Effects of Water Addition on a Catalytic Fluorination of Dienamine

Daiki Kuraoku 1, Tsunaki Yonamine 1, Genta Koja 2, Norio Yoshida 3, Satoru Arimitsu 1,* and Masahiro Higashi 4,*

1 Graduate School of Engineering and Science, University of the Ryukyus, Senbaru 1, Nishihara, Okinawa 903-0213, Japan; k198363@eve.u-ryukyu.ac.jp (D.K.); k188363@eve.u-ryukyu.ac.jp (T.Y.)
2 Center for Research Advancement and Collaboration, University of the Ryukyus, Senbaru 1, Nishihara, Okinawa 903-0213, Japan; kojagen@lab.u-ryukyu.ac.jp
3 Department of Chemistry, Graduate School of Science, Kyushu University, 744 Motooka, Nishiku, Fukuoka 819-0395, Japan; noriwo@chem.kyushu-univ.jp
4 Department of Molecular Engineering, Kyoto University, Kyoto daigaku-katsura, Nishikyo-ku, Kyoto 615-8510, Japan
* Correspondence: arimitsu@sci.u-ryukyu.ac.jp (S.A.); higashi@moleng.kyoto-u.ac.jp (M.H.)

Abstract: We investigate the effects of water addition on a highly stereocontrolled fluorination of dienamine generated by α-branched enals and 6′-hydroxy-9-amino-9-deoxy-epi-quinidine with N-fluorobenzenesulfonimide (NFSI) in the presence of Brønsted acid both experimentally and theoretically. It is experimentally found that water addition to organic solvent significantly shortens the reaction time whereas excessive water addition decreases the enantiomeric excess. The results calculated with three-dimensional reference interaction site model self-consistent field (3D-RISM-SCF) method are in good agreement with the experimental ones. It is revealed that the shortness of reaction time is caused by the reactant destabilization and that the decrease in enantiomeric excess is due to the difference of hydration free energy between two transition states.

Keywords: water addition; dienamine; fluorination; 3D-RISM-SCF; solvent effect

1. Introduction

In organic synthesis, water addition into organic solvents often enhances the reactivity and the selectivity [1–6]. For example, in triple catalytic cross-coupling reaction developed by MacMillan group, addition of 40 equiv. of water into DMSO solvent increases product yield from 42% to 85% [3]. Maruoka et al. found that water addition for asymmetric aldol reactions with a chiral organocatalyst greatly increases the enantiomeric excess up to ~90% ee [4]. Although such effects of water addition on the reactivity and selectivity are widely known, the understanding of detailed mechanisms is still limited.

Recently, we developed the asymmetric fluorination of dienamines generated by α-branched enals and 6′-hydroxy-9-amino-9-deoxy-epi-quinidine in the presence of NFSI and Brønsted acid [7,8], and the reaction gave the excellent α-regioselectivity and enantioselectivity regardless of substituents on R1 and R2; thus, both electron-withdrawing and electron-donating groups on the aromatic ring of R1 have no significant influence, and all maintain excellent enantioselectivity over 90% ee. On the other hand, in the cases of benzyl group on R1 or ethyl on R2, the reactions gave slightly lower enantioselectivities, 81% and 77%, respectively (Scheme 1) [9]. During the optimization of the reaction condition, the addition of 40 equiv. of water was found effectively to shorten the reaction time maintaining the high stereoselectivity. In the previous density functional theory (DFT) investigation,
we revealed the non-classical hydrogen bonding between C(sp\(^2\))–H and the counterion of the conjugate base of Brønsted acids is the key to stabilize the transition state for the major enantiomer, R-isomer determined by single-crystal X-ray diffraction analysis; however, the detailed role of water was remained unclear.

**Scheme 1.** α-Fluorination of α-branched enals.

Therefore, in the present study, we investigate the deep insight of the effects of water addition on this fluorination reaction both experimentally and theoretically. First, changes in reaction time and enantiomeric excess by the amount of water are examined experimentally. Then the effects of water addition are analyzed by means of three-dimensional reference interaction site model self-consistent field (3D-RISM-SCF) method [10]. The 3D-RISM-SCF method is a combination of quantum chemical method and integral equation theory of liquids, which enables us to obtain complete ensemble average of solvation structure around a solute described quantum chemically. This type of approach has been successfully applied to chemical reactions not only in pure solvents but also in mixed solvents [11–14]. The free energy change due to the water addition is examined by decomposition analysis.

**2. Results and Discussion**

First, we investigated the effect of water on the fluorination reaction experimentally. We liked to run the reaction in DMF instead of NMP to obtain a comparable data of the later DFT calculation. Furthermore, we examined the effect of Brønsted acids for the enantioselectivity if we can find the simple acids give similar enantioselectivity so that we can reduce the computational time; thus, we examined three kinds of Brønsted acids, BINOL-based phosphoric acid (B1), p-TsOH (B2) and MsOH (B3) with amine catalyst (A1) in DMF as the solvent at room temperature without water (Scheme 2). Interestingly, the enantioselectivities of the corresponding product were obtained at 72% with B1, 78% with B2 and 75% with B3, therefore Brønsted acids, B2 and B3 can be the good surrogates for B1.

**Scheme 2.** Effect of Brønsted acid on the enantioselectivity.

Based on this control experiment, we examined the effect of water using B2 as the Brønsted acid at \(\text{−25 }^\circ\text{C}\). The reaction without water did not consume the starting aldehyde even after 300 h; however, adding some water to DMF solution gave a shorter reaction time, as shown in Figure 1. On the other hand, the enantioselectivity outcomes were not changed until 40 equivalents of water was added, while a large excess of water addition decreases the enantiomeric excess drastically.
Next, we investigated the effects of water addition with the 3D-RISM-SCF method. Our previous study showed that two catalysts, quinuclidine and counterion of Brønsted acid, control the regio- and stereoselectivity at the transition state (TS) in a concerted manner (Figure 2) [9]. The fluorination proceeds at the α position because of the quinuclidine located close to the α position whereas the counterion preferentially stabilizes the TS-R configuration connecting to the major product due to the nonclassical CH hydrogen bonds. Following these results, we here considered four transition states, TS-R-A\(^-\), TS-S-A\(^-\), TS-R, and TS-S, in which “\(^-\)” indicates the counterion is bound with the TS substrate.

The calculated activation free energies with respect to volume fraction of water, x(H\(_2\)O), are shown in Figure 3. In pure DMF solution, x(H\(_2\)O) = 0, the activation free energies of TS-R-A\(^-\), TS-S-A\(^-\), TS-R, and TS-S are 18.76, 20.70, 41.12, and 39.44 kcal/mol, respectively. The activation free energy of TS-R-A\(^-\) is the lowest and ~2 kcal/mol lower than that of TS-S-A\(^-\). In contrast, the activation free energy of TS-R is higher than that of TS-S. This result indicates that the TS-R configuration is more stabilized with the counterion, which is consistent with our previous result [9]. It is noted that the counterion strongly interacts with the TS substrate in pure DMF solution. The free energy differences between with and without the counter anion are ~20 kcal/mol.
As a result, the activation free energies of the two TSs giving the minor product, TS- and TS-, the total free energy difference largely contributes to the free energy difference and solvation free energy. Figure 4b shows how these two terms contribute to the decrease in solvation separately. As a result, the free energy difference is caused by a large destabilization of the reactant. In addition, the TS complexes with the reactant and TSs become unstable by water addition because they have many nonpolar parts. The lesser polar reactant is more destabilized than the TSs. Therefore, the decrease in activation free energy is caused by a large destabilization of the reactant. In addition, the TS complexes with the reactant and TSs become unstable by water addition because they have many nonpolar parts. The lesser polar reactant is more destabilized than the TSs. Therefore, the decrease in activation free energy is caused by a large destabilization of the reactant. In addition, the TS complexes with the reactant and TSs become unstable by water addition because they have many nonpolar parts.

Furthermore, the free energy differences between with and without the counter anion are also decreased, indicating that the interaction between the TS substrate and counterion becomes weaker. As a result, the activation free energies of the two TSs giving the minor product, TS-S-A- and TS-R, approach to that of TS-R-A-. This result qualitatively agrees with the experimental result of shorter reaction time by water addition. It is also found that the free energy differences between the four TS states are monotonically decreased. In pure aqueous solution, x(H2O) = 1, the activation free energies of TS-R-A-, TS-S-A-, TS-R, and TS-S are 12.35, 12.43, 14.84, and 13.63 kcal/mol, respectively. Although the TS-R-A- state has the lowest free energy in all the solutions, the free energy difference between TS-R-A- and TS-S-A- is gradually decreased and only ~0.1 kcal/mol in pure aqueous solution. Furthermore, the free energy differences between with and without the counter anion are also decreased, indicating that the interaction between the TS substrate and counterion becomes weaker.

To investigate the effects of water addition in more detail, we calculated the free energy changes of reactant and TSs with respect to the volume fraction of water (Figure 4a). It is found that both the reactant and TSs become unstable by water addition because they have many nonpolar parts. The lesser polar reactant is more destabilized than the TSs. Therefore, the decrease in activation free energy is caused by a large destabilization of the reactant. In addition, the TS complexes with the reactant, TS-R-A- and TS-S-A-, become more unstable than the TSs without counterion, TS-R and TS-S, because the cationic TS-R and TS-S and anionic counterion can strongly interact with water solvent separately. As a result, the free energy differences between with and without the counter anion are decreased as the ratio of water is increased.

In the 3D-RISM-SCF theory, free energy of a solute is described as the sum of solute internal energy and solvation free energy. Figure 4b shows that how these two terms contribute to the decrease in free energy difference between TS-R-A- and TS-S-A-. When the volume fraction of water is small, the solvation free energy largely contributes to the free energy difference. As the ratio of water is increased, the positive solvation and negative solute contributions cancel each other, resulting in that the total free energy difference becomes nearly zero.

**Figure 3.** Calculated activation free energies with respect to volume fraction of H2O. The activation free energies of TS-R-A-, TS-S-A-, TS-R, and TS-S are shown in red, orange, purple, and blue lines, respectively.
We analyzed the differences in the TS-x(H₂O) configuration interacts with water molecules more strongly. It is noted that the MeSO⁻ larger than those at the TS-x(H₂O) configuration. Reducing the binding energy difference between MeSO⁻ and its components. We found that the interactions between MeSO₃⁻ counterion and solvent molecules particularly contribute to reducing the binding energy difference. Figure 5 shows the radial distribution functions (RDFs) for the S or O (MeSO₃⁻)-H (H₂O) distance at x(H₂O) = 0.3. The first peaks at the TS-S-A⁻ configuration are larger than those at the TS-R-A⁻ configuration, indicating that the MeSO₃⁻ counterion at the TS-S-A⁻ configuration interacts with water molecules more strongly. It is noted that the MeSO₃⁻ counterion at the TS-R-A⁻ configuration preferentially stabilizes the TS-R substrate and thus interacts with water solvent only weakly. Therefore, we concluded that the decrease in activation free energy difference between TS-R-A⁻ and TS-S-A⁻ by water addition is due to the difference of the interactions between MeSO₃⁻ counterion and water molecules.

Table 1. Analysis of the binding energy difference between TS-R-A⁻ and TS-S-A⁻ in the two solutions, x(H₂O) = 0.0 and x(H₂O) = 0.3 (in kcal/mol).

|          | ΔE₉bind(x(H₂O) = 0.0) | ΔE₉bind(x(H₂O) = 0.3) | ΔΔE₉bind ¹ |
|----------|----------------------|-----------------------|------------|
| aldehyde | -0.99                | -0.82                 | -0.18      |
| amine    | 1.49                 | 3.37                  | -1.88      |
| F atom   | -1.22                | -2.45                 | 1.23       |
| counterion | 2.81               | 1.00                  | 1.82       |
| sum      | 2.09                 | 1.10                  | 0.99       |

¹ ΔΔE₉bind = ΔE₉bind(x(H₂O) = 0.0) - ΔE₉bind(x(H₂O) = 0.3).
1.5 equiv.) was added to the reaction mixture at 0 °C. Purity was determined by a high performance liquid chromatography (HPLC) performed on JASCO.

Next, to a solution of the corresponding fluorinated aldehyde in DCM (1.0 mL, 0.1 M) we re added AcOH (0.5 mmol, 5.0 equiv.) and benzylamine (0.5 mmol, 5.0 equiv.), and the whole solution was stirred for 1 h at the room temperature.

The reaction was quenched with sat. NaHCO3 and extracted with Et2O. The combined organic phase was washed with brine and dried over MgSO4. The organic solution was filtered and concentrated under the reduced pressure. The residue was purified by a silica gel flash chromatography eluting with hexane/ethyl acetate = 7/1 to obtain the corresponding fluorinated aldehyde to determine yield of the product.

We employed the 3D-RISM-SCF method [10] for the free energy calculations of solutes in the mixture of water and DMF. Notably, several previous studies showed that water molecules directly involve organic reactions [5,6]. However, we here assumed that the added water acts as a solvent because the enantiomeric excess is only gradually changed by water addition. It is noted that the hydrogen-bond interactions between substrate and water molecules are properly treated in the framework of 3D-RISM-SCF method. To reduce computational cost, MeSO3H (B3) was used as

**Figure 5.** Calculated RDFs for the O or S (MeSO3−)-H (H2O) distances at x(H2O) = 0.3 volume fraction at TS-R-A− and TS-S-A−. The RDFs for the O-H distances at TS-R-A− and TS-S-A− and the S-H distances at TS-R-A− and TS-S-A− are shown in red, blue, orange, and purple, respectively.
Brønsted acid. The density functional theory with M06-2X functional and 6-31G(d,p) basis set was employed for the electronic structure calculation of solutes. The solute geometries were taken from our previous study [9], in which the geometries were optimized at the SMD(MMF)-M06-2X/6-31G(d,p) level. The thermal corrections to the free energy were also added at the SMD(MMF)-M06-2X/6-31G(d,p) level. The Lennard-Jones parameters for solutes were taken from the general AMBER force field (GAFF) [15]. The simple point charge (SPC) model [16] with modified hydrogen parameters ($\sigma = 1.0$ Å and $\epsilon = 0.056$ kcal/mol) and six-interaction site optimized potential computational model (CS2) [17] were used for water and DMF, respectively. The volume fractions of water, $x(H_2O)$, were set from 0.0 (pure DMF) to 1.0 (pure water) at intervals of 0.1, where we assumed that the volumes of water and DMF in mixed solution are identical to those of pure solvents. The Kovalenko-Hirata closure [18] was used to solve the 3D-RISM equation. The temperature was set at 300 K. The grid points in the 3D-RISM-SCF calculations were $128 \times 128 \times 128$ with a spacing of 0.5 Å. All the calculations were performed with a modified version of the GAMESS version 18 AUG 2016 (R1) program package [19], where the 3D-RISM-SCF method have been implemented [20].

4. Conclusions

In this article, we investigated the effects of water addition on the highly stereocontrolled fluorination both experimentally and theoretically. The experimental results are in good agreement with the calculated ones. It was revealed that the shortness of reaction time is caused by the reactant destabilization and that the decrease in enantiomeric excess is due to the difference of solute-solvent interaction between two transition states.

As noted in the Introduction, adding water into organic reaction systems often improves reactivity and selectivity. Notably, it is difficult to investigate such effects with a simple polarized continuum model, though it is widely used for the analysis of solvation effects. As the present study demonstrated, the 3D-RISM-SCF method is expected to be a powerful tool for studying the effects of water addition.

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**Sample Availability:** Samples of all the compounds are available from the authors.