Total Synthesis of Wasabidienones B₁ and B₀ via SIBX-Mediated Hydroxylative Phenol Dearomatization

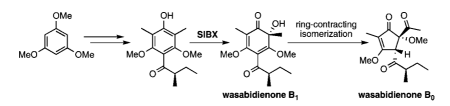
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ABSTRACT



The first total synthesis of the natural nondimerizing *o*-quinol (+)-wasabidienone B_1 was achieved from commercially available 1,3,5-trimethoxybenzene. The key dearomatizing transformation was efficiently accomplished via a hydroxylative phenol dearomatization reaction using the stabilized λ^5 -iodane reagent IBX (SIBX). (+)-Wasabidienone B_1 was then converted into its congener (-)-wasabidienone B_0 via an improved thermally induced ring-contracting isomerization reaction.

o-Quinols (i.e., 6-alkyl-6-hydroxycyclohexa-2,4-dienone derivatives) are seldom found as constituting motifs of natural products because of their high intrinsic reactivity and, notably, their propensity toward dimerization via Diels–Alder processes.¹ Among the very few examples of nondimerizing *o*-quinols are the hop constituents called humulones² and a series of fungal metabolites referred to as wasabidienones **1a**–**c** and **2a**,**b** (Figure 1). These fungal polyketides were isolated in the 1980s by Soga and co-workers from a potato culture of *Phoma wasabiae* Yokogi,³ a fungus responsible for the blackleg disease causing widespread destruction among cruciferous crops such as rape, cabbage, and wasabi

(Japanese horseradish). Some of these metabolites were also later isolated from *Phoma lingam*^{4a} and *Aspergillus viridinutans*^{4b} cultures. *Aspergillus parasiticus* and *P. lingam* were also reported to produce, respectively, aspersitin (**2c**), i.e., the C-8 epimer of wasabidienone B₂ (**2b**),^{4c} and phomaligin A (**2d**).^{4a} These fungal metabolites all feature an *o*-quinoltype cyclohexa-2,4-dienone core. They differ from one

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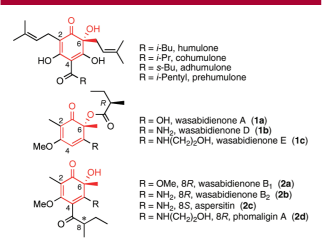


Figure 1. Examples of natural nondimerizing o-quinols.

another by the nature of their C-5 substituent and the positioning of their 2-methylbutanoyl side chain. In wasabidienones A (1a),^{3a-d} D (1b),^{3a} and E (1c),^{3e} this chiral unit acylates the tertiary alcohol function of the *o*-quinol moiety, while in wasabidienones B₁ (2a)^{3a,4b} and B₂ (2b),^{3a,4a} aspersitin (2c),^{4c} and phomaligin A (2d),^{4a} it is connected to the C-4 position of the cyclohexa-2,4-dienone core (Figure 1).

Efforts in the total synthesis of these naturally occurring *o*-quinols have, to the best of our knowledge, only resulted in that of wasabidienone A (**1a**)^{5a} and in that of (+)-aspersitin (**2c**).^{5b} In both cases, installation of the 2-methylbutanoyl moiety was achieved by Friedel–Crafts acylation using racemic 2-methylbutanoic acid. Sato and co-workers next developed an acyl rearrangement reaction from carbon to oxygen for the synthesis of **1a**, which was achieved in eight steps in ca. 4% overall yield from 1,2,3,5tetrahydroxybenzene.^{5a} In their six-step synthesis of (+)-**2c** obtained in ca. 5% overall yield from dimethylphloroglucinol, Büchi and co-workers relied on a dearomatizing Wessely oxidative acetoxylation (i.e., lead tetraacetate in acetic acid) of a phenolic intermediate to install the required oxygen atom at C-6.^{5b}

Oxygenating phenol dearomatization methods indeed stand out among the most straightforward tactics to access *o*quinols, as long as they can efficiently target the desired *o*-carbon center. Besides the Wessely oxidation, several other oxygenative dearomatizing reagents have been examined over the last 50 years to generate these systems from 2-alkylphenols.^{1a,6} More recently, the λ^5 -iodane 2-iodoxybenzoic acid (IBX) and its stabilized nonexplosive version (SIBX)⁷ were revealed as particularly appropriate reagents to promote hydroxylative phenol dearomatization (HPD) in a strictly ortho-selective manner.^{1b,8} For example, we previously applied the SIBX-mediated HPD reaction to the synthesis of natural *o*-quinol-derived [4 + 2] cyclodimers [e.g., (\pm) -biscarvacrol,^{1b,8a} (\pm) -grandifloracin,^{8a} and (+)-aquaticol^{8b}] in one step from phenolic precusors. Herein, we report the first application of this HPD reaction to the synthesis of the natural nondimerizing *o*-quinol (+)-wasabidienone B₁ (**2a**), followed by its conversion into (-)-wasabidienone B₀ (**14**).

Our choice of performing this HPD reaction on a phenolic substrate to access the title compounds may find some biomimetic justification in view of investigations on the biosynthetic origin of the C-6 oxygen atom in other o-quinolbased natural products such as the humulones^{9a,b} and dimerizing sorbicillinoids.^{9c,d} The synthesis began with the multigram-scale preparation of the known symmetric phenol 4^{10a} in four steps in 52% overall yield from commercially available 1,3,5-trimethoxybenzene (3) (Scheme 1, see the Supporting Information). Installation of the butyryl side chain was next envisioned through nucleophilic addition of a C-4 lithiated derivative of 4 on 2-methylbutanal (8). Unfortunately, lithiation conditions applied to a O-silvlated derivative^{10b} of **4** failed to generate the desired C-4 carbanion. We then relied on a standard halogen-metal exchange protocol and prepared bromobenzenes 7a,b in high yields from 4 via either bromination and silvlation or benzylation and bromination (Scheme 1). Coupling of these bromides with commercially available aldehyde (\pm) -8 or its (S)enantiomer, efficiently prepared by oxidation of commercial (S)-2-methylbutan-1-ol,¹¹ was achieved upon treatment with tert-butyllithium in pentane at low temperature, which led to 1.5:1 diastereomeric mixtures (unassigned) of benzylic alcohols 9. Of particular note is that these coupling conditions were nonepimerizing when using (S)-8. Debrominated benzene byproducts were successfully recycled in the preparation of **7a**,**b** (see the Supporting Information).

Benzylic alcohols (\pm) -**9a** and (S)-**9a**,**b** were next cleanly oxidized in excellent yields into their corresponding ketones (\pm) -**10a** and (S)-**10a**,**b**. This oxidation was performed in a 9:1 (v/v) mixture of THF/DMSO at room temperature for 1.5 h using SIBX, which we previously also unveiled as a

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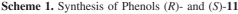
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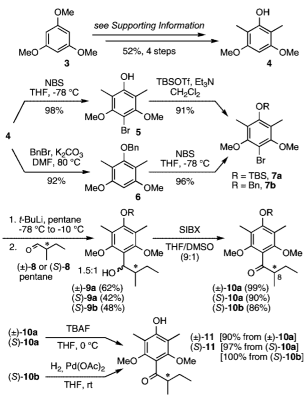
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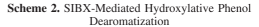
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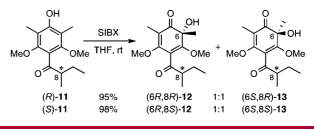




safe and efficient alternative to the use of IBX or the Dess-Martin reagent.^{7a} Again, no epimerization at the C-8 carbon center was observed under these conditions. Phenol (\pm) -11^{5b} was then obtained in a 90% yield through TBAFmediated desilylation of ketone (\pm) -10a; subsequent chiral HPLC resolution furnished both pure enantiomers (*R*)- and (*S*)-11 (see the Supporting Information). Phenol (*S*)-11 was also prepared in nearly quantitative yield via either desilylation of (*S*)-10a or hydrogenolytic debenzylation of (*S*)-10b (Scheme 1).

The key HPD reaction was next performed individually on (R)- and (S)-11 in THF at room temperature by adding SIBX (2.25 equiv, i.e., 1.10 equiv of IBX). The resulting suspension was stirred for 15 h, after which time a simple filtration, followed by an aqueous basic workup, afforded clean 1:1 diastereomeric mixtures of the corresponding o-quinols in 95% and 98% yields (Scheme 2). Separation by semipreparative HPLC furnished pure (6R, 8R)-12 and (6S,8R)-13 on the one hand and (6R,8S)-12 and (6S,8S)-13 on the other hand (see the Supporting Information). NMR analyses and optical rotation measurements of all four o-quinols led to the unambiguous identification of (6R, 8R)-12 as the targeted (+)-wasabidienone B_1 (2a, $[\alpha]^{20}$ _D +122.2).^{4b} The (6S,8R)-13, (6R,8S)-12, and (6S,8S)-13 stereoisomers were respectively identified as the non-natural (-)-6-epi-, (+)-8-epi-, and (-)-ent-wasabidienone B₁. These nondimerizing o-quinols could be stored as dry oils for several days at -20 °C without any noticeable degradation. However, they were found much less stable in solution, as

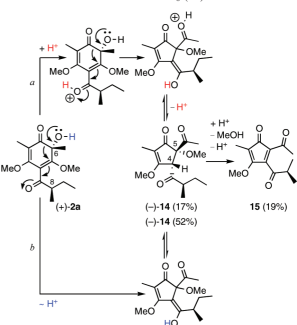




observed during ¹³C NMR analysis in CDCl₃ over several hours (vide infra). This instability was more pronounced for the 8*S*- than for the 8*R*-compounds and could explain the loss of material observed during the HPLC purification of the 8*S*-stereoisomers (see the Supporting Information).

We were satisfied with these results but yet intrigued by the instability of the synthetic wasabidienone B₁ (**2a**) and its stereoisomers. As alluded to above, (+)-**2a** indeed began to evolve slowly but noticeably into some other species in solution in CDCl₃ (ca. 0.08 M). After standing 10 days at room temperature, purification of the resulting mixture by standard silica gel chromatography afforded two compounds in 17 and 19% yield, respectively. Full characterization of the faster-eluting compound led to the identification of wasabidienone B₀ (**14**, $[\alpha]^{20}$ _D -296.7),¹² the naturally occurring ring-contracted isomer of **2a**.^{12,4b} ¹H NMR and mass analyses of the second compound indicated the cyclopentadienone structure **15**, resulting from methanol elimina-

Scheme 3. Isomerization of Wasabidienone B_1 (2a) into Wasabidienone B_0 (14)^{*a*}



 a Key: (a) CDCl₃, rt, 10 days, then SiO₂ column; (b) benzene, 80 °C, 5 days, then HPLC (OJ-H column).

tion from 14 (Scheme 3, path *a*). These data led us to reassess some earlier observations made by Soga and co-workers on the slow conversion over 20 h in refluxing benzene of (+)-2a into (-)-14 (33% yield).¹² We observed that wasabidienone B₁ (2a) is stable in freshly distilled benzene (ca. 0.02 M), even heated up to 60 °C, as evidenced by ¹H NMR monitoring in C₆D₆. Heating at refluxing temperature (80 °C) for 5 days is necessary to reach nearly complete conversion of 2a into 14 (Scheme 3, path *b*), which was purified in 52% on a cellulose-based chromatographic support (Chiralcel OJ-H column, see the Supporting Information). No elimination of methanol was observed under these conditions. It is probably the presence of some residual acidity in CDCl₃ that had previously promoted the formation of 15 (Scheme 3, path *a*).

In summary, the ortho-selective SIBX-mediated hydroxylative phenol dearomatization (HPD) reaction was successfully applied as a key (and biomimetic) step in the synthesis of the nondimerizing *o*-quinols (+)-wasabidienone B₁ (**2a**) and its three non-natural stereoisomers. The total synthesis of the fungal metabolite (+)-**2a** was thus achieved for the first time in 10 steps and ca. 5% overall yield from 1,3,5trimethoxybenzene. The conversion of (+)-**2a** into (-)wasabidienone B₀ (**14**) was performed by a thermally induced ring-contracting isomerization in an improved yield of 52%. This work demonstrates the applicability of this HPD reaction in the synthesis of *o*-quinolic natural products. The next challenge will consist in setting up an asymmetric version of this regioselective reaction. Our initial efforts in this avenue relied on substrate-controlled accesses to *o*-quinols in nonracemic forms.¹³ The convenient use and efficacy of SIBX described therein for mediating the HPD reaction have since prompted us to focus on the development of a reagent-controlled reaction using chiral iodanes. Results will be reported in due course.

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Supporting Information Available: Experimental procedures, NMR spectra, and HPLC traces for all new compounds. This material is available free of charge via the Internet at http://pubs.acs.org.

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