

Enantioselective Phenolic α -Oxidation Using H_2O_2 via an Unusual **Double Dearomatization Mechanism**

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Supporting Information

ABSTRACT: Feedstock aromatic compounds are compelling low-cost starting points from which molecular complexity can be generated rapidly via oxidative dearomatization. Oxidative dearomatizations commonly rely heavily on hypervalent iodine or heavy metals to provide the requisite thermodynamic driving force for overcoming aromatic stabilization energy. This article describes oxidative dearomatizations of 2-

(hydroxymethyl)phenols via their derived bis(dichloroacetates) using hydrogen peroxide as a mild oxidant that intercepts a transient quinone methide. A stereochemical study revealed that the reaction proceeds by a new mechanism relative to other phenol dearomatizations and is complementary to extant methods that rely on hypervalent iodine. Using a new chiral phasetransfer catalyst, the first asymmetric syntheses of 1-oxaspiro [2.5] octa-5,7-dien-4-ones were reported. The synthetic utility of the derived 1-oxaspiro [2.5] octadienones products is demonstrated in a downstream complexity-generating transformation.

1. INTRODUCTION

Oxidative dearomatizations of feedstock arenes, including phenols, are useful in delivering functionalized, complex organic building blocks. These processes often rely on excess, and in some cases costly, hypervalent iodine or heavy metalbased (i.e., lead and bismuth) reagents which can give rise to hazardous byproducts;² this characteristic may counterbalance or overshadow the benefit of using inexpensive feedstock precursors. Reactions using catalytic or heavy metal-free conditions with benign oxidants, such as oxygen or hydrogen peroxide, have seldom been explored, especially in asymmetric fashion.3 Hydrogen peroxide (H2O2) is especially appealing as an oxidant due to its high efficiency, abundance, and favorable byproduct profile (i.e., H₂O).⁴

A recent report from these laboratories employed the Adler-Becker oxidation as the initiating step in a cascade sequence for the synthesis of highly functionalized heterocycles.5 The Adler-Becker oxidation utilizes stoichiometric sodium metaperiodate (NaIO₄), a hypervalent iodine species, to convert 2-(hydroxymethyl)phenols 1, also referred to as salicyl alcohols, into racemic, dearomatized 1-oxaspiro[2.5]octa-5,7-dien-4-ones 4 (spiroepoxydienones) (Scheme 1a),6 a motif which is readily found in a number of biologically active natural products (Scheme 1c). While these oxidation products have been broadly deployed due to their functionalizable dienone motif and proclivity to participate in a variety of cycloaddition reactions,8 the lack of access to enantioenriched spiroepoxydienones limits their applicability.

The favorable attributes of the (hydroxymethyl)phenol dearomatization and the interest in enantioenriched spiroepoxydienones led to the development of the hypothesis outlined in Scheme 1b. The reaction design imagines an enabling and underexplored intersection between transient quinone methides and mechanistically validated asymmetric nucleophilic epoxidations using basic H₂O₂.

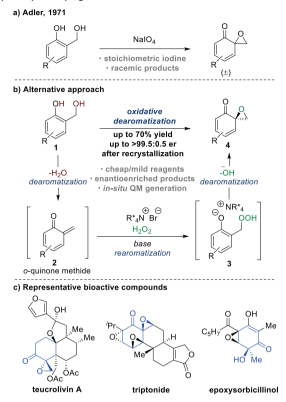
ortho-Quinone methides 2 (oQMs) are dearomatized species that have frequently been employed in forming complex natural products and synthetically useful building blocks. strong driving force for rearomatization underlies the high reactivity of the enone toward [4 + 2]-cycloadditions and 1,4-conjugate additions. 12 In almost all cases, these transformations irreversibly reset the aromaticity of the resultant system, limiting further complexity-building transformations. Reactions involving oQMs resulting in isolable, dearomatized products are rare.

Because phenol 1 is formally related to its derived oQM 2 by dehydration, a key challenge to achieving the title process would be the identification of conditions that facilitate dehydrative QM formation under mild conditions: a new method of QM generation was deemed to be a prerequisite for success of the project.¹⁴ Scheme 1 moreover postulates that the reaction of a nonstabilized, 15 ephemeral QM with H₂O₂ under basic conditions would initially re-establish aromaticity affording hydroperoxide 3 but concurrently set the stage for heterolytic O-O bond cleavage induced by engagement at the phenolic α -carbon, thereby breaking aromaticity for the second time in the sequence and creating the spiroepoxide substructure (4). Employing an asymmetric ion-pairing phase-transfer catalyst with phenoxide 3 could selectively facilitate the O-O bond cleavage, affording the enantioenriched spiroepoxydienone.

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Scheme 1. Oxidative Dearomatization of 2-(Hydroxymethyl)phenols



2. RESULTS AND DISCUSSION

Considering this hypothesis, we converted primary alcohol 1a¹⁶ to its unstable monoacetate 5 in low yield. When phenol 5 was subjected to KOH and H₂O₂ in MeCN at −5 °C, the desired spiroepoxydienone 4a was observed (Table 1, entry 1). Diacetate 6 was prepared in 70% yield and exhibited better stability than 5. Diacetate 6 in turn gave 4a in somewhat higher yield relative to the preparation from the monoacetate 5 (entry 2). The efficiency of quinone methide formation could be affected by the rates of both the phenolic deacylation (loss of

Table 1. Identification of an Optimal Activating Group for Quinine Methide Formation

 R^{1}) and expulsion of $(-)OR^{2}$; the electronic characteristics of both groups should be critical. To accelerate both steps, the more electron-deficient mono- and dichloroacetate analogues were prepared and exhibited drastically improved intermediate and product yields (entries 3 and 4). Bis(trifluoroacetate) 9 performed at a modest level (entry 5); consequently, bis(dichloroacetate) analogue 8 was selected for deployment with additional phenols.¹⁷ Dichloroacetate merits some consideration of the molecular mass "sacrificed", but the attractiveness of this acid chloride as a dehydrating agent stems in no small part from its cheap access on scale, high yields (>90%), and wide applicability and the fact that bis-(dichloroacetates) 8 are stable and often exist as easily handled white solids.

Using the optimized racemic conditions identified in Table 1, alkyl-substituted 2-(hydroxymethyl)phenols afforded the highest yields of the desired spiroepoxide products (Table 2,

Table 2. Racemic Scope of Oxidative Dearomatization Using Bis(dichloroacetates) of 2-(Hydroxymethyl)phenols^a

^aReactions performed with 1.0 equiv of 8 and 3.0 equiv of both H₂O₂ and KOH in MeCN ($[8]_0 = 0.05$ M). Yields refer to isolated yields. ^b4 equiv of KOH was used. ^cProduct isolated as dimer. ^dSlow addition of a solution of 8 and H₂O₂ over the course of 1 h. ^e9 equiv of H₂O₂ was used. ^fDetermined by ¹H NMR spectroscopic analysis.

4a-c, i-k). Alkyl groups promote the formation of QMs while reducing the rate of detrimental dimerization processes.¹⁸ Electron-withdrawing substituents are reported to inhibit QM formation¹⁸ and lead to oligomerization under basic conditions;¹⁹ however, when using difluorophenol 8e, the desired product 4e was obtained (47%). In contrast, difluorophenol 1e failed to provide any discernible product when NaIO₄ was used, highlighting the complementary nature of this method relative to the Adler-Becker oxidation. Mixed alkyl and halogen substitution afforded similar yields when 9

^aIsolated yield. ^{b1}H NMR yield versus internal standard

equiv of peroxide was used (4f). Because benzylic substitution (R⁵) often promotes facile rearrangement of spiroepoxydienones to benzodioxoles,⁵ we used bicyclic substrates 8g-h to prevent rearrangement and observed good yields with excellent diastereoselectivity in 4g.

The substitution pattern around the o-spiroepoxydienone was a critical determinant of whether the product was isolated in monomeric or dimeric form. Compounds 4i–1 with no substitution at R⁴ generally favored dimerization upon isolation. Unsubstituted salicyl alcohol 4l afforded the lowest yield and resulted in a multitude of side products (i.e., oligomers, QM dimers). Substrates prone to dimerization required that bis(dichloroacetate) 8 and H_2O_2 were added over the course of 1 h to reduce excess QM accumulation in solution. Rapid addition (<1 min) of 8 and H_2O_2 to the KOH/MeCN mixture resulted in a 1:1 mixture of the dimer and chromane 10, the product of trapping of the spiroepoxydienone with excess QM in solution (Scheme 2).

Scheme 2. Observed Competition between Product Dimerization and Quinone Methide Trapping

With a mechanistically distinct phenolic oxidation in hand, we became interested in developing an asymmetric variant of the title process. Enantioselective transformations utilizing oQMs lacking methide substitution under basic conditions are limited due to their high reactivity and propensity to dimerize rapidly in solution. While asymmetric Weitz—Scheffer-type epoxidations are well established for chalcones and other β -substituted enones using cinchona alkaloid phase-transfer catalysts (PTCs), asymmetric epoxidations of enones lacking β -substitution are rare. Employing in situ-generated oQMs that lack substitution at the methide position presents a formidable challenge in controlling the stereochemistry of the resultant spiroepoxide.

Toward this end, we envisioned employing an asymmetric ion-pairing PTC with phenoxide 3 as a method for controlling the facial selectivity of heterolytic O–O bond cleavage, a mechanism closely related to phase-transfer-catalyzed α -enolate substitution reactions. Computational analyses of these enantioselective reactions have revealed tight catalyst control of the substrate and electrophile to direct the facial selectivity of the substitution. While PTC α -enolate substitution reactions frequently involve the coordination of external electrophiles, more recent examples employ internal electrophiles using cinchona alkaloid PTCs bearing a free hydroxyl group. Expression of the substrate alkaloid PTCs bearing a free hydroxyl group.

We therefore began our catalyst screening by employing various cinchona alkaloid PTCs with a free hydroxyl group. Gratifyingly, phase-transfer catalyst CN-1 with CH_2Cl_2 solvent using 30% aqueous H_2O_2 as the oxidant resulted in a 55:45 er (Table 3, entry 1). Further catalyst optimization revealed that

Table 3. Optimization of Enantioselective Oxidative Dearomatization^a

entry	cat.	mol %	[O]	T (°C)	er ^a	yield (%) ^b
1	CN-1	20	H_2O_2 (aq)	0	55:45	<10
2	CN-2	20	H_2O_2 (aq)	0	50:50	<10
3	CN-3	20	H_2O_2 (aq)	0	63:37	<10
4	CN-3	20	UHP	0	69:31	33
5	QN-1	20	UHP	0	85:15	50
6	QN-1	20	UHP	-10	87:13	55
7	QN-1	20	UHP	-20	80:20	43
8	DHQ-1	20	UHP	-10	87:13	52
9	QN-2	20	UHP	-10	85:13	70
10	QN-2	20	UHP	-40	87:13	85
11	QN-2	10	UHP	-40	87:13	86
	N + Br	✓ N C	:N-1: R = H :N-2: R = 3-OMe :N-3: R = 3,5-(CF ₃) ₂ F ₃ C	OH N Br	OMe

 $^a\mathrm{Reactions}$ were performed using 8a (0.50 mmol), KOH (1.5 mmol), UHP (1.5 mmol), and [8a] $_0$ = 0.05 M in CH $_2\mathrm{Cl}_2$. 8a was added over the course of 2.5 h. $^b\mathrm{er}$ was determined by chiral HPLC. $^c\mathrm{Isolated}$ vields.

more electron-deficient benzyl groups (i.e., CN-3) improved selectivity (entry 3). Employing urea·H₂O₂ (UHP) to limit water content gave appreciable increases in selectivities and yields. Switching to quinine as the cinchona alkaloid (QN-1) greatly improved the er while affording mediocre yields (entry 4). Cooling the reaction temperature to -20 °C resulted in lower yields and selectivities (entry 5). Dihydroquinine catalyst DHQ-1 was investigated to minimize potential in situ derivatization of the catalyst's olefin via a Diels-Alder cycloaddition with a transient quinone methide (entry 6). Disappointingly, this catalyst resulted in the same yields and selectivities as QN-1. Upon further analysis, increased catalyst loading resulted in increased side product formation. Characterization revealed putative oxazonine 11 resulting from nucleophilic attack on a generated QM by the catalyst's quinoline nitrogen followed by cleavage of the quinuclidine core (Scheme 3). To minimize the nucleophilicity of the quinoline nitrogen while simultaneously providing steric hindrance,²⁷ catalyst QN-2 with a trifluoromethyl group was developed (30% yield over 3 steps from quinine N-oxide), which considerably improved yields while maintaining selectivities (entry 7).

Using the optimized catalyst and reaction conditions, various bis(dichloroacetates) were evaluated for conversion to their derived enantioenriched epoxides (Table 4). Monomers 4a-d were all found to afford modest selectivities with high yields;

Scheme 3. Preventing Catalyst Decomposition

Table 4. Scope of Enantioselective Oxidative Dearomatization a,b,c

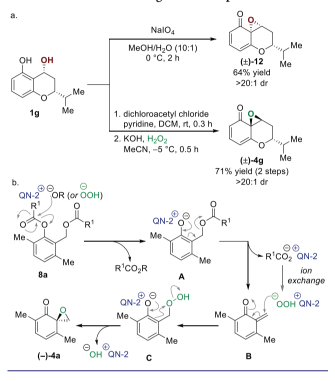
^aReactions were performed using **8** (0.50 mmol), **QN-2** (10 mol %), KOH (1.5 mmol), UHP (1.5 mmol), and [8] = 0.05 M in CH₂Cl₂, at -40 °C. ^ber was determined by chiral HPLC. ^cValues in parentheses represent recrystallized yields and enantiomeric ratios. ^a20 mol % QN-2.

however, good to excellent enantioselectivities with good mass recovery could be achieved via a single recrystallization. Similarly, dimer 4i exhibited slightly diminished selectivities that could be upgraded by a single recrystallization, affording excellent selectivities and modest recovery. It was quickly apparent that the stability and substitution pattern of the generated QM were important factors in determining the yield and enantioselectivities of the reaction. Attempts to access the enantioenriched unsubstituted dimer 4l resulted in low yields of the racemic dimer due to rapid QM oligomerization relative to epoxidation under the basic conditions. In contrast, bicyclic

substrate 4h gave excellent yields but afforded poor selectivity, although a change in the enantiodetermining step of the reaction should be noted with these β -substituted substrates.

An evaluation of the reaction mechanism was initiated by comparing the stereochemical outcome of the H_2O_2 -mediated oxidative dearomatization of $\mathbf{1g}$ to that when $NaIO_4$ was employed (Scheme 4a). $NaIO_4$ oxidation proceeded with

Scheme 4. Mechanistic Insights and Proposed Mechanism



stereoretention (12), while Weitz–Scheffer epoxidation of the planar QM intermediate proceeded with highly diastereose-lective inversion at the benzylic position (4g). Based upon this observation, we propose the initial deacylation of the phenolic dichloroacetate 8a to afford phenoxide A followed by formation of oQM B via elimination of the benzylic dichloroacetate. Conjugate addition by hydroperoxide affords the rearomatized phenoxide C that attacks the hydroperoxide to give the dearomatized epoxide C

The unique and critical role of PTC QN-2 in both QM generation and stereoselective epoxide formation is evident in this mechanism. To further understand the catalyst's role in promoting the observed stereoselectivity, the transition state of the epoxidation step between QN-2 and 3a was studied computationally using density functional theory (DFT) calculations at the level of $M062X^{28a}$ approximate functional and a compound Pople basis set. 28b,c

Upon analysis of the calculated major stereoisomer (Figure 1a), several important interactions emerge: (a) the hydroxide nucleofuge is stabilized by two significant hydrogen bonding interactions derived from the Ar–H on the electron-deficient $(CF_3)_2$ Ar ring (H···O distance 2.06 Å) and the benzylic C–H in close proximity to the ammonium cation (H···O distance 2.16 Å) as well as a weaker hydrogen bonding interaction from the C–H bond on the bridged quinuclidine (H···O distance 2.67 Å);²⁹ (b) the catalyst –OH group forms a strong hydrogen bond (H···O distance 1.83 Å) to the phenoxide of the substrate, orienting the hydroperoxide in close proximity to

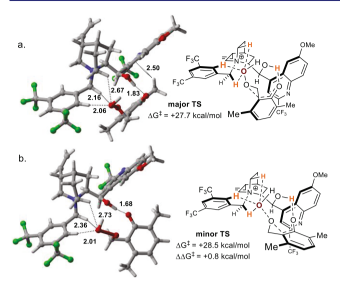


Figure 1. DFT-optimized stereodetermining transition states of QN-2 and 3a.

the three stabilizing hydrogen bond donors; and (c) the C-2 methyl group on 3a experiences an attractive $CH-\pi$ interaction with the quinoline ring (atom to plane distance of 2.50 Å). The apparent synergistic role of the R₃N⁺CH₂cationic subunits and the (F₃C)₂Ar-H is to create an unconventional trifurcated oxyanion hole to stabilize and accommodate the nascent alkoxide during O-O scission (vide infra). Such interactions for nucleofuge stabilization have been previously identified via DFT calculations but arise principally or solely from the R₃N⁺CH₂- cationic subunit.²

We were interested in comparing the stereodetermining catalyst-substrate interactions leading to the formation of the minor enantiomer to those leading to major isomer formation. These interactions were calculated to be similar to the major isomer (Figure 1b), with the distinction being a $\sim 90^{\circ}$ rotation of the substrate's aromatic ring to afford the opposite epoxide facial selectivity. The transition state leading to minor enantiomer formation was calculated to be 0.84 kcal/mol greater in energy than the major enantiomer. This higher energy transition state can be explained by (a) the loss of the attractive CH $-\pi$ interaction between the substrate C-2 methyl group and the quinoline ring and (b) the observed lengthening of the hydrogen bonding interaction between the hydroxide nucleofuge and the benzylic C-H (2.36 Å versus 2.16 Å), mitigating, in part, the stabilization of the leaving group. The increased transition state barrier due to the loss of the methyl $CH-\pi$ interaction is in accord with experimental evidence demonstrating reduced enantiocontrol with lack of alkyl substitution.

1-Oxaspiro[2.5]octa-5,7-dien-4-ones (4) are highly reactive species which readily participate in Michael additions,³ dihydroxylation,³¹ epoxide openings,³² and cycloadditions.³³ To further highlight the synthetic utility of this reaction in complexity building transformations, enantioenriched spiroepoxydienone generation was merged with subsequent basepromoted acyl-nitroso generation from 13³⁴ to realize a onepot oxidative dearomatization/acyl-nitroso Diels-Alder cycloaddition. The derived tricyclic oxazinanone 14 (Scheme 5) was obtained with excellent enantio- and diastereoselectivity after a single recrystallization. These tricyclic oxazinanones can be

Scheme 5. One-Pot Enantioselective Oxidative Dearomatization/Acyl-Nitroso Diels-Alder Cycloaddition

i. QN-2 (10 mol %), KOH (3 equiv), UHP (3 equiv), CH₂Cl₂, -40 °C, 4.5 h; then ii. 13 (1.5 equiv), -40 to 0 °C, 1.5 h

^aIsolated as an equilibrating mixture of diasteromers. ⁵ ^bValues in parentheses represent recrystallized yield, enantiomeric ratio, and diastereomeric ratio.

further elaborated to afford highly substituted cyclohexanone rings in a short number of synthetic steps.⁵

3. CONCLUSION

We have developed an enantioselective oxidative dearomatization of 2-(hydroxymethyl)phenols using H₂O₂ to afford stable, dearomatized 1-oxaspiro 2.5 octadienones employing a basepromoted in situOM activation technique. This reaction highlights the use of a mild and convenient oxidant to afford synthetically useful, dearomatized spiroepoxydienones. By using a new cinchona alkaloid-derived phase-transfer catalyst, the reaction allows for access to enantioenriched o-spiroepoxydienones which were previously inaccessible via the Adler-Becker oxidation. DFT calculations revealed a highly organized transition state involving a unique tripartite stabilization of the hydroxide leaving group leading to the observed facial selectivity. The synthetic utility of this method for rapid complexity generation has been demonstrated by preparing an enantioenriched tricyclic oxazinanone. This chemistry demonstrates the potential for complementary enantioselective dearomative processes involving quinone methides, and our laboratory is currently exploring these possibilities.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.8b13006.

CIF file giving data for compound 4i (CIF) Experimental procedures, characterization, and spectral data for all new chemical compounds as well as crystal data and data collection parameters (PDF)

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REFERENCES

- (1) (a) Vo, N. T.; Pace, R. D. M.; O'Har, F.; Gaunt, M. An Enantioselective Organocatalytic Oxidative Dearomatization Strategy. J. Am. Chem. Soc. 2008, 130, 404. (b) Jackson, S. K.; Wu, K.; Pettus, T. R. R. Sequential Reactions Initiated by Oxidative Dearomatization. Biomimicry or Artifact? In Biomimetic Organic Synthesis; Poupon, E., Nay, B., Ed.; Wiley-VCH Verlag & Co.: Weinheim, Germany, 2011; pp 723-749. (c) Roche, S. P.; Porco, J. A. Dearomatization Strategies in the Synthesis of Complex Natural Products. Angew. Chem., Int. Ed. 2011, 50, 4068.
- (2) (a) Varvoglis, A. Hypervalent Iodine in Organic Synthesis; Academic Press, Inc.: San Diego, CA, 1997. (b) Feldman, K. S. Cyclization Pathways of a (Z)-Stilbene-Derived Bis(orthoquinone monoketal). J. Org. Chem. 1997, 62, 4983. (c) Tchounwou, P. B.; Yedjou, C. G.; Patolla, A. K.; Sutton, D. J. Heavy metal toxicity and the environment. EXS 2012, 101, 133.
- (3) Dong, S.; Zhu, J.; Porco, J. A. Enantioselective Synthesis of Bicyclo [2.2.2] octenones Using a Copper-Mediated Oxidative Dearomatization/[4 + 2] Dimerization Cascade. J. Am. Chem. Soc. 2008,
- (4) (a) Goti, A.; Cardona, F. Hydrogen Peroxide in Green Oxidation Reactions: Recent Catalytic Processes. In Green Chemical Reactions; Tundo, P., Esposito, V., Eds.; Springer: Dordrecht, The Netherlands, 2008; pp 191-212. (b) Tsuji, T.; Zaoputra, A. A.; Hitomi, Y.; Mieda, K.; Ogura, T.; Shiota, Y.; Yoshizawa, K.; Sato, H.; Kodera, M. Specific Enhancement of Catalytic Activity by a Dicopper Core: Selective Hydroxylation of Benzene to Phenol with Hydrogen Peroxide. Angew. Chem., Int. Ed. 2017, 56, 7779.
- (5) Good, S. N.; Sharpe, R. J.; Johnson, J. S. Highly Functionalized Tricyclic Oxazinanones via Pairwise Oxidative Dearomatization and N-Hydroxycarbamate Dehydrogenation: Molecular Diversity Inspired by Tetrodotoxin. J. Am. Chem. Soc. 2017, 139, 12422.
- (6) (a) Adler, E.; Brasen, S.; Miyake, H. Periodate Oxidation of Phenols. IX. Oxidation of o-(omega-Hydroxyalkyl)phenols. Acta Chem. Scand. 1971, 25, 2055. (b) See the Supporting Information for a photochemical stability study of o-spiroepoxydienones.
- (7) (a) Bruno, M.; Omar, A. A.; Perales, A.; Piozzi, F.; Rodriguez, B.; Savona, G.; de la Torre, M. C. Neo-clerodane diterpenoids from Teucrium oliverianum. Phytochemistry 1991, 30, 275. (b) Kupchan, S. M.; Court, W. A.; Dailey, R. G., Jr.; Gilmore, C. J.; Bryan, R. F. Tumor inhibitors. LXXIV. Triptolide and tripdiolide, novel antileukemic diterpenoid triepoxides from Tripterygium wilfordii. J. Am. Chem. Soc. 1972, 94, 7194. (c) Sperry, S.; Samuels, G. J.; Crews, P. Vertinoid Polyketides from the Saltwater Culture of the Fungus Trichoderma longibrachiatum Separated from a Haliclona Marine Sponge. J. Org. Chem. 1998, 63, 10011.
- (8) (a) Corey, E. J.; Dittami, J. P. Total synthesis of (\pm) -ovalicin. J. Am. Chem. Soc. 1985, 107, 256. (b) Shair, M. D.; Danishefsky, S. J. Observations in the Chemistry and Biology of Cyclic Enediyne Antibiotics: Total Syntheses of Calicheamicin $\gamma_1^{\ \text{I}}$ and Dynemicin A. J. Org. Chem. 1996, 61, 16. (c) Singh, V. Spiroepoxycyclohexa-2,4dienones in Organic Synthesis. Acc. Chem. Res. 1999, 32, 324. (d) Yang, D.; Ye, X.; Xu, M. Enantioselective Total Synthesis of (-)-Triptolide, (-)-Triptonide, (+)-Triptophenolide, and (+)-Triptoquinonide. J. Org. Chem. 2000, 65, 2208.

- (9) Weitz, E.; Scheffer, A. Über die Einwirkung von alkalischem Wasserstoffsuperoxyd auf ungesättigte Verbindungen. Ber. Dtsch. Chem. Ges. B 1921, 54, 2327.
- (10) For excellent reviews on QMs, see: (a) Willis, N. J.; Bray, C. D. ortho-Quinone Methides in Natural Product Synthesis. Chem. - Eur. J. 2012, 18, 9160. (b) Bai, W.; David, J. G.; Feng, Z.; Weaver, M.; Wu, K.; Pettus, T. R. R. The Domestication of ortho-Quinone Methides. Acc. Chem. Res. 2014, 47, 3655. (c) Caruana, L.; Fochi, M.; Bernardi, L. The Emergence of Quinone Methides in Asymmetric Organocatalysis. Molecules 2015, 20, 11733. (d) Singh, M. S.; Nagaraju, A.; Anand, N.; Chowdhury, S. ortho-Quinone methide (o-QM): a highly reactive, ephemeral and versatile intermediate in organic synthesis. RSC Adv. 2014, 4, 55924.
- (11) Spence, J. T. J.; George, J. H. Total Synthesis of Peniphenones A-D via Biomimetic Reactions of a Common o-Quinone Methide Intermediate. Org. Lett. 2015, 17, 5970.
- (12) Guo, W.; Wu, B.; Zhou, X.; Chen, P.; Wang, X.; Zhou, Y.; Liu, Y.; Li, C. Formal Asymmetric Catalytic Thiolation with a Bifunctional Catalyst at a Water-Oil Interface: Synthesis of Benzyl Thiols. Angew. Chem., Int. Ed. 2015, 54, 4522.
- (13) For examples of nucleophilic addition to stable pQMs resulting in dearomatized products, see: (a) Yuan, Z.; Fang, X.; Wu, J.; Hequan, Y.; Lin, A. 1,6-Conjugated Addition-Mediated [2 + 1] Annulation: Approach to Spiro[2.5]octa-4,7-dien-6-one. J. Org. Chem. 2015, 80, 11123. (b) Ma, C.; Huang, Y.; Zhao, Y. Stereoselective 1,6-Conjugate Addition/Annulation of para-Quinone Methides with Vinyl Epoxides/Cyclopropanes. ACS Catal. 2016, 6, 6408. (c) Roiser, L.; Waser, M. Enantioselective Spirocyclopropanation of para-Quinone Methides Using Ammonium Ylides. Org. Lett. 2017, 19, 2338.
- (14) (a) Toteva, M. M.; Richard, J. P. The Generation and Reactions of Quinone Methides. Adv. Phys. Org. Chem. 2011, 45, 39. (b) Jaworski, A. A.; Scheidt, K. A. Emerging Roles of in Situ Generated Quinone Methides in Metal-Free Catalysis. J. Org. Chem. 2016, 81, 10145.
- (15) Jurd, L. Quinones and quinone-methides—I: Cyclization and dimerisation of crystalline ortho-quinone methides from phenol oxidation reactions. Tetrahedron 1977, 33, 163.
- (16) Due to the propensity of spiroepoxydienones to dimerize, 6,20 3,6-dimethylsalicyl alcohol (1) was selected as a model substrate since dimerization of 4a is slow and the monomer is stable for long periods.5
- (17) See the Supporting Information for detailed optimization of the racemic oxidative dearomatization reaction.
- (18) Weinert, E. E.; Dondi, R.; Colloredo-Melz, S.; Frankenfield, K. N.; Mitchell, C. H.; Freccero, M.; Rokita, S. E. Substituents on Quinone Methides Strongly Modulate Formation and Stability of Their Nucleophilic Adducts. J. Am. Chem. Soc. 2006, 128, 11940.
- (19) (a) Wan, P.; Hennig, D. Photocondensation of o-hydroxybenzyl alcohol in an alkaline medium: synthesis of phenol-formaldehyde resins. J. Chem. Soc., Chem. Commun. 1987, 939. (b) Chiang, Y.; Kresge, A. J.; Zhu, Y. Flash Photolytic Generation of ortho-Quinone Methide in Aqueous Solution and Study of Its Chemistry in that Medium. J. Am. Chem. Soc. 2001, 123, 8089.
- (20) For structural orientation of dimers, see: (a) Adler, E.; Holmberg, K.; et al. Periodate Oxidation of Phenols. X. Structural and Steric Orientation in the Diels-Alder Dimerization of o-Quinols. Acta Chem. Scand. 1971, 25, 2775. (b) Adler, E.; Holmberg, K.; et al. Diels-Alder Reactions of 2,4-Cyclohexadienones. I. Structural and Steric Orientation in the Dimerisation of 2,4-Cyclohexadienones. Acta Chem. Scand. 1974, 28b, 465.
- (21) For excellent examples of base-promoted asymmetric reactions involving oQMs lacking methide substitution, see: (a) Izquierdo, J.; Orue, A.; Scheidt, K. A. A Dual Lewis Base Activation Strategy for Enantioselective Carbene-Catalyzed Annulations. J. Am. Chem. Soc. 2013, 135, 10634. (b) Lee, A.; Scheidt, K. A. N-Heterocyclic carbenecatalyzed enantioselective annulations: a dual activation strategy for a formal [4 + 2] addition for dihydrocoumarins. Chem. Commun. 2015, 51, 3407. (c) Zhu, Y.; Zhang, L.; Luo, S. Asymmetric Retro-Claisen

Reaction by Chiral Primary Amine Catalysis. J. Am. Chem. Soc. 2016, 138, 3978.

- (22) Kelly, D. R.; Caroff, E.; Flood, R. W.; Heal, W.; Roberts, S. M. The isomerisation of (Z)-3-[²H₁]-phenylprop-2-enone as a measure of the rate of hydroperoxide addition in Weitz—Scheffer and Juliá—Colonna epoxidations. *Chem. Commun.* **2004**, 2016.
- (23) (a) Dolling, U. H.; Davis, P.; Grabowski, E. J. J. Efficient catalytic asymmetric alkylations. 1. Enantioselective synthesis of (+)-indacrinone via chiral phase-transfer catalysis. *J. Am. Chem. Soc.* **1984**, *106*, 446. (b) Corey, E. J.; Xu, F.; Noe, M. C. A Rational Approach to Catalytic Enantioselective Enolate Alkylation Using a Structurally Rigidified and Defined Chiral Quaternary Ammonium Salt under Phase Transfer Conditions. *J. Am. Chem. Soc.* **1997**, *119*, 12414. (c) Poulsen, T. B.; Bernardi, L.; Aleman, J.; Overgaard, J.; Jorgensen, K. A. Organocatalytic Asymmetric Direct α -Alkynylation of Cyclic β -Ketoesters. *J. Am. Chem. Soc.* **2007**, *129*, 441.
- (24) (a) de Frietas Martins, E.; Pliego, J. R., Jr Unraveling the Mechanism of the Cinchoninium Ion Asymmetric Phase-Transfer-Catalyzed Alkylation Reaction. ACS Catal. 2013, 3, 613. (b) He, C. Q.; Simon, A.; Lam, Y.-H.; Brunskill, A. P. J.; Yasuda, N.; Tan, J.; Hyde, A. M.; Sherer, E. C.; Houk, K. N. Model for the Enantioselectivity of Asymmetric Intramolecular Alkylations by Bis-Quaternized Cinchona Alkaloid-Derived Catalysts. J. Org. Chem. 2017, 82, 8645.
- (25) (a) Belyk, K. M.; Xiang, B.; Bulger, P. G.; Leonard, W. R., Jr; Balsells, J.; Yin, J.; Chen, C. Enantioselective Synthesis of (1R,2S)-1-Amino-2-vinylcyclopropanecarboxylic Acid Ethyl Ester (Vinyl-ACCA-OEt) by Asymmetric Phase-Transfer Catalyzed Cyclopropanation of (E)-N-Phenylmethyleneglycine Ethyl Ester. *Org. Process Res. Dev.* **2010**, *14*, 692. (b) Xiang, B.; Belyk, K. M.; Reamer, R. A.; Yasuda, N. Discovery and Application of Doubly Quaternized Cinchona-Alkaloid-Based Phase-Transfer Catalysts. *Angew. Chem., Int. Ed.* **2014**, 53, 8375. (c) Cullen, L. R.; Denmark, S. E. Development of a Phase-Transfer-Catalyzed, [2,3]-Wittig Rearrangement. *J. Org. Chem.* **2015**, *80*, 11818.
- (26) See the Supporting Information for a detailed optimization of the enantioselective dearomatization reaction.
- (27) (a) Johansson, C. C. C.; Bremeyer, N.; Ley, S. V.; Owen, D. R.; Smith, S. C.; Gaunt, M. J. Enantioselective Catalytic Intramolecular Cyclopropanation using Modified Cinchona Alkaloid Organocatalysts. *Angew. Chem., Int. Ed.* **2006**, *45*, 6024. (b) Attempts to only sterically inhibit rather than electronically deactivate the quinoline nitrogen by placing a phenyl group at the quinoline 2-position reduced the presence of **11** but failed to improve the yields.
- (28) (a) Zhao, Y.; Truhlar, D. G. The M06 suite of density functionals for main group thermochemistry, thermochemical kinetics, noncovalent interactions, excited states, and transition elements: two new functionals and systematic testing of four M06-class functionals and 12 other functionals. *Theor. Chem. Acc.* 2008, 120, 215. (b) Krishnan, R.; Binkley, J. S.; Seeger, R.; Pople, J. A. Self-consistent molecular orbital methods. XX. A basis set for correlated wave functions. *J. Chem. Phys.* 1980, 72, 650. (c) Rassolov, V. A.; Ratner, M. A.; Pople, J. A.; Redfern, P. C.; Curtiss, L. A. 6-31G* basis set for third-row atoms. *J. Comput. Chem.* 2001, 22, 976.
- (29) Jeffery, G. A. An Introduction to Hydrogen Bonding; Oxford University Press: Oxford, 1997.
- (30) Carroll, W. R.; Zhao, C.; Smith, M. D.; Pellechia, P. J.; Shimizu, K. D. A Molecular Balance for Measuring Aliphatic $CH-\pi$ Interactions. *Org. Lett.* **2011**, *13*, 4320.
- (31) Liotta, D. C. Branched Diepoxide Compounds for the Treatment of Inflammatory Disorders. U.S. Patent 20,100,324,133, December 23, 2010.
- (32) Cacioli, P.; Reiss, J. A. Reactions of 1-Oxaspiro[2.5]octa-5,7-dien-4-ones with nucleophiles. *Aust. J. Chem.* **1984**, *37*, 2525.
- (33) (a) Singh, V.; Lahiri, S.; Kane, V.; Stey, T.; Stalke, D. Efficient Stereoselective Synthesis of Novel Steroid-Polyquinane Hybrids. Org. Lett. 2003, 5, 2199. (b) Singh, V.; Chandra, G.; Mobin, S. Aromatics to Diquinanes: An Expeditious Synthesis of Tetramethylbicyclo[3.3.0]octane Framework of Ptychanolide. Synlett

- **2008**, 2008, 3111. (c) Jarhad, D.; Singh, V. π^4 s + π^2 s Cycloaddition of Spiroepoxycyclohexa-2,4-dienone, Radical Cyclization, and Oxidation—Aldol—Oxidation Cascade: Synthesis of BCDE Ring of Atropurpuran. *J. Org. Chem.* **2016**, *81*, 4304.
- (34) Sutton, A. D.; Williamson, M.; Weismiller, H.; Toscano, J. P. Optimization of HNO Production from *N,O*-bis-Acylated Hydroxylamine Derivatives. *Org. Lett.* **2012**, *14*, 472.