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Introduction

Controlling the regioselectivity of the bromolactonization reaction in HFIP⁺

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The halolactonization reaction provides rapid access to densely functionalized lactones from unsaturated carboxylic acids. The *endo/exo* regioselectivity of this cyclization reaction is primarily determined by the electronic stabilization of alkene substituents, thus making it inherently dependent on substrate structures. Therefore this method often affords one type of halolactone regioisomer only. Herein, we introduce a simple and efficient method for regioselectivity-switchable bromolactonization reactions mediated by HFIP solvent. Two sets of reaction conditions were developed, each forming *endo*-products or *exo*-products in excellent regioselectivity. A combination of computational and experimental mechanistic studies not only confirmed the crucial role of HFIP, but also revealed the formation of *endo*-products under kinetic control and *exo*-products under thermodynamic control. This study paves the way for future work on the use of perfluorinated solvents to dictate reaction outcomes in organic synthesis.

Halonium-promoted addition of nucleophiles to alkenes is one of the most fundamental reactions in organic chemistry, which offers widespread applications in organic synthesis.1 The intramolecular variant of this transformation is a powerful tool to construct molecular complexity by not only creating a new ring and stereogenic centres but also introducing a halide group for subsequent functionalizations.² Halolactonization is a typical example of this transformation, offering rapid access to densely functionalized lactones from acyclic unsaturated carboxylic acids. A wide range of valuable lactone analogues varying in both structures and stereochemistry could be obtained through this protocol by controlling the diastereoselectivity, enatioselectivity, and regioselectivity of the reaction (Scheme 1a).3 Traditionally, diastereoselective and enantioselective halolactonizations are usually directed by substrate structures.^{3a,b} Over the past two decades, there have been also developments of reagent-controlled enantioselective halolactonization, relying on chiral electrophilic halogenating reagents which in turn can be generated in situ through the coordination of chiral catalysts to halogenating reagents.⁴ On the other hand, regioselectivity of halolactonization has predominantly been dictated by the electronic effects of alkene substituents, with carboxylate groups intercepting halonium intermediates at the position where positive charge stabilization is most favorable. To the best of our knowledge, methods to alter this inherent regioselectivity of halolactonization or offer controllable formation of both regioisomers from one single unsaturated carboxylic acid substrate are scarce in the literature.⁵

In the last decade, hexafluoroisopropanol (HFIP) has attracted increasing attention as a reaction solvent due to its strong hydrogen-bond donating ability, low nucleophilicity, stability under redox conditions and most importantly the unique capacity to stabilize ionic reaction intermediates.6 HFIP has also demonstrated its ability to facilitate a wide range of difunctionalization reactions.7 In relevant context to this work, there have been seminal contributions from the Gulder group8 on HFIPmediated halocyclization of terpenes, and the Lebœuf and Gandon groups9 on HFIP-mediated haloamidation and halolactonization of alkenes with excellent reaction outcomes (Scheme 1b). Building upon these research studies and our prior investigations on halide-promoted addition reaction to alkenes,10 acid-promoted cyclization reactions¹¹ and HFIP-assisted Brønsted acid-catalyzed chemistry,12 we envisaged that HFIP can be used to activate bromonium sources such as NBS and promote endo-bromolactonization of readily available conjugated unsaturated carboxylic acids¹³ (Scheme 1c, upper route). On the other hand, the other bromolactonization regioisomers with smaller ring sizes are inherently more thermodynamically stable but require higher activation energies due to the lack of

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Scheme 1 (a) Halolactonization reaction; (b) HFIP-mediated halocyclization; and (c) this work: HFIP-mediated regioselectivity-switchable bromolactonization.

electronic stabilization from the aromatic ring. We believed that the presence of HFIP-activated Brønsted acid catalysts can potentially activate the reaction substrates to overcome these higher activation energy barriers, leading to the formation of the *exo*-cyclization product (Scheme 1c, lower route). This novel regioselectivity-switchable protocol would enable rapid regioselective synthesis of two different analogues of densely functionalized lactones from readily available precursors.

Results and discussion

We elected to focus on bromolactonization in this work, as brominating reagents and their corresponding products possess balanced reactivity and stability compared to chloro or iodo counterparts. We initiated this study by choosing **1a**, which was efficiently obtained by ring-opening olefination of cyclopentanone ketal according to our recent work,¹³ as the model substrate and *N*-bromosuccinimide (NBS) as a brominating reagent, which can generate ε -caprolactone **2a** and δ -valerolactone **3a** as *endo*-cyclization and *exo*-cyclization products respectively (Table 1). The bromolactonization of **1a** in HFIP happened smoothly at room temperature, selectively forming *endo*-product **2a** in 91% yield with a 13/1 regioisomer ratio (entry 1, Table 1). Both reactivity and selectivity of this HFIP-mediated *endo*-bromolactonization are in accordance with Lebœuf and Gandon work.⁹ Lowering the reaction temperature to 0 °C, with a slight decrease in yield and longer reaction time, resulted in absolute selectivity to *endo*-product **2a** (entry 2). This observation is in agreement with our initial hypothesis that *endo*-products of bromolactonization reactions are kinetically favorable while *exo*products are thermodynamically favorable. For a simple reaction setup and better reaction efficiency, we chose to carry out the reaction at room temperature in subsequent studies.

On the other hand, introducing a catalytic amount of Brønsted acids into the reaction, as predicted, shifted the selectivity towards the *exo*-product **3a** while maintaining Table 1 Optimization of HFIP-mediated regioselectivity-controllable bromolactonization



Entry ^a	Catalyst (mol%)	Solvent (v/v)	Yield of $2a^b$	Yield of $3a^b$	2a/3a ratio ^b
1		HFIP	91%	7%	13/1
2^c		HFIP	77%	Traces	>20/1
3	AcOH (10%)	HFIP	88%	10%	8.8/1
4	TFA (10%)	HFIP	67%	20%	3.4/1
5	pTSA (10%)	HFIP	Traces	89%	<1/20
6	TfOH (10%)	HFIP	Traces	86%	<1/20
7	pTSA (5%)	HFIP	Traces	87%	<1/20
8	pTSA (2%)	HFIP	Traces	79%	<1/20
9		HFIP/DCE $(1/1)$	85%	8%	10.6/1
10		HFIP/DCE $(3/7)$	85%	9%	9.4/1
11		HFIP/DCE (1/9)	71%	Traces	>20/1
12		DCE	ND	ND	_
13		TFE	29%	60%	1/2.1
14		iPrOH	ND	ND	_
15		MeNO ₂	Traces	Traces	_
16	pTSA (5%)	DCE	11%	5%	2.2/1
17	pTSA (5%)	TFE	22%	12%	1.8/1
18	pTSA (5%)	iPrOH	13%	5%	2.6/1
19	pTSA (5%)	MeNO ₂	17%	20%	1/1.2

^{*a*} Reaction conditions: **1a** (0.1 mmol), NBS (1.1 equiv.), catalyst, and solvent (v/v, 0.1 M) were stirred at room temperature for 1 h. ^{*b*} Yields and regioisomer ratios were determined by ¹H NMR using methyl benzoate as an internal standard. See pages S4–S5 in the ESI for full optimization studies. ND = not detected. ^{*c*} Reaction was carried out at 0 °C for 2 h.

excellent overall yields of both products (Table 1, entries 3-8). Interestingly, weaker acids such as acetic acid (AcOH, entry 2) and trifluoroacetic acid (TFA, entry 4) caused minor shifts in selectivity, whereas stronger acids like p-toluenesulfonic acid (pTSA, entry 5) and triflic acid (TfOH, entry 6) induced a complete alteration in selectivity toward exo-product 3a. pTSA was chosen for further studies of exo-bromolactonization as it is inexpensive and easier to handle. Reducing the amount of pTSA to 5 mol% did not change the reaction efficiency (entry 7) while the yield of *exo*-product 3a slightly decreased when employing 2 mol% of pTSA (entry 8). We also attempted to reduce the amount of HFIP in endo-bromolactonization by using DCE as a co-solvent, but both yield and selectivity slightly dropped (entries 9-11). To clarify the role of HFIP in this reaction, we carried out the reaction in DCE as a typical solvent for halocyclization chemistry, trifluoroethanol (TFE) as a weaker hydrogen-bonding donor solvent, isopropanol as the respective non-fluorinated alcohol, and nitromethane (MeNO₂) as a highly polar solvent for charge stabilization (entries 12-19). Without HFIP as solvent, reactions generally led to unsatisfactory outcomes, regardless of whether or not there was a Brønsted acid catalyst, confirming the crucial role of HFIP in the formation of both endo- and exo-products.

We subsequently explored the versatility of our newly developed method in both *endo*-bromolactonization (Scheme

2a) and *exo*-bromolactonization (Scheme 2b). A series of unsaturated carboxylic acid **1** were first subjected to our optimal conditions for *endo*-bromolactonization. Substrates with diverse electronic and steric effects (**1a**–**j**) were well tolerated under *endo*-bromolactonization conditions, yielding 7-*endo*-products **2a–j** in moderate to excellent yields. Electron-rich thiophene substrate **1k** led to poor efficiency, probably due to susceptibility to electrophilic bromination on the electron-rich aromatic ring in HFIP medium.¹⁴ Our method also demonstrated good efficiency in 6-*endo*-bromolactonization, generating 6-*endo*-product **2l** in good yield. However, 8-*endo*-bromolactonization exhibited unimpressive efficiency (products **2m** and **2n**), which can presumably be attributed to the challenging formation of eight-membered medium-sized lactones.

A quite similar trend in reaction yields was also observed when the same set of unsaturated acids **1** was subjected to our optimal conditions of *exo*-bromolactonization (Scheme 2b). In the case of $\delta_{,\epsilon}$ -unsaturated carboxylic acids, except for substrates with an electron-rich aromatic ring (**1f** and **1k**), which are sensitive towards electrophilic aromatic bromination,^{14,15} other substrates (**1a–e** and **1g–j**) exhibited a complete switch in regioselectivity to 6-*exo*-bromolactonization with moderate to excellent yields when treated with a Brønsted acid catalyst in HFIP. Absolute *exo*-regioselectivity and excellent



Scheme 2 Substrate scope of (a) *endo*-bromolactonization and (b) *exo*-bromolactonization. Reaction conditions unless otherwise noted: 1(0.2 mmol), NBS (1.1 equiv.), with/without pTSA (5 mol%), HFIP (0.1 M). Diastereomeric ratios were all >20/1 without quoted ratios in parentheses. ^[a] HFIP/DCM (1/4, 0.1 M) was used. ^[b] HFIP/DCM (1/99, 0.1 M) was used. ^[c] Reaction was stirred without catalyst for 1 h prior to adding catalyst and further stirring for 1 h. ^[d] NMR yield. ^[e] TfOH (10 mol%) was used.

yield were also observed with γ , δ -unsaturated carboxylic acid **11**, while no expected products **3m** and **3n** were detected for the case of ε , ζ -unsaturated carboxylic acids **1m** and **1n**. It should be noted that along with controllable regioselectivity, our method also offered excellent to absolute diastereoselectivity for both *endo*- and *exo*-cyclization.

Next, we extended the substrate scope of HFIP-mediated bromolactonization to other alkenoic acid scaffolds (Scheme 3). Estrone-derived $\delta_{,\epsilon}$ -unsaturated carboxylic acid **10**, obtained by modifying estrone through a three-step procedure as reported in our recent work,¹³ smoothly underwent 7-*endo*-

bromolactonization, yielding tetracyclic lactone **20** as an equimolar mixture of two diastereomers in 88% yield. Similar to substrates in Scheme 2b, the regioselectivity completely shifted to 6-*exo*-product **30**, also as an equimolar mixture of two diastereomers in excellent yield, when a catalytic amount of pTSA was used to promote the reaction (Scheme 3a). This example demonstrated a quick and efficient way for the late-stage modification of complex cyclic ketones into lactones with various ring-sizes.

As discussed earlier in Scheme 2, we believe that the selective formation of *endo*-product **2** is kinetically favored due to



Scheme 3 Expanding the substrate scope of bromolactonization: (a) rapid access to two analogues of estrone-based lactone and (b) bromolactonization on other scaffolds.

benzylic stabilization, and the selective formation of *exo*product **3** is thermodynamically supported due to generation of a more stable ring size. To support this hypothesis, we carried out some negative testing studies with two other structures often encountered in halolactonization chemistry, terminal alkenoic acid **1p** and *trans*-stilbene-type acid **1q**. Gratifyingly, both of these exclusively yielded *exo*-product **3p** and *endo*product **2q**, respectively in good to excellent yields, regardless of whether catalytic pTSA was used or not (Scheme 3b). These outcomes align well with our hypothesis, as **3p** and **2q** are both kinetically and thermodynamically favorable, owing to both benzylic stabilization and the formation of a more stable ring size. As a result, this leads to an unswitchable regioselectivity with substrates **1p** and **1q**.

We also attempted to exclude benzylic stabilization by carrying out reactions on aliphatic unsaturated acids 1r-t,



Scheme 4 Proposed reaction mechanism for the bromolactonization.

which have nearly identical electronic effects on both reactive sites. Isopropyl-substituted acids **1r** and **1s** exclusively yielded 6-*exo*-product **3r** and 5-*exo*-product **3s** in excellent yields in the presence or absence of the pTSA catalyst, respectively. Unswitchable regioselectivity and excellent yield were also recorded on a less sterically hindered methyl-substituted acid **1t**. These results confirmed the importance of the benzylic effect on the controllable regioselectivity.

To gain better insights into the reaction mechanism and effect of catalyst/solvent on the regioselectivity for the bromolactonization, we then turn our effort to density functional theory (DFT) calculations at the MN15/6-311+G(2d,2p)/SMD// M06-2X/6-31G(d,p)/SMD level of theory (see page S97 in the ESI† for computational details). The proposed reaction mechanism for the bromolactonization is shown in Scheme 4. Traditionally, this transformation is expected to take place *via* a stepwise Ad_E 2-type mechanism. The first step of this reaction is the electrophilic addition of the bromine atom to the C==C double bond of the substrate generating a reactive cyclic bromonium intermediate, which is followed by a nucleophilic addition leading to *endo-* and *exo*-cyclic products.¹⁶ Alternatively, by means of kinetics studies, NMR spectroscopy, and DFT calculations, Jackson and Borhan¹⁷ proposed that the halolactonization can take place *via* a concerted Ad_E3 -type mechanism in which the nucleophilic and electrophilic additions happen simultaneously and no ionic intermediate is generated during the reaction. Moreover, the additions of the nucleophile and electrophile can occur at the same face (*i.e.*, *syn*-addition) or opposite faces (*i.e.*, *anti*-addition) of the C==C double bond (Scheme 4).¹⁷

We first performed DFT calculations to elucidate the reaction mechanism for the bromolactonization in the aprotic DCE solvent, using **1c** (Ar = p-Br-C₆H₄-) as the model substrate. Consistent with previous experimental and theoretical studies,^{17,18} DFT calculations revealed that in DCE, the bromolactonization takes place *via* a *syn*-concerted addition pathway (Fig. 1). No transition state for the stepwise mechanism as well as *anti*-concerted addition pathway could be located. Interestingly, the activation barriers for the *syn*-additions are calculated to be fairly high, amounting to 36.8 and 39.4 kcal mol⁻¹ for **TS-1** and **TS-2**, respectively. This DFT result is in good agreement with our experimental findings (Table 1) that only a trace



Fig. 1 Optimized transition states for the concerted bromolactonization in DCE. Transition states TS-1 and TS-2 lead to *endo-* and *exo-*cyclic products, respectively.

amount of the lactonization product can be observed when this reaction is performed in DCE (*vide supra*).

DFT calculations were subsequently performed to investigate the favorable mechanistic pathway and regioselectivity for the bromolactonization in HFIP. Because of its powerful hydrogen bond (H-bond) donor ability,^{6a,d} it is possible for HFIP to form strong H-bonds with various species along the reaction course. Therefore, a mixed explicit–implicit solvation model is used, in



b. Conversion between endo- and exo-product:



Fig. 2 Experimental mechanistic studies: (a) kinetics studies and (b) conversion between *endo-* and *exo-*product. See pages S6–S9 in the ESI† for more details.



Fig. 3 Computed free energy profile for the stepwise bromolactonization in HFIP. S represents the HFIP molecule.

which explicit HFIP molecules were included in DFT calculations.^{9,12b,19} Based on our kinetic studies suggesting that the reaction order in HFIP is approximately 3 (Fig. 2a and further details in pages S6–S8 in the ESI†), three HFIP molecules were included in our computational investigations. Additionally, we have also performed calculations by involving one and two HFIP molecules (Fig. S1–S4, pages S99–S100 in the ESI†). Although there are some changes in absolute energy values, the conclusion remains similar all through our calculations, which gives a solid validation to the accuracy of the mixed explicit–implicit solvation model.

Our DFT calculations indicated that the favorable mechanism for the bromolactonization in HFIP is the stepwise pathway (Fig. 3), which is consistent with the previous calculation for the



Fig. 4 Computed free energy profile for the stepwise bromolactonization in HFIP catalyzed by TfOH. S represents the HFIP molecule.

halolactonization in protic solvent.¹⁸ The barrier height for the formation of the bridged bromonium species 5 *via* **TS**-3 is calculated to be 15.7 kcal mol⁻¹. It should be noted that we have also considered the concerted mechanism for this transformation in HFIP. However, we can only locate the transition state for the *syn*-concerted addition pathway (Fig. S5 in the ESI†), which is calculated to be 9.7 kcal mol⁻¹ higher in energy than **TS**-3. Therefore, the concerted mechanism is unlikely to occur. Additionally, the stepwise mechanism is also calculated to be the favorable pathway for the reaction of **11** in HFIP (see Fig. S6 in the ESI† for more details). This result is in agreement with the fact that HFIP has an exceptional cation stabilization ability^{6a,20} due to the low nucleophilicity and high dielectric constant, and, thus, the stepwise mechanism is supposed to be the preferable pathway.^{16b}

In the bridged bromonium intermediate 5, the natural charge of the C1 atom is higher than that of the C2 atom (Fig. 3). Therefore, the activation barrier for the nucleophilic addition from the oxygen atom of the carboxyl moiety to the C2 atom of the bromonium cation leading to the *endo*-cyclic product is calculated to be 2.6 kcal mol⁻¹ lower in energy than that for the nucleophilic addition to the C1 atom. This result is consistent with experimental data where the formation of the *endo*-product is more favorable in HFIP. Our computational study demonstrates that the regioselectivity for the bromolactonization in HFIP is determined by electronic properties and this reaction is under kinetic control.

On the other hand, our experiments demonstrated that when TfOH is used as a catalyst, the exo-cyclic product is more favorable (Table 1). By using DFT calculations, we found that in the presence of TfOH, the NBS reagent can easily be protonated lowering the LUMO energy from -1.05 eV to -2.04 eV for neutral and protonated NBS, respectively leading to the enhancement of NBS reactivity.21 When TfOH is included, the activation barrier for the electrophile addition TS-6 is calculated to be 9.2 kcal mol⁻¹ relative to **1** (Fig. 4), which is 6.5 kcal mol⁻¹ lower than that without TfOH, i.e., TS-3. In addition, TfOH can also have a great stabilization effect on the bridged bromonium species. From bromonium intermediate 8, although the nucleophilic addition generating the endo-product is kinetically more favorable, the reaction is now under thermodynamic control. The endo-product can isomerize to generate a more stable exocyclic product. This computational result is consistent with our additional control experiments, in which the endo-cyclic species 2 can transform into the exo-cyclic species 3 in the presence of TfOH or pTSA (Fig. 2b - upper route, also see page S9 in the ESI⁺ for more details). It should be noted that when the isomerization was carried out in DCE instead of HFIP (Fig. 2b - lower route), average yields of exo-product 3a with poor diastereoselectivity (in the case of TfOH) or incomplete conversion (in the case of pTSA) were observed, highlighting the vital role of HFIP in this conversion.

Conclusion

In summary, we introduce an effective and straightforward method to access two different analogues of bromolactones by

precisely manipulating the regioselectivity of the bromolactonization reaction in HFIP solvent. DFT calculations undersignificance of HFIP in both scored the endobromolactonization under kinetic conditions and exo-bromolactonization under thermodynamic conditions. This method, when combined with our recently developed ring-opening olefination of cyclic ketone ketals, offers a novel pathway for latestage modifications of cyclic ketones into functionalized lactones.

Data availability

All data are available in the main text or the ESI.†

Author contributions

Conceptualization: TAT and TVN. Experimental work: NTAP and TAT. Computational work: BKM. Supervision: TVN. Writing – original draft: TAT, BKM, and TVN. Writing – review & editing: TVN.

Conflicts of interest

The authors declare that they have no competing interests.

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