See discussions, stats, and author profiles for this publication at: https://www.researchgate.net/publication/24029672

# ChemInform Abstract: Asymmetric Oxidation of o-Alkylphenols with Chiral 2-(o-Iodoxyphenyl)-oxazolines

ARTICLE in ORGANIC LETTERS · MARCH 2009	
Impact Factor: 6.36 · DOI: 10.1021/ol8029092 · Source: PubMed	
CITATIONS	READS
66	29

#### **2 AUTHORS**, INCLUDING:



Vladimir Birman

Washington University in St. Louis

49 PUBLICATIONS 1,590 CITATIONS

SEE PROFILE



Org Lett. Author manuscript; available in PMC 2010 March 19.

Published in final edited form as:

Org Lett. 2009 March 19; 11(6): 1221–1223. doi:10.1021/o18029092.

# Asymmetric Oxidation of o-Alkylphenols with Chiral 2-(o-lodoxyphenyl)-Oxazolines

# Jagadish K. Boppisetti and Vladimir B. Birman

Department of Chemistry, Washington University, Campus Box 1134, One Brookings Drive, St. Louis, Missouri 63130

#### **Abstract**

A new class of chiral iodine (V) derivatives has been prepared. These compounds have been found to transform *ortho*-alkylphenols into *ortho*-quinol Diels-Alder dimers with significant levels of asymmetric induction.

A reaction of *o*-alkylphenols **1** with achiral oxidants, such as sodium periodate **4**,<sup>1</sup> benzeneseleninic anhydride **5**<sup>2</sup> and *o*-iodoxybenzoic acid (IBX) **6**,<sup>3</sup> produces *o*-quinols **2**, which dimerize spontaneously via a regio- and stereoselective intermolecular Diels-Alder reaction to give tricyclic products **3** (Scheme 1, Figure 1). The overall process is remarkable for the rapid increase in molecular complexity achieved in a single preparative step. Until a few years ago, enantioselective oxidative dearomatization of phenols remained unknown. In 2005, Porco et al. published a highly enantioselective oxidation of resorcinols with stoichiometric amounts of copper-spartein-dioxygen complex **7**.<sup>4a</sup> In 2008, they extended this process to *o*-alkylphenols producing o-quinol dimers with 99% ee's.<sup>4b</sup> The same year, Kita et al. developed a related process—asymmetric *ortho*-acetoxylation—using spirocyclic iodine (III) derivative **8**.<sup>5</sup> Prompted by these reports, we disclose the results of our recent studies in this area.

In connection with a total synthesis project, we sought to find new types of chiral oxidants capable of converting o-alkylphenols into o-quinol dimers in an enantioselective fashion. Inspired by the success of IBX in the racemic variant of this reaction,<sup>3</sup> we decided to explore the potential of chiral organoiodine (V) compounds. Although ortho-substituted iodoxybenzene derivatives, such as IBX  $\mathbf{6}^6$  and Dess-Martin periodinane,<sup>7</sup> are widely used in organic synthesis,<sup>8</sup> the utility of their chiral analogues remains relatively little explored. Chiral derivatives of IBX amide  $(\mathbf{9})^9$  and iodoxybenzene  $(\mathbf{10})^{10}$  (Figure 2) were synthesized and investigated by Zhdankin et al. The latter showed moderate levels of enantioselectivity in the oxidation of benzylic alcohols and thioanisole.

Iodoxybenzene derivatives *ortho*-substituted with a carbonyl, sulfonyl, or phosphonyl group, e.g., **10–13**, are known to form pseudocyclic structures in which the iodine forms a captodative bond with the adjacent oxygen atom. <sup>11</sup> Surprisingly, there have been no reports of the analogous iodoxybenzene derivatives with a *neutral* nitrogen ligand at the *ortho*-position. Given the ubiquitous use of oxazolines as chiral ligands for Lewis acids, <sup>12</sup> we decided to investigate the preparation and properties of Chiral 2-(*o*-IodoxyPhenyl)-Oxazolines, abbreviated as CIPO's (cf. **17**, Scheme 2).

Several 2-(*o*-iodophenyl)-oxazolines **16a–e** were prepared starting from chiral 2-amino alcohols **14a–e** following known general protocols. <sup>13</sup> Their oxidation with dimethyldioxirane <sup>11a,14</sup> yielded the desired CIPO's in moderate to good yields after chromatographic purification. The new compounds were obtained as white microcrystalline powders soluble in most organic solvents. The presence of the iodoxy group was evidenced by the diagnostic I=O stretches in IR (700–775 cm<sup>-1</sup> region), and the chemical shifts of the *ipso*-carbon (149 ppm) and the *ortho*-proton (8.3 ppm) in NMR. <sup>11</sup> All attempts to grow X-ray quality crystals of CIPO's have so far proved unsuccessful. Thus, the existence of a captodative bond between the oxazoline nitrogen and the iodoxy group cannot be ascertained at this point.

To our delight, reaction of 0.6 equiv. of *i*-Pr-CIPO **17a** with 1 equiv. of 2,6-dimethylphenol **1a** in chloroform produced the desired *o*-quinol dimer **3a** with an encouraging level of enantioselectivity (Table 1, entry 1). Notably, Porco's enantioselective oxidation method was reported to be unsuccessful in the case of this substrate. 4b

However, the reaction stopped at low conversions. Mindful of Barton's report on activation of the unsubstituted iodoxybenzene by trichloroacetic acid, 15 we investigated the effect of acid promoters (entries 2-5). Stoichiometric amounts of acetic acid were found to improve the reaction rate, whereas trifluoroacetic acid led to decomposition. Several solvents were screened next (entries 6–10). DME (1,2-dimethoxyethane) proved to be optimal giving 3a in 51% yield and 55% ee (entry 10). Again, addition of acetic acid was confirmed to be beneficial in this solvent (entries 10-12). At this point, all other available CIPO's were examined (entries 13-16). Only t-Bu-CIPO 17b displayed enhanced enantioselectivity compared to i-Pr-CIPO 17a and was therefore selected for further experimentation. The stoichiometry of the reaction with respect to the oxidant was initially uncertain, since in principle either one or both of the oxygen atoms of the iodoxy group could be utilized. An experiment with 2.5 equivalents of the substrate with respect to the oxidant confirmed that only one oxygen atom is utilized in the desired oxidation (entry 17). Accordingly, increasing the amount of the oxidant from 0.6 to 1.1 equivalents improved the yield of the dimer based on the substrate to 65% (entry 18). The rest of the starting material was mostly consumed by unidentified side reactions. The fate of the second oxygen atom of the iodoxy group remains unclear at this point. Only the fully reduced compounds, oxazoline 16b (54%) and amide 15b (19%) could be isolated after the reaction in entry 13.

Application of the optimized set of conditions to other dimethylphenols was investigated next. 2,5-Isomer **1b** produced the dimer (**3b**) with almost the same yield and enantioselectivity as **3a** (Scheme 3). Oxidation of 2,4-dimethylphenol **1c** attained 77% ee—the highest value observed in this study—albeit in a more modest yield. No dimer could be isolated from the reaction of 2,3-dimethylphenol **1d**, which is consistent with previous reports. <sup>3c,4b</sup> The absolute configurations of **3b** and **3c** were deduced from their signs of optical rotation. <sup>4b</sup> The configuration of **3a** was assigned by analogy.

In conclusion, we have synthesized the first examples of iodoxybenzene derivatives with chiral oxazoline groups at the *ortho*-position and applied them to the enantioselective oxidation of phenols. Although the ee values remain moderate at this point, our results demonstrate the

potential of the new class of chiral hypervalent iodine compounds in asymmetric synthesis. Further studies in this direction will be reported in due course.

### Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

#### References

- 1. Adler E, Holmberg K. Acta Chem. Scand 1971;25:2775.
- Barton DHR, Ley SV, Magnus PD, Rosenfeld MN. J. Chem. Soc., Perkin Trans 1977;1:567. [PubMed: 557487]
- 3. (a) Magdziak D, Rodriguez AA, Van De Water RW, Pettus TRR. Org. Lett 2002;4:285. [PubMed: 11796071] (b) Gagnepain J, Castet F, Quideau S. Angew. Chem., Int. Ed 2007;46:1533. (c) Lebrasseur N, Gagnepain J, Ozanne-Beaudenon A, Léger JM, Quideau S. J. Org. Chem 2007;72:6280. [PubMed: 17628111]
- (a) Zhu J, Grigoriadis NP, Lee JP, Porco JA Jr. J. Am. Chem. Soc 2005;127:9342. [PubMed: 15984841]
   (b) Dong S, Zhu J, Porco JA Jr. J. Am. Chem. Soc 2008;130:2738. [PubMed: 18266375]
- 5. Dohi T, Maruyama A, Takenaga N, Senami K, Minamitsuji Y, Fujioka H, Caemmerer SB, Kita Y. Angew. Chem., Int. Ed 2008;47:3787.
- 6. Frigerio M, Santagostino M. Tetrahedron Lett 1994;35:8019.
- 7. Dess DB, Martin JC. J. Am. Chem. Soc 1991;113:7287.
- 8. For recent reviews on hypervalent organoiodine compounds, see(a) Wirth T. Top. Curr. Chem 2003;224:1–248. (b) Zhdankin VV. Curr. Org. Synth 2005;2:121. (c) Ladziata U, Zhdankin VV. Arkivoc 2006;9:26. (d) Zhdankin VV, Stang PJ. Chem. Rev 2008;108:5299. [PubMed: 18986207]
- 9. Zhdankin VV, Smart JT, Zhao P, Kiprof P. Tetrahedron Lett 2000;41:5299.
- 10. Ladziata U, Carlson J, Zhdankin VV. Tetrahedron Lett 2006;47:6301.
- 11. (a) Zhdankin VV, Koposov AY, Litvinov DN, Ferguson MJ, McDonald R, Luu T, Tykwinski RR. J. Org. Chem 2005;70:6484. [PubMed: 16050713] (b) Macikenas D, Skrzypczak-Jankun E, Protasiewicz JD. Angew. Chem., Int. Ed 2000;39:2007. (c) Meprathu BV, Justik MW, Protasiewicz JD. Tetrahedron Lett 2005;46:5187.
- 12. For reviews, see(a) Gomez M, Muller G, Rocamora M. Coord. Chem.Rev 1999;193–195:769. (b) Desimoni G, Faita G, Jørgensen KA. Chem. Rev 2006;106:3561. [PubMed: 16967916]
- 13. Crosignani S, Young AC, Linclau B. Tetrahedron Lett 2004;45:9611.
- 14. Murray RW. Chem. Rev 1989;89:1187.
- $15.\,Barton\,DHR, Godfrey\,CRA, Morzycki\,JW, Motherwell\,WB, Stobie\,A.\,Tetrahedron\,Lett\,1982; 23:957.$

## **Acknowledgments**

 $The authors thank \ Dr. \ V. \ V. \ Zh dank in and \ Dr. \ V. \ N. \ Nemykin (University of Minnesota-Duluth) for helpful discussions and suggestions. Financial support for this study was provided in part by NIGMS (R01 GM072682).$ 

**Figure 1.** Reagents for oxidative dearomatization of *o*-alkylphenols

**Figure 2.** *o*-Substituted iodoxybenzene derivatives

OH
$$X = \begin{bmatrix} O \\ OH \\ X \end{bmatrix}$$

$$X = \begin{bmatrix} O \\ OH \\ R^2 \end{bmatrix}$$

$$X = \begin{bmatrix} (\pm)-2 \end{bmatrix}$$

$$X = \begin{bmatrix} (\pm)-2 \end{bmatrix}$$

$$X = \begin{bmatrix} (\pm)-3 \end{bmatrix}$$

$$X = \begin{bmatrix} (\pm)-3 \end{bmatrix}$$

**Scheme 1.** Transformation of *o*-alkylphenols into *o*-quinol dimers

**a**: R = i-Pr (S) **b**: R = t-Bu (S) **c**: R = Ph (R)

**d**:  $R = PhCH_2(R)$  **e**:  $R = Ph_2CH(R)$ 

**Scheme 2.** Synthesis of chiral 2-(*o*-iodoxyphenyl)-oxazolines

Asymmetric oxidation of isomeric dimethylphenols with *t*-Bu-CIPO **17b** 

NIH-PA Author Manuscript

Oxidation of 2,6-dimethylphenol 1a with CIPO's<sup>a</sup>

NIH-PA Author Manuscript

entry	oxidant	solvent	additve (equiv)	$^{9,6}_{ m vield} b$	% ee
	17a	CHCl <sub>3</sub>	none	29	30
2	17a	CHCI,	AcOH (0.25)	40	33
3	17a	CHCI,	AcOH (1.0)	47	36
4	17a	CHCI,	AcOH (3.0)	14	38
5	17a	CHCI	TFA (1.0)	dec	ND
9	17a	PhMe	AcOH (1.0)	43	53
7	17a	MeCN	AcOH (1.0)	51	43
8	17a	$CF_3CH_2OH$	AcOH (1.0)	58	17
6	17a	THF	AcOH (1.0)	36	56
10	17a	DME	AcOH (1.0)	51	55
11	17a	DME	none	29	55
12	17a	DME	AcOH (0.25)	43	56
13	17b	DME	AcOH (1.0)	43	65
14	17c	DME	AcOH (1.0)	36	46
15	17d	DME	AcOH (1.0)	36	24
16	17e	DME	AcOH (1.0)	36	32
17	17b	DME	AcOH (1.0)	26(64) <sup>C</sup>	62
18	17b	DME	AcOH (1.0)	$e^{5d}$	63

 $^{a}$ Conditions: 0.10 mmol 2,6-dimethylphenol 1a, 0.060 mmol CIPO 17a-e, 0.2 mL solvent, rt, 12 h, unless noted otherwise.

 $^{b}$  Yields are based on 1a, unless noted otherwise.

 $^{c}$  0.10 mmol 1a and 0.040 mmol 17b was used. The yield shown in parentheses is based on 17b.

 $d_{0.10 \text{ mmol } 1a}$  and 0.11 mmol 17b was used.