Catalyst-Controlled Stereoselective Barton–Kellogg Olefination
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Abstract: Overcrowded alkenes are expeditiously prepared by the versatile Barton–Kellogg olefination and have remarkable applications as functional molecules owing to their unique stereochemical features. The induced stereodynamics thereby enable the controlled motion of molecular switches and motors, while the high configurational stability prevents undesired isomeric scrambling. Bistricyclic aromatic enes are prototypical overcrowded alkenes with outstanding stereochemical properties, but their stereocontrolled preparation was thus far only feasible in stereospecific reactions and with chiral auxiliaries. Herein we report that direct catalyst control is achieved by a stereoselective Barton–Kellogg olefination with enantio- and diasterecontrol for various bistricyclic aromatic enes. Using Rh2(S-PTAD)4 as catalyst, several diazo compounds were selectively coupled with a thioketone to give one of the four anti-folded overcrowded alkene stereoisomers upon reduction. Complete stereodivergence was reached by catalyst control in combination with distinct thirane reductions to provide all four stereoisomers with e.r. values of up to 99:1. We envision that this strategy will enable the synthesis of topologically unique overcrowded alkenes for functional materials, catalysts, energy- and electron transfer, and bioactive compounds.

Introduction

The eminent Feringa motors and bistricyclic aromatic enes are overcrowded alkenes that typically adopt a characteristic anti-folded geometry upon complementary pyramidalization of the alkene carbons (Figure 1A).[1–3] Since the discovery of overcrowded alkenes,[4,5] the stereoisomerism and stereodynamics of bistricyclic aromatic enes and related systems have been studied in great detail.[6,7] When unsymmetrically disubstituted, the anti-folded structures exist as four stereoisomers in the form of two enantiomeric pairs of diastereomers (Figure 1B). Intriguingly, this isomerism results from the different alkene geometries (E or Z) together with their helicity (enantiomeric M or P) within an irreducible stereogenic unit characterized by its fourfold stereogenicity.[8]

In their pioneering work, Wynberg and Feringa uncovered that such anti-folded overcrowded alkenes exhibit particularly high configurational stability and that stereoisomer separation is feasible.[9] Because of their well-defined topology, the high thermal racemization barriers and the possibility to induce stereospecific isomerization by photochemical stimuli, they have been utilized as molecular switches and catalysts.[10,11] Enantiomerically enriched overcrowded alkenes were classically obtained by HPLC separation[12–14] of a racemate prepared by a Barton–Kellogg olefination.[15–17] When comprising stereocenters for instance in the form of a chiral auxiliary, the diastereoselective assembly of overcrowded alkenes provides an indirect stereospecific route to enantiomeric products by a range of synthetic methods, including the McMurry coupling and cascade cyclizations besides the prevalent Barton–Kellogg olefination (Figure 1C).[18–22] Indirect catalyst control was observed by Tanaka and co-workers in a rhodium-catalyzed cascade reaction with selectivity for the configuration of a stereocenter (Figure 1D).[23] Direct catalyst control over two stereoisomers was recently achieved by the Miller group in a dynamic kinetic resolution of a stereodivergent precursor by a peptide-catalyzed N-oxidation (Figure 1E).[24] Interestingly, the catalyst-controlled stereoselective synthesis of bistricyclic aromatic enes by the particularly effective Barton–Kellogg olefination has, to the best of our knowledge, not yet been described.

The Barton–Kellogg olefination essentially involves the coupling of diazo compounds with thioketones—typically in an uncatalyzed reaction—followed by the reduction of the resulting thirane intermediate.[25–26] Interestingly, a few reported examples indicated that the formation of Rh-carbenoids with Rh2(OAc)4 facilitates challenging Barton–Kellogg olefinations.[27–30] Related to the formation of the thirane intermediate, an analogous Rh2(OAc)2-catalyzed epoxidation was reported by Doyle and co-workers (Figure 1F).[31] Motivated by our interest in catalyst control over higher-order stereogenicity and the addressability of specific states of halted molecular motors,[32,33] we thus considered the feasibility of stereocontrol with the chiral dirhodium tetracarboxylate catalysts designed by the Davies group.[34–36] We specifically envisioned the possibility of an unprecedented stereoselective rhodium-catalyzed thirination[37–39] combined with a stereospecific reduction for a stereoselective Barton–Kellogg olefination. Our study was furthermore encouraged by the notable prospects for stereodivergent control to provide all four stereoisomers of anti-folded overcrowded alkenes from the same diazo- and thioketone substrates (Figure 1H). If successful, this method would circumvent

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tedious separations of enantiomers, resolutions or auxiliary methods to provide a most direct, general and stereoselective assembly of valuable overcrowded alkenes. We now disclose that the stereoselective Barton–Kellogg olefination is viable with rhodium catalysis and that a striking stereodivergence for pertinent overcrowded alkenes with fourfold stereogenicity is tractable.

Results and Discussion

We approached our first aim to achieve catalyst control in a stereoselective Barton–Kellogg olefination by developing an efficient synthesis of monosubstituted diazo substrates \( \mathbf{2} \) either by direct assembly or by diversification using cross-coupling reactions (see the Supporting Information for details). Thioketone substrate \( \mathbf{1} \) was subsequently coupled with the diazo compound \( \mathbf{2a} \) activated by dirhodium catalysts, where the outcome of the thiirane formation provided first insights into the stereoselectivity of the reactions (Table S1 in the Supporting Information). Different reductants\(^ {\text{40}} \) to convert the thiiranes to overcrowded alkenes were tested and the treatment with \( \text{P}(\text{NMMe}_2)_3 \) in toluene was thereby identified as ideal protocol. Whilst full conversion was achieved with \( \text{P}(\text{NMMe}_2)_3 \) in 3 h with excellent stereospecificity, the reduction with \( \text{PPh}_3 \) required long reaction times and led to loss of stereoisomeric enrichment (see the Supporting Information for the optimization of the reduction). The lack of diastereoselectivity in substrate-controlled reactions was also confirmed with \( \text{Rh}_2(\text{OAc})_4 \) (\( \mathbf{C}_1 \)) as catalyst, providing a 1:1 mixture of the racemic \( E \)- and \( Z \)-products (Figure 2, entry 1). In striking contrast, \( \text{Rh}_2(R\text{-DOSP})_4 \) (\( \mathbf{C}_2 \)) showed a 1:1.7 d.r. for the \( Z \)-isomer with excellent enantioselectivity (94:6 e.r.) and a 97% yield at a catalyst loading of 5 mol% (entry 2). Furthermore, deconstructing the processes by the analysis of the thiirane intermediate indicated that an identical level of stereoselectivity is already reached in the coupling step, rendering the thiirane reduction exceptionally stereospecific. Interestingly, \( \text{Rh}_2(S\text{-PTAD})_4 \) (\( \mathbf{C}_3 \)) provided an increased d.r. of 2.6:1 for the \( E \)-diastereomer with a 98:2 e.r. for the major overcrowded alkene diastereomer (entry 3, 95:5 e.r. for the minor \( Z \)-diastereomer). In comparison, the tetrachlorinated catalyst \( \text{Rh}_2(S\text{-TCPTAD})_4 \) (\( \mathbf{C}_4 \)) led to substantially lower stereocontrol (entry 4) and also dirhodium cyclopropane carboxylate catalysts \( \mathbf{C}_5 \) and \( \mathbf{C}_6 \) provided inferior selectivities (see the Supporting Information). Interestingly, \( \text{Rh}_2(S\text{-PTAD})_4 \) (\( \mathbf{C}_3 \)) was previously also identified as optimal catalyst for the stereoselective cyclopropanation of styrenes
We next varied the dimethoxyphenyl moiety to probe the P
revealing a catalyst-controlled selectivity towards P
analysis of a single crystal confirmed the
anti
4a
98:2 e.r. for the major E
Figure 2. Optimization of the catalyst-controlled stereoselective Barton–Kellogg olefination. [a] Reaction conditions: 1 (50.0 μmol, 1.0 equiv), 2a (1.5 equiv), Rh2L4, solvent, −78°C, 3 h, then P(NMe2)3 (50 equiv), toluene, 80°C or 60°C. [b] Determined for the crude product by 1H NMR
spectroscopy and confirmed by HPLC on a chiral stationary phase. [c] Determined for the isolated product by HPLC on a chiral stationary phase. [d] Yield over both steps for the isolated product. [e] Reduction at 80°C. [f] Reduction at 60°C.
with diaryldiazomethanes that bear donor and acceptor substituents to induce a differentiated alignment of the two aryl groups with respect to the rhodium carbene.[41] Using the catalyst C3, an increase of the temperature from −78°C to room temperature was found to strongly reduce stereoselectivity (see the Supporting Information). The coupling also showed a strong solvent effect and reactions in CH2Cl2 or THF both resulted in lower d.r. and e.r. values (entries 5 and 6), confirming that toluene is an ideal solvent for the catalyst-controlled stereoselective thiiranation. More dilute reactions led to a moderate improvement of diastereoc control (entry 7).
Moreover, the excellent stereospecificity and yield of the reduction remained when performed at 60°C, which allowed to further enhance the mild conditions of the method. Gratifyingly, the catalyst loading was successfully decreased to 2.5 mol% without loss of stereoselectivity, providing 3a in a d.r. of 2.7:1 and an e.r. of 99:1 for the major E-diastereomer (entry 8, 95:5 for the minor Z-isomer). When the catalyst loading was further reduced to 1 mol%, the selectivity was marginally compromised to an e.r. of 97.3 (entry 9) and the remainder of the study was therefore carried out with 2.5 mol% dirhodium catalysts.
Having identified the optimal conditions for the stereoselective catalyst-controlled Barton–Kellogg olefination (Figure 2, entry 8), the scope and limitations of the method were explored (Figure 3). The reaction scale was successfully increased from 50.0 μmol to 100 μmol without affecting selectivity, providing 4a in 95% yield with 2.7:1 d.r. and 98:2 e.r. for the major E-diastereomer. X-ray crystallographic analysis of a single crystal confirmed the anti-folded geometry and established the absolute configuration as (P-E)-4a, thus revealing a catalyst-controlled selectivity towards P-helicity. We next varied the dimethoxyphenyl moiety to probe the impact of electronic and steric alterations on the diazo building block. Interestingly, the rather small change in 4b led to a dramatic loss of stereoselectivity and also the di-ortho-methoxy-substituted 4c initially resulted in lower enantiocontrol, thus challenging the breadth of the method. Nonetheless, the catalyst Rh2[(R-BTPCP)2] (C5) restored selectivity and provided 4c in 95% yield in 4:1 d.r. with 89:11 e.r. for the major E-isomer. Since the ortho- and para- methoxy substituents exert a strong electronic influence, this higher stereoselectivity using C5 as complementary catalyst was likely observed due to the rigidity of the 2,6-substituted diazo substrate 2c. Moreover, X-ray crystallographic analysis confirmed that with Rh2[(R-BTPCP)2], the (M-E)-4c-configured product is formed. In contrast, we were pleased to find that the remainder of the scope studies could be performed with the Rh2(S-PTAD)2 catalyst (C3) under the previously optimized conditions, thus confirming the generality of the catalysi-controlled Barton–Kellogg olefination. The dichloroaryl moiety in 4d led to a notable enantioselectivity of 96:4 (E) and 95:5 (Z) when using catalyst C3. While 4-fluorophenyl substitution in 4e resulted in a lower level of stereocontrol, excellent results were achieved with 3,4,5-trifluorophenyl-substituted 4f which was obtained with 99:1 e.r. for both the E- and Z-isomers. Conversely, the phenyl-substituted overcrowded alkene 4g was obtained in 90% yield with lower enantiocontrol. Having evaluated the electronic and steric effects of various aryl substituents, sterically less-demanding substrates were examined. Remarkably, a chloro substituent was found to already ensure ample differentiation to provide bistricyclic aromatic ene 4h with 1.8:1 d.r. and 90:10 e.r. for the major E-diastereomer. Furthermore, in contrast to previous substrates, a pronounced difference of enantioenrichment of the E- and Z-diastereo-
mers was observed. Similar enantioselectivity and an improved diastereoselectivity was established when changing from chlorine to bromine substitution in $4_i$ where a 2.4:1 d.r. was measured. To our delight, also a strongly electron-withdrawing functionality was amenable and nitro-substituted $4_j$ showed excellent enantioselectivity (99:1 e.r.) for the major $P$-$E$-diastereomer. Methyl-substituted overcrowded alkene ($P$-$E$)-$4_k$, which has been reported as potential molecular switch, [7] was also obtained with 91:9 e.r. and a d.r. of 1.3:1. Notably, $4_l$ with an electron-withdrawing CF$_3$ group was accessible with similar enantioselectivity and an improved diastereomeric ratio. To systematically explore the scope and limitations of the method, also the most challenging disubstituted diazothioxanthene substrates $2_m$ and $2_n$ were employed in the stereoselective Barton–Kellogg olefination. While fluoro-dimethoxyphenyl substituted $4_m$ was obtained with a valuable level of enantiocontrol (88:12 e.r.), the reaction to brominated $4_n$ was less selective, indicating the boundaries of enantioselectivity of the method. Consistent with the stereoselectivity for acyclic rhodium diarylcarbenes, electronic effects of the substituents may impact stereocontrol by differentiating the torsion angles between the aryl groups and the rhodium carbene, albeit to a reduced extent within the cyclic structures compared to open analogues. Altogether, the comprehensive scope underscores the capacity of chiral dirhodium tetracarboxylate catalysts for diazo–thioketone couplings through Rh-carbenoids by the unique stereoselective thiiranation, whereas the breadth of the approach for distinct bistricyclic aromatic ene stereoisomers is highlighted by the remarkable stereospecificity of the thiirane reduction.

To additionally explore the stereochemical versatility of the developed stereoselective Barton–Kellogg olefination, we also aimed at stereodivergent control over all four stereoisomers of the overcrowded alkenes (Figure 4). The opposite enantiomeric pathway was expediently confirmed by using the Rh$_2$(R-PTAD)$_4$ catalyst ($en$-$C_3$), providing ($M$-$E$)-$4_a$ with the expected inverted enantioselectivity of 2:98. For the challenging redirection of diastereoselectivity, we investigated an olefin formation by means of a radical thiirane reduction. However, while the desulfurization of thiiranes to alkenes with Bu$_3$SnH and BEt$_3$ as initiator is known as particularly mild methodology, [44] it has to the best of our knowledge not been described for the synthesis of overcrowded alkenes. Nonetheless, an intriguing reversal of diastereoselectivity and an excellent stereospecificity were

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**Figure 3.** Scope and limitations of the catalyst-controlled stereoselective Barton–Kellogg olefination. [a] Reaction conditions: 1) $1$ (100 µmol, 1.0 equiv), $2$ (1.5 equiv), $C_3$ (2.5 mol%), toluene (100 mL), –78°C, 3–5 h; 2) P(NMe$_2$)$_3$, (50 equiv), toluene (10 mL), 60°C, 18 h. Yields are for the isolated product after both steps. The d.r. values were measured by $^1$H NMR spectroscopy for the crude product. The e.r. values were determined for the isolated product by HPLC on a chiral stationary phase. [b] The reaction was carried out with catalyst $C_5$. [c] The reaction was carried out with 1.0 equivalent of $2_f$. 

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found when employing the Bu₃SnH/BEt₃ system. Under optimized conditions, the diazo–thioketone coupling of 1 and 2a with Rh₂(S-PTAD)₄ (C3) followed by a radical thiirane reduction in CH₂Cl₂ at 78°C thereby provided the Z-configured overcrowded alkene (P-Z)-4a in 1:2.7 d.r. with an e.r. of 97:3. To also obtain the fourth stereoisomer, the enantioselectivity was inverted by using the Rh₂(R-PTAD)₄ catalyst (ent-C3) in combination with the radical Bu₃SnH/BEt₃ thiirane reduction, giving (M-Z)-4a in 81% yield with 3:97 e.r. for the major Z-diastereomer and 9:91 e.r. for the minor E-diastereomer.

At ambient temperature, none of the alkenes prepared in our studies showed a noticeable tendency for stereoisomerization, thus underscoring the extraordinary configurational stability of the investigated overcrowded alkenes. To determine the isomerization barriers, a sample of nearly enantiopure (P-Z)-4a (> 99:1 e.r.; 95:5 d.r.) in heptane was heated to 95°C, resulting in a stereospecific isomerization to (M-E)-4a.⁶,⁷ The process followed first-order kinetics with ΔG°‡ = 124.5 kJ mol⁻¹, reaching an equilibrium state with a (P-Z)-4a/(M-E)-4a ratio of 1:1:1. Similar P-Z/M-E isomerization was observed at room temperature under ambient light in the presence of iodine within 9 h. In contrast, the irradiation of a sample of (P-Z)-4a with a UV-A LED (370 nm, 43 W) resulted in complete isomerization between all four stereoisomers within 90 min, leading to a photostationary state (PSS) of E/Z = 45:55. With the distinct stereoisomerization processes established, we conclusively anticipate the integration of the defined overcrowded alkene stereoisomers as switchable moieties or tractable functional units for the design of molecular machinery. Having functional elements with defined starting positions controlled by stereoselective catalysis, interactions during assembly become adjustable, the analytics comprehensible and the particularly well-defined topology is expected to facilitate the construction of advanced functional molecules and materials.

**Conclusion**

The catalyst-controlled stereoselective Barton–Kellogg olefination for the high-yielding synthesis of overcrowded alkenes with fourfold stereogenicity was developed. The reaction provides access to distinct anti-folded bistricyclic aromatic enes and features a combination of a highly efficient thiranation using the Davies' Rh₂(S-PTAD)₄ catalyst and an exceptionally stereospecific thiranation. The scope of the method also comprises the possibility for divergent stereocatalysis, rendering all four stereoisomers accessible from the same substrates. Based on the generality of the Barton–Kellogg olefination and the viability of forming the central C=C double bond of the overcrowded alkene in a direct catalyst-controlled stereoselective coupling, we envision that the rational design and synthesis of various catalyst structures, bioactive compounds and functional molecular scaffolds are feasible with the method presented herein.

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Conflict of Interest

The authors declare no conflict of interest.

Keywords: Barton–Kellogg olefination · higher-order stereogenicity · overcrowded alkenes · stereodivergent catalysis · stereoselective catalysis

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