN in vacuo. However, several species from competing side reactions also formed that were difficult to separate from the desired product given the instability of diimines towards hydrolysis: pyrrole (D, 5%) from insertion of a second equivalent of alkyne;²⁹ imine from the hydroamination of 3-hexyne $(E, 6\%)$; and azobenzene $(F, 12\%)$ from direct metathesis of PhNO with leftover Ti imido.^{50,51} Using excess MeCN (10 equiv.) suppressed these competing side reactions (Fig. 4B) and led to cleaner formation of the product, with azobenzene as the predominant side-product after removal of volatiles. Conveniently, basic aqueous extraction can remove the Ti byproducts, and azobenzene can be easily removed by sublimation—avoiding the need to perform column chromatography on the sensitive products.

Based on this result, the scope of one-pot oxidative alkyne diimination with $[py_2Ticl_2(NPh)]_2$ and PhNO was examined

Fig. 4 Comparison of synthesis of 5b using (A) 1 equiv. or (B) 10 equiv. **MeCN**

(Table 2, 5a-5p). Here, the α -diimine yields are ultimately determined by the yield of in situ formed diazatitanacycle 1a-1p, as the subsequent oxidations are near-quantitative. Symmetrical internal alkynes with both alkyl and aryl substituents formed the respective α -diimines (5a–5d) in good isolated yields (51% to 63%).

Unsymmetrical internal alkyne 4e resulted in the formation of a single α -diimine isomer (5e) in a 52% yield. In contrast, α diimines 5f-5h give complex 1 H and 13 C NMR spectra, presumably due to formation of multiple imine stereoisomers. Analysis of 5f by NOESY (Fig. S53 and 54†) showed that chemical exchange occurs between each of these isomers by the presence of EXSY cross-peaks. Additionally, GC-MS revealed only one peak corresponding to the mass of 5f (Fig. S55†).

Imines are well-known to undergo rapid equilibrium between E/Z isomers through inversion (also called the lateral shift mechanism) in nonpolar solvents.⁵² Confirmation that the complex spectra of 5f–5h were a result of stereoisomer equilibration was obtained through further reaction of the isomer mixtures. For example, reaction of the 5f isomeric mixture with $ZnCl₂$ resulted in 93% yield of α -diimine adduct 6f (Fig. 5). The identities of diimines $5g$ and $5h$ were similarly confirmed via $ZnCl₂$ coordination (see ESI†).

Terminal alkynes 4j–4l were also examined. Trapping Ti \equiv NR + terminal alkyne [2 + 2] cycloadducts is challenging: terminal alkynes typically react faster to form pyrrole or alkyne trimerization products.⁵³ Nevertheless, 5k was obtained cleanly in a 68% yield, while 5j and 5l were obtained as a mixture of

^a Conditions: 0.2 mmol (0.5 equiv.) $[py_2Ticl_2(NPh)]_2$, 0.4 mmol (1 equiv.) alkyne, 4.0 mmol (10 equiv.) MeCN, 4 mL PhBr, 115 °C, 4 h, N_2 atmosphere (glovebox); then, 0.4 mmol (1 equiv.) PhNO, rt, 0.5 h. Isolated yields. ^b Mixture of stereoisomers. ^c Yield corrected for minor pyrrole impurities. α GC-FID yield (vs. 1,3,5-trimethoxybenzene standard).

stereoisomers with small pyrrole impurities (2.4% and 12% pyrrole, respectively).

1,5-Enyne 4m allows for clean formation of 5m (50%), while 1,6-enyne 4n results in formation of 5n (10%) as a mixture with competing carboamination⁴⁷ products $(6%)$. Here, the shorter linker of 4m compared to 4n prevents the intramolecular alkene insertion that would lead to carboamination. Conjugated enyne 4o yielded only trace 5o, while cyclooctyne 4p predominantly formed alkyne trimer and pyrrole, with only a trace amount of 5p.

5d was also synthesized via a telescoped in situ route from $TiCl_4$ (THF)₂, azobenzene, and Zn⁰ powder with only a moderate decrease in yield (Fig. 6). We have previously shown that this strategy is a benchtop-compatible approach to other Ti oxidative amination reactions.⁵⁴ This one-pot approach facilitates the in situ formation of diazatitanacycle intermediates such as 1 without the need for specialized equipment, making the synthesis of both α -diimines and pyrazoles³² more operationally simple.

Given that diazatitanacyclohexadiene intermediate formation is regioselective, $32,55-57$ this method could be used to prepare regioisomeric a-diimines through different combinations of alkynes, imidos, and nitrosos (Fig. 7). For example, reaction of 4k with $[py_2TiCl_2(Np-tol)]_2$ imido gives 5q (41%),

Fig. 5 Top: reaction of stereoisomeric mixture 5f with $ZnCl₂$ results in convergence to 6f. Bottom: 13 C NMR imine region of (a) 5f (C₆D₆) and (b) 6f (CDCl₃). Inset: crystal structure of 6f showing half of the asymmetric unit with cocrystallized solvent and hydrogens omitted for clarity.

Fig. 6 Telescoped one-pot synthesis of 5d from TiCl₄(THF)₂.

while its regioisomer 5r can be synthesized by using 2g and $[p_{y_2}Ticl_2(NPh)]_2$ (44%). Another regioisomer 5s can be prepared from $[py_2TiCl_2(N^tBu)]_2$ with a 50% yield of diimine stereoisomers, albeit with a small pyrrole impurity (4% yield).

Together, these reaction scopes demonstrate that a wide range of both symmetric and unsymmetric α -diimines can be accessed directly through a one-pot multicomponent reaction with a variety of Ti imidos, alkynes, and C-nitrosos.

Proposed mechanism for intermolecular diimination

A plausible mechanism and accompanying DFT calculations for the formation of 5a from 1a (IM1) and PhNO (2a) are shown in Fig. 8. The *in situ* formation of **IM1** from Ti imidos, alkynes, and nitriles through alkyne/imido $[2 + 2]$ -cycloaddition and nitrile insertion has been previously established.^{32,58-60} The formal $[4 +$ 2] cycloaddition of PhNO to IM1 occurs in a stepwise fashion. First, O-coordination of PhNO to Ti yields IM3. From IM3, the nucleophilic, electron-rich γ -carbon in the metallacycle backbone⁶¹ attacks the electrophilic N of coordinated PhNO, generating bicyclic IM4. This process can be further visualized using IBO analysis (Fig. 8a): coordination of PhNO results in a puckering of the γ -C orbital toward the N of coordinated PhNO.

Subsequently the new N–C σ -bond is formed in IM4, with simultaneous rearrangement of the Ti-N and N-O π -bonds to a new N–C π -bond and O lone pair, respectively.

The tendency of C-nitrosos to undergo Diels–Alder-type cycloaddition reactions is well known,^{62,63} but examples of this process with metallacycles are scarce.^{64,65} Titanium ß-diketiminate complexes, which differ from diazatitanacyclohexadienes by a degree of unsaturation, have also been observed to undergo $[4 + 2]$ -cycloaddition with ketenes.^{66,67}

IM4 then undergoes rate-determining $(12.6 \text{ kcal mol}^{-1})$ retro-[4 + 2] cycloaddition to extrude nitrile, forming

azaoxatitanacycle IM5. The retrocycloadditive nature of TS2 can also be visualized by IBO calculations (Fig. 8b), which show the elimination of nitrile by the breaking of C–C and Ti–N σ -bonds in IM4 and the formation of three new N–C, C–C, and Ti–N π bonds in IM5. Further IBO analysis is provided in Fig. S100.† A similar tandem $[4 + 2]$ -cycloaddition-cycloreversion process to eliminate nitriles has also been proposed in the synthesis of phosphinines.⁶⁸ IM5 then undergoes a haptotropic shift following loss of the coordinated nitrile $(IM6)$ generating η^2 .
(N.O.) bound $IM7$ Finally, N.O. bond, cleavers, results in (N,O)-bound IM7. Finally, N–O bond cleavage results in exothermic formation of a Ti=O species with the bound α -diimine product IM8. ⁶⁹ Cycloreversions of group IV heterometallacycles to yield $M = X (M = Ti, Zr; X = 0, N)$ are wellprecedented.⁷⁰–⁷⁵

Ultimately, nitrile serves as a promoter in the *in situ* reactions—first forming the key diazatitanacycle intermediate, and then being eliminated prior to product formation. Alternatively, instead of undergoing a $[4 + 2]$ -cycloaddition, a nitroso could directly insert into the $[2 + 2]$ alkyne/Ti=NR cycloadduct, bypassing the need for nitrile. Indeed, C-nitrosos undergo insertions with Ti and Zr metallacycles.^{76,77} However, a nitrilefree control reaction resulted in exclusive formation of azobenzene through metathesis of the Ti imido with PhNO (Fig. 9), making this route unlikely.^{50,51}

Fig. 8 Computed pathway for diimine formation (M06/6-311G(d,p)/SMD, 25 °C, C₆H₅Br). All free energies are referenced to **IM1** = 0.0 kcal mol⁻¹. (a) Intrinsic bond orbitals (IBOs) showing [4 + 2] cycloaddition between NO π -bond (red) orbital of coordinated PhNO and π -
backbone (green orbital) of IM3. (b) IBOs showing rearrangement of IM4 to backbone (green orbital) of IM3. (b) IBOs showing rearrangement of IM4 to IM5 via retro-[4 + 2] cycloaddition.

Fig. 9 Direct reaction of nitrosos with Ti imidos and alkynes results in formation of azobenzene.

Conclusions

In summary, we have demonstrated a one-pot multicomponent synthesis of α -diimines by the diimination of alkynes by Ti imidos and C-nitrosos. This reaction likely occurs by $[4 + 2]$ cycloaddition of a nitroso to the Ti and γ -carbon of a diazatitanacyclohexadiene intermediate, followed by two subsequent cycloreversion steps to eliminate nitrile and afford the adiimine and $Ti=O$. This is an attractive route to complex, unsymmetrical α -diimines that are difficult to obtain through classical condensation reactions, providing new strategies for ligand synthesis,^{3,18,20} and generally adding to the library of alkyne difunctionalization reactions. Efforts are ongoing to make this cycloaddition–retrocyclization strategy generalizable to other difunctionalizations. Chemical Science
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Author contributions

CWF: experimental data collection and analysis, computational analysis; DTE: computational analysis; EK: computational analysis; AJP: experimental data collection and analysis; YC: experimental data collection and analysis; IAT: project direction and all authors contributed to the writing, editing, and revision of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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