Impact of Multiple Hydrogen Bonds with Fluoride on Catalysis: Insight from NMR Spectroscopy

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ABSTRACT: Hydrogen-bonding interactions have been explored in catalysis, enabling complex chemical reactions. Recently, enantioselective nucleophilic fluorination with metal alkali fluoride has been accomplished with BINAM-derived bisurea catalysts, presenting up to four NH hydrogen-bond donors (HBDs) for fluoride. These catalysts bring insoluble CsF and KF into solution, control fluoride nucleophilicity, and provide a chiral microenvironment for enantioselective fluoride delivery to the electrophile. These attributes encouraged a "H/19F" NMR study to gain information on hydrogen-bonding networks with fluoride in solution, as well as how these arrangements impact the efficiency of catalytic nucleophilic fluorination. Herein, NMR experiments enabled the determination of the number and magnitude of HB contacts to fluoride for thirteen bisurea catalysts. These data supplemented by diagnostic coupling constants $^{1}J_{FH}$ give insight into how multiple H bonds to fluoride influence reaction performance. In dichloromethane (DCM-$d_{2}$), nonalkylated BINAM-derived bisurea catalyst engages two of its four NH groups in hydrogen bonding with fluoride, an arrangement that allows effective phase-transfer capability but low control over enantioselectivity for fluoride delivery. The more efficient N-alkylated BINAM-derived bisurea catalysts undergo urea isomerization upon fluoride binding and form dynamically rigid trifurcated hydrogen-bonded fluoride complexes that are structurally similar to their conformation in the solid state. Insight into how the countercation influences fluoride complexation is provided based on NMR data characterizing the species formed in DCM-$d_{2}$, when reacting a bisurea catalyst with tetra-n-butylammonium fluoride (TBAF) or CsF. Structure–activity analysis reveals that the three hydrogen-bond contacts with fluoride are not equal in terms of their contribution to catalyst efficacy, suggesting that tuning individual electronic environment is a viable approach to control phase-transfer ability and enantioselectivity.

INTRODUCTION

Hydrogen bonding (HB) is an important force for molecular recognition and for shaping the three-dimensional structure of molecules as large as proteins. In the past decade, hydrogen-bond donors (HBD) have also found widespread applications in asymmetric catalysis offering an alternative to Brønsted and Lewis acid catalysts. For example, dual HBD catalysts promote enantioselective nucleophilic additions through direct binding to neutral electrophiles or via the formation of chiral ion pair intermediates. In the latter scenario commonly referred to as anion-binding catalysis, an electrophile is ionized by a chiral HBD catalyst generating an ion pair, with the resulting anionic hydrogen-bonded species creating a chiral environment for the cation to react with an external nucleophile. These approaches that build on pioneering studies in non-asymmetric cases have enabled a wide range of transformations involving heteroatom-stabilized cations including acyl-Pictet–Spengler reactions, acyl-Mannich reaction, and additions to oxocarbenium ions. In further developments, HBD and Lewis acid catalysts were combined for enantioselective reactions involving less-reactive electrophiles. Specifically, chiral squaramides were used to activate silyl triflates via trilate abstraction, forming a highly Lewis acidic complex capable of generating oxocarbenium ion intermediates from acetals for enantioselective nucleophilic additions. Such a dual-catalyst system also enables activation of propargyl acetates into tertiary carbocationic intermediates that lack heteroatom stabilization for enantioconvergent catalytic $S_{N}1$ reactions. An underdeveloped manifold in chiral ion pair organocatalysis consists of using the hydrogen-bonded anion itself as the nucleophile. Hydrogen bonding is...
expected to decrease nucleophilicity, a challenge that led to
creative solutions. For example, the combination of a chiral
squaramide and pro-nucleophile bromotrimethylsilyl
(TMSBr) allowed for highly enantioselective ring-opening of
oxetanes with in situ released bromide bound to the chiral
HBD catalyst. Our laboratory disclosed an alternative
oxetanes with in situ released bromide bound to the chiral
squares via ring-opening of in situ formed meso-episulphonium or
aziridinium ions with metal alkali fluorides. Enantioen-
riched γ-fluoroamines were also within reach from achiral
azetidinium salts applying HB-PTC. For selected trans-
formations, 2 mol % of the phase-transfer catalyst was
sufficient to reach high yield and enantioselectivity. N-
Alkylation of the bisurea catalyst improved performance,
suggesting that this structural modification influences con-
formational preference and the ability of the catalyst to engage
in hydrogen bonding with fluoride.

While fluoride binding to HBD is an active area of research in
supramolecular chemistry, applications to asymmetric
catalytic fluorination have been overlooked for many years, and
data on how the number, strength, and directionality of hydrogen
bonds with fluoride influence its nucleophilicity have only recently appeared. Valuable information on the spatial
arrangement of the chiral tetrabutylammonium (TBA) N-
isopropyl bisurea–fluoride complex 2:TBAF was obtained
from a single-crystal X-ray diffraction measurement that
revealed three distinct N–H–F hydrogen-bonding inter-
actions (Figure 1C).

A 1H/19F NMR study is ideally suited to define the precise
nature of H-bonding to fluoride under conditions that simulate
catalytic fluorination. In this study, spectra were recorded in DCM-
and solid-state structure of 2:TBAF from single-crystal X-ray
diffraction.

Figure 1. (A) Proposed mechanism for hydrogen-bonding phase-
transfer catalysis (HB-PTC). (B) Asymmetric fluorination of β-
bromosulphides under HB-PTC. (C) BINAM-derived bisurea catalysts
1 and 2 and solid-state structure of 2:TBAF from single-crystal X-ray
diffraction.

RESULTS AND DISCUSSION

Conformational Properties of BINAM-Derived Bisurea Catalysts. Polar aromatic solvents such as 1,2-difluor-
benzene are optimal for fluorination under HB-PTC; chlorinated solvents such as chloroform and dichloromethane
are also effective. In this study, spectra were recorded in DCM-
d, because of its noncoordinating nature, optimal chemical
shift dispersion, and favorable temperature range. When
possible, the concentration was set to 25 mM to reflect the reaction conditions applied for enantioselective fluorination
under HB-PTC.

Initially, BINAM bisurea 1 and its N-isopropylated analogue
2 served to investigate the effect of N-alkylation on conformational change to the catalyst. A combination of
total correlation spectroscopy (TOCSY), heteronuclear single
quantum coherence spectroscopy (HSQC), and heteronuclear
multiple bond correlation spectroscopy (HMBC) ($^1$H–$^{13}$C and $^1$H–$^{15}$N) allowed for unambiguous assignment of all proton and carbon resonances of 1 and 2. Nonalkylated bisurea 1 shows uniformly sharp peaks in $^1$H and $^{13}$C NMR spectra (Figure 2, top). The chemical shifts are consistent with

electronic environments, with NH(a) proximal to electron-deficient 3,5-bis(trifluoromethyl)phenyl groups deshielded with respect to NH(b), which are positioned close to the BINAM scaffold. In contrast to 1, line broadening was observed for selected peaks in the $^1$H and $^{13}$C NMR spectra of N-isopropyl bisurea 2 (Figure 2, bottom). The resonances assigned to NH(a) and NH(c) appear at 7.1 and 6.9 ppm, respectively; NH(b) overlaps with multiple peaks in a range of common deuterated solvents (DCM-d$_2$, CDCl$_3$, tetrahydrofuran (THF-d$_4$), CD$_3$CN, dimethyl sulfoxide (DMSO-d$_6$), and MeOD) and, at first glance, was difficult to locate due to broadening (>70 Hz). In DCM-d$_2$, the chemical shift of NH(b) was estimated at ~7.6 ppm based on cross-peaks in $^1$H nuclear Overhauser effect spectroscopy (NOESY) and $^1$H–$^{15}$N HSQC; this was unexpected as NH(b) is in a less electron-deficient environment than NH(a) and NH(c). In DMSO-d$_6$, NH(a) and NH(c) are deshielded (+1.5 and +0.8 ppm, respectively) likely due to hydrogen bonding with the solvent, while NH(b) is minimally affected (+0.2 ppm). This observation alludes to NH(b) engaging in intramolecular hydrogen bonding with O═C(9) (broad $^{13}$C signal deshielded at 156.2 ppm; Figure 2, inset).

Peak line width ($\nu_{1/2}$ = half-height line width) is an important NMR observable correlating with the transverse relaxation rate $R_2$ ($R_2 = 1/T_2 = \nu_{1/2}$ Hz) and thereby with dynamic behavior of molecules in solution. Specifically, signal broadening observed for H(3′), C(3′), C(2′), C(1′), and C(9) is consistent with a dynamic conformational exchange on the NMR time scale involving the hydrogen-bond interaction NH(b)···O═C(9). The $^1$H NMR spectra of both 1 and 2 were also recorded at varying concentrations. Significant changes were observed for 1 between 2 and 64 mM, with all protons bound to carbons experiencing shielding (Δ$\delta_{H(11)} = -211$ ppb), while deshielding was observed for the NH (Δ$\delta_{NH(a)} = +300$ ppb, Δ$\delta_{NH(b)} = +227$ ppb). This pattern suggests intermolecular aggregation. In contrast to 1, the $^1$H NMR spectra of 2 display minimal variations in chemical shifts at higher concentrations, a result suggesting reluctance to form intermolecular aggregates.

Insight into the conformational preference of 2 in DCM-d$_2$ was deduced from analysis of through-space correlations by means of high-resolution $^1$H NOESY (Figure 3). Strong correlations between the isopropyl protons H(15), H(16), H(17), and NH(c) are consistent with the N-iPr urea adopting an anti–anti conformation. Nuclear Overhauser effect (NOE) between isopropyl and H(8′) indicates that the N-alkyl group is within close distance to naphthyl(I), while the NH(a) ⇔ H(11) interaction suggests that the two ureas are proximal to each other. The different orientations of the two ureas with respect to the BINAM scaffold explains the pronounced variations in chemical shifts between H(3′) and H(3) (8.47 vs. 7.56 ppm), with both positioned on the naphthyl group ortho to the urea substituent. The carbonyl group affects the chemical shifts of neighboring protons by means of anisotropic electron circulation; as a result H(3′), which is coplanar with the urea, is deshielded, whereas this is not the case for H(3) being out of plane. Together, these observations suggest that the two ureas are approximately perpendicular to each other. Magnetic anisotropy effects...
account for the shielding of the two diastereotopic methyls belonging to isopropyl. Protons 17 appear at 1.03 ppm as a sharp doublet, while H(15) pointing toward naphthyl(I) appears as a broad singlet at 0.68 ppm. Data in the solid state could not be secured because recrystallization of 2 afforded fine needles that were unsuitable for single-crystal X-ray diffraction studies.

A range of N-alkylated BINAM bisureas with various N-substituents and electronic patterns for the aryl rings were synthesized and analyzed by NMR spectroscopy in DCM-

\[\text{DCM-}d_2\] (Figure 4A). The spectra of N-alkylated analogues 3–7 all featuring two 3,5-bis(trifluoromethyl)phenyl substituents were very similar to N-isopropyl bisurea 2, implying analogous conformational preference. For this series, all NH(b) are more deshielded than NH(a)/NH(c) and appear broader; the C(9) and H(3’) signals are also broader and deshielded compared to C(9’)/H(3), respectively. When the aryl group linked to the N-alkylated urea is the less electron-deficient phenyl group (R1 = Ph, catalyst 8), an additional hydrogen-bond interaction NH(a)···O= C is observed that is consistent with increased Lewis basicity of the carbonyl oxygen (Figure 4B, top). This interaction confers conformational rigidity, giving rise to sharp proton and carbon resonances. In contrast, when the aromatic ring of the nonalkylated urea is phenyl (R1 = Ph, catalyst 9), the \(^1\)H and \(^13\)C NMR spectra are broader, suggesting the existence of a more dynamic structure likely resulting from weaker intramolecular hydrogen-bonding interactions (Figure 4B, bottom).

**Complexation of Bisureas with TBAF.** Formation of a hydrogen-bonded bisurea–fluoride complex is the cornerstone for successful enantioselective fluorinations with metal alkali fluoride under HB-PTC. Structural insight of these complexes in solution is therefore crucial for further development. The binding affinities of bisurea 1 and N-isopropyl bisurea 2 for fluoride were investigated by spectrophotometry (Figure 5).

**Figure 4.** (A) Structures of the BINAM-derived bisurea catalysts 3–13. (B) \(^1\)H NMR of 8 (top) and 9 (bottom) (DCM-

\[\text{DCM-}d_2\] 25 mM, 298 K).

**Figure 5.** Stacked UV spectra recorded for the titration of 1 (1.2 \(\mu\)M) and 2 (1.4 \(\mu\)M) with TBAF-3H\(_2\)O (0.1 mM) in DCM at 298 K. On the right, titration profiles at the \(\lambda_{\text{max}}\) (red dots) and fitting functions (red lines).\(^{13}\)

UV titrations carried out with 1 or 2 (1.2 and 1.4 \(\mu\)M, respectively) were performed in DCM by adding a solution of tetra-n-butylammonium fluoride of exact concentration quantified by \(^19\)F NMR (TBAF-3H\(_2\)O in DCM, 0.1 mM).\(^{13}\) Upon addition of an increasing amount of TBAF (0–15 equiv), UV spectra showed a bathochromic shift with the buildup of a new maximum of absorption (\(\lambda_{\text{max}}\)) at 273–275 nm. The binding constants and energies were calculated from the titration profiles at \(\lambda_{\text{max}}\) via nonlinear least-squares regression using dedicated software.\(^{16}\) Both 1:1 and 2:1 binding modes were considered, with 1:1 giving superior fitting results. A 1:1 binding mode was also consistent with the clean isosbestic points observed. Association constants in the range of \(10^6\ \text{M}^{-1}\) denote strong and similar binding affinities of 1 and 2 to fluoride (Table 1).

Next, \(^1\)H NMR titrations were performed at 3 mM concentration in DCM-

\[\text{DCM-}d_2\]. Addition of a TBAF solution (TBAF-3H\(_2\)O in DCM-

\[\text{DCM-}d_2\] 55 mM) to 1 led to line broadening...
in $^1$H NMR, indicating on–off equilibration. Sharp spectra were gradually recovered when approaching 1 equiv of added fluoride. Noticeable changes in $^1$H and $^{19}$F chemical shifts were observed between 0 and 1 equiv, after which the chemical shift of all resonances became invariant. Over the course of the titration, proton signals were shielded between 0 and 0.5 equiv of fluoride and deshielded between 0.5 and 1.0 equiv before reaching a plateau at >1 equiv. A signal assigned to NH(a) appeared in $^1$H NMR at 13.5 ppm after 1 equiv of fluoride was added, while NH(b) was observed at 7.6 ppm (Figure 6, left). Addition of TBAF to 2 also led to line broadening until quantitative binding was reached. At 1 equiv of fluoride, all NH are deshielded by $\sim$5 ppm with respect to unbound bisurea 2 (Figure 6, right). The titration of both catalysts 1 and 2 reflects a scenario not limited to 1:1 complexation. More likely, a dimeric species or higher coordinated fluorides dominate at low $F^-$ concentration, which then equilibrate toward the more stable $[UF]^-$ 1:1 complex ($U = 1$ or 2) over the course of the titration ($K_{(U_{1:1})} \gg K_{(U_{2:1})}$). The variation of chemical shift for diagnostic resonances was plotted against the concentration of added fluoride, and the data were analyzed. Initial attempts to fit the data to a 1:1 binding mode resulted in poor fit and nonsensical data. When a 2:1 binding mode was considered and $K_{(U_{2:1})}$ values secured from UV titration were taken into account, a good agreement with the experimental data was found. This allowed the calculation of 2:1 binding associations $K_{(U_{2:1})}$ ($\sim10^3$ M$^{-1}$) and binding energies for both 1 and 2 (Table 2). In our previous report on diarylurea–fluoride complexes [UF]$^-$ and [U$_2$F]$^-$, the binding constants were 250–500 M$^{-1}$ for $K_{(U_{2:1})}$ and 12 000–85 000 M$^{-1}$ for $K_{(U_{1:1})}$. These were measured in CD$_3$CN, a solvent in which hydrogen-bonding interactions are expected to be weaker.

**Table 1. Association Constants ($K_a$), log($K_a$), and Free Energies ($\Delta G$) for the Formation of 1:1 Bisurea–TBAF Complexes Derived from Bisureas 1 and 2**

| $U$ | $K_a$ (1:1) (M$^{-1}$) | log($K_a$ (1:1)) | $\Delta G$ (1:1) (kJ/mol) |
|-----|----------------------|------------------|--------------------------|
| 1   | 0.92 ± 0.02 × 10$^6$ | 5.96 ± 0.01      | $-34.01 ± 0.06$          |
| 2   | 1.43 ± 0.04 × 10$^6$ | 6.16 ± 0.01      | $-35.10 ± 0.08$          |

$^aK_a$ are calculated by nonlinear regression using DynaFit 4 and expressed as the average of two experiments.

**Table 2. Association Constant ($K_a$), log($K_a$), and Free Energies ($\Delta G$) for the Formation of 2:1 Bisurea–TBAF Complexes**

| $U$ | $K_a$ (2:1) (M$^{-1}$) | log($K_a$ (2:1)) | $\Delta G$ (2:1) (kJ/mol) |
|-----|----------------------|------------------|--------------------------|
| 1   | 600 ± 100            | 2.80 ± 0.08      | $-16.0 ± 0.5$            |
| 2   | 3100 ± 900           | 3.5 ± 0.1        | $-20.0 ± 0.7$            |

$^aK_a$ are calculated by nonlinear regression using DynaFit 4 and expressed as the average of two experiments.
Hydrogen-Bond Coupling $^{1b}J_{NH···F}$. The three NH of 2:TBAF are deshielded compared to 2 ($\Delta\delta_{313} \approx +5$ ppm), as are H(11), H(11$'$), and H(3$'$). In contrast, H(13) and H(13$'$) on the 3,5-bis(trifluoromethyl)phenyl group, as well as H(6$'$), H(7$'$), and H(8$'$) belonging to naphthyl(I), are shielded. The sharp lines in the $^1$H and $^{13}$C NMR spectra indicate that 2:TBAF is dynamically stable (Figure 7).

Each NH appears as a doublet in $^1$H NMR, and F$^-$ appears as a double–doublet doublet (ddd) in $^{19}$F NMR; the three NH collapsed into singlets in the fluorine decoupled $^1$H NMR, and the splitting of the fluoride peak disappeared when $^{19}$F NMR was acquired with proton decoupling. This signal multiplicity represents a case of scalar coupling that develops across hydrogen bonds (hydrogen-bond coupling, HBC). These couplings provide unambiguous proof of hydrogen-bonding interactions between NH and F$^-$ for the complex in solution and a measure of their strength. Because HBC results from overlap of H-bond donor and acceptor wave functions, the magnitude of the coupling depends exponentially on donor–acceptor distances, as well as geometry.18 Studies by both Grzesiek and Bax demonstrated that the empirical relationship between HBC and internuclear distances can be exploited for the structural assignment of nucleobases and proteins.19 For HBC to be observable, the hydrogen-bonding network must be rigid, which is often the case for biomolecules held together by multiple hydrogen bonds but is less common in small molecules.20 For hydrogen bonds involving the fluoride ion, pioneering work from Shenderovich and Limbach determined $^{1b}J_{NH···F}$ in $[F(HP)]_3^-$ clusters measured at ultralow temperature,21 while only a few organic molecules such as calix[4]pyrrols and amido cryptand receptors display HBC with fluoride.22 The $^1$H NMR spectrum of 2:TBAF stands out with three resonances displaying hydrogen-bond scalar couplings with fluoride clearly visible. This offers the possibility of measuring correlation spectra across the hydrogen bridge, a feature more often applied to covalently linked nuclei. The $^{19}$F chemical shift is seen on the vertical axis with the magnitude of $^{1b}J_{NH···F}$ determined from cross-peak distances (which matched those visible in the $^1$H spectrum). The three HBC constants measured for 2:TBAF (DCM-$d_2$, 3 mM, 298 K) are $^{1b}J_{NH(a)···F} = 61$ Hz, $^{1b}J_{NH(b)···F} = 51$ Hz, and $^{1b}J_{NH(c)···F} = 34$ Hz.

In DCM-$d_2$, the peak of F$^-$ at −90 ppm for 2:TBAF is deshielded compared to TBAF·$3H_2O$ which resonates at −118 ppm.24 The F$^-$ longitudinal relaxation time constant ($T_1$) would be expected to vary upon formation of a hydrogen-bonded fluoride complex because $T_1$ depends on the motions of the molecule in solution (and hence its size) as well as the proximity of neighboring protons as relaxation sources. Accordingly, $T_1$ measured for $^{19}$F via inversion recovery experiments (magnet field strength = 11.7 T) was found to decrease from 1430 ms for TBAF·$3H_2O$ to 313 ms for 2:TBAF, which is consistent with the formation of a larger, H-bonded molecular complex. The magnitude of the HBC constant for each NH inversely correlates with the NH···F internuclear distance measured from the solid-state structure of 2:TBAF (determined by single-crystal X-ray diffraction), with NH(a)···F$^-$ being the shortest HB contact with fluoride. To correlate $^{1b}J_{NH···F}$ and hydrogen-bond distances in solution, the NOE developed between fluoride and NH was investigated by $^1$H–$^{19}$F heteronuclear Overhauser spectroscopy (HOESY). NOE buildup curves were determined using mixing times between 10 and 600 ms, and NH–F distances relative to NH(a)···F$^-$ were calculated (Figure 8). The distance of NH to fluoride increased in the following order: NH(a) ≈ NH(c) < NH(b), with NH(c)···F$^-$ being 2% longer than NH(a)···F$^-$, and NH(b)···F$^-$ being 12% longer. Larger coupling constants $^{1b}J_{NH···F}$ therefore correspond to shorter HB distance. An additional NOE correlation between the ortho-aromatic proton H(11) with fluoride was observed, although of smaller...
To gain insight into the conformational preference of 2:TBAF, 1H NOESY cross-peak volumes were converted into absolute distances by calibration with the distance of the vicinal protons H(3′) and H(4′) (2.47 Å). This allowed for a quantitative mapping analysis of H−H internuclear distances for 2:TBAF (Figure 11A). Correlation H(16) ⇔ H(8′) indicates that the isopropyl C−H bond points toward naphthyl(I) with a distance between the two protons of 2.6 Å. In contrast to unbound bisurea 2, the protons belonging to the N-iPr group do not correlate with NH(c), which instead shows two NOE with H(11) and NH(b) (2.7 and 2.5 Å), suggesting that the N-isopropyl urea underwent an anti−anti conformational change upon fluoride binding. NH(a) and NH(b) belonging to the same urea display strong NOE (2.0 Å). NH(a) also presents correlation with H(11′) (2.7 Å) and H(11) (3.5 Å). These NOE interactions indicate that the three NH are in close proximity. Deshielding of H(3′) upon fluoride binding indicates more pronounced coplanarity between H(3′) and the adjacent urea carbonyl with respect to unbound 2. The internuclear distances measured by NMR are in keeping with those determined for the solid-state structure using single-crystal X-ray diffraction (Figure 11B).

Studies of Bisurea−TBAF Complexes. With a robust NMR experiment correlating NH−F−F and internuclear distances in place, we studied how fluoride positioned itself within the pocket of a range of BINAM-derived bisurea catalysts. Thirteen complexes were formed by addition of 1 equiv of TBAF:3H2O to a solution of bisurea, and the coupling constants NH−F−F were measured. For each catalyst−fluoride complex, 1H NMR, 19F NMR, and CLIP-HSQC were recorded at 273 K and at low concentration (3 mM) to minimize proton exchange with water, which we had observed to mask NH−F coupling constants. The fluoride complexation could also be responsible for the line broadening observed at a higher concentration. In 19F NMR, fluoride appears as a triplet at –86 ppm. The NH(b) pair is only slightly deshielded upon fluoride complexation, ΔδNH(b) = +1 ppm. The number of proton signals indicate that 1:TBAF

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Figure 9. Relative H−F−F distances calculated for 2:TBAF from HOESY (DCM-d6, 10 mM, 298 K, τm = 10–600 ms) and comparison with single-crystal X-ray diffraction and 1JNH−F−F−F.

Figure 10. Quantitative 1H−19F HOESY of 13:TBAF (DCM-d6, 3 mM, 298 K, τm = 30 ms).

Figure 11. (A) Structure of 2:TBAF and internuclear distances calculated from 1H NOESY (600 MHz, DCM-d6, 3 mM, 298 K, τm = 300 ms) and comparison with X-ray structure data. (B) Solid-state structure from single-crystal X-ray diffraction studies of 2:TBAF.
chains display 1H control the positioning of F, R2 = CF3, F3) resulted in 1H in DCM-2, respectively, indicating reduced interaction between NH(c) highly similar to those for NH(c) that are N-alkylated with di···ffluoride (Table 3, entries 3 and 7). Catalysts 3,5-substituted aromatic rings was examined next. Electron-poor aryl groups are expected to increase NH acidity and therefore hydrogen-bond strength with fluoride; conversely, electron-rich aromatic rings would weaken NH···F hydrogen-bonding interactions. This offers a platform to control the positioning of fluoride within the catalyst pocket, possibly tune its nucleophilicity, and influence the enantioselective delivery. Catalysts 8 (R1 = H, R2 = CF3) and 11 (R1 = F, R2 = CF3) resulted in 1H values of 39 and 44 Hz, respectively, indicating reduced interaction between NH(c) and fluoride (Table 3, entries 8 and 11). Catalysts 9 (R1 = CF3, R2 = H), 10 (R1 = CF3, R2 = Me), and 12 (R1 = CF3, R2 = F) displayed weaker hydrogen bonding between fluoride and NH(a), with 1H values increased and even exceeding 1H for 10 (Table 3, entries 9, 10, and 12). Fluoride is equally shared between NH(a) and NH(c) for 9 and 10 despite the electronic character of the rings expected to favor NH(c). Catalyst 13 (R1 = R2 = F) presents similar 1H values to 2 for NH(a) and NH(b) but a more pronounced variation in NH(c), which is reduced by 4 Hz (Table 3, entry 13).

These data are informative at various levels:

- In solution, all three NH of N-alkylated bisurea catalysts bind fluoride with the N-alkylated urea undergoing anti−anti to syn−anti isomerization. This contrasts with the non-alkylated catalyst 1, which does not engage all four NH groups in hydrogen bonding with fluoride; the two NH bound to fluoride are the ones substituted by the trifluoromethylated aryl groups, and 1H values are of large magnitude (65 Hz). Complex 1:TBAF is less dynamically stable than 2:TBAF as reflected by the broad line and unresolved 1H coupling at 3 mM concentration.

- For all catalysts, NH(a)···F is the dominant hydrogen-bonding interaction in DCM-d2, even when programmed to be weaker by tuning its electronic environment; this is likely due to more favorable geometrical arrangement for hydrogen bonding. NH(c)···F is the hydrogen bond that varies the most, and NH(b) provides the highest HB contribution with 1H values of 33−34 Hz.

- The complexes display 19F chemical shifts between −86 and −94 ppm, which represents a deshielding of ~30 ppm compared to unbound TBAF·3H2O in DCM-d2.

Complexation with CsF. Fluorinations under HB-PTC are carried out with alkali metal fluorides (MFs), requiring the formation of a soluble bisurea−MF complex as a reaction intermediate. The spectroscopic characterization of a representative bisurea−MF complex would therefore provide

![Figure 12. 1H NMR of 1:TBAF (DCM-d2, 0.58 mM, 273 K); (inset) 19F NMR of 1:TBAF showing fluoride splitting.](image)

The bisurea−MF complexes were generated by addition of 1 equiv of TBAF·3H2O (solution in DCM-d2 of known concentration) to the bisurea (3 mM in DCM-d2) and measured at 273 K. 15N DCM/DCM-d2, 0.58 mM.

Table 3. HBC Constants 1H and 19F Chemical Shifts of Bisurea−Fluoride Complexes

| entry | catalyst | R1 | R2 | 1H (Hz) | 1H (Hz) | 1H (Hz) | δ (ppm) |
|-------|----------|----|----|---------|---------|---------|---------|
| 1     | 1        | H  | CF3| 65      |         |         | 1H      |
| 2     | 2        | iPr| CF3| 60      | 50      | 33      | 90.53   |
| 3     | 3        | Me | CF3| 60      | 54      | 34      | 89.41   |
| 4     | 4        | Et | CF3| 61      | 54      | 33      | 88.79   |
| 5     | 5        | nPr| CF3| 60      | 53      | 33      | 88.79   |
| 6     | 6        | 3-Pentyl| CF3| 61      | 52      | 33      | 89.20   |
| 7     | 7        |     | CF3| 61      | 52      | 33      | 91.33   |
| 8     | 8        | iPr| H  | CF3    | 63      | 39      | 93.87   |
| 9     | 9        | iPr| CF3| 53      | 52      | 33      | 93.87   |
| 10    | 10       | iPr| CF3| 52      | 53      | 34      | 90.15   |
| 11    | 11       | iPr| F  | CF3    | 60      | 44      | 89.79   |
| 12    | 12       | iPr| CF3| F      | 59      | 51      | 91.59   |
| 13    | 13       | iPr| F  | F      | 59      | 46      | 91.17   |

The fluoride complexes were generated by addition of 1 equiv of TBAF·3H2O (solution in DCM-d2 of known concentration) to the bisurea (3 mM in DCM-d2) and measured at 273 K. 15N DCM/DCM-d2, 0.58 mM.
NMR spectrum (298–243 K) showed two signals, with the major resonating at −65 ppm and the minor resonating at −96 ppm (Figure 13 C). T$_1$ relaxation was measured by inversion recovery experiments for both fluoride resonances at room temperature (magnet field strength 11.7 T) and found to be 145 and 5 ms for the major and minor species, respectively. These T$_1$ values are substantially shorter than those of 2:TBAF (313 ms) and TBAF in DCM-$d_2$ (1430 ms), suggesting slower tumbling due to the species in solution being larger in size and/or increased contributions from local relaxation sources. 133Cs NMR shows a wide peak (Figure 13 C, inset) broadened as a result of its quadrupolar moment, as well as dynamic averaging of more than one cesium fluoride complex. The 133Cs signal is recorded at −14 ppm at room temperature, which is significantly more shielded than previously observed for the ion pair formed from Cs$^+$ and the weakly coordinating [H$_2$NB$_2$(C$_6$F$_5$)$_6$]$^{-}$ anion (17 ppm), a rare example of 133Cs NMR for a Cs$^+$ salt measured in DCM-$d_2$.²⁸

Proton diffusion experiments (1H DOSY) were performed at 243 K and revealed that the three sets of peaks assigned to isopropyl CH(16) belong to two species. The minor species had a smaller diffusion coefficient ($D = 1.9 \times 10^{-10}$ m$^2$ s$^{-1}$), denoting a larger molecular complex, while the major species displayed faster diffusion ($D = 2.4 \times 10^{-10}$ m$^2$ s$^{-1}$).¹³ These data support the presence of a 1:1 bisurea–fluoride complex as the major species and a minor nonsymmetrical 2:1 complex. 19F NMR indicated that the 1:1 to 2:1 ratio was 85:15 at room temperature.

The NH signals for both species were unambiguously identified using NOESY (ESY) and ROESY correlations (measured, respectively, at 273 and 243 K). $^{1}$H$^{}$F$^{-}$ coupling constants together with $^{1}$H$^{}$−$^{19}$F NOE gave insight into binding modes. The chemical shifts for the NH protons of the 1:1 complex (2:CsF) were comparable to those observed for 2:TBAF, with NH(a) being the most deshielded (δ$_{NH(a)}$ = 11.46 ppm), followed by NH(c) (δ$_{NH(c)}$ = 10.52 ppm) and NH(b) (δ$_{NH(b)}$ = 10.26 ppm). Complex 2:CsF displayed $^{1}$H$^{}$F$^{-}$ splitting of 52 Hz for NH(a), 37 Hz for NH(c), and 34 Hz for NH(b). These values differ from those for 2:TBAF, which shows larger HB coupling constants for both NH(a) and NH(c) (cf. 2:TBAF: $^{1}$H$^{}$F$^{-}$ splitting of 61 Hz, $^{13}$N$^{}$H$^{(c)}$F$^{-}$ splitting of 51 Hz, and $^{13}$N$^{}$H$^{(b)}$F$^{-}$ = 34 Hz, measured at 298 K). $^{13}$N$^{}$H$^{(c)}$F$^{-}$ stands out for being reduced by 14 Hz. The 2:1 complex Cs$_2$(2)F$^{-}$ shows five NH···F correlations in $^{1}$H$^{}$−$^{19}$F CLIP-HSQC and $^{1}$H$^{}$−$^{19}$F ROESY. In Cs$_2$(2)F$^{-}$, one of the two bisureas exhibited tridentate binding to fluoride ($^{13}$N$^{}$H$_{NH(a)}$F$^{-}$ = 40 Hz, $^{13}$N$^{}$H$_{NH(b)}$F$^{-}$ = 37 Hz, and $^{13}$N$^{}$H$_{NH(c)}$F$^{-}$ = 14 Hz), while the other featured two HB contacts engaging its nonalkylated urea submotif ($^{13}$N$^{}$H$_{NH(a)}$F$^{-}$ = 58 Hz and $^{13}$N$^{}$H$_{NH(b)}$F$^{-}$ = 21 Hz). NH(c)’ appeared as a shielded singlet at 5.8 ppm and did not interact with fluoride (Figure 14).

We noted important differences in fluoride chemical shifts. The observed shifts were −96 and −65 ppm for Cs$_2$(2)F$^{-}$ and 2:CsF$^{-}$, respectively. This large difference suggests no interaction between fluoride and Cs$^+$ in Cs$_2$(2)F$^{-}$; in contrast, for 2:CsF, Cs$^+$ likely forms a contact ion pair with F$^−$ in DCM-$d_2$. No data are available in the literature on fluoride chemical shifts. For Cs$_2$F$^{-}$, the fluorescence intensity of Cs$_2$F$^{-}$ is enhanced in apolar solvent likely due to Cs$^+$ screening effects of solubilizing the salt; strongly deshielded F$^−$ in DCM-$d_2$.

The crystallization of bisurea 13 in the presence of CsF gave a single crystal suitable for X-ray diffraction studies. In the solid state, 13:CsF formed columnar superstructures, where Cs$^+$ was bridged between F$^−$ and the two carbonyls of a second bisurea molecule, with each pair of bisurea being oriented in a head-to-tail arrangement (Figure 15). The complex showed 1:1 binding to fluoride, which was itself hydrogen-bonded to all three NH groups. The structure of the BINAM scaffold is highly similar to that of 2:TBAF; however, in 13:CsF NH(c)···F$^−$ showed the longest hydrogen-bonding distance (Table 4).
fluoride as well as a cation–π interaction with naphthyl. From these insights, it appears that Cs⁺ in place of TBA⁺ does not drastically affect the structure of the 1:1 catalyst–fluoride complex, apart from weakening NH(a)···F⁻ and NH(c)···F⁻ interactions. The π–cation interaction of Cs⁺ with BINAM directly contributes to the positioning of Cs⁺ and indirectly to that of fluoride and its HB interactions. An analogous cation–π contact between the positively charged episulfonium ion and naphthyl was found to be a key noncovalent interaction in the transition state (TS) leading to product formation.⁶⁻⁷

The observation that Cs⁺ interacts with fluoride in the solid state and in solution has implications for catalytic fluorination under HB-PTC because the process requires ion metathesis between Cs⁺ and the electrophile (E⁺). Also, under catalytic conditions, the formation of Cs[(2)₂:F] will depend on the rate of formation (phase transfer) and consumption (fluoride delivery) of 2:CsF. For fast fluorination involving highly reactive electrophiles, the formation of Cs[(2)₂:F] will be kinetically unfavorable. However, under conditions favoring aggregation such as high concentration, low temperature, or poorly reactive electrophiles, one would assume that Cs[(2)₂:F] is present in solution.

Further information was gained on reactivity. To a solution of 2:CsF/Cs[(2)₂:F] generated in situ by sonication of 2 and CsF in DCM-d₂, β-bromosulfolide 1a (2 equiv) was added as a solid, and the sample was briefly agitated. This resulted in the instantaneous formation of a precipitate assigned as CsBr. ¹H and ¹⁹F NMR analyses showed the presence of β-fluorosulfolide 3a formed with an enantiomeric ratio (e.r.) of 86.5:13.5, which is consistent with the enantioselectivity measured for the catalytic reaction (cf. e.r. = 88:12). This experiment unambiguously demonstrated that 2:CsF (in equilibrium with Cs[(2)₂:F]) is effective for the enantioselective fluorination of 1a, an observation that supports our mechanistic hypothesis (Figure 1A).

Impact of Multiple H Bonds to Fluoride on Catalytic Fluorination. Preliminary computational studies suggested that 2:Fs⁻ participates in the enantiodetermining step for the fluorination of meso-episulfonium ions under HB-PTC,⁶⁻⁷ striking structural similarities were found for the bisurea–fluoride anionic component featured in the TS and complex 2:TBAF in its ground state (in the solid state and in DCM-d₂). The structure and conformation of bisurea–fluoride complexes in their ground state could therefore provide valuable information to understand catalyst performance. Catalysts 1–13 were tested with the fluorination of β-bromosulfolide 1a with CsF in DCM or 1,2-difluorobenzene (1,2-DFB) at room temperature for 1.5 h (Table S). Initially, yields and enantioselectivities were determined for catalysts 3–7 bearing different N-alkyl groups and compared with the results obtained with the nonalkylated bisurea 1 and N-iPr bisurea 2 (Table S, entries 1–7). All catalysts 1–7 provided excellent yields (>95%), implying that the 3,5-bis(trifluoromethyl)-phenyl substituents ensure strong binding with fluoride and thus effective solid–liquid phase transfer for CsF. The nonalkylated BINAM urea 1 led to a lower e.r. compared to all N-alkylated analogues (entry 2, 86:14 e.r. (ΔΔΔΔG° = 1.07 kcal/mol). For catalysts 4–6, the level of enantioselectivity achieved is highly similar to that for 2 (e.r. 89.5:10.5–90:10); only the N-methylated catalyst 3 provided (S,S)-3a in slightly lower e.r. (entry 3, e.r. 88:12). N-(3-Pentyl) catalyst 7 on the other hand led to slight improvement (entry 7, 91:9 e.r. (ΔΔΔΔG° = 1.4 kcal/mol).

A distinctive cation–π interaction between Cs⁺ and naphthyl-(II) (distance of 3.328(4) Å) was also prominent. Compared with the NH···F distances of 13:TBAF determined by NMR in DCM-d₂ (Table 4), an elongation of NH(c)···F⁻ and shortening of NH(b)···F⁻ were observed. NH(a)···F⁻ displayed a similar HB distance in solution and in the solid state. The Cs···F distance for 13:CsF was 2.742(3) Å,¹⁵ and Cs⁺ displayed short-distance contacts with the two carboxyls of a neighboring bisurea, a multimeric arrangement likely reinforced by solid-state packing. In solution, the shielding for ¹³³Cs resonance was consistent with an interaction with

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**Figure 14.** NMR parameters measured for NH resonances and F⁻ of 2:CsF and Cs[(2)₂:F].

**Figure 15.** Solid-state structure of 13:CsF from single-crystal X-ray diffraction (solvent omitted for clarity).

**Table 4. Relevant Intramolecular Distances of 13:F⁻⁻⁻**

| DH···A | X-ray 13:CsF | NMR 13:TBAF |
|-------|-------------|-------------|
| d D–H (Å) | d D–A (Å) | d H–A (Å) | d H–A (Å) |
| NH(a)···F⁻ | 0.85 | 2.641(7) | 1.88 | 1.83 |
| NH(b)···F⁻ | 0.87 | 2.727(6) | 1.93 | 2.05 |
| NH(c)···F⁻ | 0.85 | 2.837(6) | 2.10 | 1.91 |
| CH(11)···F⁻ | 0.93 | 3.119(7) | 2.52 | 2.52 |

*H = HB donor, A = HB acceptor. Determined by NMR from HOESY experiments on 13:TBAF.*
The small variation in e.r. for catalysts bearing different N-alkyl groups suggests that this structural variation does not play a prominent role in differentiating between the TSs leading to one or the other enantiomer. This is corroborated by NMR conformational analyses, X-ray structures, and DFT calculations, which indicate that the N-alkyl substituent points away from the catalyst pocket that binds fluoride. These data also inform that the tridentate fluoride binding mode of N-alkylated catalysts 2–7 is beneficial over bidentate binding to ensure higher e.r. as demonstrated with the lower performance of 1, which lacks an N-alkyl group. Next, N-isopropyl bisureas 8–13 served the purpose of investigating the influence of NH-ary1 substitution on both yields and e.r. (Table 5, entries 8–13). In this series, only catalysts 12 and 13 featuring fluorine substitution on both aryl rings afforded 3a in >95% yield, offering optimal performance as a phase-transfer catalyst for CsF and the ability to release fluoride for C–F bond construction. The replacement of 3,5-bis(trifluoromethyl)phenyl for phenyl or meta-xylyl, as shown with catalysts 8–10, was detrimental as yields plummeted as low as 18%, indicating poor phase-transfer capability. The most drastic effect was encountered when the aryl group attached to NH(c) lacked electron-deficient substituents (catalyst 8). Reactivity is dependent on the solvent, with catalysts 8–10 affording higher yields in DCM and catalyst 11 being more effective in 1,2-DFB. The trends in enantioselectivity are similar in both solvents, with 1,2-DFB giving more often higher enantiomeric ratios. In this series, catalyst 13 is the least effective in terms of enantiocontrol. In all cases, no side products were detected in the crude mixture based on H NMR analysis.

The NMR data of the best-performing catalyst—TBAF complexes indicate that bisureas having stronger NH(c)···F– contribution (larger \( \Delta_{\text{NH(c)···F–}} \)) typically lead to a superior reaction outcome, suggesting that NH(c)···F– is a key interaction for achieving high yield and enantioselectivity. The striking role of NH(c) was highlighted in our original report, which presented intrinsic reaction coordinates to shed light on the mechanism of the reaction. The calculations showed that the three NH···F hydrogen bonds elongated during fluoride delivery to the epilsulphonium ion as a consequence of charge neutralization; however, the elongation of NH(c)···F– occurred faster compared to those of NH(a)···F– and NH(b)···F–.

### CONCLUSION

This investigation has unveiled important information on the conformation of BINAM-derived bisureas and their ability to bind fluoride in solution. Analysis of a set of bisureas featuring up to four NH groups enabled us to draw conclusions on the effect of multiple hydrogen bonds to fluoride on catalytic nucleophilic fluorination. The key findings are summarized hereafter.

- In DCM-d<sub>2</sub> (25 mM), N-alkylated BINAM-derived bisureas with three NH groups exist as equilibrating structures resulting from an intramolecular time-dependent hydrogen bond that involves the two urea motifs. The corresponding nonalkylated BINAM bisurea with four HB donor NHs does not display such intramolecular interaction but undergoes aggregation when the concentration increases.
- N-alkylated and alkylated bisureas bind fluoride in DCM-d<sub>2</sub>, resulting in the formation of two equilibrating species: a 2:1 bisurea–F– complex at low TBAF stoichiometry and a more stable 1:1 complex dominant when TBAF stoichiometry is >1 equiv. Conformational analysis by quantitative NOESY informed that N-alkylated bisureas underwent anti–anti to syn–anti isomerization of the N-alkylated urea upon addition of TBAF. This conformational change led to a dynamically stable 1:1 complex, allowing all
three NHs to bind fluoride, an arrangement similar to the conformation observed for this complex in the solid state. The corresponding nonalkylated bisurea binds fluoride, engaging two of its four NH groups.

- Hydrogen-bond coupling constants ($^{3}J_{\text{NH-F}}$) were observed and measured by $^{1}H-^{19}F$ CLIP-HSQC experiments, providing insight into the hydrogen-bonding network around fluoride, with distances calculated from $^{1}H-^{19}F$ NOE experiments. All N-alkylated bisurea catalysts bind fluoride with all three NHs participating in hydrogen bonding, NH(a), located on the nonalkylated urea submotif and away from naphthyl, suggesting that this electronic pattern was necessary for additional insight. Excellent reactivity was observed for all catalysts all characterized by NMR spectroscopy provided $37 \text{ Hz}$, $1\text{H}$ $37 \text{ Hz}$, and $1\text{H}$ $33 \text{–}34 \text{ Hz}$. In contrast, only two of the four NH groups of nonalkylated BINAM bisurea I displayed HB with fluoride ($^{3}J_{\text{NH-F}}=65 \text{ Hz}$); the NHs involved are those proximal to the 3,5-bis(trifluoromethyl)phenyl rings.

- A bisurea–fluoride complex prepared from CsF was characterized in solution and in the solid state. Low-temperature NMR and $^{1}H$ diffusion experiments revealed two species in equilibrium, a 1:1 bisurea–CsF complex as the main species along with a less-abundant 2:1 complex (85:15 at 298 K). $^{3}J_{\text{NH-F}}$ detection provided information on the binding mode, with the 1:1 species showing a four-centered trifurcated HB with fluoride ($^{3}J_{\text{NH(a)-F}}=52 \text{ Hz}$, $^{3}J_{\text{NH(c)-F}}=37 \text{ Hz}$, and $^{3}J_{\text{NH(b)-F}}=34 \text{ Hz}$). The 2:1 bisurea–CsF led to a six-centered hydrogen-bond network, engaging both bisureas as hydrogen-bond donors ($^{3}J_{\text{NH(a)-F}}=40 \text{ Hz}$, $^{3}J_{\text{NH(b)-F}}=37 \text{ Hz}$, $^{3}J_{\text{NH(c)-F}}=14 \text{ Hz}$, $^{3}J_{\text{NH(d)-F}}=58 \text{ Hz}$, and $^{3}J_{\text{NH(b)-F}}=21 \text{ Hz}$). Spectroscopic studies of the 1:1 complex agreed with the solid-state structure from single-crystal X-ray diffraction, indicating that the Cs$^+$ ion interacted both with naphthyl via a cation–π interaction and with F$^-$. These interactions resulted in some reorganization of the HB network in the catalytic pocket, with NH(a)...F$^-$ and NH(c)...F$^-$ interactions reduced compared to those observed in the corresponding TBAF complex. The reaction of the preformed bisurea–CsF complex with a model β-bromosulfoxide led to instantaneous formation of the expected enantioenriched β-fluorosulfide; this experiment demonstrated that this bisurea–CsF complex enabled asymmetric fluorination with fast fluoride delivery.

- Catalytic fluorinations carried out with thirteen bisurea catalysts all characterized by NMR spectroscopy provided additional insight. Excellent reactivity was observed for all catalysts with electron-withdrawing substituents on the N-aryl groups, suggesting that this electronic pattern was necessary for the catalyst to act as a phase-transfer agent for CsF. The resulting bisurea–fluoride complexes remained competent nucleophiles and provided the necessary chiral environment for enantioselective fluoride delivery. The data show that the hydrogen-bond interaction NH(c)...F$^-$ was critical to ensure good control over enantioselectivity. The nonalkylated bisurea gave the desired product in high yield, indicating that the formation of a three-centered bifurcated fluoride complex was sufficient for phase-transfer but not optimal for enantioselectivity.

In this study, we demonstrated that $^{1}H-^{19}F$ NMR spectroscopy is a powerful tool to analyze multiple H-bonding interactions with fluoride, thereby offering insight into catalyst performance for enantioselective nucleophilic fluorination under HB-PTC. Considering the importance of hydrogen-bonding interactions in catalysis, we anticipate that the analytical approach described herein would encourage further investigations with a multitude of catalyst candidates featuring multiple hydrogen-bond donor functionalities that are capable of anion binding.

## ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/10.1021/jacs.0c09832.

NMR methods and characterization, binding studies, hydrogen-bond coupling, catalyst synthesis and characterization, and NMR spectra (PDF)

Crystallographic data of 13-CsF (CIF)

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### Notes

The authors declare no competing financial interest.

Crystallographic data have been deposited with the Cambridge Crystallographic Data Centre (2031585) and can be obtained free of charge via http://www.ccdc.cam.ac.uk/data_request/cif.

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