

NMR Spectroscopy

Predicting ^{19}F NMR Chemical Shifts: A Combined Computational and Experimental Study of a Trypanosomal Oxidoreductase–Inhibitor Complex

Johannes C. B. Dietschreit, Annika Wagner, T. Anh Le, Philipp Klein, Hermann Schindelin, Till Opatz, Bernd Engels, Ute A. Hellmich,* and Christian Ochsenfeld*

Abstract: The absence of fluorine from most biomolecules renders it an excellent probe for NMR spectroscopy to monitor inhibitor–protein interactions. However, predicting the binding mode of a fluorinated ligand from a chemical shift (or vice versa) has been challenging due to the high electron density of the fluorine atom. Nonetheless, reliable ^{19}F chemical-shift predictions to deduce ligand-binding modes hold great potential for *in silico* drug design. Herein, we present a systematic QM/MM study to predict the ^{19}F NMR chemical shifts of a covalently bound fluorinated inhibitor to the essential oxidoreductase trypanothione (Tpx) from African trypanosomes, the causative agent of African sleeping sickness. We include many protein–inhibitor conformations as well as monomeric and dimeric inhibitor–protein complexes, thus rendering it the largest computational study on chemical shifts of ^{19}F nuclei in a biological context to date. Our predicted shifts agree well with those obtained experimentally and pave the way for future work in this area.

Fluorine is considered a “magic” element in medicinal and agricultural chemistry. It forms strong bonds to carbon, is the smallest biocompatible hydrogen substitute,^[1] has the ability to form hydrogen bonds, and possesses a high electronegativity. Its introduction into small molecules can increase metabolic stability and allows the fine-tuning of physico-chemical properties.^[2] It is therefore not surprising that more

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than 20% of all FDA-approved drugs and more than 30% of all agrochemicals contain fluorine.^[2] Replacing hydrogen by fluorine has been used successfully to, for example, investigate the interaction of inhibitors with proteases, explore their active site properties, and characterize inhibitors for neglected tropical diseases.^[3]

With its 100% natural abundance, high gyromagnetic ratio, and the resulting high sensitivity, the spin-1/2 nucleus ^{19}F is of particular interest for NMR studies.^[4] While practical advantages of fluorine for NMR spectroscopy have been exploited for many decades, the performance of corresponding quantum-chemical calculations for complex systems has gained momentum only lately.^[5]

Chemical shifts of compounds containing fluorine have been calculated for many decades, from small molecules in the gas phase over biological systems in solution to solid-states.^[6] The two most recent studies focusing on ^{19}F chemical shifts of biologically relevant molecules investigated crystals of fluorinated tryptophans^[7] or monofluorinated phenylalanines in a protein (Brd4).^[8] In the case of the tryptophan crystals, four molecules were used as a representation of the entire crystal. For Brd4, a quantum-mechanical/molecular-mechanical (QM/MM) setup was used with a buffer region of 4 Å and Boltzmann weighting of a few conformers. Nonetheless, the calculations differed from the measurements by between one and more than 20 ppm even after improving

[*] J. C. B. Dietschreit, Prof. Dr. C. Ochsenfeld
Theoretical Chemistry,
Department of Chemistry, University of Munich (LMU)
Butenandtstr. 7, 81377 Munich (Germany)
E-mail: christian.ochsenfeld@uni-muenchen.de
Prof. Dr. C. Ochsenfeld
Max Planck Institute for Solid State Research
70569 Stuttgart (Germany)
E-mail: c.ochsenfeld@fkf.mpg.de
A. Wagner, Prof. Dr. U. A. Hellmich
Dept. Chemistry, Section Biochemistry,
Johannes Gutenberg-Universität Mainz
55128 Mainz (Germany)
and
Centre for Biomolecular Magnetic Resonance (BMRZ),
Goethe-University Frankfurt
Max-von-Laue Str. 9, 60438 Frankfurt (Germany)
E-mail: u.hellmich@uni-mainz.de
T. A. Le, Prof. Dr. B. Engels
Institute for Physical and Theoretical Chemistry,
University of Würzburg
Emil-Fischer-Straße 42, 97074 Würzburg (Germany)

P. Klein, Prof. Dr. T. Opatz
Dept. Chemistry, Section Organic Chemistry,
Johannes Gutenberg-Universität Mainz, 55128 Mainz (Germany)
Prof. Dr. H. Schindelin
Institute of Structural Biology,
Rudolf Virchow Center for Experimental Biomedicine,
University of Würzburg, 97080 Würzburg (Germany)

Supporting information, which includes the experimental (NMR measurements) and computational (MM-MD simulation and QM/MM calculations) methods, NMR dilution curves, analysis of protein complex stability during the MD simulations, and figures highlighting the flexibility of CFT in complex with Tpx, and the ORCID identification number(s) for the author(s) of this article can be found under:
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predictions by linear regression to experimental data. Another study benchmarked different levels of quantum-chemical methods for fluorinated amino acids in implicit solvent, achieving at best a mean absolute error of 2.68 ppm with respect to the experiment.^[9] Despite the impressive progress in the field, this is not sufficient to explain subtle differences in experimental spectra. Here, we use hundreds of frames from molecular dynamics (MD) simulations to ensure proper sampling of conformers and a significantly larger buffer region in our QM/MM calculations to increase the accuracy of our results.

Methods for computing NMR parameters range from empirical programs, such as SPARTA+,^[10] to highly accurate QM calculations.^[5,11,12] When using quantum-chemical methods, it has been shown that sufficiently large QM regions are necessary when describing complex systems.^[13,14] However, the inclusion of many atoms is computationally very demanding. Thus, a plethora of methods has been devised to reduce the computational effort.^[14,15] Here, we employ rigorous linear-scaling formulations that allow us to exploit the locality of the electronic structure within density-matrix-based theories. While this strongly reduces the computational scaling, for example, for the computation of NMR chemical shifts within density-functional theory from cubic to asymptotically linear, the accuracy is numerically unchanged and fully controlled.^[5,16]

As a medically relevant test system, we selected the oxidoreductase trypanothione (Tpx), an essential enzyme of *Trypanosoma brucei*, the parasite that causes African sleeping sickness.^[17] Tpx is inhibited by covalently binding to 2-(chloromethyl)-5-(4-fluorophenyl)thieno[2,3-d]pyrimidine-4-(3*H*)-one, CFT, which efficiently kills *T. brucei*.^[18,19] CFT carries a 4-fluorophenyl moiety (Figure 1 A and Supporting Information, Figure S5). The chlorine leaving group facilitates the covalent interaction with Cys40 in the active site of Tpx.

In the asymmetric unit of our monoclinic crystals, three protein chains with two different inhibitor orientations are present (PDB: 6GXY, binding pose 1 for chains A and B, binding pose 2 for chain C, Figure 1 B).^[19] In binding pose 1, the covalently bound CFT features extensive intramolecular interactions with the protein, including T-shaped π -stacking interactions with Trp70 and a weak hydrogen bond of the CFT fluorine with the backbone-H α of Glu107. In binding pose 2, CFT is not in contact with the protein beyond the covalent bond to Cys40, and its fluorine atom is solvent exposed instead. In both, crystal and solution, CFT binding to the wild-type protein in pose 1 leads to Tpx dimerization mediated by extensive intermolecular inhibitor–inhibitor stacking and inhibitor–protein interactions.^[19] The dissociation constant for the CFT-induced Tpx dimer is approximately 5 μ M. In binding pose 2, dimerization is structurally not possible. We

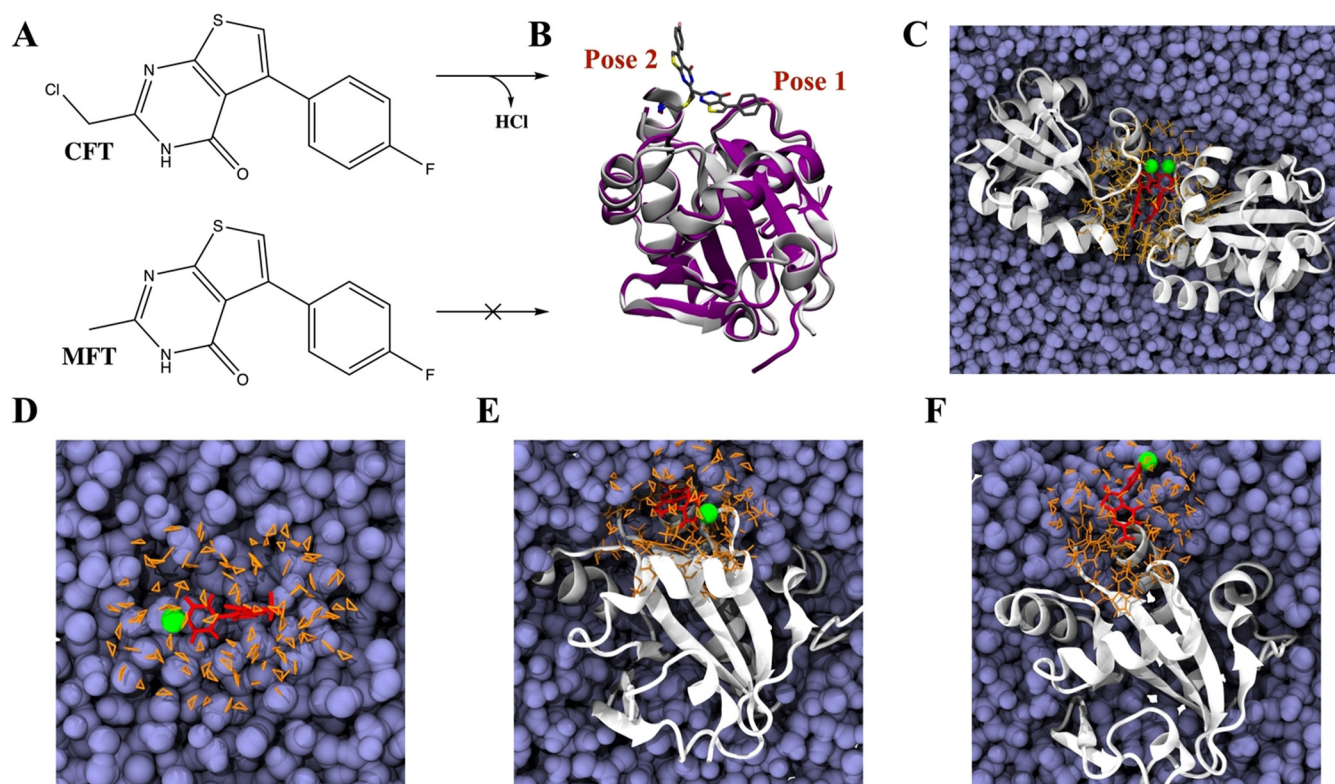


Figure 1. Interaction of *T. brucei* oxidoreductase trypanothione (Tpx) with a covalent inhibitor. A) cysteine-reactive CFT (top) and non-reactive MFT (bottom). B) Overlay of Tpx–CFT monomers in poses 1 and 2 as observed in our crystal structures (PDB: 6GXY).^[19] C–F) Depiction of the QM region and MM embedding. Tpx is shown in white, water in blue, and all atoms in the QM region as orange sticks. The inhibitor is highlighted in red with its fluorine atom as green sphere. C) shows the Tpx–CFT dimer, D) the inhibitor in solution, E) the Tpx–CFT monomer in pose 1, and F) the Tpx–CFT monomer in pose 2.

identified residue Trp39 to be crucial for dimerization. Mutation of this active-site residue to alanine (Tpx-W39A) yields a protein that can still covalently interact with CFT, but dimerization upon inhibitor binding is extremely weak (Supporting Information, Figure S1).^[19]

The ¹⁹F signals for free CFT and its unreactive analogue 2-methyl-5-(4-fluorophenyl)thieno[2,3-d]pyrimidin-4(3*H*)-one, MFT, which is missing the chlorine leaving group (Figure 1A), were measured in solution at 298 K. Both ¹⁹F chemical shifts were found to be very similar (−114.79 and −114.77 ppm, respectively). Upon binding of CFT to Tpx and the subsequent dimerization, a downfield chemical shift of approximately 0.3 ppm for the ¹⁹F signal (−114.47 ppm) and substantial line broadening are observed, in agreement with incorporation of CFT into a high molecular weight, dimeric complex (Figure 2). Hence, the simultaneous availability of ¹⁹F NMR and X-ray data for the Tpx–inhibitor system renders it exceptionally well-suited for systematic ¹⁹F chemical shift studies. At high concentrations (greater than 500 μM), the ¹⁹F chemical shift for the CFT–Tpx complex does not depend on the concentration of the protein–inhibitor complex. This suggests that under these conditions the ¹⁹F chemical shift is not influenced by the monomer/dimer equilibrium since these concentrations are far above the K_D for dimerization. Only after significantly lowering the concentration, the ¹⁹F signal starts to shift further downfield, indicating an increasing population of the monomer in exchange with the dimer

(Supporting Information, Figure S2). However, due to limitations in the signal-to-noise ratio for the ¹⁹F NMR measurements at high dilution, the concentration of the complex could not be reduced far enough below the K_D ($\ll 5 \mu\text{M}$) to reach conditions where the monomer is exclusively observed. In agreement with the above observation, the measured chemical shift for CFT bound to the W39A-mutant is shifted further downfield by an additional approximately 0.8 ppm compared to the WT complex (−113.7 ppm). This mutant does not significantly dimerize, and the distances from the inhibitor's fluorine group to the W39 C α atom and to the W39 indole group are more than 16 Å and 11.5 Å, respectively, and thus should not affect the inhibitor's ¹⁹F chemical shift. Ergo, this mutant can be used as a reference point for the ¹⁹F chemical shift of CFT bound to a Tpx monomer.

In contrast to small organic molecules, the free energy landscape of solvated biomolecules typically does not possess one deep minimum, but rather a multitude of energetically close configurations that are thermodynamically accessible at physiological temperatures. In the NMR experiment, this implies the recording of an ensemble-averaged NMR chemical shift. Therefore, many different configurations of the system have to be taken into account to compute the observed chemical shift.^[20] It has been demonstrated that inclusion of bond-length and bond-angle vibrations is often crucial for reliable chemical-shift computations.^[21,22] Furthermore, the different relative orientations of molecules have to be accounted for as well. Here, we present a robust methodology based on MM-MD sampling and subsequent QM/MM calculations of ¹⁹F NMR chemical shifts to identify CFT's binding poses relevant in solution.

Based on the crystal structure of the Tpx–CFT complex (PDB: 6GXY),^[19] we computed separate MM-MD trajectories for free CFT and MFT in solution, for the monomeric complex with CFT bound in either pose 1 or 2, and the Tpx-dimer with both protomers binding to CFT in pose 1 (see the Supporting Information for setup details). An analysis of the complex stability during the simulations and inhibitor flexibility is given in Figures S4–S8 and Table S2 in the Supporting Information.

Subsequently, the ¹⁹F chemical shifts of the inhibitor were calculated with our linear-scaling methods^[5,16] for 200 snapshots taken evenly spaced in time from each MD trajectory. All interactions of either the protein or solvent atoms with the inhibitor were included explicitly. The aim was to perform high quality ¹⁹F chemical-shift calculations on a large scale, based on an adequate description of the accessible phase space. The configurational ensembles included, for example, different solvation patterns or orientations of the inhibitor towards Tpx.

The high electron density of ¹⁹F renders its spectroscopic properties, such as NMR chemical shifts, particularly challenging to calculate.^[23] Motivated by previous studies,^[12,13] we tested KT2^[24] and B97-2^[25] together with the NMR-specific basis set pcS-1,^[26] and the necessary QM buffer size (Supporting Information, Figure S9). Both functionals show identical QM size convergence. As the radius of the QM region is increased from 4 to 5 Å, the calculated chemical shift changes by more than 0.5 ppm, highlighting the importance of

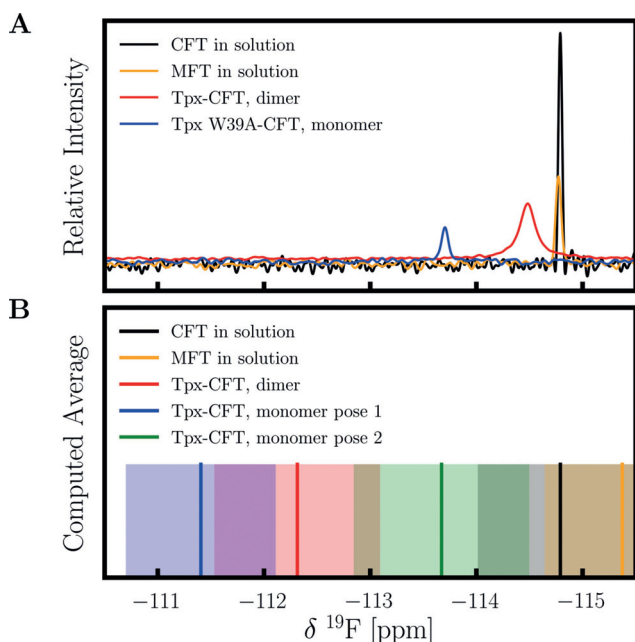


Figure 2. Comparison of experimentally measured (A) and calculated (B) ¹⁹F NMR shifts. A) We compare CFT and MFT in solution, CFT in the dimeric complex and the W39A monomeric mutant. In the experiment, only one peak could be found for the monomeric protein. B) The computed averages are shown as vertical lines with the SEM indicated by an area shaded with decreased saturation. The calculations distinguish between two poses observed in the crystal structure (Figure 1). Importantly, the calculations and the experiment reveal the same ordering of peaks and indicate that pose 2 does not exist in solution.

a sufficiently large QM sphere. A 7 Å QM buffer region around the inhibitor was found to be necessary to obtain size-converged shifts. Hence, we performed QM/MM-NMR calculations with KT2/pcS-1 for a 7 Å QM region. For the dimeric wild-type Tpx-CFT complex, this region (Figure 1C) includes more than 1000 atoms. In combination with the 200 sampling points taken from each of the individual MD trajectories, this makes it one of the largest QM-based ¹⁹F NMR chemical-shift calculations reported so far.^[7,8,27] The QM size and the combined number of calculations in this context are unprecedented and are at the frontier of what is currently possible for such large biological systems. The embedded QM regions are shown in Figure 1C–F. A detailed description is given in the Supporting Information (Supporting Information, Table S3).

It is important to note that the values of the calculated ¹⁹F shifts for individual MD frames are scattered over a large shift range of about 60 ppm (Supporting Information, Figure S10). This is, however, not unexpected due to bond-length and bond-angle vibrations in the MD simulations.^[21] The vibrations causing these distributions, especially those of the C–F bond, are fast processes on the spectroscopic time scale, and thus are not observed experimentally. We use the experimentally measured ¹⁹F signal of free CFT in solution as reference for our calculated values, as relative shieldings are much more accurate.

As expected, our calculated ¹⁹F chemical shifts correctly predict that the chemical shifts for free CFT and MFT are very similar to each other. Importantly, the predicted chemical shift for the inhibitor bound to Tpx is calculated to be downfield shifted compared to free CFT. An even more pronounced downfield shift compared to free CFT is predicted for CFT bound to monomeric Tpx in binding pose 1. This is in excellent agreement with what is observed experimentally for CFT bound to the monomeric W39A Tpx-mutant and in our dilution experiments (Supporting Information, Figures S1 and S2). In contrast, for the monomeric complex in pose 2, a chemical shift is calculated that lies between free CFT and the dimeric complex. Thus, our calculations qualitatively predict the correct order of the ¹⁹F chemical shifts for CFT in the different states, as well as the true direction of chemical-shift changes induced by protein binding and complex dissociation. They further suggest that binding pose 2 of CFT observed in chain C of the crystal structure is not relevant in solution, as one would then expect, for CFT bound to the monomeric W39A mutant, a ¹⁹F signal with a chemical shift in between those of free CFT and CFT bound to dimeric wild-type Tpx. This agrees well with the extended degree of solvent exposure of the inhibitor in this binding pose (Supporting Information, Figure S8), thus rendering it more similar to the free inhibitor. However, our calculations overestimate the chemical-shift differences between the different states. Already the calculated chemical shift difference between free CFT and free MFT (0.58 ppm) is larger than the measured one (0.02 ppm). This pattern continues for the other pairwise chemical-shift differences (CFT vs. CFT-WT: 0.32 ppm/2.48 ppm; Dimer vs. Monomer (pose 1): 0.77 ppm/0.97 ppm). Nevertheless, the computed trend allows us to discriminate between the different struc-

tures observed experimentally and to assign the measured shifts to a given conformer.

The accuracy of the prediction could be further increased by using a higher level of theory, a larger basis set, or more accurate dynamics (QM/MM-MD instead of MM-MD) improving the description of bond lengths, vibrations, and non-covalent interactions, which would entail, however, significantly higher computational costs.

Our study underlines the usefulness of ¹⁹F NMR for the investigation of complex protein–inhibitor interactions, showcases current computational possibilities, and illustrates the power of predicting ¹⁹F NMR chemical shifts in a complex biological system as a prerogative for further biomedical applications and drug design.

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Conflict of interest

The authors declare no conflict of interest.

Keywords: African sleeping sickness · covalent inhibitors · NMR spectroscopy · quantum chemistry · structural biology

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