

A Stereoselective Intramolecular Halo-Etherification of Chiral Enamides in the Synthesis of Halogenated Cyclic Ethers

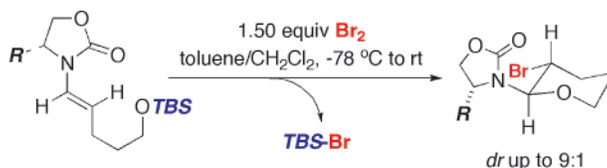
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ABSTRACT



A stereoselective halo-etherification of chiral enamides is described here. This work provides an approach to halogen containing cyclic ethers and reveals further mechanistic insights to the chemistry of chiral enamides.

With elegant advances in metal-catalyzed *N*-alkenylations,^{1–4} chiral enamides should emerge as versatile building blocks for developing stereoselective synthetic methods.^{5–12} We

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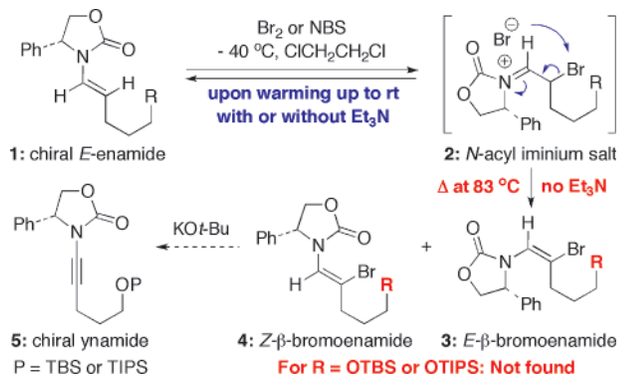
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encountered an interesting phenomenon involving enamides when we attempted years ago to transform enamides **1** to chiral ynamides¹³ via a sequence of bromination–elimination of the intermediate β -bromo-enamides **3** and **4** [Scheme 1].¹⁴ Two intriguing observations were made. First, the bromination behaved differently from a standard bromination of olefins. It was reversible with or without an amine base, and

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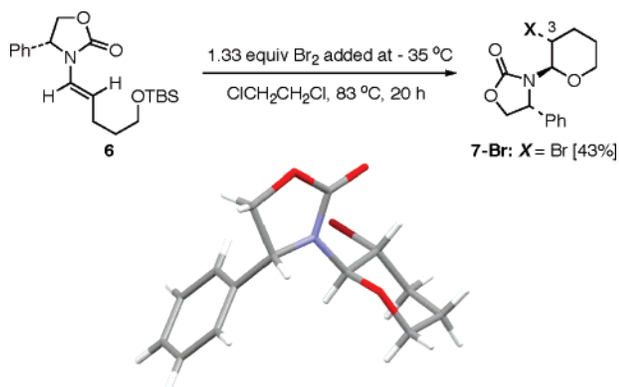
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Scheme 1. An Arrested Bromination–Elimination of Enamides

3 and **4** were obtained only if the reaction was heated at ≥ 80 °C without base. Although upon its addition the bromine color disappeared rapidly in a colorimetric titration manner, the color returned upon warming to rt. Second, when R is a TBSO or TIPSO group, the bromination led to a completely different product, thereby failing to access ynamides **5** via this protocol. While we suspected the *N*-acyl iminium salt intermediate **2** to be responsible for the reversibility issue [see blue arrows], we recently resolved the mystery product in the second observation. We reported here a stereoselective halo-etherification of chiral enamides in the synthesis of halogen containing cyclic ethers.

When chiral enamide **6**¹⁵ was subjected to bromination conditions in which 1.33 equiv of Br₂ was added at -35 °C and the resulting mixture was heated at 83 °C in ClCH₂-CH₂Cl for 20 h, pyran **7-Br** was isolated in 43% yield as a single isomer. Its relative stereochemistry was unambiguously assigned via single-crystal X-ray structure [Scheme 2]. No β-bromo-enamides related to **3** and **4** were found,

Scheme 2. Observation of a Desilylative Bromo-Etherification

and pyran **7-Br** implies a bromine-promoted desilylative cyclization had taken place instead.

Given that halogen-induced etherifications^{16,17} involving enamides, especially chiral enamides,^{18,19} have only been sparsely explored, we examined this reaction in greater detail. As shown in Table 1, we were able to establish that (1) the

Table 1. Temperatures and Choice of Activations

entry	activation	solvent	temp [°C]	time [h]	yield [%] ^a	pyrans	dr ^b	8 ^c [%]
1	Br ₂	toluene ^d	-78 to rt	12	92	7-Br	6.9:1	0
2		CH ₂ Cl ₂	-78	2	63	7-Br	6.9:1	0
3		CH ₂ Cl ₂	-45 ^e	2	73	7-Br	7.0:1	0
4		CH ₂ Cl ₂	-20 ^e	2	82	7-Br	7.0:1	0
5		CH ₂ Cl ₂	0 ^e	2	81	7-Br	7.0:1	0
6		CH ₂ Cl ₂	rt ^e	1	82	7-Br	6.9:1	0
7	NBS	CH ₂ Cl ₂	-78 to 40	24	NR	7-Br		0
8	NBS + TBAB ^f	CH ₂ Cl ₂	-45 to rt	6	42	7-Br	1:1.2	0
9	NCS	toluene ^d	-78 to 60	48	NR	7-Cl ^g		0
10	NCS + TBAC ^h	CH ₂ Cl ₂	-45 to rt	24	54	7-Cl	6.8:1	30 ⁱ
11	I ₂ ^j	CH ₂ Cl ₂	-78 to 40	24	30	7-I ^k	1:1.1	33 ^l
12	NIS	CH ₂ Cl ₂	-78 to 40	24	NR	7-I		0
13	ICl	CH ₂ Cl ₂	-78 to rt	2	70	7-I	6.0:1	0

^a Isolated yields. NR: no reactions. ^b Ratios assigned using ¹H and/or ¹³C NMR. ^c X = H at C3. See Scheme 2 for the structure. ^d Br₂ was added as a 0.5 M solution in CH₂Cl₂: toluene:CH₂Cl₂ is 3:1. ^e Br₂ was added at -78 °C. The reaction was stirred at the temperature and time indicated after the addition. ^f TBAB: tetra-*n*-butyl ammonium bromide; NBS:TBAB:6 = 1.5:1.5:1. NBS and TBAB were premixed at -78 °C for 10 min. ^g X = Cl at C3. ^h TBAC: tetra-*n*-butyl ammonium chloride; NCS:TBAC:6 = 1.5:1.5:1. NCS and TBAC were premixed at -78 °C for 10 min. ⁱ dr for **8** = 2:1. ^j 4 Å MS was used. ^k X = I at C3. ^l dr for **8** = 3:1.

bromo-etherification could take place readily at much lower temperatures with best yields obtained between -20 °C and rt with the diastereomeric ratio being independent of tem-

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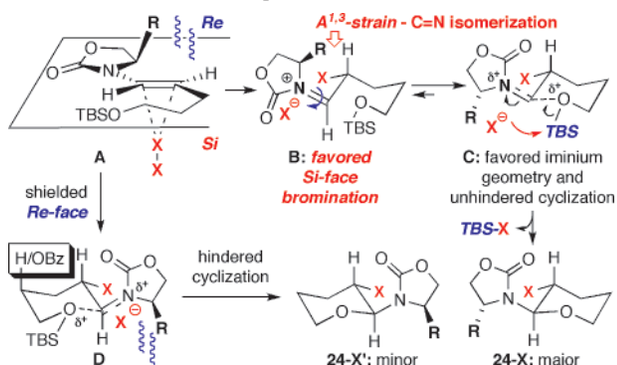
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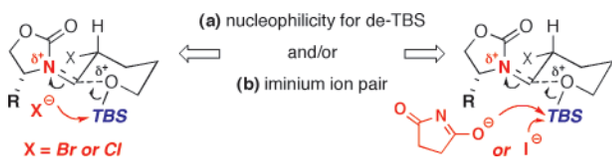
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Scheme 4. A Proposed Stereochemical Model

hindered cyclization through intermediate **D**. Support for this model arises from the halo-etherification in which the minor isomer **14-Br'** [Figure 1] was enriched relative to the reaction leading to **7-Br/7-Br'**. The only difference is the presence of the OBz group [see the box]. This is in fact consistent with related cation intermediates where an OR substituent would prefer the axial position.^{24,25} This preference can only be accommodated in intermediate **D** [not shown but the OBz group would be equatorial in intermediate **C** en route to **14-Br**].

In addition, the observations that NXS did not work and I_2 was sluggish suggest that an effective desilylative cyclization through intermediate **C** likely depends upon not solely the nucleophilicity of the halide anion [versus succinimido anion: see Figure 3] since I^- should be more nucleophilic

**Figure 3.** Nucleophilicity and the nature of iminium ion pairs in C.

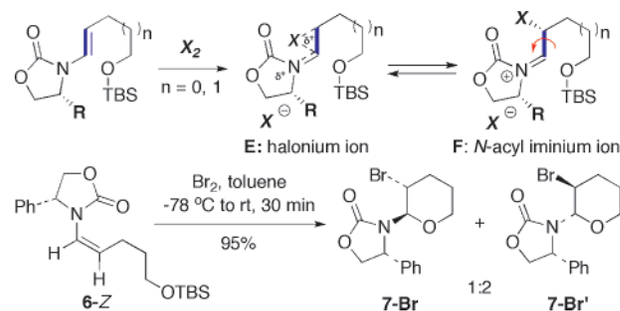
than Br^- or Cl^- . Thus, we believe the proximity of the anion, or the “tightness” of the iminium ion pair, is also very important. In this regard, as the TBSO group approaches the iminium carbon, Br^- or Cl^- being a much tighter counter-

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anion to the iminium cation than I^- provides a more facile desilylation.

Another question concerns the involvement of the halonium intermediate **E** [Scheme 5] and its significance versus

Scheme 5. Bromination of a Z-Enamide

N-acyl iminium ion **F**. The isolation of **19-Cl-cis**, arriving from its respective *E*-enamide, implies that halonium cation **E** is likely not important because isomerization of the C–C bond [in blue] is required to give **19-Cl-cis**. A more definitive answer came when we brominated **6-Z**^{3,15} and only obtained *trans*-bromo pyrans **7-Br** and **7-Br'** in 95% yield with a 1:2 ratio²⁶ in favor of the latter isomer. We detected no corresponding *cis*-products. The fact that the olefin geometry is not preserved again supports a free rotation of the C–C bond [in blue] occurring prior to the desilylative cyclization.

We have described here a stereoselective halo-etherification of chiral enamides, leading to the synthesis of halogenated cyclic ethers, which are prevalent among natural products. While this work provides a stereoselective entry to chiral secondary halides, these halo-etherifications also provide mechanistic insight to the chemistry of chiral enamides. Applications of this new synthetic method are currently underway.

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Supporting Information Available: Experimental details, characterization data, X-ray structural analysis, and NMR spectral for all new compounds. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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(26) This reversal in ratio is consistent with the mechanistic model in which the same favored *Si*-face bromination of **6-Z** would lead to an *N*-acyl iminium intermediate that is slow to cyclize, whereas the hindered *Re*-face bromination gives the respective *N*-acyl iminium intermediate that is more favored for the desilylative cyclization.