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FLORIDA INTERNATIONAL UNIVERSITY

Miami, Florida

THEORETICAL STUDY OF CHLOROPEROXIDASE CATALYZED CHLORINATION OF BETA-CYCLOPENTANEDIONE AND ROLE OF WATER IN THE CHLORINATION MECHANISM

A dissertation submitted in partial fulfillment of the

requirements for the degree of

DOCTOR OF PHILOSOPHY

in

CHEMISTRY

by

Cassian D'Cunha

2011

To: Dean Kenneth Furton College of Arts and Sciences

This dissertation, written by Cassian D'Cunha, and entitled Theoretical Study of Chloroperoxidase Catalyzed Chlorination of beta-Cyclopentanedione and Role of Water in the Chlorination Mechanism, having been approved in respect to style and intellectual content, is referred to you for judgment.

We have read this dissertation and recommend that it be approved.

Alexander Mebel

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Date of Defense: November 9, 2011

The dissertation of Cassian D'Cunha is approved.

Dean Kenneth Furton College of Arts and Sciences

Dean Lakshmi N. Reddi University Graduate School

Florida International University, 2011

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DEDICATION

I dedicate this dissertation to my parents. To my mother, my role model, for her constant

love and support, and in loving memory of my father.

ACKNOWLEDGMENTS

I would first like to thank my committee members. Their help, encouragement and patience during the various stages of my dissertation have been most appreciated. I would like to thank Dr. Xiaotang Wang for being particularly helpful during the design phase of my dissertation. I would like to thank Dean Kenneth Furton for taking time from his extremely busy schedule to serve on my committee. I would like to thank Dr. Bernard Gerstman for taking an interest in my research and inviting me to present my research to his research group on several occasions. I would like to thank Dr. Alexander Mebel for being supportive and patient, and for answering the many questions I had regarding quantum mechanics approaches. I would especially like to thank Dr. David Chatfield, my major professor. He has been very supportive, encouraging and patient, and I sincerely appreciate the time I spent under his guidance. I cannot thank him enough for the time he took to discuss research, guide me through my dissertation and make resources available for my research. His mentorship was invaluable and much appreciated.

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I would like to thank my friends and roommates over the years. They have been my family away from home while I was pursuing my degree. I would also like to thank the Martinez family that has all but adopted me and made me feel welcome in their home.

V

ABSTRACT OF THE DISSERTATION

THEORETICAL STUDY OF CHLOROPEROXIDASE CATALYZED CHLORINATION OF BETA-CYCLOPENTANEDIONE AND ROLE OF WATER IN THE CHLORINATION MECHANISM

by

Cassian D'Cunha

Florida International University, 2011

Miami, Florida

Professor David Chatfield, Major Professor

Chloroperoxidase (CPO) is a potential biocatalyst for use in asymmetric synthesis. The mechanisms of CPO catalysis are therefore of interest. The halogenation reaction, one of several chemical reactions that CPO catalyzes, is not fully understood and is the subject of this dissertation.

The mechanism by which CPO catalyzes halogenation is disputed. It has been postulated that halogenation of substrates occurs at the active site. Alternatively, it has been proposed that hypochlorous acid, produced at the active site via oxidation of chloride, is released prior to reaction, so that halogenation occurs in solution. The freesolution mechanism is supported by the observation that halogenation of most substrates often occurs non-stereospecifically. On the other hand, the enzyme-bound mechanism is supported by the observation that some large substrates undergo halogenation stereospecifically.

The major purpose of this research is to compare chlorination of the substrate β cyclopentanedione in the two environments. One study was of the reaction with limited hydration because such a level of hydration is typical of the active site. For this work, a purely quantum mechanical approach was used. To model the aqueous environment, the limited hydration environment approach is not appropriate. Instead, reaction precursor conformations were obtained from a solvated molecular dynamics simulation, and reaction of potentially reactive molecular encounters was modeled with a hybrid quantum mechanical/molecular mechanical approach. Extensive work developing parameters for small molecules was pre-requisite for the molecular dynamics simulation.

It is observed that a limited and optimized (active-site-like) hydration environment leads to a lower energetic barrier than the fully solvated model representative of the aqueous environment at room temperature, suggesting that the stable water network near the active site is likely to facilitate the chlorination mechanism.

The influence of the solvent environment on the reaction barrier is critical. It is observed that stabilization of the catalytic water by other solvent molecules lowers the barrier for keto-enol tautomerization. Placement of water molecules is more important than the number of water molecules in such studies. The fully-solvated model demonstrates that reaction proceeds when the instantaneous dynamical water environment is close to optimal for stabilizing the transition state.

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LIST OF ACRONYMS

ACRONYM	FULL NAME
ABNR	Adopted Basis Newton Raphson
ACM	Adiabatic Connection Method
AGLC	Alpha-D-Glucose
ANEB	Adaptive Nudged Elastic Band
AXYS	Alpha-D-Xylopyranos
BXYS	Beta-L-Xylopyranos
BARB	Beta-L-Arabinose
BNAG	N-Acetyl-Glucosamine
CC	Coupled Cluster
CI-NEB	Climbing Image-Nudged Elastic Band
СРО	Chloroperoxidase
CPD	1,3 Cyclopentanedione
DFT	Density Functional Theory
DHP	Dehaloperoxidase
ESP	Electrostatic Potential
GGA	Generalized Gradient Approximation
GTO	Gaussian Type Orbitals
GRMS	Root Mean Square Gradient
HF	Hartree-Fock
IC	Internal Coordinate
IRC	Intrinsic Reaction Coordinate

LCAO	Linear Combination of Atomic Orbitals
LDA	Local Density Approximation
MCD	Monochlorodimedone
MD	Molecular Dynamics
MEP	Minimum Energy Path
МК	Merz-Kollman
MM	Molecular Mechanics
МО	Molecular Orbitals
MP	Moller-Plesset
NEB	Nudged Elastic Band
NMR	Nuclear Magnetic Resonance
PCA	Pyroglutamic Acid
PCM	Polarizable Continuum Model
PES	Potential Energy Surface
QM	Quantum Mechanics
QM/MM	Quantum Mechanics/Molecular Mechanics
RMSD	Root Mean Square Deviation
RPATH	Replica Path
SD	Steepest Descent
STO	Slater Type Orbitals
TS	Transition State
VDW	van der Waals

1. INTRODUCTION

1.1.Chloroperoxidase

Shaw and Hager [1] reported chlorination activity when extracts of a marine fungus, *Caldariomyces fumago*, were used to react with β -ketoadipic acid to give δ -chlorolevulinic acid. Another observation made was that the main metabolite of this organism was a chlorinated species, 2,2-dichloro-1,3-cyclopentanediol, clearly indicating that there was an enzyme or group of enzymes in the organism that facilitated chlorination. It was later discovered by the same group that there were at least four enzymes in the mold extract. The four enzymes identified were an amyloltic enzyme, a catalase, glucose oxidase and another enzyme that they termed Chloroperoxidase (CPO) to denote the activity of the enzyme [2].

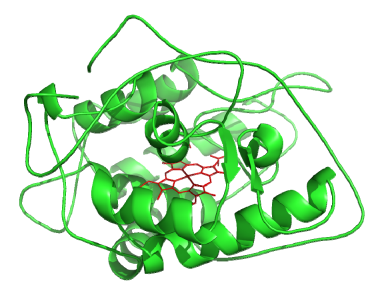


Figure 1: Chloroperoxidase (CPO)

Chloroperoxidase is a heme containing metalloprotein enzyme. It shares this feature with several other globular enzymes like myoglobin, haemoglobin, cytochrome P-450 and horseradish peroxidase. One of the main characteristics distinguishing CPO from most heme enzymes is that the proximal ligand bound to the heme is a cysteine residue [3]. It shares this feature with cytochrome P-450, but a main distinguishing feature between the two is that the distal site of the porphyrin group is charged or polar in CPO and hydrophobic in cytochrome P-450.

Since being discovered more than 50 years ago, CPO has been studied extensively because of its versatility, ecological significance and potential applicability in the pharmaceutical industry [4].

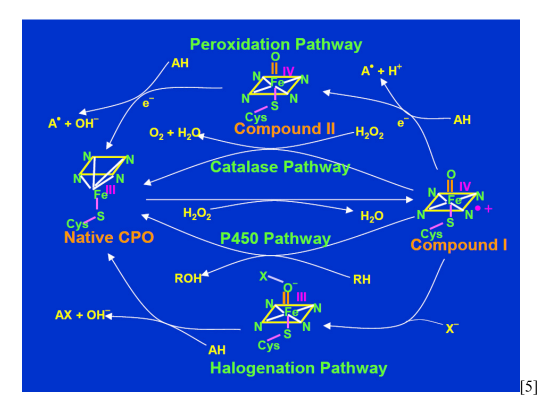


Figure 2: Reaction pathways for different reactions CPO catalyzes

Chloroperoxidase is known to catalyze several different types of oxidation reactions, summarized in Figure 2. All of these reactions begin with activation of CPO by an oxygen donor. The most commonly used reagent for activating the enzyme is hydrogen peroxide. The unreactive form of the enzyme, called the resting state, has the heme iron in the +3 oxidation state (ferric). Peroxide activates the resting state to a highly reactive state with the iron formally in the +5 oxidation state (perferryl). The active form of the enzyme is called compound I. In fact, it has been shown that a better description of compound I has the iron in the +4 oxidation state (ferryl) and a delocalized radical cation on the porphyrin ring.

1.1.1. Catalytic Reactions of CPO

Perhaps the most important of CPO's catalytic reactions is the oxygen insertion reaction, one form of which is epoxidation. Chloroperoxidase has been shown to catalyze the epoxidation of a number of organic substrates. These epoxidation reactions are highly enantiospecific, which makes CPO a pharmaceutically significant enzyme with substantial biotechnological potential. It is this potential combined with the ease of manufacture and availability of CPO that has driven the study of this reaction over the past several years. Much has been learned about the reaction mechanism of this reaction. Modification of the histidine residue at the distal site inhibits the epoxidation reaction, suggesting that the reaction occurs near the distal site of the heme. Other catalytic reactions compete with the epoxidation reaction. The most important competitor is the catalase reaction, which involves the dismutation of hydrogen peroxide to give water and an oxygen atom. The catalase reaction is much faster than the epoxidation reaction and depletes the hydrogen peroxide. The active site near the heme is only accessible via two narrow channels. One possible reason that the catalase reaction can out-compete the epoxidation reaction is that the substrates for the epoxidation reaction have to travel through the narrow channel to reach the active site. One line of research is aimed at controlling or inhibiting the catalase activity. It was discovered that there are different optimal pH ranges for different substrates and that pH 5.5 - 6.0 diminishes the catalase activity while not affecting the epoxidation reaction greatly. There is, however, a risk of deactivating CPO at the high end of this pH range. Using an alkyl peroxidase instead of hydrogen peroxidase as the oxygen donor slows down the catalatic activity, but the alkyl peroxide tends to lower the stability of Chloroperoxidase. Other mutation studies are being performed to discover motifs that would reduce catalase activity while maintaining or improving the epoxidation turnover rates.

Chloroperoxidase also catalyzes the dehydrogenation of substrates that is characteristic of heme peroxidase enzymes [6, 7]. The mechanism for this reaction is not completely understood. It is thought that for most, though not necessarily all, substrates the peroxidase reaction occurs at the surface of the enzyme and not near the heme active site. Like the epoxidation reaction, the peroxidase reaction also competes with the catalase reaction.

Another reason CPO is extensively studied is that it shares structural and functional features with several important enzymes, most importantly cytochrome P-450 and the thyroid enzyme involved in the iodonization process. Chloroperoxidase is more

4

easily produced and activated than these enzymes. Structure and reactivity can be studied using CPO, and the findings can be extrapolated to these other enzymes to better understand how they function.

The most prominent of CPO's catalytic reactions is the chlorination reaction, which has ecological implications [8-14]. In addition to the chlorination reaction, CPO also catalyzes other halogenation reactions including those of iodine and bromine, but not fluorine. The enzymatic chlorination reaction occurs at low pH. There was early concern [9] that the global prevalence of organohalogens was mainly due to anthropogenic contributions. However, several experiments have been able to show that there is significant contribution from biological activity to the global organohalogen burden. Given that CPO is easily accessible and found in nature, it was the enzyme of choice used by many research groups to demonstrate the possibility of enzymatic catalysis of chlorination of organic compounds found in plant litter, fluvic acid, lignin structures and humic substances found in soil samples. There are also examples of plant matter still attached to trees that contains organohalogens originating from degradation, presumably from microbial or fungal attacks [13]. The source of the halogen and oxidizer could be the plant itself, which tends to produce hydrogen peroxidase in cell walls [13]. There are several organisms like fungi, algae and bacteria that produce similar enzymes that would explain the presence of organohalogens in soils, surface water and volatile halogenated anisoles found on surface of oceans [14].

1.1.2. Previous Work on Mechanism of CPO-catalyzed Halogenation Reaction

Given the versatility and potential economic significance of CPO, there is little wonder that it has been studied extensively over the years. However, its most prominent reaction, chlorination, is still not completely understood. One postulated mechanism involves catalytic formation of HOCl at the active site of CPO followed by release into solution, where it halogenates substrates [4, 15-21]. The other postulated mechanism involves the chlorination of substrates at the heme active site [22-30]. Studies have been performed to support either of these two postulates, but it has been difficult for a consensus to be reached.

The non-active site hypothesis is based on the observation that most of the chlorinated substrate products show no enantiomeric excesses; that some substrates are too large to access the active site; and that HOCl and Cl₂, which can chlorinate any of the substrates, are observed in solution in chlorination reactions involving CPO.

The active site hypothesis is based on the following experiments. (1) There is a preference for certain substrates over others when the reaction is carried out using a mixture of substrates. For example, the preference for thiourea and methionine over monochlorodimedone (MCD) is 50:1 and 30:1, respectively, for the enzymatic reaction. By comparison, the preference for thiourea over MCD is 2:1 when HOCl is directly added to the reaction mixture in the absence of CPO, and 3:1 when Cl₂ is directly added. There is no preference for methionine over MCD using either HOCl or Cl₂ directly [24]. (2) The rates of chlorination of some substrates are higher for the reactions in the presence of CPO than for reactions involving chemical controls [23]. (3) Woggon et al.

developed synthetic active-site analogs of CPO for which the formation of stable intermediates with OCI⁻ and HOCl bound to the heme was demonstrated. These were presumed to be the intermediates through which the enzyme chlorinates the substrates [27-31]. (4) Some steady state kinetic studies suggest that chlorination via the free HOCl mechanism is unlikely. The studies show that the production of HOCl and Cl₂ at the enzyme active site and the subsequent release into solution is a slow process that may be considered as competing with the chlorinating reaction [22, 23]. (5) Some substrates give different product distributions in the enzyme catalyzed reaction and the chemical control reaction. For example, while no appreciable stereoselectivity is observed for most chlorinated substrates, for 2methyl-4-propyl-cyclopentane-1,3-dione it was reported that the enzyme catalyzed reaction results in a product, 2-chloro-2-methyl-4-propyl-cyclopentane-1,3-dione, with a 40:60 ratio of the racemic diastereomers, while the chemical control gives a 55:45 ratio of the racemic diastereomers [26]. (6) There is also a reported case of stereoselective halohydration of glycals and steroids using CPO [32-35].

Thus the experimental evidence regarding the mechanism of enzymatic halogenation by CPO is inconclusive. Some consider the general lack of enantiospecificity to be the most persuasive evidence and conclude that halogenation occurs outside the active site. However, the many observations favoring active-site halogenation leave the case far from clear. The bulk of the work presented in this dissertation is aimed at contributing to the resolution of this dispute.

1.1.3. Substrate Used to Study Reaction

In the research presented here, I employ a variety of theoretical approaches to study the chlorination mechanism of CPO, and I choose a particular model substrate, 1,3 cyclopentanedione (CPD), for consideration. This substrate was chosen for several reasons. First, CPD is a cyclic β -diketone, and it is known that cyclic β -diketones, β -keto acids, and substituted phenols make good halogen acceptors. Second, 2,2-dichloro-1,3cyclopentanediol was found to be the main metabolite of CPO [1]. CPD has been shown to be catalyzed to 2,2-dichloro-1,3-cyclopentanedione [36]. CPD Third, CPO and another cyclic β -diketone, monochlorodimedone (MCD), are the compounds most widely used for monitoring the halogenation activity of CPO spectroscopically. The intermediate species in the reaction of CPD to the dichloro metabolite is a monochloro species, 2chloro-1,3-cyclopentanedione [36]. It is easy to observe the conversion of CPD to the metabolite because CPD and the intermediate have absorption maxima at 242 and 248 nm, respectively, with a high extinction coefficient while the metabolite has an absorption maximum at 277 nm with a very low extinction coefficient [37]. Monochlorodimedone is more commonly used because it is readily available, and it is easy to observe the conversion from MCD to dichlorodimedone [37]. For computational studies, though CPD is preferable to MCD because it contains fewer heavy atoms and is hence less demanding of computational resources. Finally, a crystal structure of CPO with CPD at its active site has been reported [38]. This structure will be useful for future studies of halogenation at the active site.

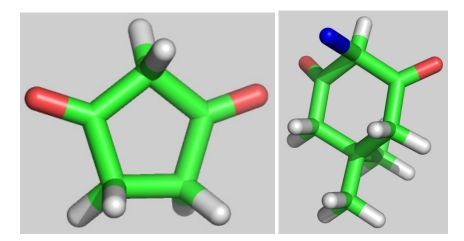


Figure 3: Substrates - CPD and MCD

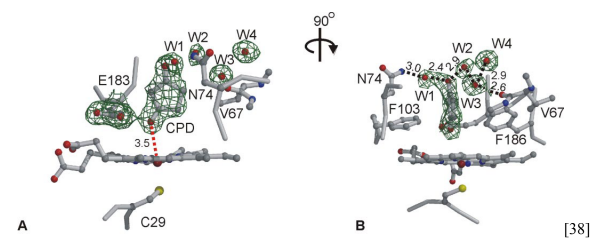


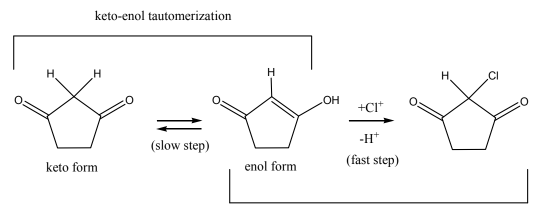
Figure 4: CPO active-site showing CPD and water molecules

1.2. Chlorination Mechanism of CPO

It is well known that ketones in aqueous solution can undergo an internal rearrangement in which a hydrogen is transferred from the α -carbon to the keto oxygen and the double bond shifts from C=O to C=C. The resulting species is called an enol because it contains both –ene (carbon-carbon double bond) and alcohol (OH) groups.

The two forms are called tautomers, and the process of interconversion is tautomerization. In aqueous solution, both tautomers are present in equilibrium. An alternative nomenclature for the tautomers of β -diketones is diketo and keto-enol.

The overall process of the chlorination reaction, with keto-enol tautomerization as the first step, is depicted in Figure 5. The mechanisms of the individual steps are shown in Figure 6 and Figure 7. The chlorination (or more generally, the halogenation) of ketones begins with tautomerization to the enol form. Electrophilic attack of a chloronium (Cl⁺) on the carbon-carbon double bond of the enol form, followed by transfer of the hydroxy proton to a solvent water, yields the chlorinated product. The tautomerization step can be either acid- or base-catalyzed, as shown in Figure 6. I consider acid catalysis because CPO-catalyzed chlorination is carried out at low pH (pH 3-4).



chloronium addition

Figure 5: Chlorination of CPD

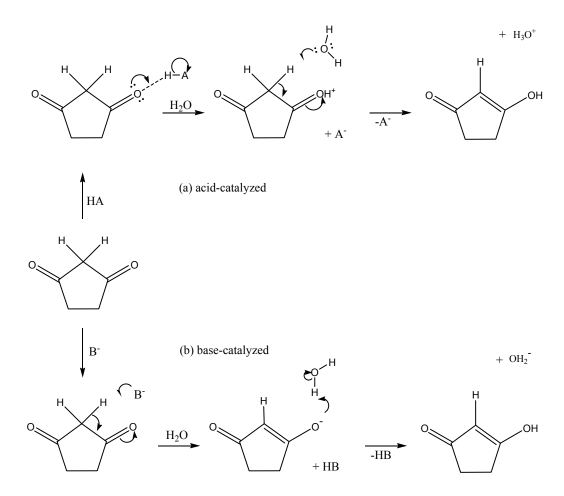


Figure 6: Mechanism of keto-enol tautomerization: (a) acid catalyzed, (b) base catalyzed

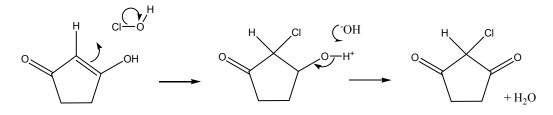


Figure 7: Mechanism of chlorination step

For the overall process of the chlorination reaction shown in Figure 5, the second step is expected to be fast and barrierless. The mechanism of this step is depicted in Figure 7. I modeled the chlorination step in a limited hydration environment, with 14

water molecules, using the quantum mechanics approach explained in section 2.4. The data in Figure 9 verifies that the reaction shown in Figure 8 is nearly barrierless and hence a kinetically fast reaction. Therefore, I focus the study on the rate-limiting step which is the acid catalyzed keto-enol tautomerization.

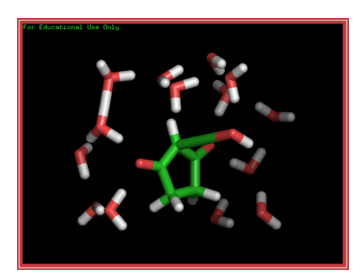


Figure 8: Second step of the chlorination mechanism in a limited hydrated environment

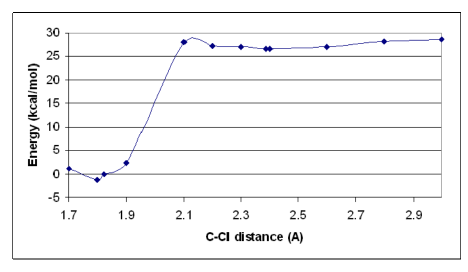


Figure 9: Second step of the chlorination mechanism depicting barrierless reaction

1.3. Previous theoretical studies of keto-enol tautomerization

Solvation plays a critical role in many reactions including keto-enol tautomerization and will be the focus of my studies on the chlorination mechanism of CPD. Considerable prior theoretical work has been done on the role of water in keto-enol tautomerization. Here this prior work is summarized.

The equilibrium between the keto and enol tautomers in solution has been studied with implicit solvent methods, with reasonable success, reproducing logK, where K is the equilibrium constant, to within $\sim \pm 0.2$ in many cases [39-41]. However, such methods are not sufficient for modeling the kinetics of tautomerization in aqueous solution because the mechanism explicitly involves water molecules.

Calculations determine the barrier for keto-enol tautomerization via intramolecular proton transfer to be generally in the range 60-85 kcal/mol, depending on the compound. This is much larger than the experimental reaction barriers [42-44]. Inclusion of a single explicit catalytic water lowers the barrier by 30-40 kcal/mol for typical molecules. The mechanism is a two-step process with the water first abstracting a proton from the alpha carbon, and subsequently donating a proton to the keto oxygen. Ma et al. determined that a second catalytic water lowers the tautomerization barrier for 3-methyl-5-hydroxyisoxazole a further 8 kcal/mol [43]. Freitag et al. showed that for malonaldehyde, a third explicit water with a solvating but not catalytic role can lower the barrier another 7 kcal/mol [44]. Further solvating waters had decreasing incremental effects. These studies demonstrate that inclusion of explicit water molecules is crucial

for modeling the kinetics of keto-enol tautomerization, but the degree of explicit solvation required for an accurate model is not known.

Cucinotta et al. applied an ab initio metadynamics treatment to keto-enol tautomerization of acetone in aqueous solution [45]. They found that a single water lowers the free energy barrier from 57.7 to 46.1 kcal/mol, and a solution environment of 28 waters lowers the barrier further to 38.5 kcal/mol. In solution, a Grotthuss mechanism involving four water molecules in a proton transfer chain was observed.

With the exception of the metadynamics work, all the studies focused on optimized structures with up to two catalytic waters, and in one case with up to three further waters that were solvating but not catalytic. It is unknown whether this degree of hydration is sufficient. More fundamentally, the assumption is made that barriers for fully optimized structures involving multiple waters can accurately predict experimental However, such structures might represent reaction events so activation energies. extremely rare that they do not predict the observed kinetics well. I address these issues with two studies on CPD. In the first study, I systematically increase the number of explicit waters in fully optimized structures and determine the barrier and the reaction energy as functions of the number of waters. I find that the barrier to reaction continues to decrease as the number of waters increases beyond five. In the second study, I use classical molecular dynamics to generate solution environments representative of 300 K. I then determine reaction barriers from snapshot structures using a hybrid QM/MM method, allowing one or two water molecules to be explicitly active. I find that the instantaneous environment has a significant effect on the barriers. This result, ultimately,

is consistent with Cucinotta's metadynamics work [45] but provides complementary detail on the nature of the influence of water on this keto-enol tautomeric reaction.

The energies of the stationary points (reactant, product and transition state) were calculated in different environments as described above. I start with the simplest system consisting of only the CPD and HOCl molecules, with or without a single catalytic water, in the gas phase. I then focus most of the study on systems consisting of CPD, HOCl and one reactive/catalytic water molecule in different environments.

1.4. Solvation

Based on previous theoretical work on keto-enol tautomerization it is clear that solvation effects are quite substantial and critical, as they influence the reaction barrier strongly. Both at the active site and in aqueous solution, water molecules are present and will interact with the substrate during reaction. Water itself, in fact, plays a catalytic role in the reaction. Furthermore, water molecules not directly involved in the breaking and forming of chemical bonds may also influence the reaction strongly through hydrogen bonding, in addition to less directional dielectric effects that water exerts. Thus we will find that the way in which water is incorporated into the reaction model is critical. The most basic way to represent solvent in theoretical calculations, as a dielectric continuum, turns out to be entirely inadequate in this case.

In reaction at the active site, a limited number of waters are available to interact with the substrate. This is apparent from the crystal structure shown in Figure 4, which identifies four waters quite near a CPD molecule bound at the active site. These four waters are resolved in the X-ray structures because they have strong stabilizing interactions with neighboring moieties. In reality, there may be other waters in addition to these four, but the total number is limited by the size and shape of the active site pocket. Thus in the context of the active site, it is misleading to speak of solvation because the number of waters is limited. In this work, I will refer to such a water environment as "limited hydration." In contrast I will refer to the water environment of CPD dissolved directly in aqueous solution as "full solvation."

The central work in this dissertation reports on two investigations, one of the chlorination reactions in "limited hydration" conditions and the other chlorination reaction in "full solvation" conditions. Quite different approaches are used for the two investigations. The limited hydration work connects with previous theoretical work reported on keto-enol tautomerization, which is a critical step in the reaction (see below). Keto-enol tautomerization is a fundamental concept in organic chemistry, but the influence of the water environment on the reaction is still not fully understood. My work on limited hydration suggests that previous work on keto-enol tautomerization needs to be understood in a more restricted way than that work made clear. The energetics worked out in the previous work is representative of the particular limited hydration structures used, but not representative of limited hydration in general and certainly not representative of full solvation.

My work on the reaction in "full solvation" at near-physiological temperature leads to a paradoxical observation. The dominant reaction events have energetic barriers very similar to those encountered under extremely limited hydration conditions, namely a single water that assumes a catalytic role. This similarity in barriers turns out to be a consequence of equivalent degrees of reactant- and transition-state stabilization by other waters in the full-solvation calculations. The full-solvation barriers are quite different from the barriers for limited hydration with several waters present.

The work on the chlorination reaction presented here is restricted to the various water environments just described. This turned out to be rich and difficult undertaking in itself. Reaction of CPD situated within the enzyme CPO at the active site is a subject for future work. The work presented here demonstrates that inclusion of hydrating waters will be critical for modeling the reaction at the active site. Implications of this work extend beyond CPO, as many chemical reactions are likely to be sensitive to the nature of the limited-hydration environment found at an active site.

The rest of this dissertation is organized as follows. Chapter 2 contains an outline of theoretical methods used (content on basic quantum mechanics and molecular mechanics is from standard computational chemistry texts [46, 47]). Part of the work presented is based on molecular mechanics models. This necessitated the development of such models for CPD and other molecules used. This is an involved process of parameterization, and it is detailed in Chapter 3. Chapter 4 presents the research on the chlorination reaction in limited hydration conditions representative of the active site environment. Chapter 5 presents the research on the chlorination reaction in fully solvated conditions representative of aqueous solution. Chapter 6 presents the major conclusions regarding the halogenation mechanism. Chapter 7 briefly describes contributions to three other research projects. The appendix contains the most important scripts used for running the calculations.

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2. METHODS

2.1. Molecular Mechanics

The term Molecular Mechanics (MM), which was first introduced in 1970, describes the application of classical mechanics in the determination of molecular structures at equilibrium. In the simplest MM treatment of molecules, the atoms are treated as balls and the bonds as springs connecting them. These springs follow Hooke's law, which is a representation of a harmonic oscillator. The application of classical mechanics to the motion of these balls (or atoms) bound by springs (or bonds) is the basis of molecular mechanics. Thinking of molecules as being made up of balls and springs dates back to as early as 1930 and was first suggested by Andrews [48, 49].

It might seem as if the laws of quantum mechanics are ignored with this treatment of molecules. In reality, though, the force fields are based on experimental and quantum mechanical calculations, so quantum mechanics is incorporated in the force field. The MM treatment of molecules works quite well as long as the structures do not deviate too much from their respective equilibrium structures. The harmonic oscillator approximation of the bonds often provides sufficiently accurate results. However, bond breaking or forming cannot be modeled using molecular mechanics.

Quantum Mechanics (QM) cannot be used to model large molecules such as proteins, which number in the thousands of atoms due to computational costs. The MM models use a simplistic representation of atoms in comparison to quantum mechanics to allow one to study systems with large numbers of atoms with reasonable accuracy. Molecular mechanics is used extensively in the study of biomolecular systems like docking for drug design, protein folding, binding constants and reaction precursors.

One of the main objectives of a good force field is transferability from molecule to molecule. For example, we would like to be able to apply the force field derived from one molecule to another similar molecule to correctly predict the minimum energy structure for the new molecule. This key development in MM, where it was possible to transfer force constants between alkanes was first made by Snyder and Schachtschneider [50].

The functional forms used for developing force fields are described below. The various types of interactions in a macromolecule can be classified as bonded interactions and non-bonded interactions. The potential energy contributions from the various interactions are combined to give the total potential energy of the macromolecule. Each contribution is described separately below.

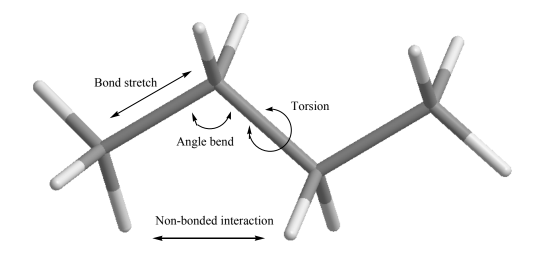


Figure 10: Bonded and non-bonded interactions in MM

The program I use for the MM model is CHARMM [51]. CHARMM is a molecular simulation program developed to study large, many-atom structures usually encountered in biological systems, for example proteins, peptides, nucleic acids, carbohydrates and organic drug-like molecules or ligands in solution, crystal or membrane environments. CHARMM provides computational tools to perform molecular energy minimization, molecular dynamics, analysis and path sampling. It can be interfaced with several QM software packages to study systems that require the hybrid QM/MM method explained in section 2.5.

2.1.1. Bonded Interactions

Bond Stretching

To determine the potential energy contribution from bond stretching we consider the ball and spring model. The force due to the spring is given by Hooke's Law

$$F_s = -k_s (R - R_e)$$

The potential energy is given by

$$U(r_{AB}) = -\int F(r_{AB})dr$$

Applying a Taylor expansion we obtain

$$U(r) = U(r_{eq}) + \frac{dU}{dr}\Big|_{r=r_{eq}}(r-r_{eq}) + \frac{1}{2!}\frac{d^2U}{dr^2}\Big|_{r=r_{eq}}(r-r_{eq})^2 + \frac{1}{3!}\frac{d^3U}{dr^3}\Big|_{r=r_{eq}}(r-r_{eq})^3 + \dots$$

We label the equilibrium bond distance r_{eq} . We can, for convenience, consider the minimum of the functional form to be zero. Therefore, the first and second terms in the Taylor's expansion are zero. The simplest contribution then is from the first non-zero term which is the quadratic term. This approximation may be sufficient for small stretching, but as the bond stretches far from the equilibrium values the potential energy contribution becomes infinitely positive, which is unrealistic, and the harmonic approximation fails. This failure is inherent in truncating the Taylor's expansion, so the logical thing would be to include more terms. When the next cubic term is included, it causes an unintended divergence to negative infinity as the bond is stretched further. Therefore, the quartic term also has to be included.

Alternately, the functional form of the Morse potential can be used which describes the bond stretch rather accurately.

$$U(r_{AB}) = D_{AB} [1 - e^{-\alpha_{AB}(r_{AB} - r_{AB,eq})}]^2$$

However, the exponential term in the Morse potential is computationally less efficient for calculation than the truncated polynomial terms from the harmonic oscillator. In most force fields some variation of the following bond stretching term is used

$$U(r_{AB}) = \frac{1}{2} [k_{AB} + k_{AB}^{(3)}(r_{AB} - r_{AB,eq}) + k_{AB}^{(4)}(r_{AB} - r_{AB,eq})^2 + \dots](r_{AB} - r_{AB,eq})^2$$

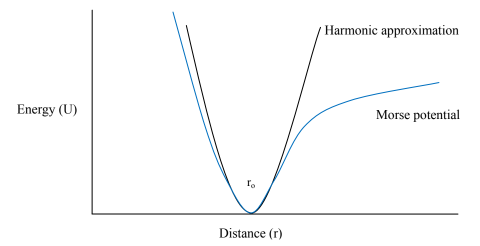


Figure 11: Bond stretch energy functions

Angle Bending

As with bond stretching, angle bending or bond angle deformation can be modeled using the harmonic approximation about the equilibrium bond angle. It has an important contribution to the total potential energy as evidenced by vibrational spectroscopy data that show energy variation attributed to bond bending. The same considerations as for bond stretching apply to the functional form, so a generally useful function form is

$$U(\theta_{ABC}) = \frac{1}{2} [k_{ABC} + k_{ABC}^{(3)}(\theta_{ABC} - \theta_{ABC,eq}) + k_{ABC}^{(4)}(\theta_{ABC} - \theta_{ABC,eq})^2 + \dots](\theta_{ABC} - \theta_{ABC,eq})^2$$

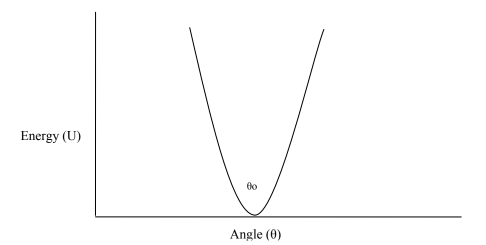


Figure 12: Angle bending energy function

Torsions

Torsional motion is periodic, so its contribution to the potential energy is also periodic. It is conveniently modeled using a cosine function, or more generally a sum of cosines to allow for irregularity in the periodic function. Energy contribution due to dihedral (torsional) motion involving four atoms is given by the following expression.

$$U(\boldsymbol{\omega}_{ABCD}) = \frac{1}{2} \sum_{\{j\}_{ABCD}} V_{j,ABCD} [+(-1)^{j+1} \cos(j\boldsymbol{\omega}_{ABCD} + \boldsymbol{\psi}_{j,ABCD})]$$

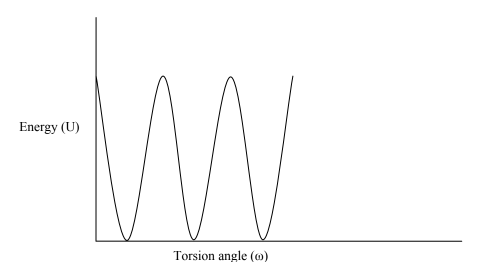


Figure 13: Torsion energy function

Improper torsions

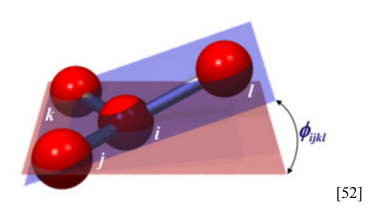


Figure 14: Improper torsion angle

Improper torsions are a special situation that involves four atoms, but unlike the case with regular torsions, for improper torsions there is a central atom bonded to the three other atoms. If we define a plane by the central atom and two other atoms that are bonded to it, then the angle the fourth atom creates to that plane is the improper torsion angle. Improper torsion terms are included only when the other energy terms are

insufficient to maintain correct geometries. Specifically, improper torsion terms are used to maintain puckering in structures. Improper torsional energy contributions are only accounted for by CHARMM if there are explicit parameters for them for a specific molecule. Improper torsion terms are not included in the energy by default like the other bonded interaction terms. The CHARMM topology file must specify that a particular improper torsion term be included.

2.1.2. Non-bonded Interactions

As stated earlier, one of the main objectives in developing a force field is the transferability of the force field to similar groups of molecules. During the initial development of force fields it was observed that 'transferability' was attainable by including non-bonded interactions, at least of the nearest neighbors. This is why non-bonded interaction contributions are critical and have to be included. The following two types of non-bonded interactions are included in most force fields, while some force fields include additional interactions.

Van der Waals Interactions

Van der Waals force is the attractive forces experienced by non-bonded neutral atoms. As neutral atoms gets in close vicinity to each other they experience attractive forces due to the induced dipoles. These short distance intermolecular forces are weaker than a chemical bond, but are important when atoms get close enough like in liquids and solids and are responsible for properties like surface tension and capillary action.

In the simplest case we have two atoms, represented as hard spheres, each with a characteristic radius. We also assume they have no charge. At infinite separation, there is no interaction and hence the contribution to the potential energy is zero. As they approach each other the two spheres eventually touch. At this point they cannot approach each other and the energy contribution tends to infinity. Realistically, we know from QM that this is a simplistic case and in fact as the atoms approach each other there is an attractive force due to the electronic correlation, also called dispersion. As they come closer, at some very short distance, the energy contribution tends to rise exponentially to infinity. The attractive energy is inversely proportional to r^6 . This dispersion interaction between atoms is usually modeled by the Lennard-Jones potential and is given below.

$$U(r_{AB}) = 4\varepsilon_{AB} \left[\left(\frac{\sigma_{AB}}{r_{AB}} \right)^{12} - \left(\frac{\sigma_{AB}}{r_{AB}} \right)^{6} \right]$$

It would be more accurate to represent the repulsion with an exponential term instead of the inverse r^{12} term. But, exponential calculations are more difficult on the computer in comparison to a polynomial term, which is why the inverse r^{12} term is used instead. This is an empirical result and used for computational convenience.

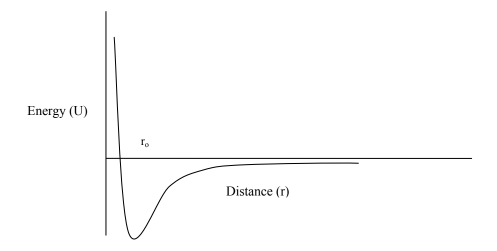


Figure 15: Lennard-Jones potential for van der Waals interaction

Electrostatic Interactions

Electrostatic interactions are the energy contribution that arises from non-bonded interactions between atoms that have a charge associated with them. Again, if we consider the simplest case, we assume a fixed point charge associated with an atom. We then apply Coulomb's Law to determine the energy contribution which is given by the following equation.

$$U_{AB} = \frac{q_A q_B}{\varepsilon_{AB} r_{AB}}$$

We know that as atoms move the effective atomic charges will respond to the environment, that is to say, atomic systems are polarizable. When permanent charges are assigned, the polarizability is neglected. However, the fixed permanent charges used are generally capable of reproducing several experimental observable or QM computed properties quite accurately. Development of polarizable force fields is an active topic of research and there are several groups working on developing polarizable force fields. However, the community has not yet embraced any particular polarizable force field.

The calculations of non-bonded interactions are the most time consuming part of the energy evaluations since it would mean calculation for every pair of atoms in the system ($N^2/2$ interactions, where N is the number of atoms). These interactions are distant dependent and two atoms that are quite far from each other may have minimal contribution to the total energy. An implementation consideration then is how far we want to calculate non-bonded interactions. In interest of saving computation time a truncation scheme is used. A cutoff distance is set so that non-bonded interactions are only calculated if an atom is within a certain distance from another atom. However, this introduces a discontinuity in the energy evaluations. Consider for example, two atoms that are exactly at the cutoff distance when the energy contribution is calculated. At the next step, they move just beyond the cutoff distance from each other, thus the new energy contribution calculated is precisely zero. The small step would cause a drastic difference in the energy evaluation. To avoid this abrupt discontinuity, a switching function is used to gradually exclude the non-bonded interactions instead of abruptly. For example, if the cutoff was set at 12 Å, one can use a switching function to gradually exclude the nonbonded interactions from 12 Å to 14 Å instead of abruptly setting it to zero at 12 Å. The preferred method of treating electrostatic interactions is the Particle Mesh Ewald summation method. It is a method to calculate the electrostatic interactions in an infinite lattice or under periodic boundary conditions (explained in section 2.3.3).

2.1.3. Solvation

Modeling gas phase reactions are relatively simple. However, all biochemical processes occur in the condensed phase and the chemical reaction we study occurs in solution. Therefore, it is critical to model a solvation environment around the molecule of interest. A solvation environment model can be broadly classified as an implicit model or explicit model.

In an implicit solvent model, also called the continuum solvation model, the environment is like a continuous medium with a dielectric constant representing the solvent and properties consistent with those of the solvent. To represent water as the extended solvent environment we use the dielectric constant of water ($\varepsilon = 78.4$). This kind of representation of the solvent has the advantage that the computation overhead is low and used with QM model. I used the Polarizable Continuum Model (PCM) method as implemented in Gaussian98 [53]. The PCM model is a widely used implicit solvent model that represents the environment of the system by an dielectric continuum having the dielectric constant of the solvent. However, an implicit model is only used when the condensed phase does not play a critical role in the reaction environment.

An explicit solvent model includes explicit solvent molecules. Adding explicit water molecules can get prohibitively expensive in the quantum mechanics model. At best it is feasible to include a single solvation shell. However including explicit water molecules is quite feasible in the molecular mechanics model which is routinely done while studying large biomolecular systems. I use both types of explicit solvated environments.

2.2. Geometry optimization

One of the key motivations of MM is to find minimum energy structures or equilibrium structures that are representative of realistic molecules. Locating minimum energy structures constitutes a mathematical problem and the general field of study is called optimization theory. One of the priorities is to facilitate geometry optimization by simplifying the functional forms without compromising accuracy. One such example is using polynomial terms instead of exponential terms to facilitate computation. There are several algorithms that have been developed that are used in achieving this goal. Obviously, some are more efficient that others and in most cases it is best to use a combination of different algorithms.

Consider a molecule with N atoms. This molecule has 3N degrees of freedom, of which six are translational and rotational (or five if it is a linear molecule). The remaining 3N - 6 (or 3N - 5) degrees of freedom are vibrational degrees of freedom and therefore, there are 3 N - 6 independent variables that describe them which are also called internal coordinates.

All these variables can be written as a matrix

$$q = \begin{pmatrix} q_1 \\ q_1 \\ q_3 \\ \dots \\ q_p \end{pmatrix}$$

The simplest way to optimize would be to optimize the energy for one variable at a time. This is called a line search, but when we have several variables and if they are correlated this approach becomes inefficient.

Another approach would be to proceed in a direction that minimizes the total energy with respect to all the variables at the same time. The first derivative matrix, also called the gradient and shown below, is needed. The optimization process proceeds by altering the coordinates so as to modify the structure in the negative direction of g(q) until a local minimum is reached. This is a first order approach.

$$g(q) = \begin{pmatrix} \frac{\partial U}{\partial q_1} \\ \frac{\partial U}{\partial q_2} \\ \dots \\ \frac{\partial U}{\partial q_p} \end{pmatrix}$$

Steepest Decent methods, which was introduced by Wiberg in 1965 [54], and Conjugate Gradient methods, which were introduced by Fletcher and Reeves [55], are examples of first derivative methods. Such methods get us close to the minimum quickly, but an inherent drawback of such an approach is the convergence. It can sometimes be hard for such algorithms to converge to the minimum. One usually starts an optimization process with such an algorithm to get close to the minimum. Then one uses a more sophisticated method, usually a second order method, to finally converge to the minimum.

Second order methods make use of the second derivative of the energy function, also called the Hessian.

$$H(q) = \begin{pmatrix} \frac{\partial^2 U}{\partial q_1} & \frac{\partial^2 U}{\partial q_1 \partial q_2} & \dots & \frac{\partial^2 U}{\partial q_1 \partial q_p} \\ \frac{\partial^2 U}{\partial q_2 \partial q_1} & \frac{\partial^2 U}{\partial^2 q_2} & \dots & \frac{\partial^2 U}{\partial q_2 \partial q_p} \\ \dots & \dots & \dots \\ \frac{\partial^2 U}{\partial q_p \partial q_1} & \frac{\partial^2 U}{\partial q_p \partial q_2} & \dots & \frac{\partial^2 U}{\partial^2 q_p} \end{pmatrix}$$

If we substitute in the Taylor's expansion series of energy we obtain

$$U(q) - U(q_{eq}) = \sum_{i=1}^{p} \xi_{i} (\frac{\partial U}{\partial q_{i}})_{\xi_{i}=0} + \frac{1}{2} \sum_{i=1}^{p} \sum_{j=1}^{p} \xi_{i} \xi_{j} (\frac{\partial^{2} U}{\partial q_{i} \partial q_{j}})_{\xi_{i}=0}, \xi_{j}=0} + \dots$$

$$U(q) - U(q_{eq}) = \xi^T g + \frac{1}{2} \xi^T H \xi + \dots$$

Second order methods tend to converge to the minimum quickly if they are close to the minimum to begin with. However, they are computationally more expensive because of the necessity of calculating the Hessian, especially if needed at every step. In practice, it is necessary to calculate the Hessian for the first step when using a second order method, but some algorithms use an approximate Hessian for subsequent steps to make the

calculation computational efficient. However, in some sensitive cases it is essential to calculate the Hessian for every step, which is the basis for the Newton-Raphson method.

The minimization algorithms used in this project were a combination of first order and second order methods. Minimization was initiated using Steepest Descent followed by the Adopted Basis Newton Raphson (ABNR) method. ABNR is a quasi-Newton-Raphson method.

2.3. Molecular Dynamics

The most popular application of Molecular Mechanics is optimization of a structure to find its minimum energy or an equilibrium structure using the laws of classical mechanics. While this is quite useful, it is even more interesting to predict molecular properties computationally that may or may not be feasible to measure experimentally. One technique that is used to do this computationally is called Molecular Dynamics (MD).

2.3.1. Phase Space

In order to describe MD, I will first discuss the concept of phase, and then show that MD is a way to sample phase space in order to compute average properties. The state of a macromolecule can be completely described using classical laws by specifying the position and momenta of all the particles that make up the macromolecule. A system made up of N particles would need 6N coordinates to describe the macromolecule, 3N coordinate to describe position along the x, y and z axes and 3N coordinates to describe their corresponding momenta along the respective axes. The resulting 6N dimensional space is called the 'phase space' of the macromolecule. At any given instant in time, the macromolecule occupies one point in phase space. Over time, a system maps out a trajectory in phase space. Molecular dynamics is a technique used to construct the phase-space trajectory of a macromolecule, given a set of initial conditions.

Phase space point

X = (q, p)

 $q = (x_1, y_1, z_1, x_2, y_2, z_2, \dots)$

$$p = (p_{x,1}, p_{y,1}, p_{z,1}, p_{x,2}, p_{y,2}, p_{z,2}, \dots)$$

Properties measured are averages over the states of the system. Phase-space is extremely large and encompasses every possible state of the system, even those that the system is unlikely to sample, like unrealistic high energy states. In other words, phase space includes states that have a very low probability of being visited in reality. Thus phase-space is essentially a large wasteland that mostly contains states that are unlikely to contribute much to the expectation or average value of a particular property. Hence, it makes sense to use available computational resources efficiently by only mapping out trajectories that concentrate on regions of phase space the system would be most likely to sample. These regions would likely be near local minima or equilibrium structures. For this reason, most MD simulations are preceded by short energy minimization simulations. (Another reason MD simulations are preceded by a short energy minimization is that most structure coordinates come from x-ray crystallography. The crystal structures usually contain bad contacts between atoms with artificially high forces. If we start the MD simulation with these high forces, the trajectory may be pushed away from the local minima into high-energy, improbable regions of phase space.)

Experimental measurements of properties are averages over states of a system, as mentioned above. Often the system of interest is microscopic or submicroscopic, yet the experimental conditions include a macroscopic sample containing many "copies" of the system. For example in a solution containing a mole of enzyme molecules in which case Avogadro's number of enzyme molecules is present, each capable of sampling a different state of the enzyme. In this case, an experimental measurement of some property of the enzyme will represent an ensemble average over many copies. (Experimental measurements will also be time averaged if the measurement takes longer than the characteristic time associated with the property in question, but the important point here is the ensemble-average nature of most experiments). An MD trajectory provides a sampling over time of a single copy (or small number of copies) of the system. The connection between simulation and experiment rests on the assumption that the time-average over the trajectory of a single copy of the system will give the same results as the ensemble average over many copies of the . This assumption is the ergodic theorem. It

allows us to make a connection between simulation and experiment, provided the simulation is long enough for sufficient sampling of phase space.

An interesting aspect of a phase-space point is that the location of the next point can easily be determined from the current position and momentum. The relationship between two positions and the momentum over some time interval is given by

$$q(t_2) = q(t_1) + \int_{t_1}^{t_2} \frac{p(t)}{m} dt$$

The relationship between momenta and acceleration is given by

$$p(t_2) = p(t_1) + m \int_{t_1}^{t_2} a(t) dt$$

The acceleration a(t) is given by

$$a = \frac{F}{m}$$

The acceleration can be calculated from the forces acting on each particle via the force field. Hence, we can map out the trajectory in phase-space in the forward and reverse directions once we have a single phase-space point.

For non-analytical systems the above expression can be re-written using Euler's approximation in the limit $\Delta t \rightarrow 0$. We get the same result if we use the Taylor's expansion truncated at second order.

$$q(t + \Delta t) = q(t) + \frac{p(t)}{m} \Delta t$$

$$p(t + \Delta t) = p(t) + ma(t)\Delta t$$

2.3.2. Verlet-leapfrog algorithm

On the basis of the above general discussion of phase space we now consider the algorithm used to map the trajectory of the system through phase space. One way to map out the trajectory is by using the Verlet algorithm. Consider time steps of Δt in the forward and reverse direction.

The expression for position in the forward direction

$$q(t + \Delta t) = q(t) + (\frac{dq}{dt})_t \Delta t + \frac{1}{2!} (\frac{d^2 q}{dt^2})_t (\Delta t)^2 + \dots$$

The expression for position in the reverse direction

$$q(t - \Delta t) = q(t) - (\frac{dq}{dt})_t \Delta t + \frac{1}{2!} (\frac{d^2 q}{dt^2})_t (\Delta t)^2 + \dots$$

If we add the two expressions we get

$$q(t + \Delta t) = 2q(t) - q(t - \Delta t) + \frac{1}{2}a_t(\Delta t)^2$$

$$a_t = \left(\frac{d^2 q}{dt^2}\right)_t$$

We can see that to get the next position, after a time interval of Δt , all we need are the current and previous position and the acceleration, which we can get from the force field.

$$a = \frac{F}{m}$$

$$F = -\frac{dV}{dq}$$

The Verlet algorithm allows us to propagate through phase-space with no explicit reference to velocity or momentum. However, in most cases it is useful to run simulations at a particular temperature or gradually to be able to change temperature. Temperature is directly related to random velocity, so one needs to be able to manipulate velocity directly in order to control the temperature of the simulation. The leapfrog algorithm, a modification to the Verlet algorithm, was developed to couple the propagation of position and velocity directly.

$$q(t + \frac{\Delta t}{2} + \frac{\Delta t}{2}) = q(t + \frac{\Delta t}{2}) + \frac{dq(t + \frac{\Delta t}{2})}{dt} \frac{\Delta t}{2} + \frac{1}{2!} \frac{d^2 q(t + \frac{\Delta t}{2})}{dt^2} (\frac{\Delta t}{2})^2 + \dots$$

$$q(t + \frac{\Delta t}{2} - \frac{\Delta t}{2}) = q(t + \frac{\Delta t}{2}) - \frac{dq(t + \frac{\Delta t}{2})}{dt} \frac{\Delta t}{2} + \frac{1}{2!} \frac{d^2 q(t + \frac{\Delta t}{2})}{dt^2} (\frac{\Delta t}{2})^2 + \dots$$

Subtracting we get

$$q(t + \Delta t) = q(t) + v(t + \frac{\Delta t}{2})\Delta t$$

$$v(t + \frac{\Delta t}{2}) = \frac{dq(t + \frac{\Delta t}{2})}{dt}$$

For v(t) we obtain

$$v(t + \frac{\Delta t}{2}) = v(t - \frac{\Delta t}{2}) + a(t)\Delta t$$

From the above two expressions we see that for half a time step ($\Delta t/2$) we evaluate velocity and then for the second half (Δt) we evaluate the new position. The new

position is calculated based on the current position at time t and velocity at time step $\Delta t/2$. The velocity at time step $\Delta t/2$ in turn is based on the previous velocity at time step $-\Delta t/2$ of the particle and the current acceleration of the particle at time t.

An important consideration that has to be made is the time interval Δt used for computation. If the time step is too large we will deviate from the true phase-space trajectory, since we take a step in the direction tangent to the true trajectory i.e. constant acceleration. However, if we take time steps that are too small, we will end up with very short trajectories through time that will sample too small a region of phase-space to be useful. Practically, a time step one or two orders of magnitude smaller than the highest frequency motion generally yields a sufficiently accurate trajectory. The highest frequency motion in atomic systems is the vibrational motion of bonds between a heavy atom and a hydrogen atom, which has a period of about 10⁻¹⁴ s. In practice, time steps of 0.5-1.0 fs are recommended for most purposes.

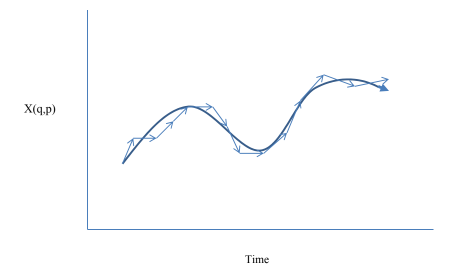


Figure 16: Trajectory in phase space

Another option to allow the use of larger time steps is to freeze the highest frequency motion. This can be done by constraining the heavy atom – hydrogen bond length. The keyword in CHARMM to constrain the heavy atom – hydrogen atom bond is called SHAKE. The next highest frequency motion would then be a heavy atom – heavy atom vibration with a frequency typically a factor of 2 - 5 smaller in magnitude. This approach can only be used if the property being studied is independent of those degrees of freedom which are removed. In addition to increasing the time step by a factor of two 2 (time step of 2 fs), reduction in the number of degree of freedom provides some saving in time and integration stability.

2.3.3. Periodic Boundary Conditions

A molecule being modeled is placed in a box of finite size. This model presents a technical difficulty. The size of the box cannot be very large because of the computational cost of including a large number of atoms is great, especially if the box is filled with water. If the box is too small then the surface effects of the box may affect the chemistry of the system. Beyond the edge of the box is essentially vacuum. Both these effects are unrealistic.

One way to handle this situation is to use Periodic Boundary Conditions (PBC). The system is placed in a unit cell, usually a cubic or truncated octahedral unit cell. The main unit cell is then surrounded by identical images. The long-distance non-bonded interaction energies are calculated between the unit cell and the images that are within the cutoff distance. Each atom in the main unit cell has identical images surrounding the unit cell that follows the trajectory of the main unit cell particle. During the course of the MD simulation, if for example, a water molecule from the main unit cell moves out of the unit cell an identical image water molecule enters the main unit cell. This image water molecule then becomes part of the main unit cell when the atom list is updated. Therefore, PBC conserves mass, the number of particles and total energy. An important consideration is the size of the main unit cell. Ideally, the size of the main unit cell should be the total length of the molecule being modeled plus two times the largest cutoff distance used for non-bonded interactions. When the size of the unit cell is defined in this way, no two solute atoms will interact with each other directly and no solvent molecule will see two copies of a given solute atom.

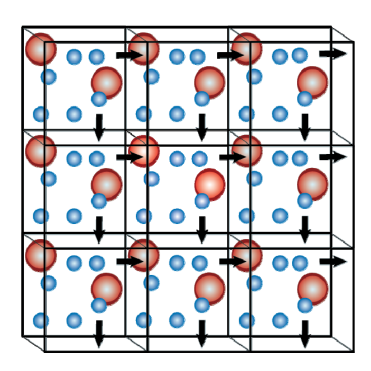


Figure 17: Schematic representation of periodic boundary condition

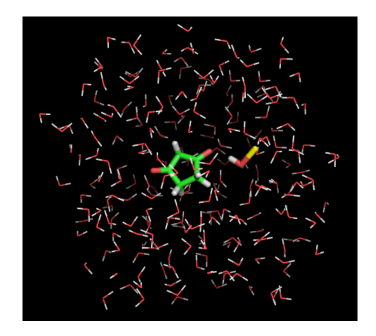


Figure 18: Solvated CPD and HOCl in a truncated octahedron box of water

2.4. Quantum Mechanics

To study reactions that involve the formation of new bonds or breaking of bonds, one needs to solve the equations of Quantum Mechanics (QM) applied to the electronic structure of molecules. The QM model is described by the time-independent Schrödinger equation:

$$\hat{H}\psi = E\psi$$

where ψ is the wavefunction describing the system, \hat{H} is the Hamiltonian operator and E is the total energy of the system. The Hamiltonian operator for the time-independent Schrödinger equation is given by the following:

$$\hat{H} = -\sum_{i} \frac{\hbar^{2}}{2m_{e}} \nabla_{i}^{2} - \sum_{k} \frac{\hbar^{2}}{2m_{k}} \nabla_{k}^{2} - \sum_{i} \sum_{k} \frac{e^{2}Z_{k}}{r_{ik}} \nabla_{i}^{2} + \sum_{i < j} \frac{e^{e}}{r_{ij}} + \sum_{k < l} \frac{e^{e}Z_{k}Z_{l}}{r_{il}}$$

where i and j label electrons, and k and l label nuclei. The first two terms describe the kinetic energy of electrons and nuclei, respectively, and the last three terms describe the potential energy of interaction electrons with nuclei, of electrons with each other, and of nuclei with each other, respectively. To simplify the above Hamiltonian the Born-Oppenheimer approximation is applied, which decouples the correlated motion of electrons with nuclei. The nuclei are assumed fixed in space while the energy of the electrons surrounding the nuclei is calculated. The above Hamiltonian is then simplified to the following (nuclear-nuclear repulsion energy can be added back in after the electronic energy is calculated):

$$\hat{H} = -\sum_{i} \frac{\hbar^2}{2m_e} \nabla_i^2 - \sum_{i} \sum_{k} \frac{e^2 Z_k}{r_{ik}} \nabla_i^2 + \sum_{i < j} \frac{e^e}{r_{ij}}$$

Most approaches to solving the Schrödinger equation begin by expressing the wavefunction as a sum over basis functions:

$$\boldsymbol{\psi} = \sum_{i=1}^{N} a_i \phi_i$$

where ϕ_i is a basis function and a_i is a coefficient. When the basis functions are appropriately chosen and spin is correctly accounted for, the correct solution for ψ can in principle be determined with an appropriate methodology.

2.4.1. Theoretical Methods

It is not practical to solve the Schrödinger equation exactly for any but the simplest of systems which include only one electron, for example a hydrogen molecule ion, because of the electron-electron repulsion term. Hence, there are several approximations made to solve systems of interest. First, the basis set expansion above must be truncated at a finite number of orbitals. In most approaches, the basis functions in the expansion above are themselves expanded further:

$$\phi_i = \sum_{j=1}^M a_j \varphi_j$$

The functions φ_i are termed atomic orbitals, and the functions ϕ_i are termed molecular orbitals. When either expansion is finite (which must be the case for practical applications), the solution will in principle be approximate. A molecular orbital constructed as a linear combination of atomic orbitals is called a linear combination of atomic orbitals (LCAO).

One of the most basic methods for solving the Schrödinger equation, the Hartree-Fock (HF) method, makes the approximation that each electron is influenced by an average

static potential of the remaining electrons in the system and an iterative 'self-consistent field' (SCF) method is used to determine the wavefunction. The HF molecular orbitals consist of Slater Determinant wavefunctions that include the exchange effects of the electrons. However, the HF method neglects most aspects of electron correlation, and this approximation may yield large errors in properties calculated from the wavefunctions. More sophisticated and robust methods need to be used, but HF often provides a practical stepping stone to a more exact solution.

There are a large number of methods for including more aspects of electron correlation than HF theory does. Some of the most widely used methods are Moller-Plesset (MP*n*) perturbation theory and Coupled Cluster (CC) theory. The MP*n* method depends on the perturbation order (*n*) correction used. However, these methods do not scale too well with the number of basis set and are computationally very expensive. For example, the CCSD (CC with single and double excitations) scales at the order of N⁶ and MP2 (with second-order perturbation correction) scales at the order of N⁵, where N is the number of basis sets. All of the methods described thus far are called ab initio methods because they do not make reference to any empirical parameters, and thus construct a solution from first principles.

Another alternative method is Density Functional Theory (DFT), which is different from the wavefunction theory discussed so far. The DFT approach accounts for electron correlation and offers equal or comparable accuracy to MP2 methods with much better computational efficiency. Scaling of DFT is of the order N³, but for practical purposes (discussed later in this section) it scales to the order of N⁴. Density Functional

Theory scales substantially better than the other methods., For these reasons, I use DFT for my calculations.

Density functional theory was first introduced in 1927 by Thomas and Fermi, but the approximations made were inaccurate. The theory was later legitimatized by the proof of two theorems, the Hohenberg-Kohn existence theorem and the Hohenberg-Kohn variational theorem in 1964. The most difficult part of DFT is not having an analytical form for the exchange correlation energy, and for this reason some approximations are made. One is to use parameters that are empirically optimized. Hence, DFT is sometimes referred to as a semi-empirical method, but in fact the functionals used are not parameterized to the extent that semi-empirical molecular orbital theory is. Density functional theory makes an approximation in order to calculate the energy density, a component of the exchange correlation energy. One form of the approximation, called local density approximation (LDA), treats the electron density as a uniform electron gas and is single valued at every position. Because of this approximation we can expect limitations in accuracy. One way to improve the exchange correlation functional is to allow it to depend not only on the local electron density but also on the extent on the gradient of the local density. This approximation is called the generalized gradient approximation (GGA).

The most popular GGA exchange functional is due to Becke (B); it incorporates a single empirical parameter. The most popular GGA correlation functional is LYP. The functional I use, B3LYP, is a hybrid functional that uses B3 with a three parameter scheme for the exchange functional and an adiabatic connection method (ACM) and LYP

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for the correlation functional. ACM methods are called hybrid methods because they incorporate HF and DFT exchange terms.

I chose the B3LYP/6-31+G** level of theory and basis set because it has been shown to be sufficiently accurate for cases like my system, and it is sufficiently fast. The software packages I used for the QM calculations were Gaussian98 and Gaussian09 [56].

2.4.2. Basis sets

Choosing an appropriate basis set is very important for achieving an accurate solution to the electronic structure problem. In practice, the atomic orbitals in the second expansion above are themselves further expanded in terms of other functions. I will briefly discuss these. It is computationally efficient to represent atomic orbitals in terms of Gaussian type orbitals (GTOs) [46]. While GTOs are computationally convenient, they have issues with representing the radial portion of the orbital. The best way to deal with this drawback of GTOs is to form linear combination of GTOs. Basis sets are made up of linear combinations of primitive GTOs called contracted orbitals. Using primitive and contracted orbitals gives more flexibility in manipulating the coefficients so as to have a proper radial representation of the orbital. It is found that the most efficient (accuracy vs. computational efficiency) way to create a minimally realistic representation of an atomic orbital is in terms of three contracted orbitals. The nomenclature used to describe this kind of representation is "STO-3G basis set."

For the accuracy that I need, larger basis sets than the STO-3G are needed. The most important considerations for increasing the basis set size are described next. It is known that the valence orbitals play a much more major role in reactions than the core orbitals. Hence, there is more to be gained by having flexibility in the valence basis functions than in the core basis functions. This led to the concept of a 'split-valence' basis set, in which the core orbitals are represented by a single contracted basis function and the valence orbitals are represented by several contracted basis functions. I use the 6-31G basis set in my QM calculations. The '6' in the basis set representation indicates that the core orbitals are represented by one contracted orbital consisting of six primitive orbitals. The other two numbers, '31', indicate that the valence orbitals are represented by two contracted basis functions each. The first contracted basis function includes three primitive functions and the second contracted function includes one normalized primitive function. In addition polarization and diffuse functions can be augmented to the basis set, which are described below.

Polarization functions

Several molecular properties and especially geometries are sensitive to polarization effects. Atomic orbitals can be accurately represented with s, p functions, etc. This is because atomic orbitals, like these functions, are centered about the atom nucleus. Molecular orbitals, on the other hand, are often extended in space and are expanded in terms of LCAOs with several atom nuclei. For this reason it is necessary for the molecular orbital to have more flexibility than atomic orbitals.

One way to provide this flexibility is to include an additional quantum number of higher angular momentum than the valence orbitals in the form of a GTO basis function. This means that elements in row two like carbon and oxygen. have polarization functions that include d-type GTOs; for hydrogen atoms the polarization functions includes p-type GTOs. The nomenclature used in this case is either a '*' or, equivalently, (d) at the end of the basis set specification for row two heavy atoms; and '**' or, equivalently, (d,p) when polarization functions are to be used for both second row atoms and hydrogen atoms. In my calculations I use the following basis set specification and polarization functions: 6-31G**.

Diffuse functions

Electrons in some molecular complexes tend to be more spatially diffuse than in atomic orbitals. To provide the flexibility needed to represent this spatial diffuseness, the basis function is augmented with diffuse functions when needed. The nomenclature used to indicate the addition of diffuse functions is a '+' added to the basis set specification for additional heavy atom s and p functions and a '++' to indicate diffusion functions of s type for hydrogen atoms in addition to heavy atom diffuse functions. In my calculations I use diffuse function for only heavy atoms; 6-31+G**.

2.5. Quantum Mechanics / Molecular Mechanics

As described in previous chapters MM is useful for modeling large systems. However, it cannot be used to model reactions that involve bond-making or bondforming. Instead we use a QM model to model chemical reactions. The disadvantage to the QM model is that it does not scale well as the number of heavy atoms increases and therefore the limiting factor is the size of the system. We are restricted to using the QM model for only small systems, or for reactions of large systems, one has to ignore atoms not near the reaction site. In many cases, though, these atoms may significantly influence the reaction even though their role may not be as important as the ones near the reaction site. These atoms may influence the reaction via long range non-bonded interactions (electrostatic or van der Waals) with the reactive atoms or, in the case of macromolecules, by influencing the conformation of the system as a whole.

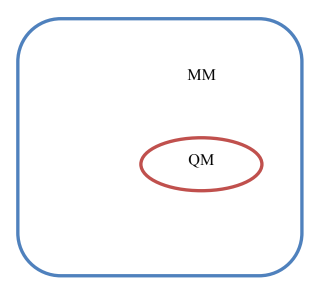


Figure 19: Schematic of QM and MM regions

Fortunately, larger systems can often be treated by using a combination of the MM and QM approaches. This kind of a hybrid method is denoted QM/MM. One can model reactive site atoms that are deemed crucial to the reaction using QM. The other atoms of the large system can then be modeled using MM instead of ignoring them. The development and application of QM/MM methods have grown rapidly over the past few years.

For a QM/MM model, the total Hamiltonian is written as

$$H_{Total} = H_{QM} + H_{MM} + H_{QM/MM}$$

This equation has a deceptively simple appearance. The details of $H_{QM/MM}$ can become quite complicated, depending on the system. The range of factors that can arise is described below, but as in earlier chapters emphasis is given to factors arising in the research reported here.

The first thing that comes to mind when we consider QM/MM systems is the boundary between the QM section and MM section and whether the boundary intersects a bond or not. It is a little simpler if the boundary does not intersect a bond. If it does, there are a few ways it is handled. One way is to use a link atom, usually hydrogen, to cap off the dangling bond in the QM section. Another way is to use a frozen orbital. Sometimes a different methodology, ONIOMM, is used. In my system, the boundary does not intersect any bond, so I did not have to deal with any of these issues.

Even though my system does not have to deal with the issue of the QM/MM boundary severing bonds, there are still other matters that need to be considered and

choices to be made. One is the coupling or polarization between the two regions. There are three possible interactions. Both regions can be unpolarized, only the QM region is polarized and the third option is having both regions polarized.

My model, incorporates polarization of the QM region by the MM region, but not polarization of the MM region. The MM atoms are represented as fixed point charges. The Hamiltonian for such a system is

$$H_{\underline{QM}/\underline{MM}} = \sum_{i}^{\text{solute electrons}} \sum_{m}^{\underline{MM} \text{ atoms}} \frac{q_{m}}{r_{im}} + \sum_{k}^{\text{solute nuclei}} \sum_{m}^{\underline{MM} \text{ atoms}} [\frac{Z_{k}q_{m}}{r_{km}} + 4\mathcal{E}_{km}(\frac{\sigma_{km}^{12}}{r_{km}^{12}} - \frac{\sigma_{km}^{6}}{r_{km}^{6}})]$$

Another issue that needs to be considered is the interaction of atoms in both regions that are very close to the boundary. MM model represents atoms as point charges, while the QM atoms have electronic and nuclear representation of atoms. The MM point charges near the boundary can have large unrealistic influences on the electronic representation of the QM atoms near the boundary. They can distort the QM electron density and create forces acting on QM nuclei. One way to deal with this situation is to blur the point charges in the MM region that are close to the boundary by Gaussian functions so that the charges are spread or 'blurred' and do not have such drastic, unphysical effects on QM atoms close to the boundary. The CHARMM program includes such an option, and I used it in the work reported here.

The QM software I use to interface with CHARMM to implement the hybrid QM/MM model is Q-Chem 3.2 [57].

2.6. Chain-of-replica Methods

A reaction mechanism is better understood by studying its reaction path. A reaction path is the relative motion taken by atoms during a chemical reaction. We usually think of a reaction path as beginning with reactants, proceeding through a highenergy structure called a transition state, and terminating in products. In complicated cases, there may also be significant intervening structures called intermediates. For any chemical reaction, there are many possible reaction paths. Particular significance is given to that reaction path that passes through the lowest-energy transition state and for which the path from the transition state toward either reactants or products follows the direction of steepest descent; this is called the Minimum Energy Path (MEP) or Steepest Descent (SD) path. The significance of the MEP is that in a macroscopic system of many reacting molecules, the vast majority of reactive events follow a trajectory close to the MEP, and consequently the energetics of the MEP dominates the kinetics. Thus the MEP is often predictive of rate constants and reaction mechanisms, and so most analyses are based on the MEP. In mathematical terms, reactants, products and transition states (TS), and also intermediates if they exist, are stationary points because for them the gradient of the potential energy surface is zero. Clearly, the development of methods to identify MEPs, and especially the stationary points, is very important in understanding reaction mechanisms.

There are broadly three classifications of methods used to identify the reaction path. The first are eigenvector following methods, of which the most popular method is the Intrinsic Reaction Coordinate (IRC) method [58, 59]. This method relies on first

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identifying the transition state, for which by definition the Hessian possesses exactly one negative eigenvalue, and then following the eigenvector corresponding to the negative eigenvalue in the forward and reverse directions. This process eventually leads to the reactant and product structures. While it is robust, there are several drawbacks to this method. It is sometimes very difficult to find the TS structure if the potential energy surface is complex or the saddle point is quite flat. Locating the TS is also a very expensive method in terms of scaling. Memory scales as N^2 and time scales as N^3 , where N is the number of degrees of freedom. The size of the system restricts the use of this method. It is not practical to use in large systems such as the one studied here.

Another method which is quite intuitive is the reaction coordinate following method. If we know the reaction coordinate that drives a particular reaction, we just follow it and it should connect the three stationary points. The reaction coordinate following method, however, is impractical for all but the simplest of systems, since it is very difficult to know the reaction coordinate in advance, especially for reaction mechanisms that involve large numbers of atoms. Another issue with this method is that one needs to check the hysteresis to make certain that the same path is mapped out in the forward and reverse directions.

The method I use is from a group of methods called chain-of-replica methods [60-71] that are fairly new compared to the other methods. Its development is quite significant because it allows one to study reaction mechanisms of large systems, and in most cases only first derivatives need to be calculated. In some cases the results can be refined using a second derivative method for geometry optimization, but even then, chain-of-replica methods are more feasible for large systems than are other methods. Another significant advantage of chain-of-replica methods is that one only needs the reactant and product structures as input, whereas the IRC method also requires a fully optimized transition state. In fact, chain-of-replica methods are efficient for locating transition states in high-dimensional systems where the usual transition-state geometry optimization methods may be impractical. There is also no need to know the reaction coordinate in advance which can be a very difficult task with a system having many degrees of freedom. A further benefit of this method is that it can take advantage of today's parallel computers. Since individual points on the path are only weakly coupled, they can be refined on different processors. The other methods are sequential and cannot make use of parallel computers as efficiently.

The chain-of-replica method was originally proposed by Elber and Karplus [67]. It was based on work by Empedocles who in 1969 had proposed using a line integral for searching a reaction coordinate.

$$S = \frac{1}{L} \int_{R_A}^{R_B} V(R) dl(R)$$

where S is the function to be optimized, R_A and R_B are position vectors representing the coordinates of reactant and product structures, V(R) is the potential energy and dl(R) is the line element along the length of the path, L, between R_A and R_B . V(R) and dl(R) are functions of the N-dimensional position vector representing coordinates, where N is the number of degrees of freedom.

Elber and Karplus approximated the above line integral by a discretized version.

$$S = \frac{1}{L} \sum_{j=0}^{M} V(R_j) \Delta l_j + \text{ constraints}$$

where M is the number of intermediate discretized grid points and ΔI_j is the interval between each point. Constraints were included to keep the individual points equidistant from each other. This is because, during the process of minimization, if the ΔI_j were widely different the equation above would no longer be a good approximation of the line integral. Later additional repulsion constraints were added so that the points did not all pool into a local minimum. In the methods I used that are described below, the distance ΔI between coordinate sets representing points on the path is defined as the root mean square difference (RMSD) between the atomic coordinates of the two sets. The definition of ΔI includes overall translation and rotation of the coordinate sets to achieve a best fit in a least squares sense. The total length of the path, L, is the sum of the distances ΔI between points on the path so defined. Thus, L is a function defined in RMSD space, $L(r_1, r_2, ..., r_{M-1})$, where r_i is an RMSD of coordinates just defined..

The formalism described above forms the basis for all chain-of-replica methods. Individual methods differ mainly in the way the points are generated, the minimization protocol used, and most importantly the kind of constraints imposed to control the optimization.

The methods I used for include the Replica Path method and the Nudged Elastic Band (NEB) method with the Climbing Image algorithm to obtain the TS structure (CI-NEB). I also used the Adaptive Nudged Elastic Band (ANEB) method for a single test case. All these methods used are implemented in CHARMM, the software I used for my calculations.

2.6.1. Replica Path Method

The easiest and most common way to create initial replicas or images is by linear interpolation of the reactant and product structures. The replica path method, like most modern chain-of replica methods, only replicates certain atoms that are deemed important in the reaction being studied instead of the entire system. The un-replicated part of the system is still present and identical across all replicas. The un-replicated part can and will affect the replicas and respond, if needed, but it will do so the same way for all replicas along the MEP. This, quite conveniently, sets up for use with the QM/MM model. It allows the focus of the study and resources on the active site of the molecule. In the geometry optimization calculation of the replicas the un-replicated atoms were kept fixed.

The main aspect of replica path that sets it apart from the other chain-of-replica methods is that the method uses a harmonic penalty function to constrain/restrain the points on the reaction path. The penalty function applied is directly proportional to the root mean square deviation (RMSD) in the atomic coordinates of two neighboring replicas or images. During the evaluation of the RMSD the deviation because of rotation and translation are avoided. An advantage of using this RMSD space to define the restraints is that there is no need to deal with effects of rigid body rotation and translation which would otherwise affect the minimization process.

Below are the penalty functions and other keywords used with replica path as implemented in CHARMM.

$$E_{rms} = \frac{1}{2} \sum_{i=1}^{N} K_{rms} (r_i - \bar{r})^2$$

where E_{rms} is the penalty function used to force the replicas or images to be approximately equidistant from each other. N is the number of replicas or images along the pathway. The optimal value of the force constant K_{rms} should result in a path step variation of no greater than 15 – 20 %. The distance between path points i and i+1 in RMSD-space is denoted r_i , and the average over all pairs of consecutive path points is \bar{r} :

$$r_i = RMSD_{best fit}(i, i+1)$$

$$\overline{r} = \sum_{i=1}^{N} \frac{r_i}{N}$$

One artifactual difficulty sometimes encountered is a tendency for the reaction path to become highly curved so that points fall into each other, resulting in small values of the constraint energy but giving unrealistic paths. To avoid this problem, an additional restraint, E_{anele} , is used to keep the reaction path realistically linear:

$$E_{angle} = \frac{1}{2} \sum_{i=1}^{N} K_{angle} [COSMAX - \cos(\Theta)_i]^2 \text{ if } COSMAX > \cos(\Theta)_i$$

$$E_{angle} = 0 \qquad \qquad \text{if } COSMAX \le \cos(\Theta)_i$$

where Θ is the (i - 1) - i - (i + 1) angle in RMSD-space. COSMAX is a value between 0.9 and 0.98, with the value 0.98 making the pathway more linear and rigid. I use the value 0.9 in my calculations. K_{angle} is the force constant used for the restraint. The recommended range is 100 – 500 kcal/mol. I used the value 100 kcal/mol in the calculations reported here. Below are the definitions of the most important keywords used in the CHARMM's replica path facility.

RMAX: Maximum distance the replicas are allowed to be separated before switching on the force constant KMAX instead of K_{rms} in the penalty function E_{rms} .

KMAX: The force constant to be used when the distance between replicas exceeds RMAX and is usually greater than or equal to K_{rms} .

WEIGHT: This keyword causes the values in the main weighting array are to be used. This allows one to place more emphasis on atoms that play a major role in the reaction, so that when the RMSD is calculated to constraint the replicas more emphasis is placed on those atoms. ROTATION and TRANSLATION: Keywords that turn on the RMSD best-fit procedure between neighboring replicas, thus preventing the rigid-body rotations and translations from effecting the minimization. Without the RMSD best-fit procedure the atoms would adjust unnecessarily by rotating and/or translating in order to meet the constraint requirement.

BLOCK: The utility in CHARMM that calculates the total potential energy at each point along the minimum energy path. The total potential energy is the sum of the interaction energy between atoms in the un-replicated region plus the interaction energy between atoms in an individual replica plus the interaction energy between atoms in a single replica and the atoms in the un-replicated region. The BLOCK utility has to be appropriately setup so that the inter-replica interactions are ignored and the interaction matrix is appropriately scaled. The scaling factor for the interaction energy between atoms in the un-replicated region is set at 1.0. The scaling factor for the interaction energy between atoms in an individual replica is the inverse of the number of replicas created. The scaling factor for the interaction energy between atoms in a single replica and the atoms in the un-replicated region is the inverse of the number of replicas created. The scaling factor for the interaction energy between atoms in a single replica

2.6.2. Nudged Elastic Band Method

The NEB method is similar to the replica path method and uses the same keywords in CHARMM. The implementation, however, is different. The NEB method was originally proposed to address a concern about replica path. In replica path, the physical interatomic interactions and the extra spring restraints can interact, potentially causing deviations from the true MEP. Such deviations are especially likely to occur in steep regions of the reaction path, such as are often encountered near a transition state. While replica path method implements the penalty function based on the RMSD as described above, NEB uses a force projection scheme in which the physical force due to interactions acts only perpendicular to the path, and spring-constraint forces act only along the path. The spring forces keep the replicas separated from each other and the potential forces acting perpendicular to the path 'nudge' the pathway closer to the MEP. This is accomplished with the following set of equations.

$$F_i^{NEB} = F_i^{\perp} + F_i^{S|}$$

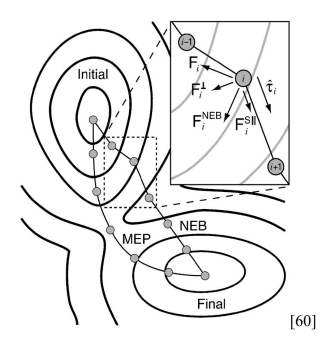


Figure 20: NEB

where F_i^{\perp} is the component of the force due to the potential perpendicular to the band given by

$$F_i^{\perp} = -\nabla(R_i) + \nabla(R_i) \bullet \hat{\tau}_i \hat{\tau}_i$$

where $\hat{\tau}_i$ is the unit vector in the tangential direction of the higher energy neighboring replica, and $F_i^{S\parallel}$ is the spring force parallel to the band and is given by

$$F_i^{S||} = k(|R_{i+1} - R_i| - |R_i - R_{i-1}|)\hat{\tau}_i$$

2.6.3. Climbing Image – Nudged Elastic Band Method

A problem often encountered in the methods described above is that the actual transition state may be difficult to reach, and the resolution of replicas near the transitions state may be poor. Poor resolution is particularly problematical if the energy barrier is narrow in comparison to the length of the minimum energy path. What is usually noticed in such situations is that the replicas tend to slide away from the transition state and cause 'corner-cutting' around the transition state.

The climbing image algorithm (CI-NEB) is a modification of the NEB method. The main difference is the treatment of the replica that has the highest energy in the pathway after a few minimization steps. That point is not subjected to the spring forces and is driven towards the transition state instead of being restricted by the spring forces. The total force acting on that particular point is given by

$$F_{i_{\text{max}}} = -\nabla E(R_{i_{\text{max}}}) + 2\nabla E(R_{i_{\text{max}}}) \mid_{\parallel}$$

$$F_{i_{\max}} = -\nabla E(R_{i_{\max}}) + 2\nabla E(R_{i_{\max}}) \bullet \hat{\tau}_{i_{\max}} \hat{\tau}_{i_{\max}}$$

This modification to the NEB does not add any additional computational time. The replica moves along the elastic band up the potential energy surface to the transition state.

2.6.4. Adaptive Nudged Elastic Band Method

The ANEB method [61] is a variation of the NEB method. The traditional NEB method can sometimes have difficulty approaching the transition state structure due to poor convergence on difficult and flat potential energy surfaces. The focus of the ANEB method is to increase the resolution of points or replicas near the transition state and to acquire a good estimate of the reaction barrier. Just like the regular NEB method the initial setup requires two local minima representing the reactant and product structures. The initial replicas or images may be created by linear interpolation as in the NEB method and minimized until a reasonable convergence criterion is met. At this point ANEB deviates from the regular NEB method. New fixed points are selected instead of the reactant and product structures in the previous step. The new fixed points selected are

the ones on either side of the point or replica with the maximum energy. Presumably, these points encompass the actual transition state. New images are then created between these points as in regular NEB and minimized. This procedure is continued until the transition state is found. Hence, the resolution of points or replicas is adaptively increased near the transition state. Another benefit of the ANEB method is that fewer replicas are needed at each iteration.

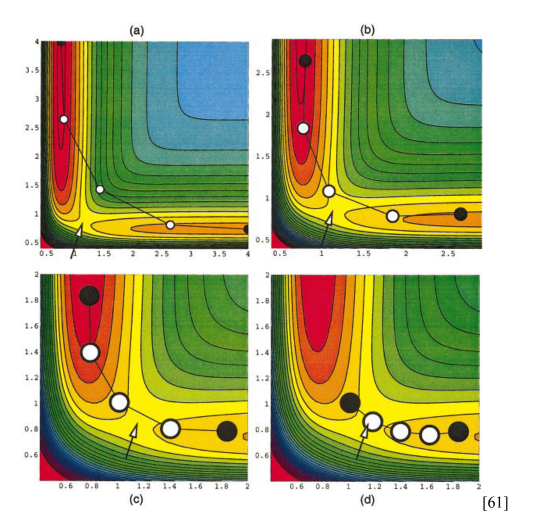


Figure 21: ANEB

3. PARAMETERIZATION

3.1. Principles

As discussed earlier in the Molecular Dynamics and QM/MM sections, programs running MD calculations require parameters that need to be read at the beginning of the script. The default parameters provided by the software are adequate for most systems or molecules. However, in some rare cases one has to develop one's own set of parameters if the software does not provide parameters for the molecule being studied.

Parameters are provided in two files called the topology file and the parameter file. The topology file contains information such as the names of molecules, atom names and atom types (defined below) partial charges on individual atoms, atom connectivity (bonds), and if present, improper dihedral angles, acceptor or donor atoms, internal coordinates and patch information (primarily need for protein molecules). The molecule name is used by the script to build or make reference to the molecule and hence has to be unique with a maximum of four characters. The connectivity information lets the software know which atoms are connected and if they share a single, double or triple bond. Acceptor or donor atom information is used for hydrogen bond analysis. A data structure for internal coordinates is set up to reserve memory in case internal coordinate manipulations are needed or in order to build Cartesian coordinates from internal coordinates. Patch information is used to connect fragments or segments and in some cases it tells the program how to handle a residue at the ends of a segment. The parameter file contains information such as the atom type, Van der Waals constants, force constants for bonds, angles, dihedrals and impropers and their respective equilibrium values. This information is used in energy calculations as described earlier. In cases where there are no parameters available for the system or molecule(s) being studied one has to develop one's own parameters. Depending on the software being used there are different protocols. One cannot mix and match parameters from different types of software. For example, one cannot use some parameters from Amber and some from CHARMM. All the parameters have to be from either Amber or all from CHARMM for internal consistency. Hence, it is very important to be consistent in using the correct protocol. Alexander MacKerell has become the primary developer of parameters for the CHARMM suite of force fields. I use the CHARMM parameter set and therefore follow the protocol outlined by MacKerell et al. to develop the parameters for my system [72].

There are two ways to go about developing parameters for atoms or molecules for use in CHARMM. The first and simplest way is called "by analogy." One identifies fragments of the molecule being parameterized that are similar to fragments of molecules already present in the default CHARMM parameter files. If it is possible to identify a complete set of fragments that when combined yield the complete molecule, only a few parameters need to be developed. These are for connecting the individual fragments. The analogy method cannot be used if the default parameter set lacks the needed fragments. In that case, one must develop all parameters for the entire molecule from the ground up. If the parameter set contains only a few of the needed fragments, it is generally observed that developing all parameters from the ground up yields the best parameterization. Below is a flowchart showing the process by which parameters are developed. Following it, the procedures used to develop parameters for the systems of interest to us are described in detail.

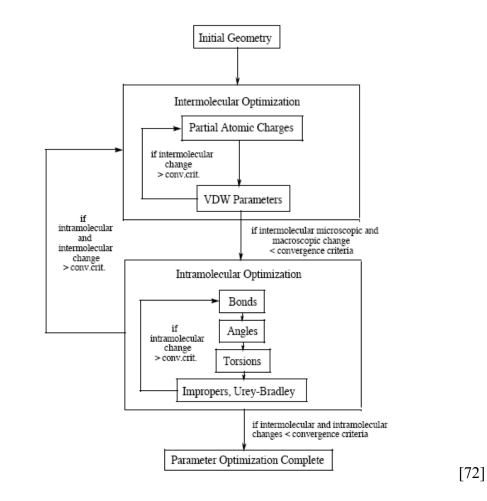


Figure 22: Parameterization protocol flow chart

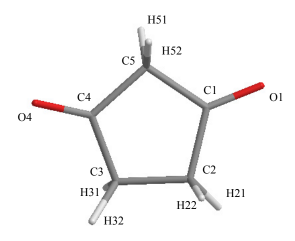


Figure 23: Cyclopentanedione (CPD). Atom names are shown

The molecule CPD is directly involved in the chlorination reaction being studied. Therefore highly accurate parameters are needed and I choose to develop all of them from the ground up. I only need to develop parameters for the diketo form of CPD and not the ketoenol form, since I only need the parameters to perform MD simulations involving the diketo form to identify "unique encounter reaction precursors," as explained in chapter 5. I will actually begin with a set of parameters determined by analogy, but I subsequently refine them. First, though, I must define a few terms.

3.1.1. Atom type and atom name

In Figure 23, CPD is shown with a unique "atom name" for each atom. In force fields, in addition to a unique "atom name," each atom is also assigned an "atom type." "Atom type" is a technical term that groups atoms according to properties. Parameters are defined in terms of atom types, not in terms of elements or individual atoms (i.e.

"atom names"). Thus the first step in parameter development is to decide which set of atoms will be assigned to the same atom type. A given element may, in different contexts, have quite different properties. Therefore, for example, an oxygen in a keto group is assigned to one atom type, while an oxygen in a hydroxyl group is assigned to another. In principle, almost every environment is unique, and so one could define a very large number of atom types. This would make force fields extraordinarily complicated and costly. In practice, a middle ground is sought in which atoms in quite different environments are assigned different atom types, while those in similar environments are assigned the same.

In the case of CPD, it is reasonable to identify two carbon atom types, one oxygen atom type and one hydrogen atom type. I look for similar atom types in the CHARMM general force field parameter file. The carbon, oxygen and hydrogen atom types are similar to their respective atoms type in 2-pyrrolidinone - C1 and C4 of type CG2R53; C2, C3 and C5 of type CG3C52; O1 and O4 of type OG2D1; H21, H22, H31, H32, H51, H52 of type HGA2. The VDW parameters are not expected to change significantly and so I use the same values without refinement, as recommended by Mackerel.[72] The reason I look for similar atom types already present in the default parameter files is to avoid creating new atom types in the original topology and parameter files if possible. I could just as easily create completely new atom types since I modify all the parameters (except the VDW parameters since it is not expected it to change significantly) for CPD. It is worth noting that some thought must be given while deciding whether it is best to create new atom types or using existing atom types. If one uses existing atom types, like I did, then the new parameters will overwrite old parameters, if they already exist, in the parameter file. So, care must be taken that one does not intend using the old parameters for other molecules. In this case, some parameters are in fact overwritten, but in the context of my work I do not require the old parameters so it is safe for us to use the old atom types.

Atom Name	Atom Type	Description
C1	CG2R53	5-membered ring carbon bound to a keto oxygen, e.g. in 2-pyrrolidinone
O1	OG2D1	carbonyl oxygen; e.g. carboxylic acids
C2	CG3C52	5-membered ring carbon bound to aliphatic hydrogen e.g. in proline
H21	HGA2	aliphatic hydrogen
H22	HGA2	aliphatic hydrogen
C3	CG3C52	5-membered ring carbon bound to aliphatic hydrogen e.g. in proline
H31	HGA2	aliphatic hydrogen
H32	HGA2	aliphatic hydrogen
C4	CG2R53	5-membered ring carbon bound to a keto oxygen, e.g. in 2-pyrrolidinone
O4	OG2D1	carbonyl oxygen; e.g. in carboxylic acids

C5	CG3C52	5-membered ring carbon bound to aliphatic hydrogen e.g. in proline
H51	HGA2	aliphatic hydrogen
H52	HGA2	aliphatic hydrogen

Table 1: CPD Atom Names and Atom Types

3.1.2. Partial atomic charges

The next step in the protocol is to assign initial guess partial charges to the different atoms. Each atom may in principle be assigned a different partial charge (i.e. the atom type does not determine the partial charge), but in general atoms in similar environments are assigned the same charge. Following the protocol of MacKerell et al. I start by assigning a +0.09 charge to all hydrogen atoms. The next step is to assign charges to the other atoms so that the total charge is zero, since the system is neutral. I determine an initial guess by calculating the charges using a quantum mechanical method. MacKerell et al. recommend using the HF/6-31G* level of theory to optimize the geometries and electronic wave functions of all molecules. This provides internal consistency for CHARMM parameter development. They recommend using the Merz-Kollman method derives atomic charges from a least-squares fit to the electrostatic potential (ESP) calculated at a large number of points around the molecule of interest. The Merz-Kollman method differs from similar methods in the choice of points where the ESP is calculated.

The MK scheme uses grid points on several layers around the molecule with different scaling factors for each layer. Keeping in mind the symmetry of the molecule and the fact that aliphatic hydrogen atoms need to have a +0.09 charge, I assign initial guess charges for all the atoms.

The next step is calculating target data for use in optimizing the partial atomic charges. These are obtained from quantum mechanical calculations. Two types of target data are calculated. One target datum is the dipole moment. The others pertain to the interaction between water molecules and CPD. In particular, the interaction energy and the optimal interaction distance are evaluated. An evaluation is made for each of the solvent-exposed atoms of CPD. This is accomplished by placing a single water molecule in close proximity to the atom under consideration, and performing the calculations described below. The partial atomic charges are then adjusted iteratively until the MM calculations well reproduce the dipole moment and water interactions obtained from QM calculations.

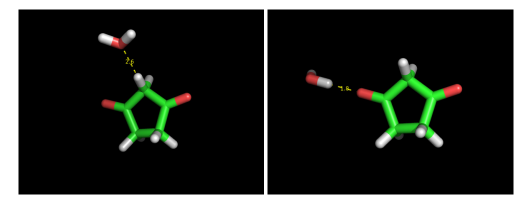


Figure 24: Two example representations of water interacting with atoms in CPD I begin with two structures; one optimized using QM and the other optimized

using the CHARMM program for 200 minimization steps using the Conjugant Gradient

algorithm and another 50 minimization steps using the Newton-Raphson algorithm. The QM optimized structure is copied in the comparison coordinate set and then the CHARMM optimized structure is aligned or oriented so that the dipole moments can be calculated and compared.

To get the other target data for water interaction energies and optimal hydrogen bond distances I first read in the QM optimized structure and build a TIP3P water molecule 1.6 Å from one of the atoms. The TIP3P water molecule is a classical water model that I use in the solvation study. The energy is set at an arbitrary maximum value of 1000 kcal/mol. The distance between the water molecule and the atom is gradually decreased by 0.01Å and the corresponding energy calculated. This is done until the minimum interaction energy is located. I follow the procedure for every periphery atom, with a single water molecule, one at a time. This data is then compared to the target data which I obtain from the QM calculation.

As mentioned above, the CHARMM General Force Field is an additive force field. One of the inherent disadvantages to an additive force field is the way electrostatic interactions are calculated. Coulomb's law is used to calculate the electrostatic interactions and uses fixed point charges on atoms to do so. Using fixed point charges is a major simplification. It is well known that the electron density or distribution around an atom and in this case atomic charges, changes as a function of the electrostatic field around the molecule. So, as the atoms move during simulation, their atomic charge should change or respond to its environment. This is ignored in an additive force field. To compensate for the lack of polarizability the atomic charges are 'tweaked' or 'enhanced', especially for molecules that are polar and in condensed phase. CPD is a polar molecule; hence, the target data acquired from QM calculations are 'enhanced'. One of the major developers of the CHARMM force fields, MacKerell et al., has outlined the modifications that need to be made to the target data. I follow these recommendations and make the following changes. The target interaction energies for the QM optimized structure are scaled by a factor of 1.16. The target optimal hydrogen bond distances are increased by about 0.2 Å. The target dipole moment is scaled by 0.8-0.9.

The only parameters I can change at this step are the partial atomic charges on CPD. CPD is a symmetric molecule so we expect atoms like C1 and C4 to have the same charge. Likewise, C2 and C3 would have the same charge and also O1 and O4. The aliphatic hydrogen atoms need to have an atomic charge of +0.9, as described earlier. Therefore, there are only four atomic charges I am able to manipulate, including C1 (or C4), C2 (or C3), O1 (or O4) and C5. Minor changes were made to these parameters and then the empirical data was compared to the target data. Several such iterations were done to approach the target data. The approach used to achieve this goal was to have the least possible deviation from the different types of target data like the dipole moment, water interaction energies and their respective minimum energy distance for all the atoms in CPD.

Ideally, the recommended level of agreement, for water interaction energies should be within 0.2 kcal/mol of the target values. The dipole moment should be overestimated by 20 to 50% in comparison to the QM calculated dipole moment. The best partial charges I was able to obtain are listed in Table 2.

Atom Name	МК	Initial CHARMM	Final CHARMM
C1	0.56	0.55	0.48
C2	-0.18	-0.18	-0.19
C3	-0.16	-0.18	-0.19
C4	0.55	0.55	0.48
C5	-0.45	-0.36	-0.2
01	-0.46	-0.46	-0.46
O4	-0.46	-0.46	-0.46
H21	0.07	0.09	0.09
H22	0.08	0.09	0.09
H31	0.07	0.09	0.09
H32	0.07	0.09	0.09
H51	0.15	0.09	0.09
H52	0.15	0.09	0.09

Table 2: CPD Partial atomic charge

The interaction energies with water molecules, the respective hydrogen bond distances with the water molecule and the dipole moments are listed in Table 3. Note: the QM energies are scaled by 1.16.

	QM (kcal/mol)	CHARMM (kcal/mol)	∆ (CHARMM – QM) (kcal/mol)
O1 - HOH	-4.1479	-4.62955	-0.48165
O4 - HOH	-4.1479	-4.6454	-0.4975
H21 - OHH	-3.0084	-2.15067	0.85773
H22 - OHH	-2.5196	-1.68288	0.83672
H31 - OHH	-2.5196	-1.68288	0.83672
H32 - OHH	-3.0084	-2.15067	0.85773
H51 - OHH	-3.29	-1.84931	1.44069
H52 - OHH	-3.29	-1.84931	1.44069
		AVG Difference	0.661391
		RMS Difference	0.707352

Table 3: CPD – H₂O Interaction Energies

	QM (Å)	CHARMM (Å)	∆ (CHARMM – QM) (Å)
01 - НОН	2.11	1.83	-0.28
O4 - HOH	2.11	1.83	-0.28
H21 - OHH	2.47	2.64	0.17
H22 - OHH	2.47	2.66	0.19
H31 - OHH	2.47	2.66	0.19
H32 - OHH	2.47	2.64	0.17
H51 - OHH	2.34	2.63	0.29
H52 - OHH	2.34	2.63	0.29

Table 4: CPD – H₂O Interaction Distances

QM (D)	CHARMM (D)
1.8625	1.8152

Table 5: Dipole moments

3.1.3. Bond and angle terms

The next step is the optimization of intra-molecular bond and angle parameters. The functional forms for these terms are shown below.

$$\sum_{bonds} K_b (b-b_0)^2 + \sum_{angles} K_{\theta} (\theta-\theta_0)^2$$

Bond and angle parameters are optimized at the same time. There are two values needed for bond and angle parameters, the equilibrium values and the force constants. The equilibrium values can easily be obtained from a corresponding QM optimization of the molecule. The MP2/6-31G* level of theory is recommended for this purpose. Ideally, deviations of up to 0.03 Å for equilibrium bond distances and up to 3° for equilibrium angles in comparison to QM calculations is acceptable. The optimization of the force constants, however, is quite involved and was possibly the longest and most tedious part of the optimization process for CPD. CHARMM uses a module called MOLVIB, which is a general-purpose vibrational analysis program used for small molecules that contain fewer than 50 atoms.

The MOLVIB module is very structured and the syntax has to be closely followed. The MOLVIB module is also used to get the target data, but the target force matrix for the vibrational analysis is obtained from a QM calculation of a frequency calculation. The force matrix from the QM (Gaussian98) output is then copied into the MOLVIB with the target frequencies scaled by a factor of 0.89. The force constants are adjusted so that they best match the target data. The goal is to reproduce the frequencies and the contributions of the internal degrees of freedom. Lower frequencies represent motions involving heavy atoms and, therefore, motion that deviates the most from its optimal geometry. Hence, it is a priority to reproduce the low frequencies more accurately. The high frequency motion that represent stretching motion involving hydrogen atom are trivial and in most cases not important. It is preferable to produce vibrational frequencies that are slightly lower than the target values. The reasoning is that it is better to have a more flexible molecule instead of a more rigid molecule.

As a rough guideline the empirical frequencies calculated should be, on average, within 5% of the corresponding QM calculated frequencies. "Apart from these basic guidelines, it is difficult to objectively quantify the quality of the parameterization." "It is often difficult to unambiguously correlate the QM and MM frequencies due to multiple local internal valence coordinate contributions to each frequency. These contributions will often differ between the QM and MM spectra." [72].

Freq	quencies	(cm^{-1}) and	% cont	cribution (from	QM calculat	ed force	e constants)
	Freq		olo		olo		olo	
1	33.8	t-5RNGsy	84.					
2	95.9	t-5RNGas	83.					
3	370.0	d-C1=01	34.	d-C4=04	34.			
4	387.3	w-C4=O4	30.	w-C1=01	30.	r-C5H2	25.	
5	444.8	w-C1=01	32.	w-C4=04	32.			
6	499.7	d-C4=04	40.	d-C1=01	40.			
7	552.8	d-5RNGas	48.					
8	572.6	d-5RNGsy	49.	r-C5H2	16.			
9	683.1	r-C3H2	19.	r-C2H2	19.			
10	775.3	sC3-C4	29.	sC1-C2	29.			
11	790.4	sC4-C5	27.	sC1-C5	27.			
12	870.2	r-C5H2	30.					
13	884.7	r-C2H2	28.	r-C3H2	28.	t-C5H2	27.	
14	977.1	sC2-C3	65.					
15	1104.6							
16	1122.3	t-C5H2	57.					
17	1128.5	t-C3H2	34.	t-C2H2	34.			
18	1157.3							
19	1204.8	w-C5H2	76.					
20	1224.7	t-C3H2	21.	t-C2H2	21.			
21	1255.1	w-C3H2	29.	w-C2H2	29.			
22	1271.1	w-C2H2	39.	w-C3H2	39.			
23	1387.2	sc-C5H2	100.					
24	1432.7	sc-C2H2	50.	sc-C3H2	50.			
25	1448.3	sc-C3H2	49.	sc-C2H2	49.			
26	1695.2	sC4=04	45.	sC1=01	45.			
27	1725.5	sC1=01	44.	sC4=04	44.			
28	2943.2	sC3-H3sy	40.	sC2-H2as	29.	sC2-H2sy	20.	
29	2945.3	sC3-H3sy	45.	sC2-H2sy	29.	sC2-H2as	21.	
30	2951.1	sC5-H5sy	99.					
31	3000.9	sC5-H5as	100.					
32	3015.4	sC3-H3as	40.	sC2-H2sy	29.	sC2-H2as	21.	
33	3022.5	sC3-H3as	45.	sC2-H2as	29.	sC2-H2sy	21.	

 Table 6: QM calculated Frequencies (cm⁻¹) and % contribution

	Frequen	cies (cm-1) and	% contribu	tion (from empir	ical d	calculated	l force
				con	stants	:)			
	Freq		olo		00		0 0		
1	41.6	t-5RNGsy	90.						
2	112.9	t-5RNGas	84.						
3	371.6	d-C4=04	33.	d-C1=01	33.				
4	404.6	w-C4=O4	40.	w-C1=01	40.				
5	435.0	w-C4=O4	41.	w-C1=01	41.				
6	507.9	d-5RNGas	47.						
7	514.6	d-C4=04	26.	d-C1=01	26.				
8	571.5	d-5RNGsy	62.						
9	747.5	sC3-C4	25.	sC1-C2	25.	d-5RNGas	17.		
10	755.8	sC4-C5	15.	sC1-C5	15.				
11	788.0	r-C2H2	32.	r-C3H2	32.	r-C5H2	17.		
12	895.6	r-C5H2	65.						
13	925.5	r-C3H2	39.	r-C2H2	39.				
14	984.2	sC2-C3	68.						
15	1094.7	t-C5H2	93.						
16	1138.1	t-C3H2	43.	t-C2H2	43.				
17	1155.8	t-C3H2	42.	t-C2H2	42.				
18	1159.4	w-C5H2	30.						
19	1240.4	w-C3H2	24.	w-C2H2	24.				
20	1272.1	w-C5H2	38.						
21	1307.6								
22	1385.9	w-C2H2	20.	w-C3H2	20.				
23	1507.0	sc-C3H2	47.	sc-C2H2	47.				
24	1518.7	sc-C5H2	42.	sc-C3H2	26.	sc-C2H2	26.		
25	1523.8	sc-C5H2	45.	sc-C3H2	22.	sc-C2H2	22.		
26	1729.1	sC4=04	36.	sC1=01	36.				
27	1747.8	sC1=01	32.	sC4=04	32.				
28	2797.5	sC5-H5sy	99.						
29	2827.6	sC5-H5as	100.						
30	2849.1	sC3-H3sy	49.	sC2-H2sy	45.				
31	2855.6	sC2-H2sy	51.	sC3-H3sy	50.				
32	2873.5	sC3-H3as	50.	sC2-H2as	45.				
33	2884.9	sC2-H2as	51.	sC3-H3as	50.				

Table 7: MM calculated Frequencies (cm⁻¹) and % contribution

3.1.4. Dihedral and improper terms

The last two sections of the parameter optimization are the dihedral and improper angles and force constants. The torsions are one of the degrees of freedom that are optimized while optimizing the bond and angle force constants. The dihedrals that have no hydrogen are fine-tuned next. The optimization of dihedrals is designed to best reproduce the QM potential energy surface (PES). The QM calculations use the MP2 level of theory and the 6-31G* basis set for the target data.

All the dihedrals are held fixed, while the one under consideration is restrained to a certain value and the energy calculated after a short minimization. The dihedral under consideration is then restrained to a new value, while still keeping the other dihedrals fixed, and the energy is again calculated after a short minimization. This is done for a set of several restrained values for the dihedral under consideration till a PES is mapped out. The short minimization carried out consists of 50 conjugate gradient algorithm steps and 100 Newton-Raphson algorithm steps. For example, the improper angle is first restrained to -45.0 degrees, the system minimized and energy calculated. The improper is angle is then restrained to -44.0 degrees, system minimized and energy calculated. This is done until a PES is obtained for the improper angle 'scan' in 1.0 degree increments from -45.0 degrees to +45.0 degrees. Similarly, I get the PES for all the dihedrals I intend to optimize, one at a time. The regular dihedral angles were restrained from -90.0 degrees to +90.0 degrees in 3.0 degree increments. Emphasis is placed on reproducing low energy regions and low barriers, since high barriers will not be crossed during typical room temperature MD simulations. I get a similar PES from QM calculations for the target data. Just like the other parameter optimization sections I try to reproduce the target data by adjusting the dihedral parameters.

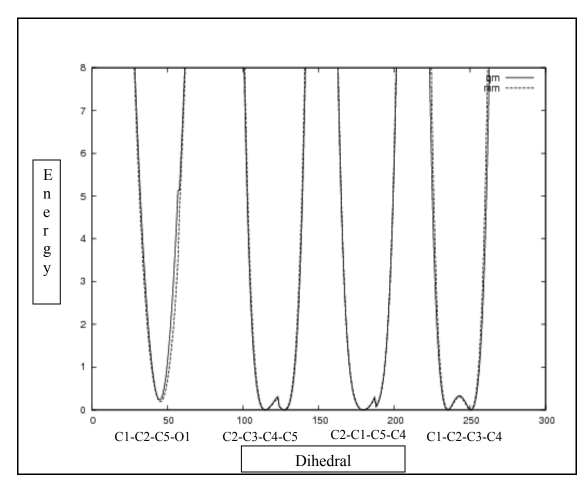


Figure 25: Dihedral potential energy

Topology a	nd para	meter fil	les for CP	D									
* Stream file with compound to be parameterized.													
*													
read rtf	card	apper	nd										
* Topolo	gy of	comp	ound to	b be j	parame	eter	ized.						
*													
36 1													
RESI CPI)		0.00										
GROUP													
ATOM C1													
ATOM 01													
ATOM C2													
ATOM H21													
ATOM H22													
ATOM C3 ATOM H31													
ATOM H31 ATOM H32													
ATOM H32 ATOM C4													
ATOM 04													
ATOM C5													
ATOM H51													
ATOM H52													
BOND C1	C2		C2	C3		C3	C4		C4	C5	C5	C1	
BOND C2	H21		C2	H22									
BOND C3	H31		C3	H32									
BOND C5	H51		C5	H52									
DOUBLE (1 C	1	C4	04									
IMPR C1	C2 C5	01											
IMPR C4	C5 C3	04											
ACCEPTOR													
ACCEPTOR	04												
IC C1	C2	C3	C4	0	.0000		0 00	0 0		0 00	0.0000		
IC C2	C3	C4	C5		.0000		0.00			0.00	0.0000		
IC C3	C4	C5	C1		.0000			0.0		0.00	0.0000		
IC C5	C2	*C1	01		.0000		0.00			0.00	0.0000		
IC C1	C3	*C2	H21		.0000		0.00	120.0		0.00	0.0000		
IC C1	C3	*C2	H22		.0000		0.00	240.0		0.00	0.0000		
IC C2	C4	*C3	H31	0	.0000		0.00	120.0		0.00	0.0000		
IC C2	C4	*C3	H32	0	.0000		0.00	240.0		0.00	0.0000		
IC C3	C5	*C4	04	0	.0000		0.00	120.0		0.00	0.0000		
IC C4	C1	*C5	H51	0	.0000		0.00	120.0		0.00	0.0000		
IC C4	C1	*C5	H52	0	.0000		0.00	240.0		0.00	0.0000		
PATC FI	RS NC	NE LA	ST NON	E									
END													
<u>. </u>													

Table 8: CHARMM Topology File for CPD

read param card flex append * Parameters needed for CPD BONDS CG3C52 HGA2 307.00 1.1000 CG2R53 CG3C52 285.00 1.5300 CG3C52 CG3C52 187.00 1.5300 CG2R53 OG2D1 691.00 1.2350 ANGLES HGA2 CG3C52 HGA2 39.50 106.00 5.40 1.802 CG2R53 CG3C52 HGA2 41.50 111.00 CG3C52 CG3C52 HGA2 36.00 111.40 22.53 2.179 CG2R53 CG3C52 CG3C52 50.00 106.50 CG3C52 CG2R53 CG3C52 93.00 109.50 CG2R53 CG3C52 CG2R53 75.00 106.50 CG3C52 CG2R53 OG2D1 70.00 122.00 DIHEDRALS CG2R53 CG3C52 CG3C52 CG2R53 0.0000 3 0.000 CG2R53 CG3C52 CG2R53 CG3C52 0.1000 3 0.000 CG3C52 CG3C52 CG2R53 CG3C52 0.000 3 0.000 OG2D1 CG2R53 CG3C52 CG2R53 0.0078 3 172.550 OG2D1 CG2R53 CG3C52 CG2R53 0.9000 1 0.000 OG2D1 CG2R53 CG3C52 CG3C52 0.0078 3 160.564 OG2D1 CG2R53 CG3C52 CG3C52 0.9000 1 0.000 OG2D1 CG2R53 CG3C52 HGA2 0.0000 3 0.000 OG2D1 CG2R53 CG3C52 HGA2 0.0000 1 0.000 CG3C52 CG2R53 CG3C52 HGA2 0.1000 3 0.000 CG2R53 CG3C52 CG3C52 HGA2 0.1000 3 0.000 IMPROPERS CG2R53 CG3C52 CG3C52 OG2D1 90.0000 0 0.000 END

Table 9: CHARMM Parameter File for CPD

3.2. HOCl



Figure 26: Hypochlorous acid (HOCl). Atom names are shown

Parameters for HOCl were also developed from the ground up, using the procedure described in detail for CPD. Developing parameters for HOCl was simpler than for CPD since HOCl has only three atoms and three atom types. Obviously, no dihedral optimization is needed and only two bond parameters and one angle parameter in addition to the partial atomic charges are needed. The van der Waals parameters are taken directly from similar atom types in the existing parameter file and not modified, as for CPD and as recommended. The finalized parameters are given in Appendix. Table 11 and Table 12 below compares QM and MM observables.

Atom Name	Atom Type	Description
CL	CLGA1	Chlorine e.g. Mono and dichloro ethane

OG	OG311	hydroxyl oxygen
HG1	HGP1	polar hydrogen

 Table 10: HOCl Atom Names and Atom Types

	QM (kcal/mol)	CHARMM (kcal/mol)	∆ (CHARMM – QM) (kcal/mol)
H - OHH	-9.6396	-6.61081	3.02879
O - HOH	-2.76436 -3.6361		-0.87174
CI - OHH	-2.6294	-0.588162	2.041238
		AVG Difference	1.39943
		RMS Difference	1.65579

Table 11: HOCl – H₂O Interaction Energies

	QM (kcal/mol)	CHARMM (kcal/mol)	∆ (CHARMM – QM) (kcal/mol)
H - OHH	1.86	1.86	0
O - HOH	1.97	1.97	0
CI - OHH	2.92	3.53	0.61

Table 12: HOCl – H₂O Interaction Distances

QM (D)	CHARMM (D)	
1.8253	1.96711	

Table 13: Dipole moments

*Topology file for HOCl
* Stream file with compound to be parameterized.
*
RESI HOCL 0.00
! GROUP
ATOM HG1 HGP1 0.38
ATOM OG 0G311 -0.49
ATOM CL CLGA1 0.11
BOND CL OG OG HG1
DONO HG1 OG
DONO CL OG
ACCE OG
PATC FIRS NONE LAST NONE
END

Table 14: CHARMM Topology File for HOCl

* read param card flex append				
* Parameters needed for HOCl				
*				
DONDO				
BONDS				
CLGA1 OG311	428.00	1.71739		
ANGLES				
CLGA1 OG311	HGP1 50.	.00 102.637		
END				
21.2				

Table 15: CHARMM Parameter File for HOCl

3.3. CPO sugars

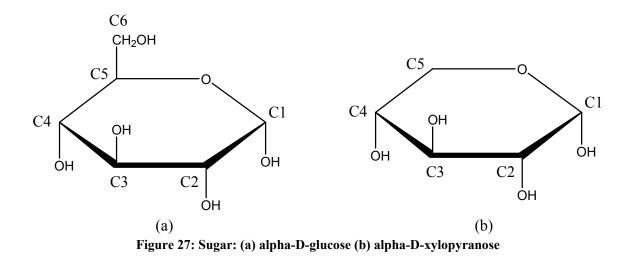
Sugar parameters were developed using the analogy method, for two reasons. First, the CHARMM force field contains parameters for some related sugars, and these parameters are expected to be largely transferable to the other sugars present in Chloroperoxidase. Second, the sugars themselves are located at the periphery of CPO, far from the active site where reaction takes place. Thus calculations of the reaction barrier for epoxidation at the active site or chlorination, if it occurs at the active site, are not expected to be very sensitive to the sugar parameters, and approximate parameters would be sufficient under such circumstances.

Application of the analogy method involves minor modification of atoms or atom types that would not significantly affect the energetic. This requires some chemical intuition. The process by which sugar parameters for addition to the default CHARMM topology file is described below.

3.3.1. Alpha-D-xylopyranose (AXYS)

Alpha-D-xylopyranose is similar to alpha-D-glucose (AGLC), which is already present in the CHARMM sugar topology file. All that is required to construct AXYS is to remove a small group of atoms from AGLC and replace them with hydrogen atoms. There are a few important details, so I describe the procedure carefully below.

I make a new entry for AXYS in the topology file. The AGLC section in the topology file is copied for AXYS, keeping only the atoms that are common to both molecules. Then the two extra hydrogen atoms are added to complete the AXYS molecule. The following atoms from AGLC are deleted: C6, H61, H62, O6, HO6. The atoms H51 and H52 are added. Partial atomic charges must be adjusted slightly so that the individual charges remain consistent and the overall charge is zero. In the process of adjusting charges, the aliphatic hydrogen atoms are given charges of ± 0.09 as recommended by MacKerell. Carbon atom C5 has its charge adjusted to ± 0.250 to accommodate overall charge neutrality.



Adjustments are made in the 'BOND' section of the topology file to provide information about which atoms are bonded. The donor ('DONO'), acceptor ('ACCE') and internal coordinate (IC) table sections are also modified. This is not absolutely necessary, though, because the information they provide can be included in a CHARMM input script if required.

3.3.2. Beta-L-xylopyranose (BXYS)

The molecule BXYS and AXYS are similar. Therefore all parameters for BXYS are the same as for AXYS; only the coordinates or stereochemistry are different. An α or β sugar are defined by the stereochemistry of the first carbon (C1) with respect to the last chiral carbon atom (C4). If they have the opposite stereochemistry the sugar is define as α and if they both have the same stereochemistry the sugar is defined as β .

We do not need to consider the orientation or the cartesian coordinates in the topology file (they are gotten from the Protein Data Bank (PDB) or a coordinate (CRD) file). Consequently, all that is needed is to create an entry for BXYS and copy all the parameters directly from AXYS.

3.3.3. Beta-L-arabinose(BARB)

Like BXYS, BARB is similar to AXYS. Essentially, only the coordinates differ between BARB and AXYS; the parameters are again identical. A new entry BARB is created with the AXYS parameters copied in.

3.3.4. N-Acetyl-Glucosamine (BNAG)

Like AXYS, BNAG is similar to a sugar already in the parameter set, alpha-Dglucose (AGLC). Thus parameterization of BNAG requires a small modification of AGLC parameters. However, in this case the hydroxyl group attached to C2 is replaced by –NHCOCH3 group. The new group is split in to three sub-groups NH, CO and CH3, each with a net partial charge of zero.

3.3.5. Pyroglutamic acid (PCA)

Pyroglutamic acid is an uncommon amino acid and is a terminal amino acid in CPO. It is similar to a more common amino acid, glutamic acid. The terminal hydroxyl is missing in PCA and the delta-carbon in glutamic acid is connected to the backbone nitrogen to form a ring structure in PCA.

3.4. Patches

Patches are tools used in CHARMM when individual residues or molecules are required to be connected to others. These connections can be made explicitly in the CHARMM input script, but if this needs to be done regularly and often, it is worth creating a patch to do so. A patch usually contains two residues that are to be connected. Often some atoms need to be deleted; bond, angle and perhaps dihedral terms need to be added; and in some cases partial charges of the atoms need to be adjusted to maintain the overall charge deletion of atoms.

In the macromolecule CPO there are several sugar molecules that are connected to each other and also to the enzyme. To accomplish this in the model, several patches were created in the topology file. They are to be invoked in the input script when the entire enzyme is built. We also need a patch to connect the heme to the proximal ligand cysteine (CYS). An example of one patch is given below, and the others are given in the Appendix.

PRES SSO4 0.000	! patch between MAN-MAN(C1-O4)
DELETE ATOM 101	
DELETE ATOM 1HO1	
DELETE ATOM 2HO4	
BOND 1C1 2O4	
GROU	
ATOM 1C1 CC3162	0 240
ATOM 1H1 HCA1	0.090
ATOM 1C5 CC3163	0.080
ATOM 1H5 HCA1	0.090
ATOM 105 0C3C61 -	-0.500
GROU	
ATOM 2C4 CC3161	0.360
ATOM 2H4 HCA1	0.090
ATOM 204 0C311 -	-0.450

Table 16: Patch to connect two mannose sugar molecules

3.5. Conclusions

In summary, I developed a set of molecular mechanics parameters for two molecules used in the study of the CPO halogenation mechanism and a set of sugars used in the study of the CPO epoxidation mechanism. My modeling work uses the CHARMM parameter set and therefore, to be consistent, I followed the protocol outlined by MacKerell et al. to develop parameters. I developed parameters for CPD and HOCl from the ground up because precise parameters were needed. The parameters developed were tested and met the requirements outlined in the protocol by MacKerell et al [72].

I used the less time-consuming analogy method to develop parameters for sugars. The sugar structures are very similar to other sugars already in the default CHARMM parameter set, and these sugars are not close to the enzyme active site and do not require a high level of accuracy. For these reasons I expect the analogy method to be satisfactory in this case. Patches were developed to create heme–cysteine, sugar–sugar and sugar– amino acid covalent bonds.

4. STUDY OF CPD DIKETO → KETO-ENOL TAUTOMERIZATION WITH LIMITED HYDRATION

4.1. Gas phase / Intramolecular proton transfer

I begin for reference with the intra-molecular hydrogen transfer process, in which a hydrogen is transferred from the alpha-carbon to one of the keto oxygens of CPD. As expected the adiabatic reaction barrier is quite high, 63.9 kcal/mol. Reactant, transition state and product structures are shown in Figure 28. Single-point calculations using the PCM method to model solvation implicitly do not alter the barrier heights significantly. The PCM barrier is 64.6 kcal/mol.

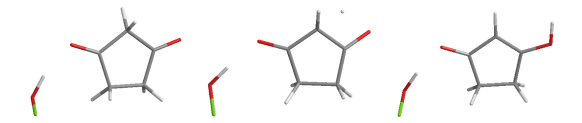


Figure 28: Intramolecular gas phase reaction. Stationary points. HOCl present to play stabilizing role representative of acidic conditions

4.2. Intermolecular proton transfer with one catalytic water molecule

When one catalytic water molecule is included, the reaction barrier decreases to 27.7 kcal/mol. Reactant, transition state and product structures are shown in Figure 29.

The water-catalyzed mechanism is a two-step process, with nearly complete abstraction of the alpha hydrogen by the water preceding transfer of a hydrogen from the water to the keto oxygen. As with intra-molecular hydrogen transfer, the implicit solvation model PCM method single-point calculation did not alter the barrier height significantly. The PCM barrier for the intermolecular hydrogen transfer is 26.8 kcal/mol.

The ketoenol product is slightly more stable that the diketo reactant in both the uncatalyzed and the single-water-catalyzed reactions (Table 17); PCM-modeled solvation increases the degree of product stabilization.

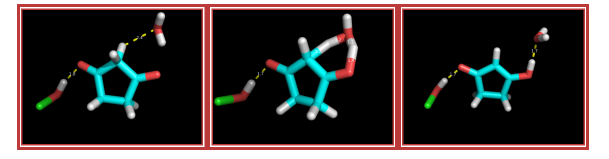


Figure 29: Intermolecular gas phase reaction. Stationary points. HOCl present to play stabilizing role representative of acidic conditions

4.3. Water clusters, fully QM optimized

Previous work on related systems has shown that additional explicit waters can lower the reaction barrier significantly [43-45]. Therefore I systematically evaluated the effect of additional hydration on the reaction barrier. To do so, I constructed clusters of water molecules surrounding the CPD/HOCl system and determined the adiabatic reaction barriers, treating the entire system at the QM level. Systems with multiple waters possess a large number of degrees of freedom and great flexibility. Therefore the potential energy surface (PES) has a high dimensionality and many local minima. Meaningful, interpretable results require careful placement of waters. My approach was to begin with the stationary points of the single-water system and add water molecules incrementally. The water molecules were placed in highly favorable positions for forming hydrogen bonds. For reactants, the waters were added incrementally until the hydrogen bonding positions around the keto oxygens, the HOCl and the catalytic water were saturated, that is, until these hydrophilic atoms had one full solvation shell of explicit waters. I attempted to maintain similar degrees of hydration of these three moieties as I increased the level of hydration. To increment the degree of hydration, one or two new waters were added to a previously minimized reactant structure, and this structure was fully minimized. The same procedure was applied to the corresponding product and transition state structures. With this procedure, trends in barrier heights and reaction energies exhibited a systematic relation to level of hydration. When incremental waters were not carefully placed, however, trends were not systematic and were difficult to interpret (data not shown).

The results are shown in Table 17 and Figure 34. The overall reaction energy, ΔE , decreases from -3.1 kcal/mol for a single catalytic water to -17.2 kcal/mol for eight waters. Eight explicit waters are sufficient to model the overall reaction energetics, as ΔE is nearly unchanged beyond eight waters. The reaction barrier exhibits a stronger dependence on level of hydration, and in a way that is critically dependent on the placement of incremental waters. The barrier height is 27.7 and 27.6 kcal/mol with one or three explicit waters, respectively. Waters two and three were placed so as to hydrate

the HOCl, having little effect on the reaction barrier. However, the barrier decreases rapidly as more explicit waters are added. Waters four and five were placed so as to hydrate the catalytic water, and thus had a substantial effect on the barrier. For systems with more than five water molecules, I was unable to fully optimize the transition states. This difficulty probably stems from the large number of highly flexible degrees of freedom and consequently quite flat PES in the vicinity of stationary points.

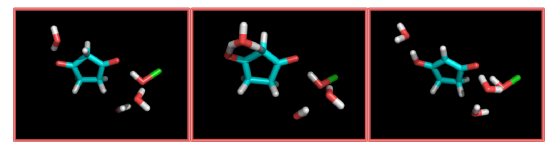


Figure 30: Stationary points of system with 3 explicit water molecules

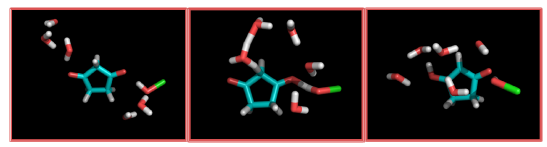


Figure 31: Stationary points of system with 5 explicit water molecules

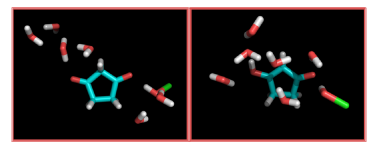


Figure 32: Stationary points of system with 6 explicit water molecules

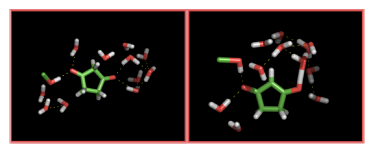


Figure 33: Stationary points of system with 11 explicit water molecules

For each stationary point at each hydration level, I also performed single-point energy calculations using the PCM method. The reaction energies changed by up to 8.3 kcal/mol. However, the barrier heights changed by no more than 1.6 kcal/mol. This finding is not consistent with the results described above, which demonstrate that adding explicit waters can significantly alter the barrier heights, at least for waters added to the first hydration shell of a hydrophilic moiety such as a keto oxygen. I conclude that an implicit water environment is not capable of accurately assessing the energetics of this reaction, in which specific directional solvent-solute interactions, and also solventsolvent interactions in the vicinity of the solute, are important. It represents the barrier for fully optimized structures.

Incremental explicit water molecules placed in the manner described significantly stabilize all structures (reactants, transition states and products) with respect to infinitely separated species. Stabilization is generally greatest for the transition state, with the result that the reaction barrier decreases with increasing level of hydration. With five explicit water molecules, the forward reaction (diketone to ketoenol) is significantly lower. To make a physical connection, such structures might be thought of fully optimized and stable conformation that one might expect to see in a protective channel in an enzyme such as the major channel of CPO. This stable network of water molecules is present in

the crystal structure reported by Kuehnel et. al. [38]. While this method does allow us to model a highly stable environment like the enzyme major channel of CPO, it is not representative of kinetics in solution at 300 K.

	in vacuo		PCM (ε=78.4)
# H ₂ O's	ΔE (kcal/mol)	$\Delta E^{TS,f}$ (kcal/mol)	ΔE (kcal/mol)	$\Delta E^{TS,f}$ (kcal/mol)
0	-1.6	63.9	-2	64.6
1	-3.1	27.7	-6.3	26.8
3	-5.1	27.6	-5.6	
5	-12.7	15.5(22)	-4.4	17.1
6	-11.6			
8	-17.2		-11.8	
10	-16.7			
11	-17.0			

Table 17: Reaction energies and barrier heights (kcal/mol) for reaction in water clusters

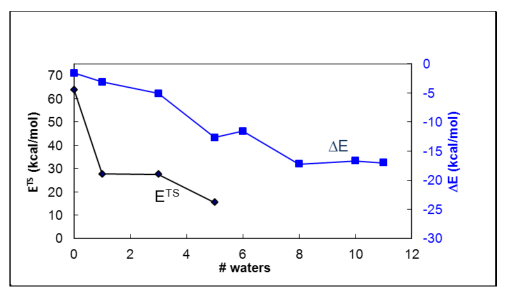


Figure 34: Reaction energies and barrier heights for reaction in water clusters

It is interesting to note that the water molecules significantly affect the barrier height only if they participate directly in a proton transfer, acting as a catalyst or if they help stabilize the catalytic water. For example, in the cluster with three waters only one water molecule is involved in the proton transfer and acts as a catalyst. The other two waters are only involved in hydration. The reaction barrier, 27.6 kcal/mol, is almost the same as for the single-water example (27.7 kcal/mol). When two more water molecules were added in close vicinity to the catalytic water the barrier height dropped significantly. The reaction barrier dropped from 27.7 kcal/mol to 15.5 kcal/mol.

I also used the IRC method [58, 59] to obtain the respective reactant and product structures from the transition states. The same reactant and product structures were obtained as before, except for the system with five water molecules. The IRC barrier is given in parentheses in Table 17 for the system with five water molecules.

4.4. Conclusions

The calculations demonstrate that the barrier height is quite sensitive to the particular pattern of an elaborate hydrogen bonded structure. Thus for reaction at the active site of CPO, it is likely that the placement of water molecules, and possibly also charged side chains, is critical for achieving the lowest barrier to reaction. The barrier height is observed to be most sensitive to the placement of water molecules near the first catalytic water and less sensitive to waters located elsewhere. No proton transfer is observed from HOCl to CPD as might have been expected to initiate or accompany abstraction of the alpha hydrogen as H^+ .

The energies of reaction leveled off as the number of waters increased beyond five, but the barrier heights did not. Thus previously reported theoretical calculations on keto-enol tautomerization, which involved 1-4 waters, are unlikely to have achieved reaction barriers representative of solution conditions, which was in most cases the stated goal. More waters, and probably a different approach altogether, are needed to realistically model the barriers in full solvation conditions.

The location of transition states with many water molecules present is very difficult due to the high dimensional, flat potential energy surface. Therefore extending this approach to larger clusters was not feasible. It is emphasized that these structures are what one might expect to see in a protective channel in an enzyme such as the major channel of CPO and unlikely to be representative of liquid water at room temperature. In order to determine the barrier height representative of full solvation conditions, I employed a different approach, which is described in the next chapter.

5. STUDY OF CPD DIKETO → KETO-ENOL TAUTOMERIZATION UNDER FULL SOLVATION CONDITIONS

To represent the solvated environment with explicit solvent molecules, I used Molecular Dynamics (MD) run at 300 K to obtain snapshots of reaction precursor structures. A QM/MM approach was then used to obtain an adiabatic reaction path and barrier height for each precursor structure. The replica path and Climbing Image Nudged Elastic Band (NEB) methods were used to determine the reaction paths.

5.1. MD simulation and selection of frames

For a reaction to occur in solution, the reactive molecules need to be favorably positioned and oriented with respect to each other. Such positioning, sometimes called a 'rare event' occurrence, is not likely to occur frequently on the nanosecond timescale. I therefore chose frames from the ten ns MD simulation that represent such 'rare event' occurrences. The MD provides the randomization of the water molecules present in a solvated environment.

When the HOCl and CPD approach each other in solution, they remain close for a duration on the order of a few ps, trapped by a solvent cage. I selected only one frame for each unique encounter, choosing frames at least ten ps apart. Ten ps represents the residence time for a hydrogen bond. Favorable conformations were selected such that the HOCl was near enough to CPD to form a hydrogen bond with one of the keto oxygens,

and a water molecule was positioned ready to accept a proton from the alpha carbon of CPD and to donate a proton to the other keto oxygen. The selection process thus used the following criteria: The distance between one of the keto oxygens of CPD and the hydrogen of HOCl is less than or equal to 2.4 Å (1). There exists a water molecule whose oxygen is within 2.4 Å of the alpha carbon of CPD (2) and one of whose hydrogens is within 3.2 Å of the keto oxygen not used in the first criterion (3). Statistics regarding the trajectory frames satisfying these criteria are shown in Figure 36. A representative selected structure is shown in Figure 35. From each unique encounter, the best frame was selected for further consideration. The best frame was taken as the one with the smallest value for criterion (1); in the case of a tie, the smallest value for criterion (2) was determining; and if a tie still remained, the smallest value for criterion (3) was decisive.

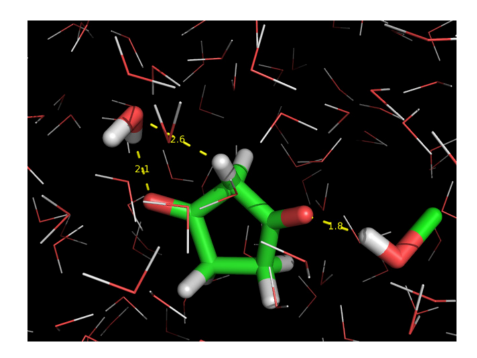


Figure 35: Representative rare event frame favorable for reaction, chosen from the MD simulation according to the selection criteria. Atoms treated at the QM level in subsequent QM/MM calculations are represented with thick sticks, and MM atoms are represented with thin lines

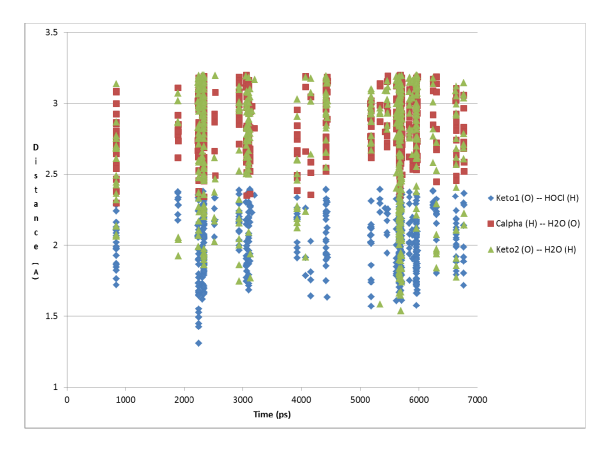


Figure 36: Trajectory frames satisfying the selection criteria. Each vertical group of data points represents one encounter, i.e., all frames lie within 10 ps of the best frame selected. Frames were saved every 20 steps (every 0.02 ps). 617 frames satisfied the selection criteria out of a total of 350,000 saved trajectory frames (0.18%)

5.2. Determination of minimum energy path

I determined the reaction barriers for the selected frames using a QM/MM model. I treated the CPD, HOCl and a single water molecule, which participates in the reaction catalytically, at the QM level and the rest of the water molecules at the MM level. All the MM waters were kept fixed while the QM atoms were flexible. Thus the barriers calculated are adiabatic with respect to the flexible degrees of freedom, in a fixed solvent environment representative of a liquid solution at 300 K. Physically, this represents a reaction that takes place much faster than solvent response. This is clearly not what happens in reality but provides a well-defined limiting behavior for analysis. Each frame represents a different solvent environment consistent with a temperature of 300 K. Boltzmann weighted averaging of barriers across a sufficient number of frames can provide an effective activation barrier for the limiting physical process considered. The MEP was determined with chain-of-replicas methods as described earlier.

5.2.1. Determining MEP using Replica Path Method

I used the Replica Path Method as the method of choice to map out the minimum energy path. In this method, a chain of conformations, or replicas, representing stages of reaction between reactants and products is created. The distance between pairs of conformations is defined using an RMSD metric. Simultaneous optimization of all replicas with a constraint to maintain spacing between the replicas yields an MEP. Optimization with the RPATh facility in CHARMM required 500 steps to achieve the convergence criterion. The convergence criterion was an average root mean square gradient (GRMS) value of less than 0.06 kcal/mol/ Å between successive 100-cycle iterations. A typical optimization is shown in Figure 38. For frame 1 Table 18 shows the energy level convergence criteria for every 100 optimization cycle, Table 19 shows the gradient convergence criteria for every 100 optimization cycle and Table 20 shows that the RMSD deviations are within limit (i.e. less than 20% RMSD deviation from neighboring replicas or images) to validate the parameters used in RPATH. For the

optimizations, the end points were fixed. Consequently the minima, which correspond to reactants and products, occur at intermediate replicas. The reaction path obtained with RPATh results in approximately equal spacing between replicas on the PES. Thus the transition state is not located with precision. To obtain an accurate transition state, I refined the reaction path using the nudged elastic band method with the climbing image option (CI-NEB).

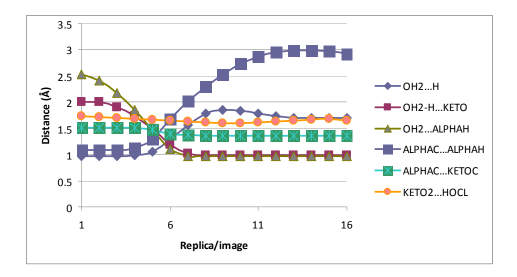


Figure 37: Frame 1: Distance change showing the hydrogen abstraction and enolization from reactant to product as the reaction proceeds in the vicinity of the transition state

	Optimization steps (SD + ABNR)				
Replicas/Images	50+100	50+200	50+300	50+400	50+500
1	21	21	21	21	21
2	2	2	2	1	1

3	3	2	2	2	2
4	4	4	6	9	7
5	9	10	19	<mark>28</mark>	<mark>23</mark>
6	20	26	<mark>30</mark>	18	22
7	<mark>35</mark>	<mark>27</mark>	12	5	6
8	21	12	5	2	2
9	11	6	2	1	1
10	7	4	2	1	1
11	5	3	1	1	1
12	3	3	1	1	1
13	2	2	1	1	1
14	1	1	0	0	0
15	0	0	0	0	0
16	15	15	15	15	15

 Table 18: Frame 1: Normalized energy level for each replica for every 100 ABNR optimization cycles. The maximum energy for each optimization cycle is highlighted

Optimization Steps (SD + ABNR)	Avg. RMS Gradient
50+100	2.094
50+200	0.388
50+300	0.165
50+400	0.121
50+500	0.065

Table 19: Frame 1: Average RMS Gradient for every 100 ABNR optimization steps

Replicas/Images	RMSD	%RMSD deviation
1	0.123	2.70%
2	0.122	3.70%
3	0.124	1.80%
4	0.13	3.00%
5	0.147	16.60%
6	0.13	3.30%
7	0.126	0.00%

8	0.125	1.00%
9	0.125	1.00%
10	0.125	0.80%
11	0.124	1.50%
12	0.123	2.40%
13	0.123	2.30%
14	0.123	2.80%
15	0.123	2.80%

Table 20: Frame 1: RMSD and % RMSD deviation from neighboring replicas

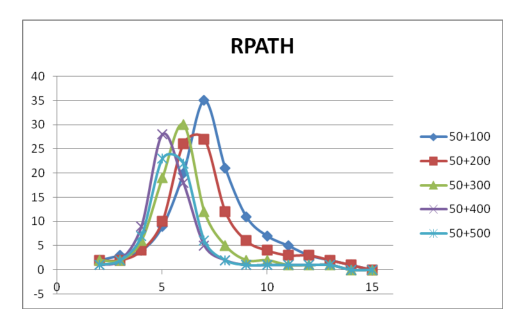


Figure 38: Optimization of Replica Path Method (SD+ABNR)

5.2.2. Locating transition state using Climbing Image Nudged Elastic Band

I use CI-NEB to locate the transition state once I have the MEP mapped out by the Replica Path Method. This option causes a path point near the transition state to climb in energy toward the transition state, while all other path points are energy minimized in directions orthogonal to the MEP. A typical optimization is shown in Figure 39. Further details are given in the supplemental information.

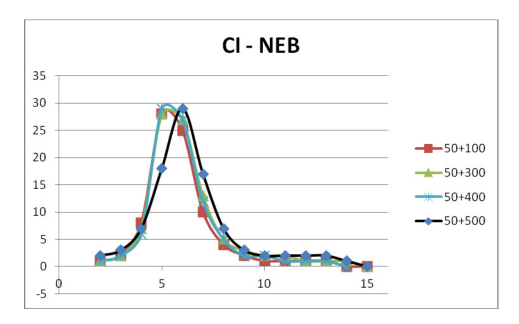


Figure 39: Optimization of CI-NEB to locate Transition state (SD+ABNR)

Table 21 gives the barrier height for each of the selected frames. There is a substantial variability in barrier height of 25-44 kcal/mol. Applying Boltzmann weighting to the barrier heights for the ten frames in Table 21, I obtain an effective activation energy (E_a) of 25 kcal/mol. Some important distances are tabulated in x, y, and table for reactant transition state and product respectively. To investigate the cause of the variation

in barrier heights, I visually analyzed each frame, focusing on hydrogen-bonding interactions likely to be important for reactants and transition states. These are summarized in Table 25.

Frame	ΔE
1	27
2	30
3	36
4	41
5	44
6	42
7	35
8	38
9	25
10	30

Table 21: Barrier heights (kcal/mol) for reaction for selected frames from solvated MD simulation

Fromo	OH2	OH2-H	OH2	ALPHAC	ALPHAC	KETO2
Frame	 H	KETO	 ALPHAH	 ALPHAH	KETOC	HOCL
1	0.972	1.995	2.379	1.093	1.512	1.721
2	0.968	2.823	3.201	1.100	1.522	1.792
3	0.974	2.107	2.490	1.099	1.521	1.675
4	0.974	2.104	2.675	1.095	1.522	1.942
5	0.979	2.385	3.165	1.098	1.501	2.321
6	0.972	2.651	2.915	1.096	1.523	1.786
7	0.973	2.201	2.517	1.092	1.511	2.024
8	0.97	3.42	2.22	1.10	1.52	1.72
9	0.98	3.15	2.20	1.10	1.52	1.88
10	0.972	1.796	2.247	1.096	1.526	1.714

Table 22: Reactant Distances for all the frames

Frame	OH2 H	OH2-H KETO	OH2 ALPHAH	ALPHAC ALPHAH	ALPHAC KETOC	KETO2 HOCL
1	1.11	1.38	1.22	1.47	1.44	1.64
2	1.13	1.39	1.28	1.53	1.44	1.64

3	1.14	1.34	1.23	1.49	1.44	1.71
4	1.24	1.23	1.27	1.47	1.44	1.95
5	1.15	1.41	1.26	1.66	1.41	3.01
6	1.2	1.26	1.28	1.47	1.44	1.65
7	1.18	1.3	1.22	1.47	1.43	1.92
8	1.11	1.42	1.23	1.54	1.44	1.73
9	1.02	1.67	1.19	1.52	1.44	1.77
10	1.16	1.33	1.27	1.5	1.45	1.66

Table 23: Transition state Distances for all the frames

Frame	OH2	OH2-H	OH2	ALPHAC	ALPHAC	KETO2
Tame	 H	KETO	 ALPHAH	ALPHAH	KETOC	HOCL
1	1.71	0.99	0.97	2.93	1.37	1.66
2	1.69	0.99	0.96	2.48	1.37	1.61
3	2.00	0.98	0.97	3.07	1.36	1.66
4	3.18	0.97	0.97	3.29	1.35	1.82
5	2.26	0.98	0.97	3.16	1.35	2.66

6	1.94	0.98	0.97	3.07	1.36	1.63
7	3.48	0.98	0.97	2.51	1.36	1.87
8	4.04	0.97	0.97	3.00	1.36	1.68
9	1.74	0.99	0.97	2.76	1.36	1.77
10	1.95	0.98	0.97	4.62	1.37	1.59

Table 24: Product Distances for all the frames

	Frame 1		Frame 9		Frame6	
	Reactant	TS	Reactant	TS	Reactant	TS
HBond	1	1	0	0	1	2
RMSD	0.07	0.21	0.38	0.30	0.29	0.52

 Table 25: Important hydrogen bonding interactions for reactants, transition states and products calculated for selected frames

To gain further insight into the cause of the variability in barrier heights, I recalculated the MEP, allowing nearby waters to respond to the progress of reaction. The MM water molecules within 2.6 Å of the QM atoms were made flexible, with the remaining MM waters still fixed. Distinguishing the treatment of water molecules in this way is easily achieved in the replica path implementation in CHARMM, as it allows for

the replicated system to be larger than the QM system. When this test was performed on one of the frames with a high barrier, frame 6, the barrier height decreased by 7 kcal/mol. The flexible MM water molecules underwent significant motion to stabilize the transition state structure, but little motion in the reactant structure, which was already solvent stabilized. When the test was performed on low-barrier structures, frames 1 and 9, there was little change in the barrier height.

The hydrogen bond analysis parameters had a cutoff distance of 4.0 Å and an angle of 150.0 degrees between acceptor and donor atoms. Only hydrogen bonds between the QM solvation shell and the CPD, HOCl and the catalytic water were analyzed.

The RMSD was calculated for the flexible solvation shell with respect to the fixed shell in the reactant and transition state structures for frames 1, 9 and 6.

Returning to the original reaction barrier calculations without the additional solvent flexibility, I can now trace the variability in barrier heights to the degree to which surrounding solvent waters stabilize the reactants. In no cases were transition states stabilized, but in some reactants were. In reality, it is unlikely that relatively massive solvent oxygens can move on the time scale of the proton transfer processes. Thus solvent stabilization of transition states is unlikely to occur, and the original set of barrier height calculations is more representative of the actual physical process than the calculations in which surrounding waters are allowed to move during the proton transfer processes.

5.3. Conclusions

I find that the solvating water environment creates a wide range of barriers for potentially reactive events. In my sampling of ten frames, these range from 25 to 44 kcal/mol. In view of Boltzmann weighting, the lowest of these are dominant and the effective barrier is 26 kcal/mol, which compares with the barrier of 28 kcal/mol for gasphase reaction with a single catalytic water. Thus, surprisingly, the barrier in a fully solvated system and the barrier with only a single water present are nearly the same. Close analysis reveals that this is not because other waters do not play a significant role in reaction. Rather it is because of a balancing effect. In low-barrier reaction events, which are mainly responsible for the observed rate constant, surrounding waters stabilize the transition state and the reactants to nearly the same extent. This is not true of most potentially reactive encounters; more often, surrounding waters stabilize reactants to a greater extent. Assuming that proton transfer occurs faster than water molecules can reorient, those same solvating waters are usually ill-positioned to stabilize the transition state and thus have a net effect of retarding rather than promoting reaction. In a small percentage of cases, though, the surrounding waters are positioned so as to lower the energies of both reactants and the transition state. Thus I find that transition state stabilization by solvent water is critical for reaction in solution. Transition state stabilization is also thought to be an important mode via which enzymes catalyze reactions. Thus it appears that enzymes and aqueous solution may have similar ways of catalyzing keto-enol tautomerization.

6. OVERALL CONCLUSIONS

The goal of this work was to elucidate the chlorination mechanism of chloroperoxidase using CPD as a model substrate, and to shed light on which of two possible reaction pathways is more likely: reaction with free HOCl in aqueous solution or reaction at the enzyme active site. I show that the rate limiting step for the chlorination of CPD is the diketo-ketoenol tautomerization step and that the solvent environment plays a crucial role in the reaction. I model the hydration environments of the two possible pathways and calculate the reaction barriers for both. I observe that a limited and optimized (active-site-like) hydration environment leads to a lower energetic barrier than the fully solvated model representative of the aqueous environment at room temperature. This result indicates that the stable water network near the active site and the access channel is likely to facilitate the chlorination mechanism. Confirmation of this predication could be obtained from calculations of the reaction barrier with the enzyme active site explicitly modeled, including active site residues and hydrating waters, using a QM/MM approach. This, however, is beyond the scope of the current project.

I determined the nature of the influence of the solvent environment on the reaction barrier. I observe that stabilization of the catalytic water by other solvent molecules is critical for keto-enol tautomerization. When studying a limited-hydration environment, placement of water molecules is more important than the number of water molecules. Careful placement of hydrating water molecules saves a significant amount of computational time in such studies. Previous theoretical limited-hydration studies have been accepted as a tractable method for determining effective reaction barriers in

solution. My work demonstrates that the limited-hydration approach is inadequate for determining effective reaction barriers for keto-enol tautomerization. In particular, I demonstrate that previous theoretical studies of keto-enol tautomerization in a limited-hydration environment, which used five or fewer waters, are not representative of a solution environment. My modeling of the reaction in a fully-solvating environment demonstrates that reaction proceeds when the instantaneous dynamical water environment is close to optimal for stabilizing the transition state.

7. OTHER SYSTEMS STUDIED/CONTRIBUTED TO

This chapter describes my contributions to three other projects carried out in the Chatfield research group. Section 7.1 describes work on dehaloperoxidase, an enzyme bearing similarity to chloroperoxidase, the subject of my main project. Section 7.2 contains the preliminary results of an MD study of the solution dynamics of a small, highly cross-linked peptide, α -conotoxin GI. Section 7.3 contains work done on the CPOcatalyzed epoxidation reaction. My major contribution to the studies described in sections 7.1 and 7.3 was the development of parameters for molecular mechanics modeling, including MD simulation. This work was a natural application of the expertise I gained in parameter development working on parameters for molecules involved in the CPOcatalyzed halogenation reaction, as explained in chapter 3. My contribution to the project described in section 7.1 also involved carrying out MD simulations using techniques and scripts I developed for my main project. The entire project described in section 7.2 was carried out primarily by me. This work in section 7.2 was a preliminary project carried out before beginning the CPO-catalyzed halogenation work that is focus of this dissertation. In carrying out the work described in section 7.2, I became familiar with molecular modeling techniques that I was able to apply to the other system studied. My contribution to each project is detailed in the subsequent sections.

7.1. DHP

I made contributions to an MD study of the enzyme dehaloperoxidase (DHP), which is found in the marine worm Amphitrite ornata. The enzyme has a globin fold and can bind dioxygen as can most globins, but its primary function is to catalyze the dehalogenation of p-halophenols and related compounds. Thus DHP can function either as a dehalogenase or as a globin. The mechanism via which the enzyme switches between the two modalities is of interest. An active-site residue, His55, which can adopt multiple conformations, has been thought to play a major role in the switching between modalities. Molecular dynamics simulations of wildtype DHP and two mutants, H55V, H55R, with and without a p-iodophenol ligand present, were performed.

My contribution to this project was development of parameters for the piodophenol ligand, much of the simulation setup, and the running of several of the native and mutated enzyme simulations. Project design and analysis of the results were carried out by other members of the research group. A very brief summary of the simulation method and analysis to date is described in the next paragraph.

A 10-ns simulation was performed and analyzed for each system. Hydrogen bonds between the ligand, Tyr38 and His55 were found to highly stabilize the ligand at the active site. The ligand was found to block access to the heme iron, for most of the 10ns simulation. When His55 was mutated to Val or Arg, the p-iodophenol ligand was no longer maintained in a position where it blocks access to the heme iron. The simulations also provided insight into the external binding site of the substrate.

7.2. α-Conotoxin GI

I began work on this project at the outset of my Ph.D. The project was framed as a joint MD/NMR study of motional motifs, with an ultimate goal of contributing to the development of MD-based methods for interpreting the results of dynamical NMR. For practical reasons relating to difficulties in obtaining the labeled peptide for the NMR portion of the study, the project was not completed.



Figure 40: Cone snail

The peptide α -conotoxin GI belongs to a family of small multiply cross-linked peptide toxins produced by marine mollusks known as cone snails. The slow-moving snails use the toxins to hunt their prey, which are usually fish. Its venom needs to act fast

in order to bring down its prey given the snails' limited mobility in comparison to its prey, which is one of the reasons its venom has evolved to be highly potent. Depending on the species, the conotoxins range in length from 13-30 residues, making them among the smallest of peptide toxins (snake venom toxins, by comparison, tend to be in the 70-residue range). Conotoxins act on the nervous system, serving to paralyze the snails' prey. For example, the α -conotoxins bind to the nicotinic acetylcholine receptor (nAChR), to Ca-gated voltage channels.

Conotoxins are notable for their specificity of binding. For example, the α conotoxins of one species of cone snail are often inactive against the nAChR of the prey
of another species of cone snail. The specificity of conotoxins are achieved through
conservation of a scaffold based on patterns of cross-linking combined with a remarkable
variability of loop composition. For this reason, conotoxins are of interest as possible
lead compounds for drug development. The α -conotoxins are of particular interest
because their binding properties may help to map out the nAChR, whose threedimensional structure is thus far unknown.

The α -conotoxin GI was one of the first-isolated conotoxins and has been widely studied. Its sequence is ECCNPACGRHYSC, and there are disulfide bonds between residues 2 and 7, and 3 and 13. The three-dimensional structure has been characterized by X-ray crystallography and solution-state NMR. Two regions of secondary structure have been identified. In all structures, residues 4-8 have been described as 3-10 helical. Residues 9-13 have been described as a β -turn. In the case of the X-ray structure, this was on the basis of both backbone dihedral angles and hydrogen bonds. For the NMR structures, secondary structure assignment was based mainly on dihedral angles.

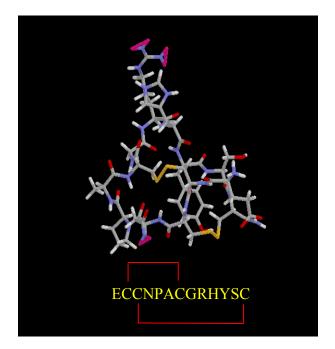


Figure 41: α-conotoxin GI

Notably, a recent NMR study identified two distinct conformers present in a 70:30 ratio (referred to hereafter as the major and minor conformers) on the basis of simulated annealing with the NMR structural restraints. It may be important to understand whether the conformers interconvert and whether one or both is active in binding to nAChR.

Multiple molecular dynamics simulations were performed on a 13-residue, doubly cross-linked peptide toxin, α -conotoxin GI, in each of two solution conformations identified by NMR. The two conformers remained distinct on the 100-ns simulation time scale, supporting the hypothesis that only one of the conformers is biologically active. The secondary structure as judged by backbone dihedral angles was preserved during simulation, but backbone hydrogen bonds were almost entirely absent. This observation suggests that binding at the active site probably involves hydrogen bonds to the peptide

backbone. The backbone exhibited unusual, concerted motions of dihedral angles in nonconsecutive residues, which may have implications for the binding free energy.

Below is a set of figures that bring out the salient features of the simulation. Figure 42 shows overlays of the several structures obtained from NMR structural data on peptide in solution using simulated annealing. The backbone and most sided chains have well defined conformations, but several side chains are clearly mobile. From this structure, two primary backbone conformations were determined. These are shown in Figure 43 and are called the major and minor structures. Figure 44 depicts the root mean square deviation (RMSD) in the backbone coordinates for ten 10-ns simulations of each structure (major and minor). The RMSD for both structures are shown together on each panel. It can be seen that the structures remain distinct on the 10-100 ns time scale.

Figure 45 depicts correlation functions, $C_i(t)$, for motion of backbone N-H bond vectors for all residues of each of the two structures. The correlation functions are defined by

$C_t(t) = \langle P_2(\hat{u}_t(0) \cdot (\hat{u}_t(t)) \rangle$

where $P_2 = \frac{8}{2}x^2 - \frac{1}{2}$ is a second Legendre polynomial and \hat{u}_i is a unit vector in the direction of the N-H bond vector. The asymptotic value of $C_i(t)$ as time gets large is the order parameter, S². Small values of S² correspond to relatively unrestricted motion, and values of S² close to one correspond to highly restricted mobility. The two sets of data (major and minor) have very different asymptotic values, but this is because the data were treated differently. The trajectory frames for the major structures were reoriented

before calculating the correlation functions, while those for the minor structures were not. When the trajectory frames for the minor structures are reoriented to remove overall translation and rotation (data not shown), the asymptotic behavior of the correlation functions is very similar to what is seen for the major structures in Figure 45. Values of S^2 by residue are depicted with a color code for both structures and also for the crystal structure (obtained from X-ray B factors) in Figure 46. The crystal structure shows generally more motional restriction (larger average value of S^2) as a consequence of crystal packing; the simulation environment is aqueous solution.

Maps showing the pattern of N-H backbone bond vector motion are shown in Figure 47. These are a major finding of the work. The N-H bond vectors display differing modes of motion, with some exhibiting motion within a cone, others elongated mobility, and yet others bimodal sampling. This indicates that interpretation of NMRderived order parameters in terms of motional models must be made with care.

Finally, Figure 48 shows that some backbone dihedrals angles exhibit correlated motions. When interpreting order parameters in terms of conformational entropy, this correlated motion will need to be taken into account.

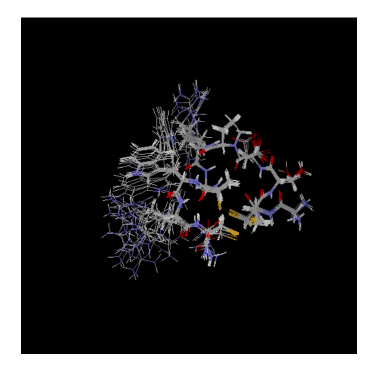


Figure 42: NMR structure showing stable backbone

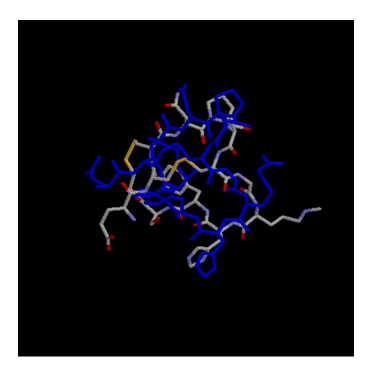


Figure 43: distinct structures in solution – Major and Minor

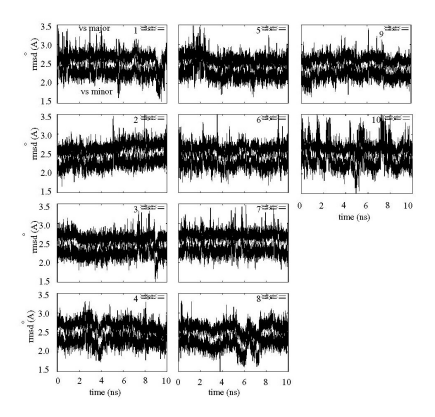
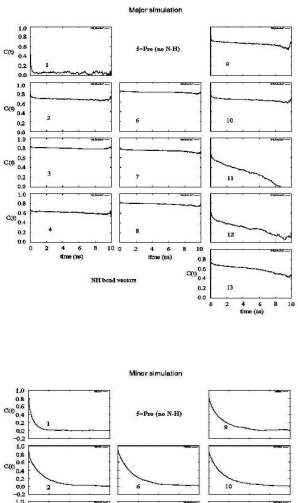


Figure 44: Simulation of major and minor conformation for 10x10 ns simulation demonstrating that the two structures are stable and distinct



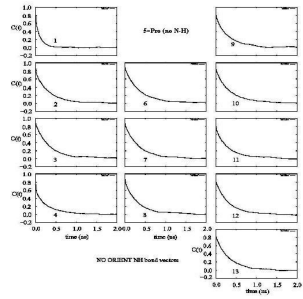


Figure 45: N-H correlation functions. Trajectory frames were reoriented for Major conformation but not for Minor conformation (S² values shown elsewhere are based on reoriented data.)

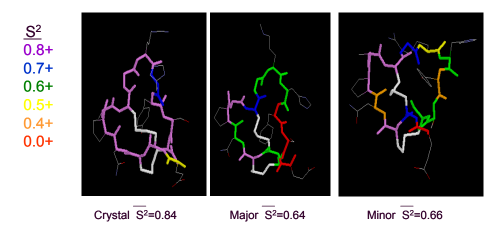


Figure 46: Backbone NH order parameter

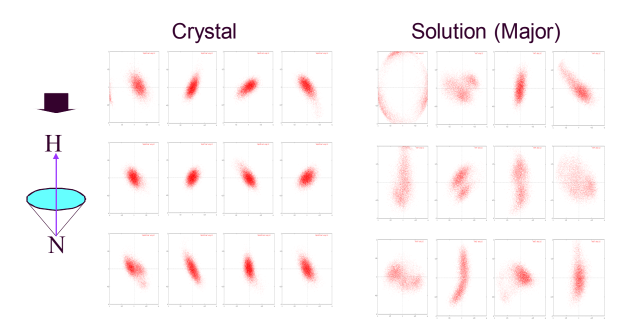
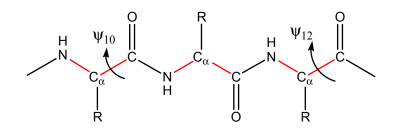


Figure 47: NH bond vector orientation and sampling



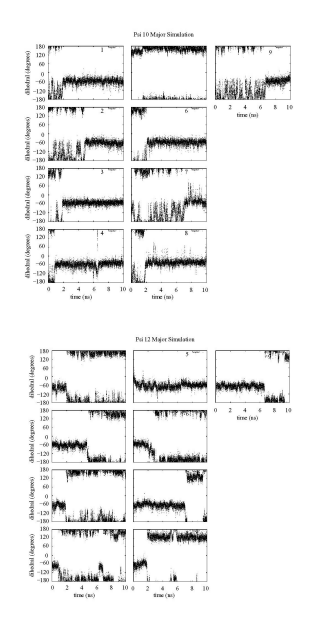


Figure 48: Dihedral (Ψ) of residue 10 and 12 of the major conformation

7.3. Chloroperoxidase epoxidation

I made contributions to an MD study of CPO aimed at elucidating the mechanism of the epoxidation reaction, which takes place at the active site. My contribution to the project was development of parameters for the sugar molecules and help with development of parameters for the substrate, cis- β -methylstyrene. In addition, I created patches to connect the heme to the proximal ligand, cysteine, and to connect sugar molecules to each other and to amino acids with covalent bonds. Project design and analysis of the results were carried out by other members of the research group, primarily Alexander Morozov. Below is the abstract from a paper published on this work [74]:

Molecular dynamics simulations of an explicitly solvated cis- β methylstyrene /chloroperoxidase-Compound I complex are performed to determine the cause of the high enantiospecificity of epoxidation. From the simulations, a two-dimensional free energy potential is calculated to distinguish binding potential wells from which reaction to 1S2R and 1R2S epoxide products may occur. Convergence of the free energy potential is accelerated with an adaptive biasing potential. Analysis of binding is followed by analysis of 1S2R and 1R2S reaction precursor structures in which the substrate, having left the binding wells, places its reactive double bond in steric proximity to the oxyferryl heme center. Structural analysis of binding and reaction precursor conformations is presented. We find that i) a distortion of Glu183 is important for CPO-catalyzed epoxidation as was postulated previously based on experimental results; ii) the free energy of binding does not provide significant differentiation between structures leading to the respective epoxide enantiomers; iii) CPO's enantiospecificity toward cis-β-methylstyrene is likely to be caused by a specific group of residues which form a hydrophobic core surrounding the oxyferryl heme center.

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APPENDICES

Appendix A: Computer scripts

A1 Scripts used in the study of limited-hydration CPD keto - enol tautomerization

# 1	B3LYI	P/6-31	G* opt						
Opt	timi:	zation	of read	ctant	structure	(CPD+	HOCl+1H20)	
0	1								
С									
С	1	cc2							
С	2	cc3	1	ccc3					
С	3	cc4	2	ccc4	1	dih4			
С	4	cc5	3	ccc5	2	dih5			
0	5	006	4	0006		dih6			
0	4	oc7		occ7	2	dih7			
h	4	hc8		hcc8		dih8			
h		hc9		hcc9		dih9			
h		hc10		hcc10		dih10			
h		hc11		hcc11		dih11			
h		hc12		hcc12		dih12			
h		hc13		hcc1		dih13			
h		ho14		hoc14		dih14			
h		ho15		hocl		dih15			
0		oc16		occle		dih16			
h		ho17		hoc1'		dih17			
0		oh18		ohol		dih18			
cl	18	clo19	17	cloh	19 16	dih19)		
CC			1.545410						
CC			1.50280	5					
CC			105.470						
CC4			1.458564	4					
CCO			112.500						
dil			9.623	-					
CC:			1.553092	2					
CCO			103.369						
dil			-20.776	2					
OC			1.197888	5					
oco dil			124.984 158.289						
oc'			3.01253	7					
00			106.729	,					
dil			62.621						
UII			02.021						

hc8	1.117716
hcc8	107.667
dih8	85.649
hc9	1.091939
hcc9	116.089
dih9	-151.414
hc10	1.092588
hcc10	113.544
dih10	127.397
hc11	1.095078
hcc11	111.498
dih11	-110.412
hc12	1.096916
hcc12	109.711
dih12	133.326
hc13	1.099154
hcc13	108.637
dih13	-111.324
ho14	0.970819
hoc14	109.561
dih14	-160.888
ho15	0.970299
hoc15	136.088
dih15	60.908
oc16	1.220000
occ16	120.000
dih16	180.000
ho17	1.500000
hoc17	179.000
dih17	180.000
oh18	0.950000
oho18	
	179.000
dih18	180.000
clo19	1.700000
cloh19	109.471
dih19	180.000

Table 26: Optimization script for reactant (or product) structure of CPD + HOCl + explicit water molecule

```
# B3LYP/6-31G* opt=ModRedundant
Scan to locate structure close to transition state
0 1
 С
     1 cc2
 С
     2 cc3
                  1 ccc3
 С
                                 1 dih4
     3 cc4
                  2 ccc4
 С
                                 2 dih5
                  3 ccc5
С
     4 cc5
                                 1 dih6
 0
     3 0C6
                  2 occ6
                  4 occ7
                                 3 dih7
 0
     5 oc7
```

o 4 oc8	3 occ8	2 dih8
h 4 hc9	3 hcc9	2 dih9
h 4 hc10	3 hcc10	2 dih10
h 1 hc11	2 hcc11	3 dih11
h 1 hc12	2 hcc12	3 dih12
h 2 hc13	3 hcc13	4 dih13
h 2 hc14	3 hcc14	4 dih14
h 6 ho15	3 hoc15	2 dih15
h 8 ho16	4 hoc16	3 dih16
h 8 ho17	4 hoc17	3 dih17
o 15 oh18	6 oho18	3 dih18
cl 18 clo19		6 dih19
cc2	1.544633	
cc3	1.508992	
ccc3	104.626	
cc4	1.423838	
ccc4	112.514	
dih4	15.495	
cc5	1.514990	
ccc5	106.402	
dih5	-10.465	
0C6	1.280959	
0006	124.332	
dih6 -	164.197	
oc7	1.211892	
occ7	124.205	
dih7	174.516	
oc8	2.559776	
0CC8	123.858	
dih8	87.988	
hc9	1.282261	
hcc9	115.146	
dih9	94.779	
hc10	1.093171	
hcc10	117.758	
dih10 -	147.959	
hc11	1.096993	
hcc11	111.438	
dih11	103.568	
hc12	1.092699	
hcc12	114.102	
dih12 -	135.021	
hc13	1.094783	
hcc13	110.746	
dih13	139.816	
hc14	1.099688	
hcc14	108.489	
dih14 -	103.953	
ho15	1.700000	
hoc15	114.173	
dih15	1.528	
ho16	0.979979	
hoc16	92.849	
dih16 -	143.860	

ho17	0.974744
hoc17	119.979
dih17	104.314
oh18	0.950000
oho18	179.000
dih18	180.000
clo19	1.700000
cloh19	109.471
dih19	180.000
8 9 1.32 S !	5 -0.1
6 15 1.72 S	5 -0.1

Table 27: Script for scan of potential energy surface to locate structure close to the transition state

```
# B3LYP/6-31G* opt=(TS,CalcFC) IOp(1/8=10)
Transition state optimization
  1
0
 С
     1 cc2
 С
 С
     2 cc3
                  1 ccc3
                                 1 dih4
 С
     3 cc4
                  2 ccc4
                                 2 dih5
                 3 ccc5
 С
     4 cc5
                 3 occ6
                                 2 dih6
 0
     4 oc6
     1 oc7
                 2 occ7
                                 3 dih7
 0
                                 3 dih8
     6 008
                 4 0008
 0
                                 4 dih9
cl
     8 clo9
                 6 cloo9
                                 3 dih10
 0
    5 oc10
                  4 occ10
   10 ho11
                  5 hoc11
                                 4 dih11
h
h
    5 hc12
                  4 hcc12
                                 3 dih12
h
     2 hc13
                 3 hcc13
                                 4 dih13
     2 hc14
                 3 hcc14
                                 4 dih14
h
     3 hc15
                  2 hcc15
                                 1 dih15
h
                                 1 dih16
h
     3 hc16
                  2 hcc16
h
     6 ho17
                  4 hoc17
                                 3 dih17
                                 4 dih18
h
   10 ho18
                  5 hoc18
                                 4 dih19
h
   10 ho19
                  5 hoc19
cc2
            1.524179
cc3
            1.542551
            104.650
ccc3
cc4
            1.527352
ccc4
            104.750
dih4
            -18.017
cc5
            1.453226
ccc5
            110.078
dih5
            17.393
            1.242614
oc6
occ6
            124.139
```

Cullob -162.499 cc7 1.25.202 dih7 -158.458 co8 2.330560 coc8 110.574 dih8 3.363 clo9 1.725564 clo9 104.618 dih9 103.668 occ10 2.501081 occc10 131.864 dih10 89.754 hol1 1.220000 hoc11 14.063 dih11 30.876 hcc12 1.093236 hcc12 1.093236 hcc13 1.098553 hcc13 1.098553 hcc14 1.979 dih13 98.138 hcc14 1.093351 hcc15 1.091907 hcc15 1.091907 hcc16 11.551 dih16 98.877 hc16 1.092098 hcc16 113.730 dih16 98.071 hc17 1.220000 hc217 13.730 dih18 -136.840 hc18<	dih6	-162.496	
occ7125.202dih7-158.458 $oo8$ 2.330560 $ooc8$ 110.574dih83.363clo91.725564clo09104.618dih9103.668 $occ10$ 2.501081 $occ10$ 131.864dih1089.754hol11.220000hoc1114.063dih1130.876hcc12116.890dih1130.876hcc131.098553hcc131.098553hcc141.5038dih15-139.992hc151.091907hcc15115.884dih1698.877hol71.220000hcc17113.730dih173.751hol81.002126hoc1880.741dih18-136.840hol90.973841hoc191.2.608			
$\begin{array}{llllllllllllllllllllllllllllllllllll$			
co8 2.330560 coc8 110.574 dih8 3.363 clo9 1.725564 cloo9 104.618 dih9 103.668 oc10 2.501081 occ10 131.864 dih10 89.754 ho11 1.220000 hoc11 1.093236 hcc12 1.093236 hcc13 1.098553 hcc13 1.098553 hcc13 1.098553 hcc14 15.038 dih14 -139.992 hc15 1.091907 hcc16 1.15.884 dih15 -139.659 hc16 1.098098 hcc16 111.551 dih16 98.877 ho17 1.220000 hcc17 113.730 dih17 3.751 ho18 1.002126 hoc18 80.741 dih18 -136.840 ho19 0.73841			
ooc8 110.574 dih8 3.363 clo9 1.725564 clo09 104.618 dih9 103.668 oc10 2.501081 occ10 131.864 dih10 89.754 hol1 1.220000 hoc11 14.063 dih11 30.876 hc12 1.093236 hcc12 116.890 dih12 -145.272 hc13 1.098553 hcc13 110.979 dih13 98.138 hc14 1.093351 hcc14 115.038 dih14 -139.992 hc15 1.091907 hcc16 11.551 dih15 -139.659 hc16 1.098098 hc16 1.098098 hc17 1.220000 hc217 113.730 dih17 3.751 ho18 1.002126 hoc18 80.741 dih18			
dih8 3.363 clo9 1.725564 clo9 104.618 dih9 103.668 oc10 2.501081 occ10 131.864 dih10 89.754 hol1 1.220000 hoc11 14.063 dih11 30.876 hc12 1.093236 hcc12 116.890 dih12 -145.272 hc13 1.098553 hcc13 110.979 dih13 98.138 hc14 1.093351 hcc15 1.5038 dih14 -139.992 hc15 1.091907 hcc15 115.884 dih15 -139.659 hc16 1.098098 hcc16 11.551 dih16 98.877 ho17 1.220000 hc17 13.730 dih18 1.002126 hce18 80.741 dih18 -136.840 ho19 0.973841 hc19 1.2.608			
clo9 1.725564 cloo9 104.618 dih9 103.668 oc10 2.501081 occ10 131.864 dih10 89.754 hol1 1.220000 hoc11 14.063 dih11 30.876 hc12 1.093236 hcc12 116.890 dih12 -145.272 hc13 1.098553 hcc13 110.979 dih13 98.138 hc14 1.093351 hcc14 115.038 dih14 -139.992 hc15 1.091907 hcc15 115.884 dih15 -139.659 hcc16 111.551 dih16 98.877 ho17 1.220000 hoc17 113.730 dih17 3.751 ho18 1.002126 hcc18 80.741 dih18 -136.840 hc19 0.973841 hoc19 112.608			
$ \begin{array}{c} cloo9 & 104.618 \\ dih9 & 103.668 \\ oc10 & 2.501081 \\ occ10 & 131.864 \\ dih10 & 89.754 \\ hol1 & 1.220000 \\ hoc11 & 14.063 \\ dih11 & 30.876 \\ hc12 & 1.093236 \\ hcc12 & 116.890 \\ dih12 & -145.272 \\ hc13 & 1.098553 \\ hcc13 & 110.979 \\ dih13 & 98.138 \\ hc14 & 1.093351 \\ hcc14 & 115.038 \\ dih14 & -139.992 \\ hc15 & 1.091907 \\ hcc15 & 115.884 \\ dih15 & -139.659 \\ hc16 & 1.098098 \\ hcc16 & 111.551 \\ dih16 & 98.877 \\ hol7 & 1.220000 \\ hoc17 & 113.730 \\ dih17 & 3.751 \\ hol8 & 1.002126 \\ hoc18 & 80.741 \\ dih18 & -136.840 \\ hc19 & 0.973841 \\ hoc19 & 112.608 \\ \end{array} $			
$ \begin{array}{llllllllllllllllllllllllllllllllllll$			
oc102.501081 $occ10$ 131.864dih1089.754hol11.220000hoc1114.063dih1130.876hc121.093236hcc12116.890dih12-145.272hc131.098553hcc14110.979dih1398.138hc141.093351hcc151.091907hcc151.5.884dih15-139.659hc161.098098hcc16111.551dih1698.877hol71.220000hoc17113.730dih173.751hol81.002126hoc1880.741dih18-136.840hoc19112.608			
occ10131.864dih10 89.754 hol1 1.220000 hocl1 14.063 dih11 30.876 hcl2 1.093236 hcc12 116.890 dih12 -145.272 hc13 1.098553 hcc13 110.979 dih13 98.138 hc14 1.093351 hcc14 115.038 dih14 -139.992 hc15 1.091907 hcc16 11.9869 hc16 1.098098 hcc16 11.551 dih16 98.877 ho17 1.220000 ho217 113.730 dih17 3.751 ho18 1.002126 hoc18 80.741 dih18 -136.840 ho19 0.973841 hoc19 112.608			
$\begin{array}{llllllllllllllllllllllllllllllllllll$			
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hocl1 14.063 dih11 30.876 hcl2 1.093236 hcc12 116.890 dih12 -145.272 hc13 1.098553 hcc13 110.979 dih13 98.138 hc14 1.093351 hcc14 115.038 dih14 -139.992 hc15 1.091907 hcc15 115.884 dih15 -139.659 hc16 1.098098 hcc16 11.551 dih16 98.877 ho17 1.220000 hoc17 113.730 dih17 3.751 ho18 1.002126 hoc18 80.741 dih18 -136.840 ho19 0.973841 hoc19 112.608			
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hc12 1.093236 hcc12 116.890 dih12 -145.272 hc13 1.098553 hcc13 110.979 dih13 98.138 hc14 1.093351 hcc14 115.038 dih14 -139.992 hc15 1.091907 hcc15 115.884 dih15 -139.659 hc16 1.098098 hcc16 111.551 dih16 98.877 ho17 1.220000 hcc17 113.730 dih17 3.751 ho18 1.002126 hoc18 80.741 dih18 -136.840 ho19 0.973841 hoc19 112.608			
hcc12 116.890 dih12 -145.272 hc13 1.098553 hcc13 110.979 dih13 98.138 hc14 1.093351 hcc14 115.038 dih14 -139.992 hc15 1.091907 hcc16 115.884 dih15 -139.659 hc16 1.098098 hcc16 111.551 dih16 98.877 ho17 1.220000 hoc17 113.730 dih17 3.751 ho18 1.002126 hoc18 80.741 dih18 -136.840 ho19 0.973841 hoc19 112.608			
$\begin{array}{llllllllllllllllllllllllllllllllllll$			
hc13 1.098553 hcc13 110.979 dih13 98.138 hc14 1.093351 hcc14 115.038 dih14 -139.992 hc15 1.091907 hcc15 115.884 dih15 -139.659 hc16 1.098098 hcc16 111.551 dih16 98.877 ho17 1.220000 hcc17 113.730 dih17 3.751 ho18 1.002126 hoc18 80.741 dih18 -136.840 ho19 0.973841 hoc19 112.608			
hcc13 110.979 dih13 98.138 hc14 1.093351 hcc14 115.038 dih14 -139.992 hc15 1.091907 hcc15 115.884 dih15 -139.659 hc16 1.098098 hcc16 111.551 dih16 98.877 ho17 1.220000 hcc17 113.730 dih17 3.751 ho18 1.002126 hoc18 80.741 dih18 -136.840 ho19 0.973841 hoc19 112.608			
dih1398.138hc141.093351hcc14115.038dih14-139.992hc151.091907hcc15115.884dih15-139.659hc161.098098hcc16111.551dih1698.877hc171.220000hcc17113.730dih173.751hc181.002126hcc1880.741dih18-136.840hc190.973841hcc19112.608			
hc14 1.093351 hcc14 115.038 dih14 -139.992 hc15 1.091907 hcc15 115.884 dih15 -139.659 hc16 1.098098 hcc16 111.551 dih16 98.877 ho17 1.220000 hoc17 113.730 dih17 3.751 ho18 1.002126 hoc18 80.741 dih18 -136.840 ho19 0.973841 hoc19 112.608			
hcc14 115.038 dih14 -139.992 hc15 1.091907 hcc15 115.884 dih15 -139.659 hc16 1.098098 hcc16 111.551 dih16 98.877 ho17 1.220000 hoc17 113.730 dih17 3.751 ho18 1.002126 hoc18 80.741 dih18 -136.840 ho19 0.973841 hoc19 112.608			
dih14-139.992hc151.091907hcc15115.884dih15-139.659hc161.098098hcc16111.551dih1698.877ho171.220000hcc17113.730dih173.751ho181.002126hoc1880.741dih18-136.840ho190.973841hoc19112.608			
hc15 1.091907 hcc15 115.884 dih15 -139.659 hc16 1.098098 hcc16 111.551 dih16 98.877 ho17 1.220000 hoc17 113.730 dih17 3.751 ho18 1.002126 hoc18 80.741 dih18 -136.840 ho19 0.973841 hoc19 112.608			
dih15 -139.659 hc16 1.098098 hcc16 111.551 dih16 98.877 ho17 1.220000 hoc17 113.730 dih17 3.751 ho18 1.002126 hoc18 80.741 dih18 -136.840 ho19 0.973841 hoc19 112.608		1.091907	
dih15 -139.659 hc16 1.098098 hcc16 111.551 dih16 98.877 ho17 1.220000 hoc17 113.730 dih17 3.751 ho18 1.002126 hoc18 80.741 dih18 -136.840 ho19 0.973841 hoc19 112.608	hcc15	115.884	
hc16 1.098098 hcc16 111.551 dih16 98.877 ho17 1.220000 hoc17 113.730 dih17 3.751 ho18 1.002126 hoc18 80.741 dih18 -136.840 ho19 0.973841 hoc19 112.608			
hcc16 111.551 dih16 98.877 ho17 1.220000 hoc17 113.730 dih17 3.751 ho18 1.002126 hoc18 80.741 dih18 -136.840 ho19 0.973841 hoc19 112.608		1.098098	
ho171.220000hoc17113.730dih173.751ho181.002126hoc1880.741dih18-136.840ho190.973841hoc19112.608	hcc16		
hoc17113.730dih173.751ho181.002126hoc1880.741dih18-136.840ho190.973841hoc19112.608	dih16	98.877	
dih173.751ho181.002126hoc1880.741dih18-136.840ho190.973841hoc19112.608	ho17	1.220000	
ho181.002126hoc1880.741dih18-136.840ho190.973841hoc19112.608	hoc17	113.730	
hoc1880.741dih18-136.840ho190.973841hoc19112.608	dih17	3.751	
dih18 -136.840 ho19 0.973841 hoc19 112.608	ho18	1.002126	
ho19 0.973841 hoc19 112.608	hoc18	80.741	
hoc19 112.608	dih18	-136.840	
	ho19	0.973841	
dih19 117.714	hoc19	112.608	
	dih19	117.714	

Table 28: Transition state optimization script

```
# B3LYP/6-31G* SCRF=PCM
Optimization of reactant/product
0 1
c
c
c 1 cc2
c 2 cc3 1 ccc3
c 3 cc4 2 ccc4 1 dih4
```

c 4 cc5 3 ccc5 2 dih5	
o 2 oc6 3 occ6 4 dih6	
o 5 oc7 4 occ7 3 dih7	
o 7 008 5 00C8 4 dih8	
cl 8 clo9 7 cloo9 5 dih9	
o 6 0010 2 00c10 3 dih10	
h 1 hc11 2 hcc11 3 dih11	
h 1 hc12 2 hcc12 3 dih12	
h 3 hc13 2 hcc13 1 dih13	
h 3 hc14 2 hcc14 1 dih14	
h 4 hc15 3 hcc15 2 dih15	
h 4 hc16 3 hcc16 2 dih16	
h 10 ho17 6 hoo17 2 dih17	
h 10 ho18 6 hoo18 2 dih18	
h 8 ho19 7 hoo19 5 dih19	
cc2 1.522321	
cc3 1.532052	
ccc3 110.030	
cc4 1.538389	
ccc4 106.565	
dih4 -1.736	
cc5 1.528754	
ccc5 106.324	
dih5 9.117	
oc6 1.215396	
occ6 124.391	
dih6 179.531	
oc7 1.219052	
occ7 125.165	
dih7 167.039	
008 2.763081	
00C8 108.786	
dih8 -5.875	
clo9 1.725303	
cloo9 103.543	
dih9 86.440	
0010 2.895279	
ooc10 99.285	
dih10 153.329	
hc11 1.096825	
hccl1 111.853	
dih11 -131.969	
hc12 1.100739	
hcc12 109.515	
dih12 111.013	
hc13 1.095062	
hcc13 109.525	
dih13 -125.427	
hc14 1.096736	
hcc14 107.722	
dih14 119.290	
hc15 1.098302	
hcc15 112.200	
dih15 -107.804	

hc16	1.093686	
hcc16	115.025	
dih16	131.078	
ho17	0.975757	
hoo17	18.144	
dih17	173.629	
ho18	0.968553	
hoo18	114.146	
dih18	113.727	
ho19	0.991861	
hoo19	9.666	
dih19	170.874	

 Table 29: Single point energy calculation script using implicit solvent environment (PCM), of gasphase optimized structure of reactant (or product) of CPD + HOCl + explicit water molecule

```
#HF/6-31G* NoSymm opt POP=(MK,DIPOLE)
Calc for initial partial atomic charge quess and target dipole
0 1
 С
 С
         1 cc2
  c 2 cc3
                                  1 ccc3
  c 3 cc4
                                2 ccc4
                                                               1 dih4

      c
      3
      cc4
      2
      ccc4
      1
      dih4

      c
      4
      cc5
      3
      ccc5
      2
      dih5

      o
      2
      oc6
      3
      occ6
      4
      dih6

      o
      5
      oc7
      4
      occ7
      3
      dih7

      h
      3
      hc8
      2
      hcc8
      6
      dih8

      h
      3
      hc9
      2
      hcc9
      6
      dih9

      h
      4
      hc10
      3
      hcc10
      2
      dih10

      h
      4
      hc11
      3
      hcc11
      2
      dih11

      h
      1
      hc12
      2
      hcc12
      6
      dih12

      h
      1
      hc13
      2
      hcc13
      6
      dih13

cc2
                      1.521729
cc3
                      1.528426
                     109.193
ccc3
cc4
                    1.530854
                    105.445
ccc4
                       19.512
dih4
                    1.528425
cc5
ccc5
                   105.445
-23.418
dih5
006
                     1.221537
                    125.081
0006
                  -160.564
dih6
                     1.221538
oc7
                      125.082
occ7
dih7
                   -160.561
                     1.097780
hc8
                     107.519
hcc8
dih8
                       80.770
hc9
                     1.092772
                   110.168
hcc9
dih9
                      -35.822
                    1.092771
hc10
                   115.075
hcc10
dih10
                  -145.032
hc11
                     1.097780
hcc11
                     111.148
                       92.796
dih11
hc12
                      1.096378
hcc12
                      111.170
dih12
                       -66.825
hc13
                 1.096379
```

hcc13	111.260
dih13	51.984

Table 30: Parameterization script (QM): Initial partial atomic guess and target dipole calculation

#HF/6-32	1G* Opt	
Calc for	r water - CPD(H12)	interaction distance and energy
0 1		
C C	cc2	
	cc3 1 ccc3	
	cc4 2 ccc4	1 dih4
	cc5 3 ccc5	
	oc7 4 occ7 hc8 2 hcc8	6 dih8
	hc9 2 hcc9	
	hc10 3 hcc10	
	hc11 3 hcc11	
	hc12 2 hcc12 hc13 2 hcc13	
	oh14 1 ohc14	
	ho15 12 hoh15	
h 14 1	hol6 12 hoh16	1 dih16
cc2	1.521729	
cc3	1.528426	
ccc3	109.193	
cc4 ccc4	1.530854 105.445	
dih4	19.512	
cc5	1.528425	
ccc5 dih5	105.445	
oc6	-23.418 1.221537	
0006	125.081	
dih6	-160.564	
oc7 occ7	1.221538 125.082	
dih7	-160.561	
hc8	1.097780	
hcc8	107.519	
dih8 hc9	80.770 1.092772	
hcc9	110.168	
dih9	-35.822	
hc10	1.092771	
hcc10 dih10	115.075 -145.032	
hc11	1.097780	

c11 1	11.148
h11	92.796
12 1	096378
c12 1	11.170
h12 -	66.825
13 1	096379
c13 1	11.260
h13	51.984
14 2	.100000
c14 1	79.000
h14 1	80.000
15 0	.950000
h15 1	09.471
h15 1	80.000
16 0	.950000
h16 1	09.471
h16 3	00.000
	h11 12 1 c12 1 h12 - 13 1 c13 1 h13 1 h14 2 c14 1 h14 1 h15 1 h15 1 h16 0 h16 1

Table 31: Parameterization script (QM): Target water-CPD (H12) interaction energy and distance calculation

```
* From tutorial on A. MacKerell's website
* Modified by Cassian
* Water interactions
ioformat extended
set pdbdir .
!### Set Variables ###
set residue cpd
set zeed 1 C1 1 C2 1 C3 ! seed to create molecule
set frist none ! first capping residue
set lsat none ! last capping residue
set bombover 0 !needs to be set to -1 for 3-membered rings
set ffsuffix cpd
set stage 5 !4=do everything with mp2 geometry for h2o inter
            !5=do everything with charmm geometry for h2o inter
!### Things that usually should not be changed. ####
set dontmini 0 ! Do not minimize main conformation. Should normally
not be used.
system "./do remas"
open unit 10 read form name top all22 prot.rtf
read rtf card unit 10
close unit 10
open unit 10 read form name par_all22_prot.prm
read param card flex unit 10
```

```
close unit 10
open unit 10 read card name top all36 cgenff remas@ffsuffix.rtf
read rtf card unit 10 append
close unit 10
open unit 10 read card name par all36 cgenff remas@ffsuffix.prm
read para card flex unit 10 append
close unit 10
if @?str eq 1 stream @str
stream toppar dum nobel gases.str
read sequence card
* @residue
*
1
@residue
bomlev @bombover
generate @residue first @frist last @lsat setup warn
bomlev 0
! For testing and debugging purposes
   open unit 30 write form name test.psf
   write psf card unit 30
   close unit 30
ic param
  ic seed @zeed
  ic build
   ic purge
   ic print
! Write out initial structure for testing and debugging purposes
   open unit 30 write form name @residue init.pdb
   write coor pdb unit 30
   close unit 30
!minimize structure and calculate dipole moment
if @dontmini eq 1 goto skip min
   MINI CONJ nstep 200 nprint 20 inbfrg 1000 cutnb 999.
   MINI NRAP nstep 50 tolgrd 0.00001
   prnlev 3
   coor dipole oxyz select segid @residue end
!save minimized geometry
  coor print
   open unit 30 write form name @residue min.crd
   write coor card unit 30
   open unit 30 write form name @pdbdir/@residue_min.pdb
   write coor pdb unit 30
   close unit 30
label skip min
if @stage lt 4 stop
!save coordinates in comparison set
```

```
coor copy comp
ic save
ic fill
ic print
!water interactions
if @stage eq 4 then
 open unit 50 write form name @residue water constr.ene
else
 open unit 50 write form name @residue water.ene
endif
!variables for averages etc.
set count 0
set asum 0.0
set sum 0.0
set sum2 0.0
!read QM optimized coordinates into the comparison set
!and align charmm optimized structure to allow for the
!components of the dipole moment to be checked.
open unit 20 read form name @residue_mp2.crd
if @stage eg 4 then
 read coor card unit 20
else
 read coor card unit 20 comp
 coor orient rms
 coor orient mass noro
 coor print
 open unit 30 write form name @residue min orient.pdb
 write coor pdb unit 30
endif
coor dipole oxyz select segid @residue end
! For testing and debugging purposes
open unit 30 write form name test.pdb
write coor pdb unit 30
* @residue
! Dipole values below from QM calculation for comparison
echu 50
echo @residue water interaction
if @stage eq 4 echo ######## USING MP2 GEOMETRY!!! #########
echo QM dipole (Debye)
echo mp2/6-31g*: X=
                       1.4581
                                 Y=
                                       -0.4891
                                                   Z =
                                                          0.0853
Tot=
        1.5404
echo hf/6-31g*: X=
                       1.5098
                                Y=
                                       0.1997
                                                  Z =
                                                          1.1617
Tot=
        1.9153
echo empirical dipole: ?xdip ?ydip ?zdip ?rdip
echo
ic delete sele all end
!1) C3H32...OHH 90.0 degrees
```

```
set d 1.60 ! initial interaction distance
read sequence card
* water
*
1
tip3
generate tip3 first none last none setup warn noangle nodihedral
join @residue tip3 renumber
read sequence card
* dummy
+
2
dum dum
generate dum first none last none setup warn noangle nodihedral
join @residue dum renumber
!preparation of IC table for surface of interest
ic edit
dihe 1 H31 1 C3 1 H32
                          3 dum
                                   0.0
dihe 1 C3 1 H32 3 dum 4 dum 0.0
dihe 4 dum 3 dum 1 H32 2 oh2 180.0
end
ic fill preserve
ic edit
dihe 3 dum 1 H32 2 oh2 2 h1 90.0
dihe 3 dum 1 H32 2 oh2 2 h2 -90.0
bond 1 H32 3 dum 1.
bond 3 dum 4 dum 1.
bond 1 H32 2 oh2 @d !varied
bond 2 oh2 2 h1 0.9572
bond 2 oh2 2 h2 0.9572
angl 1 C3 1 H32 3 dum 90.0
angl 1 H32 3 dum 4 dum 90.0
angl 3 dum 1 H32 2 oh2 90.0
angl 1 H32 2 oh2 2 h1 127.74
angl 1 H32 2 oh2 2 h2 127.74
end
ic print
ic build
coor print
update cutnb 999.0 ctofnb 998.0 ctonnb 997.0
set u 1000. ! minimum energy
```

```
label loop 1
inter -
  sele resn @residue end -
  sele resn tip3 end
!check for lowest energy and store data
set n @d
set o ?elec
set x ?vdw
set m ?ener
if @u lt @m goto done 1
set u @m
ic edit
bond 2 oh2 1 H32
                   @d
end
!initialize water coordinates
coor init sele resn tip3 .or. resn dum end
ic build
prnlev -5
incr d by 0.01
if @d lt 4.5 goto loop_1
label done_1
prnlev 5
delete atom sele type dum end
open unit 30 write form name @residue wat1.pdb
write coor pdb unit 30
* @residue
*
!accumulate statistics
calc count = @count + 1
set ai -3.0084 ! From QM calc and scaled by 1.16 for neutral
compound
set aidist 2.47
calc diff = @m - @ai
calc sum = @sum + @diff
calc diff2 = abs(@diff*@diff)
calc sum2 = @sum2 + @diff2
calc asum = @asum + abs(@diff)
calc distdiff = @n - @aidist
! write ab initio and empirical information to unit 50
```

```
echo 1) C3H32..OHH 90.0 degrees
echo a.i. @ai @aidist
echo emp. @m @o @x @n
echo ene diff: @diff dist diff: @distdiff
echo
delet atom sele resn tip3 .or. resn dum end
!2) C3H31...OHH 90.0 degrees
set d 1.60 ! initial interaction distance
read sequence card
* water
1
tip3
generate tip3 first none last none setup warn noangle nodihedral
join @residue tip3 renumber
read sequence card
* dummy
*
2
dum dum
generate dum first none last none setup warn noangle nodihedral
join @residue dum renumber
!preparation of IC table for surface of interest
ic edit
dihe 1 H32 1 C3 1 H31 3 dum
                                 0.0
           1 H31 3 dum 4 dum
dihe 1 C3
                                 0.0
dihe 4 dum 3 dum 1 H31 2 oh2 180.0
end
ic fill preserve
ic edit
dihe 3 dum 1 H31 2 oh2 2 h1
                               90.0
dihe 3 dum 1 H31 2 oh2 2 h2 -90.0
bond 1 H31 3 dum 1.
bond 3 dum 4 dum 1.
bond 1 H31 2 oh2 @d !varied
bond 2 oh2 2 h1 0.9572
bond 2 oh2 2 h2 0.9572
angl 1 C3 1 H31 3 dum 90.0
angl 1 H31 3 dum 4 dum 90.0
angl 3 dum 1 H31 2 oh2 90.0
angl 1 H31 2 oh2 2 h1 127.74
angl 1 H31 2 oh2 2 h2 127.74
end
```

```
ic print
ic build
coor print
update cutnb 999.0 ctofnb 998.0 ctonnb 997.0
set u 1000. ! minimum energy
label loop_2
inter -
  sele resn @residue end -
  sele resn tip3 end
!check for lowest energy and store data
set n @d
set o ?elec
set x ?vdw
set m ?ener
if @u lt @m goto done 2
set u @m
ic edit
bond 2 oh2 1 H31 @d
end
!initialize water coordinates
coor init sele resn tip3 .or. resn dum end
ic build
prnlev -5
incr d by 0.01
if @d lt 4.5 goto loop 2
label done 2
prnlev 5
delete atom sele type dum end
open unit 30 write form name @residue wat2.pdb
write coor pdb unit 30
* @residue
*
!accumulate statistics
calc count = @count + 1
set ai -2.5196 ! From QM calc and scaled by 1.16 for neutral
compound
set aidist 2.47
```

```
calc diff = @m - @ai
calc sum = @sum + @diff
calc diff2 = abs(@diff*@diff)
calc sum2 = @sum2 + @diff2
calc asum = @asum + abs(@diff)
calc distdiff = @n - @aidist
! write ab initio and empirical information to unit 50
echo 2) C3H31..OHH 90.0 degrees
echo a.i. @ai @aidist
echo emp. @m @o @x @n
echo ene diff: @diff dist diff: @distdiff
echo
delet atom sele resn tip3 .or. resn dum end
13) C2H22...OHH 90.0 degrees
set d 1.60 ! initial interaction distance
read sequence card
* water
1
tip3
generate tip3 first none last none setup warn noangle nodihedral
join @residue tip3 renumber
read sequence card
* dummy
*
2
dum dum
generate dum first none last none setup warn noangle nodihedral
join @residue dum renumber
!preparation of IC table for surface of interest
ic edit
dihe 1 H21
           1 C2 1 H22 3 dum
                                  0.0
dihe 1 C2 1 H22 3 dum 4 dum
                                  0.0
dihe 4 dum 3 dum 1 H22
                         2 oh2 180.0
end
ic fill preserve
ic edit
dihe 3 dum 1 H22 2 oh2 2 h1 90.0
dihe 3 dum 1 H22 2 oh2 2 h2 -90.0
bond 1 H22 3 dum 1.
bond 3 dum 4 dum 1.
bond 1 H22 2 oh2 @d !varied
```

bond 2 oh2 2 h1 0.9572 bond 2 oh2 2 h2 0.9572 angl 1 C2 1 H22 3 dum 90.0 angl 1 H22 3 dum 4 dum 90.0 angl 3 dum 1 H22 2 oh2 90.0 angl 1 H22 2 oh2 2 h1 127.74 angl 1 H22 2 oh2 2 h2 127.74 end ic print ic build coor print update cutnb 999.0 ctofnb 998.0 ctonnb 997.0 set u 1000. ! minimum energy label loop 3 inter sele resn @residue end sele resn tip3 end !check for lowest energy and store data set n @d set o ?elec set x ?vdw set m ?ener if @u lt @m goto done_3 set u @m ic edit bond 2 oh2 1 H22 @d end !initialize water coordinates coor init sele resn tip3 .or. resn dum end ic build prnlev -5 incr d by 0.01 if @d lt 4.5 goto loop_3 label done_3 prnlev 5 delete atom sele type dum end open unit 30 write form name @residue wat3.pdb

```
write coor pdb unit 30
* @residue
!accumulate statistics
calc count = @count + 1
set ai -2.5196 ! From QM calc and scaled by 1.16 for neutral
compound
set aidist 2.47
calc diff = @m - @ai
calc sum = @sum + @diff
calc diff2 = abs(@diff*@diff)
calc sum2 = @sum2 + @diff2
calc asum = @asum + abs(@diff)
calc distdiff = @n - @aidist
! write ab initio and empirical information to unit 50
echo 3) C2H22..OHH 90.0 degrees
echo a.i. @ai @aidist
echo emp. @m @o @x @n
echo ene diff: @diff dist diff: @distdiff
echo
delet atom sele resn tip3 .or. resn dum end
!4) C2H21...OHH 90.0 degrees
set d 1.60 ! initial interaction distance
read sequence card
* water
*
1
tip3
generate tip3 first none last none setup warn noangle nodihedral
join @residue tip3 renumber
read sequence card
* dummy
*
2
dum dum
generate dum first none last none setup warn noangle nodihedral
join @residue dum renumber
!preparation of IC table for surface of interest
ic edit
dihe 1 H22
           1 C2
                    1 H21
                            3 dum
                                    0.0
dihe 1 C2 1 H21 3 dum 4 dum
                                   0.0
```

```
dihe 4 dum 3 dum 1 H21 2 oh2 180.0
end
ic fill preserve
ic edit
dihe 3 dum 1 H21 2 oh2 2 h1
                                 90.0
dihe 3 dum 1 H21 2 oh2 2 h2 -90.0
bond 1 H21 3 dum 1.
bond 3 dum 4 dum 1.
bond 1 H21 2 oh2 @d !varied
bond 2 oh2 2 h1 0.9572
bond 2 oh2 2 h2 0.9572
angl 1 C2 1 H21 3 dum 90.0
angl 1 H21 3 dum 4 dum 90.0
angl 3 dum 1 H21 2 oh2 90.0
angl 1 H21 2 oh2 2 h1 127.74
angl 1 H21 2 oh2 2 h2 127.74
end
ic print
ic build
coor print
update cutnb 999.0 ctofnb 998.0 ctonnb 997.0
set u 1000. ! minimum energy
label loop 4
inter -
 sele resn @residue end -
  sele resn tip3 end
!check for lowest energy and store data
set n @d
set o ?elec
set x ?vdw
set m ?ener
if @u lt @m goto done 4
set u @m
ic edit
bond 2 oh2 1 H21 @d
end
!initialize water coordinates
coor init sele resn tip3 .or. resn dum end
ic build
prnlev -5
incr d by 0.01
```

```
if @d lt 4.5 goto loop 4
label done 4
prnlev 5
delete atom sele type dum end
open unit 30 write form name @residue wat4.pdb
write coor pdb unit 30
* @residue
*
!accumulate statistics
calc count = @count + 1
set ai -3.0084 ! From QM calc and scaled by 1.16 for neutral
compound
set aidist 2.47
calc diff = @m - @ai
calc sum = @sum + @diff
calc diff2 = abs(@diff*@diff)
calc sum2 = @sum2 + @diff2
calc asum = @asum + abs(@diff)
calc distdiff = @n - @aidist
! write ab initio and empirical information to unit 50
echo 4) C2H21..OHH 90.0 degrees
echo a.i. @ai @aidist
echo emp. @m @o @x @n
echo ene diff: @diff dist diff: @distdiff
echo
delet atom sele resn tip3 .or. resn dum end
!5) C5H52...OHH
                  0.0 degrees
set d 1.60 ! initial interaction distance
read sequence card
* water
*
1
tip3
generate tip3 first none last none setup warn noangle nodihedral
join @residue tip3 renumber
read sequence card
* dummy
*
2
dum dum
```

generate dum first none last none setup warn noangle nodihedral join @residue dum renumber !preparation of IC table for surface of interest ic edit dihe 1 H51 1 C5 1 H52 3 dum 0.0 dihe 1 C5 1 H52 3 dum 4 dum 0.0 dihe 4 dum 3 dum 1 H52 2 oh2 180.0 end ic fill preserve ic edit dihe 3 dum 1 H52 2 oh2 2 h1 90.0 dihe 3 dum 1 H52 2 oh2 2 h2 -90.0 bond 1 H52 3 dum 1. bond 3 dum 4 dum 1. bond 1 H52 2 oh2 @d !varied bond 2 oh2 2 h1 0.9572 bond 2 oh2 2 h2 0.9572 angl 1 C5 1 H52 3 dum 90.0 angl 1 H52 3 dum 4 dum 90.0 angl 3 dum 1 H52 2 oh2 90.0 angl 1 H52 2 oh2 2 h1 127.74 angl 1 H52 2 oh2 2 h2 127.74 end ic print ic build coor print update cutnb 999.0 ctofnb 998.0 ctonnb 997.0 set u 1000. ! minimum energy label loop 5 inter sele resn @residue end sele resn tip3 end !check for lowest energy and store data set n @d set o ?elec set x ?vdw set m ?ener if @u lt @m goto done 5 set u @m ic edit bond 2 oh2 1 H52 @d

end

```
!initialize water coordinates
coor init sele resn tip3 .or. resn dum end
ic build
!prnlev -5
incr d by 0.01
if @d lt 4.5 goto loop_5
label done 5
prnlev 5
delete atom sele type dum end
open unit 30 write form name @residue wat5.pdb
write coor pdb unit 30
* @residue
*
!accumulate statistics
calc count = @count + 1
set ai -3.2900 ! From QM calc and scaled by 1.16 for neutral
compound
set aidist 2.34
calc diff = @m - @ai
calc sum = @sum + @diff
calc diff2 = abs(@diff*@diff)
calc sum2 = @sum2 + @diff2
calc asum = @asum + abs(@diff)
calc distdiff = @n - @aidist
! write ab initio and empirical information to unit 50
echo 5) C5H52..OHH 90.0 degrees
echo a.i. @ai @aidist
echo emp. @m @o @x @n
echo ene diff: @diff dist diff: @distdiff
echo
delet atom sele resn tip3 .or. resn dum end
!6) C5H51...OHH 0.0 degrees
set d 1.60 ! initial interaction distance
read sequence card
* water
*
1
tip3
generate tip3 first none last none setup warn noangle nodihedral
```

```
join @residue tip3 renumber
read sequence card
* dummy
*
2
dum dum
generate dum first none last none setup warn noangle nodihedral
join @residue dum renumber
!preparation of IC table for surface of interest
ic edit
dihe 1 H52
           1 C5 1 H51
                         3 dum
                                  0.0
dihe 1 C5 1 H51 3 dum 4 dum
                                 0.0
dihe 4 dum 3 dum 1 H51 2 oh2 180.0
end
ic fill preserve
ic edit
dihe 3 dum 1 H51 2 oh2 2 h1 90.0
dihe 3 dum 1 H51 2 oh2 2 h2 -90.0
bond 1 H51 3 dum 1.
bond 3 dum 4 dum 1.
bond 1 H51 2 oh2 @d !varied
bond 2 oh2 2 h1 0.9572
bond 2 oh2 2 h2 0.9572
angl 1 C5 1 H51 3 dum 90.0
angl 1 H51 3 dum 4 dum 90.0
angl 3 dum 1 H51 2 oh2 90.0
angl 1 H51 2 oh2 2 h1 127.74
angl 1 H51 2 oh2 2 h2 127.74
end
ic print
ic build
coor print
update cutnb 999.0 ctofnb 998.0 ctonnb 997.0
set u 1000. ! minimum energy
label loop_6
inter -
 sele resn @residue end -
  sele resn tip3 end
!check for lowest energy and store data
set n @d
set o ?elec
```

```
set x ?vdw
set m ?ener
if @u lt @m goto done 6
set u @m
ic edit
bond 2 oh2 1 H51 @d
end
!initialize water coordinates
coor init sele resn tip3 .or. resn dum end
ic build
!prnlev -5
incr d by 0.01
if @d lt 4.5 goto loop 6
label done 6
prnlev 5
delete atom sele type dum end
open unit 30 write form name @residue wat6.pdb
write coor pdb unit 30
* @residue
!accumulate statistics
calc count = @count + 1
set ai -3.2900 ! From QM calc and scaled by 1.16 for neutral
compound
set aidist 2.34
calc diff = @m - @ai
calc sum = @sum + @diff
calc diff2 = abs(@diff*@diff)
calc sum2 = @sum2 + @diff2
calc asum = @asum + abs(@diff)
calc distdiff = @n - @aidist
! write ab initio and empirical information to unit 50
echo 6) C5H51..OHH 90.0 degrees
echo a.i. @ai @aidist
echo emp. @m @o @x @n
echo ene diff: @diff dist diff: @distdiff
echo
delet atom sele resn tip3 .or. resn dum end
17) 01...OHH 0.0 degrees
```

```
set d 1.60 ! initial interaction distance
read sequence card
* water
*
1
tip3
generate tip3 first none last none setup warn noangle nodihedral
join @residue tip3 renumber
read sequence card
* dummy
*
3
dum dum dum
generate dum first none last none setup warn noangle nodihedral
join @residue dum renumber
!obtain the inner angle.
!quick 3 1 12
!calculate the outer angle
!calc oangle = (360.0 - ?thet) / 2.
ic edit
dihe 1 C2
          1 C5 1 C1
                         3 dum 180.0
dihe 3 dum 1 C1
                  1 01
                        4 dum 0.0
dihe 1 C1 1 01
                4 dum 5 dum 0.0
dihe 5 dum 4 dum 1 O1
                         2 h1 180.0
end
ic fill preserve
ic edit
dihe 4 dum 1 01 2 h1 2 oh2 0.0
dihe 1 01 2 h1 2 oh2 2 h2 180.0
!dihe 5 dum 2 h1 2 oh2 2 h2 0.0
dihe 5 dum 2 h1 2 oh2 2 h2 -90.0
bond 1 C1 3 dum 1.
bond 1 01 4 dum 1.
bond 4 dum 5 dum 1.
bond 1 01 2 h1 @d !varied
bond 2 oh2 2 h1 0.9572
bond 2 oh2 2 h2 0.9572
angl 1 C5 1 C1 3 dum 127.9075
angl 1 C1 1 O1 4 dum 90.0
angl 1 01 4 dum 5 dum 90.0
angl 4 dum 1 01 2 h1 90.0
angl 1 01 2 h1 2 oh2 180.0
angl 2 h1 2 oh2 2 h2 109.47
end
ic print
ic build
```

```
coor print
update cutnb 999.0 ctofnb 998.0 ctonnb 997.0
set u 1000. ! minimum energy
label loop_7
inter -
 sele resn @residue end -
  sele resn tip3 end
!check for lowest energy and store data
set n @d
set o ?elec
set x ?vdw
set m ?ener
if @u lt @m goto done 7
set u @m
ic edit
bond 2 h1 1 01 @d
end
!initialize water coordinates
coor init sele resn tip3 .or. resn dum end
ic build
!prnlev -5
incr d by 0.01
if @d lt 4.5 goto loop 7
label done 7
prnlev 5
delete atom sele type dum end
open unit 30 write form name @residue wat7.pdb
write coor pdb unit 30
* @residue
*
!accumulate statistics
calc count = @count + 1
set ai -4.1479 ! From QM calc and scaled by 1.16 for neutral
compound
set aidist 2.11
calc diff = @m - @ai
calc sum = @sum + @diff
calc diff2 = abs(@diff*@diff)
```

```
calc sum2 = @sum2 + @diff2
calc asum = @asum + abs(@diff)
calc distdiff = @n - @aidist
! write ab initio and empirical information to unit 50
echo 7) O1..HOH 0.0 degrees
echo a.i. @ai @aidist
echo emp. @m @o @x @n
echo ene diff: @diff dist diff: @distdiff
echo
delet atom sele resn tip3 .or. resn dum end
18) 04...OHH 0.0 degrees
set d 1.60 ! initial interaction distance
read sequence card
* water
*
1
tip3
generate tip3 first none last none setup warn noangle nodihedral
join @residue tip3 renumber
read sequence card
* dummy
٦
dum dum dum
generate dum first none last none setup warn noangle nodihedral
join @residue dum renumber
!preparation of IC table for surface of interest
ic edit
dihe 1 C5
           1 C3
                  1 C4
                         3 dum 180.0
dihe 3 dum 1 C4
                  1 04
                         4 dum 0.0
dihe 1 C4
           1 04
                  4 dum 5 dum 0.0
dihe 5 dum 4 dum 1 04
                         2 h1 180.0
end
ic fill preserve
ic edit
dihe 4 dum 1 04
                 2 h1 2 oh2
                               0.0
          2 h1 2 oh2 2 h2 180.0
dihe 1 04
!dihe 5 dum 2 h1 2 oh2 2 h2 0.0
dihe 5 dum 2 h1 2 oh2 2 h2 90.0
bond 1 C4 3 dum 1.
bond 1 04 4 dum 1.
bond 4 dum 5 dum 1.
```

```
bond 1 O4 2 h1 @d !varied
bond 2 oh2 2 h1 0.9572
bond 2 oh2 2 h2 0.9572
angl 1 C3 1 C4 3 dum 127.9075
angl 1 C4 1 O4 4 dum 90.0
angl 1 04 4 dum 5 dum 90.0
angl 4 dum 1 04 2 h1 90.0
angl 1 04 2 h1 2 oh2 180.0
angl 2 h1 2 oh2 2 h2 109.47
end
ic print
ic build
coor print
update cutnb 999.0 ctofnb 998.0 ctonnb 997.0
set u 1000. ! minimum energy
label loop_8
inter -
 sele resn @residue end -
  sele resn tip3 end
!check for lowest energy and store data
set n @d
set o ?elec
set x ?vdw
set m ?ener
if @u lt @m goto done_8
set u @m
ic edit
bond 2 h1 1 04 @d
end
!initialize water coordinates
coor init sele resn tip3 .or. resn dum end
ic build
!prnlev -5
incr d by 0.01
if @d lt 4.5 goto loop_8
label done 8
prnlev 5
delete atom sele type dum end
```

```
open unit 30 write form name @residue wat8.pdb
write coor pdb unit 30
* @residue
!accumulate statistics
calc count = @count + 1
set ai -4.1479 ! From QM calc and scaled by 1.16 for neutral
compound
set aidist 2.11
calc diff = @m - @ai
calc sum = @sum + @diff
calc diff2 = abs(@diff*@diff)
calc sum2 = @sum2 + @diff2
calc asum = @asum + abs(@diff)
calc distdiff = @n - @aidist
! write ab initio and empirical information to unit 50
echo 8) 04..HOH 0.0 degrees
echo a.i. @ai @aidist
echo emp. @m @o @x @n
echo ene diff: @diff dist diff: @distdiff
echo
delet atom sele resn tip3 .or. resn dum end
!calculate statistics from all water calculations
!average difference
calc avediff = @sum/@count
calc avediff2 = @sum2/@count
calc rmsdif = sqrt(@avediff2 - (@avediff*@avediff))
!AAE; average absolute error
calc aae = @asum/@count
echo avediff, rmsdiff, average absolute error
echo @avediff @rmsdif @aae
stop
```

Table 32: Parameterization script (MM): Water – CPD dipole and optimal interaction energies and distances

%CHK=CPD_OPT_FREQ #MP2/6-31G* NOSYMM OPT=TIGHT POP=(MK,DIPOLE) DENSITY=CURRENT Initial opt 0 1 C -0.01119 0.01187 -0.18971 0 -0.74465 -0.97376 -0.20661

-			
С	1.51541	-0.02896	-0.12172
Н	1.91154	-0.52385	-1.02065
Н	1.84573	-0.63344	0.73519
С	1.98602	1.42895	-0.02109
Н	2.80533	1.65720	-0.71746
Н	2.35965	1.66407	0.98651
С	0.76424	2.29273	-0.32771
0	0.77831	3.51602	-0.44209
С	-0.48850	1.45196	-0.17369
Н	-1.19878	1.64493	-0.98696
Н	-0.99637	1.68132	0.77091
%CHK #MP2		MM GEOM=ALLC	CHECK FREQ IOP(7/33=1)
	calc for fo ribution	rce constant	s to get target normal mode freq and
0 1			
1			

Table 33: Frequency calculation script for force constants to get target normal mode frequency and percent contribution

```
* MOLVIB analysis of Gaussian second derivatives
* NOTE: The MOLVIB module is very sensitive to the format
*
ioformat extended
MOLVIB NDI1 33 NDI2 33 NDI3 44 NATOM 13 NOTO
GFX
PRNT
       1
DIM
      33
           33
                44
CART
   0.001835 0.015210
                        -0.040459
                                    12.000
   -0.705793 -0.955884
                          0.179520
                                    16.000
   1.522114 -0.022283 -0.193538
                                     12.000
             -0.349330
                          -1.217897
   1.743134
                                      1.000
                                      1.000
   1.950783
              -0.765002
                          0.483782
   1.990997
               1.416118
                          0.040270
                                     12.000
               1.727522
                          -0.561949
                                    1.000
   2.848032
   2.255480
              1.580291
                          1.092989
                                      1.000
   0.780126
             2.294838
                          -0.272340
                                    12.000
   0.826424
               3.487778
                          -0.531000
                                     16.000
   -0.482586
               1.448084
                          -0.207391
                                     12.000
   -1.083215
               1.559459
                          -1.117822
                                      1.000
   -1.120590
                                      1.000
               1.752238
                          0.630754
```

IC									
1	1	2			!1		R(1,2)	C1 = 01	
1	1	3	0	0	!2	R2	R(1,3)	C1 - C2	
1	1	11	0	0	!3	R3	R(1,11)	C1 - C5	
1	3	4	0		!4		R(3,4)	C2 - H21	
1	3	5	0		!5		R(3,5)	C2 - H22	
1	3	6			.5 !6		R(3,6)	C2 - C3	
			0						
1	6	7	0		!7		R(6,7)	C3 - H31	
1	6	8	0				(6,8) C3 - 1		
1	6	9	0		!9		10(0))	C3 - C4	
1	9	10	0	0	!10	R10	R(9,10)	C4 = O4	
1	9	11	0	0	!11	R11	R(9,11)	C4 - C5	
1	11	12	0				R(11,12)	C5 - H51	
1		13	0	0			R(11,13)	C5 - H52	
2	2	1	3			A1		01 - C1 - C2	1b2
2	2	1	11				A(2,1,11)	01 - C1 - C5	1b1
2	3	1	11				A(3,1,11)	C2 - C1 - C5	rla4
2	1	3	4			A4		C1 - C2 - H21	2b3
2	1	3	5		!18	A5	A(1,3,5)	C1 - C2 - H22	2b4
2	1	3	6	0	!19	A6	A(1,3,6)	C1 - C2 - C3	rla5 gamma
2	4	3	5	0	!20	A7	A(4,3,5)	H21 - C2 - H22	
2	4		6			A8		H21 - C2 - C3	2b1
2	5	3	6			A9		H22 - C2 - C3	2b2
2	3	6	7			A10		C2 - C3 - H31	3b3
2									
	3	6	8			A11		C2 - C3 - H32	3b4
2	3	6	9			A12		C2 - C3 - C4	rlal gamma
2	7	6	8			A13		H31 - C3 - H32	
2	7	6	9	0	!27	A14	A(7,6,9)	H31 - C3 - C4	3b1
2	8	6	9	0	!28	A15	A(8,6,9)	H32 - C3 - C4	3b2
2	6	9	10	0	!29	A16	A(6,9,10)	C3 - C4 - O4	4b1
2	6	9	11	0		A17		C3 - C4 - C5	rla2
2	10	9		0		A18		04 - C4 - C5	4b2
2		11	9			A19		C1 - C5 - C4	rla3 gamma
									-
2		11				A20		C1 - C5 - H51	5b1
2		11				A21		C1 - C5 - H52	5b2
2		11				A22			5b3
2		11		0	!36	A23			5b4
2	12	11	13	0	!37	A24	A(12,11,13)	H51 - C5 - H52	5 alpha
3	1	3	11	2	!38	I1	I(1,3,11,2)	C1-C2-C5-O1	C1
3		11			!39		I(9,11,6,10)		C4
4	11	1	3	6	!40		D(11,1,3,6)	C5-C1-C2-C3	rlt4
4	1	3	6	9	.10 !41		D(1,3,6,9)	C1-C2-C3-C4	r1t5
	1 3	6							
4			9		!42		D(3, 6, 9, 11)	C2-C3-C4-C5	r1t1
4	6	9	11		!43		D(6,9,11,1)	C3-C4-C5-C1	r1t2
4		11	1		!44		D(9,11,1,3)	C4-C5-C1-C2	r1t3
UMA	ΑT		0	1	L	0	! ro	w normalization	
:	1 1	1 1	1.			2	2 1.	3 3 1.	
4	1 4	1 3	1.			4	5 1.		
	5 4		1.			5	4 -1.		
			1.						
			1.			7	8 1.		
	3		1.				8 -1.		
						8			
	9 9		1.			10	10 1.		
1:			1.						
12	2 12	2 2	1.			12	13 1.		
		_		_	_				

13 12 1.	13 1	13 -1.					
14 19 1.	14 2	20 5.					
15 17 1.	15 3	18 -1.	15	21 1.	. 15	22 -1	
16 17 -1.	16	18 -1.	16	21 1.	. 16	22 1	
17 17 -1.	17			21 1.		22 -1	
			17	Δ⊥ ⊥.	• 1/	22 I	•
18 25 1.	18 2		1.0	0 - 1	1.0	00 1	
19 23 1.		24 -1.		27 1.		28 -1	
20 23 -1.	20 2	24 -1.	20	27 1.	. 20	28 1	•
21 23 -1.	21 2	24 1.	21	27 1.	. 21	28 -1	•
22 32 1.	22 3	37 5.					
23 33 1.	23	34 -1.	23	35 1.	. 23	36 -1	
24 33 1.		34 1.		35 -1.		36 -1	
25 33 1.		34 -1.		35 -1.		36 1	
	25 .	54 -1.	25	55 -I.	. 25	30 I	•
26 38 1.							
27 15 1.	27 :	14 -1.					
28 39 1.							
29 29 1.	29 3	31 -1.					
30 25 1.		30 -0.81	30	32 0.	.31 30	16 0	.31
30 19 -0.8			20				
		22 1 01	21	16 1	01 71	10 1	10
31 30 -1.3				16 -1.			.12
32 42 0.3		43 -0.81	32	44 1.	. 32	40 -0	.81
32 41 0.3	31						
33 42 -1.8	31 33 4	43 1.12	33	40 -1.	.12 33	41 1	.81
-1							
PED 0	15						
1 sC1=01							
2 sC1-C2							
3 sC1-C5	5						
4 sC2-H2	lsy						
5 sC2-H2	-						
6 sC2-C3							
8 sC3-H3							
9 sC3-C4							
10 sC4=04	Ł						
11 sC4-C5	5						
12 sC5-H5							
13 sC5-H	-						
15 r-C2H2							
16 w-C2H2	2						
17 t-C2H2	2						
18 sc-C3H	12						
19 r-C3H2							
20 w-C3H2							
21 t-C3H2							
22 sc-C5H							
23 r-C5H2	2						
24 w-C5H2	2						
25 t-C5H2	2						
26 w-C1=0							
28 w-C4=0							
29 d-C4=0							
30 d-5RN0	Bsy						
L	-						

31 d-5RNGas 32 t-5RNGsy 33 t-5RNGas -1 FMAT 1 1 0 ! Gaussian FX matrix is in input orientation Force constants in Cartesian coordinates: 1 2 3 4 5 1 0.643611D+00 2 0.236491D+00 0.805451D+00 3 -0.770802D-01 -0.108522D+00 0.263264D+00 4 -0.318400D+00 -0.293306D+00 0.705900D-01 0.332572D+00 5 -0.296109D+00 -0.506491D+00 0.983692D-01 0.344365D+00 0.553658D+00 6 0.692472D-01 0.967589D-01 -0.871856D-01 -0.832153D-01 -0.115810D+00 7 -0.169439D+00 0.318536D-01 0.368375D-02 -0.296714D-01 -0.215520D-01 8 0.173909D-01 -0.881021D-01 -0.136055D-02 -0.421041D-01 0.352002D-02 9 0.348225D-02 0.172581D-02 -0.893633D-01 0.108227D-01 0.510280D-02 10 -0.733210D-02 0.801079D-02 0.366843D-01 -0.464062D-03 -0.372434D-03 11 0.634848D-03 -0.134744D-02 0.116934D-02 -0.106853D-02 -0.118202D-03 12 0.140842D-02 -0.321319D-03 -0.252530D-02 -0.487130D-04 -0.259966D-03 13 -0.134787D-01 0.252618D-01 -0.242071D-01 0.493770D-02 -0.129191D-02 14 0.203779D-02 -0.184045D-02 -0.148578D-02 -0.123142D-02 0.667746D-03 15 0.554168D-02 -0.403406D-02 0.287438D-02 -0.873699D-03 -0.698227D-03 16 -0.166763D-01 -0.325877D-01 -0.642492D-02 -0.776981D-02 -0.149683D-03 17 -0.184391D-01 -0.620067D-02 0.154042D-02 0.268947D-02 0.124631D-02 18 0.460571D-03 0.600984D-02 0.266769D-03 0.368527D-02 0.520298D-04 19 -0.122688D-02 -0.256346D-03 0.676854D-04 -0.841820D-04 -0.325315D-03 20 0.286359D-02 -0.684261D-02 -0.673492D-03 -0.348106D-02 0.497678D-04 21 -0.141565D-02 -0.179596D-02 0.108578D-03 -0.811366D-03 -0.251657D-03 22 -0.350794D-03 -0.209015D-03 -0.858540D-03 -0.188155D-03 -0.230343D-03 23 0.164796D-02 -0.142444D-02 -0.853379D-03 -0.795777D-03 -0.281608D-05 24 0.415373D-03 -0.469615D-03 -0.338904D-03 -0.286168D-03 0.536667D-04 25 0.265346D-01 -0.248586D-01 0.217737D-02 -0.553549D-02 0.167721D-01

26 -0.252890D-01	-0.388526D-01	0.467323D-02	0.335264D-02	-
0.236797D-02 27 0.489794D-02				
	0.3/8506D-02	-0.114/41D-02	-0.113154D-02	
0.149518D-02		0 0105530 04	0 0601600 00	
28 -0.143017D-01	0.560882D-02	-0.8195/3D-04	0.263160D-02	-
0.722220D-02	0 (100500 00		0 1041460 00	
29 -0.788183D-02	0.613259D-02	-0.467272D-03	0.104146D-02	-
0.538102D-02				
30 0.683863D-03	-0.618144D-03	0.202353D-02	0.149466D-03	
0.642444D-03				
31 -0.116618D+00	0.389450D-01	-0.452435D-02	0.211636D-01	-
0.305527D-01				
32 0.512841D-01	-0.151697D+00	0.522296D-02	-0.688964D-02	-
0.485713D-01				
33 -0.346672D-02	0.468933D-02	-0.921110D-01	0.109782D-02	
0.117843D-01				
34 -0.554912D-02	0.173462D-02	-0.572823D-02	0.242513D-03	-
0.154103D-02				
35 0.162405D-01	-0.133538D-02	0.300695D-01	-0.107804D-02	
0.747642D-03				
36 0.315346D-03	-0.355828D-04	-0.854985D-03	0.874142D-04	-
0.112599D-03				
37 -0.677248D-02	0.331252D-02	0.570218D-02	0.565655D-03	-
0.179078D-02				
38 0.191282D-01	-0.745061D-02	-0.276817D-01	-0.149373D-02	
0.304232D-02				
39 -0.449006D-02	0.282816D-02	0.498922D-02	-0.659496D-04	-
0.367584D-03				-
0.367584D-03	0.282816D-02 7	0.498922D-02 8	-0.659496D-04 9	-
0.367584D-03 6 10	7			-
0.367584D-03 6 10 6 0.493274D-01	7			-
0.367584D-03 6 10 6 0.493274D-01 7 0.863396D-02	7 0.458667D+00	8		-
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0.367584D-03 6 10 6 0.493274D-01 7 0.863396D-02 8 0.944539D-02 9 0.170653D-01	7 0.458667D+00 -0.671308D-01 0.172014D-01	8 0.571449D+00 -0.344504D-01	9 0.578994D+00	-
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0.367584D-03 6 10 6 0.493274D-01 7 0.863396D-02 8 0.944539D-02 9 0.170653D-01 10 -0.479862D-02 0.615450D-01 11 -0.115974D-02 0.160573D-01 12 0.126200D-02 0.577158D-01 13 0.290698D-02 0.399346D-02 14 -0.128320D-03 0.449950D-02 15 0.263988D-04 0.142046D-01 16 0.258439D-02 0.241314D-02 17 -0.173542D-02 0.535128D-02	7 0.458667D+00 -0.671308D-01 0.172014D-01 -0.580467D-01 0.174177D-01 0.499845D-01 -0.912681D-01 0.731191D-01 -0.732942D-01 -0.927224D-01 -0.328830D-01	8 0.571449D+00 -0.344504D-01 0.203512D-01 -0.685688D-01 -0.715961D-01 0.729754D-01 -0.176164D+00 0.121059D+00 -0.335632D-01 -0.184450D+00	9 0.578994D+00 0.603948D-01 -0.722820D-01 -0.286280D+00 -0.734666D-01 0.120416D+00 -0.161957D+00 -0.103106D-01 -0.185659D-01	- - -
0.367584D-03 6 10 6 0.493274D-01 7 0.863396D-02 8 0.944539D-02 9 0.170653D-01 10 -0.479862D-02 0.615450D-01 11 -0.115974D-02 0.160573D-01 12 0.126200D-02 0.577158D-01 13 0.290698D-02 0.399346D-02 14 -0.128320D-03 0.449950D-02 15 0.263988D-04 0.142046D-01 16 0.258439D-02 0.241314D-02 17 -0.173542D-02 0.535128D-02 18 0.701632D-03	7 0.458667D+00 -0.671308D-01 0.172014D-01 -0.580467D-01 0.174177D-01 0.499845D-01 -0.912681D-01 0.731191D-01 -0.732942D-01 -0.927224D-01 -0.328830D-01	8 0.571449D+00 -0.344504D-01 0.203512D-01 -0.685688D-01 -0.715961D-01 0.729754D-01 -0.176164D+00 0.121059D+00 -0.335632D-01 -0.184450D+00	9 0.578994D+00 0.603948D-01 -0.722820D-01 -0.286280D+00 -0.734666D-01 0.120416D+00 -0.161957D+00 -0.103106D-01 -0.185659D-01	-
0.367584D-03 6 10 6 0.493274D-01 7 0.863396D-02 8 0.944539D-02 9 0.170653D-01 10 -0.479862D-02 0.615450D-01 11 -0.115974D-02 0.160573D-01 12 0.126200D-02 0.577158D-01 13 0.290698D-02 0.399346D-02 14 -0.128320D-03 0.449950D-02 15 0.263988D-04 0.142046D-01 16 0.258439D-02 0.241314D-02 17 -0.173542D-02 0.535128D-02	7 0.458667D+00 -0.671308D-01 0.172014D-01 -0.580467D-01 0.174177D-01 0.499845D-01 -0.912681D-01 0.731191D-01 -0.732942D-01 -0.927224D-01 -0.328830D-01	8 0.571449D+00 -0.344504D-01 0.203512D-01 -0.685688D-01 -0.715961D-01 0.729754D-01 -0.176164D+00 0.121059D+00 -0.335632D-01 -0.184450D+00	9 0.578994D+00 0.603948D-01 -0.722820D-01 -0.286280D+00 -0.734666D-01 0.120416D+00 -0.161957D+00 -0.103106D-01 -0.185659D-01	- - -
0.367584D-03 6 10 6 0.493274D-01 7 0.863396D-02 8 0.944539D-02 9 0.170653D-01 10 -0.479862D-02 0.615450D-01 11 -0.115974D-02 0.160573D-01 12 0.126200D-02 0.577158D-01 13 0.290698D-02 0.399346D-02 14 -0.128320D-03 0.449950D-02 15 0.263988D-04 0.142046D-01 16 0.258439D-02 0.241314D-02 17 -0.173542D-02 0.535128D-02 18 0.701632D-03	7 0.458667D+00 -0.671308D-01 0.172014D-01 -0.580467D-01 0.174177D-01 0.499845D-01 -0.912681D-01 0.731191D-01 -0.732942D-01 -0.927224D-01 -0.328830D-01 -0.258670D-02	8 0.571449D+00 -0.344504D-01 0.203512D-01 -0.685688D-01 -0.715961D-01 0.729754D-01 -0.176164D+00 0.121059D+00 -0.335632D-01 -0.184450D+00 -0.211970D-01	9 0.578994D+00 0.603948D-01 -0.722820D-01 -0.286280D+00 -0.734666D-01 0.120416D+00 -0.161957D+00 -0.103106D-01 -0.185659D-01 -0.857305D-01	- - -
$\begin{array}{c} 0.367584D-03 \\ & 6 \\ 10 \\ & 6 & 0.493274D-01 \\ & 7 & 0.863396D-02 \\ & 8 & 0.944539D-02 \\ & 9 & 0.170653D-01 \\ & 10 & -0.479862D-02 \\ 0.615450D-01 \\ & 11 & -0.115974D-02 \\ 0.615450D-01 \\ & 12 & 0.126200D-02 \\ 0.577158D-01 \\ & 13 & 0.290698D-02 \\ 0.399346D-02 \\ & 14 & -0.128320D-03 \\ 0.449950D-02 \\ & 15 & 0.263988D-04 \\ 0.142046D-01 \\ & 16 & 0.258439D-02 \\ 0.241314D-02 \\ & 17 & -0.173542D-02 \\ 0.535128D-02 \\ & 18 & 0.701632D-03 \\ 0.140985D-01 \\ & 19 & 0.570686D-04 \\ 0.627639D-03 \\ \end{array}$	7 0.458667D+00 -0.671308D-01 0.172014D-01 -0.580467D-01 0.174177D-01 0.499845D-01 -0.912681D-01 0.731191D-01 -0.732942D-01 -0.927224D-01 -0.328830D-01 -0.258670D-02	8 0.571449D+00 -0.344504D-01 0.203512D-01 -0.685688D-01 -0.715961D-01 0.729754D-01 -0.176164D+00 0.121059D+00 -0.335632D-01 -0.184450D+00 -0.211970D-01 0.749229D-03	9 0.578994D+00 0.603948D-01 -0.722820D-01 -0.286280D+00 -0.734666D-01 0.120416D+00 -0.161957D+00 -0.103106D-01 -0.185659D-01 -0.857305D-01 0.153965D-02	- - -

0.801433D-03	
	-0.183888D-02 0.372903D-02 -
0.303484D-03	
22 0.325049D-03 -0.152622D-02 0.214924D-03	-0.728631D-03 -0.639794D-02
	-0.641852D-02 -0.327574D-01 -
0.371775D-02	-0.0410520-02 -0.5275740-01 -
	0.329583D-03 -0.165535D-02 -
0.517449D-03	0.020002 00 0.2000002 02
25 -0.230640D-02 0.106298D-01	0.523737D-02 0.417508D-02 -
0.137768D-02	
26 0.425099D-03 0.185893D-01	-0.343851D-01 0.828822D-02
0.231544D-03	
27 0.175799D-02 0.648766D-02	-0.873153D-03 0.114493D-02
0.504702D-04	
28 0.877512D-03 -0.500167D-02	0.242211D-02 -0.359245D-02
0.329929D-03	-0.709010D-03 0.284934D-02 -
0.359325D-04	-0.7090100-05 0.2049540-02 -
30 -0.723913D-03 -0.388804D-02	0.474336D-03 -0.111159D-03
0.157395D-03	
31 0.567157D-02 -0.165118D-01	0.235584D-01 -0.297146D-02 -
0.159781D-02	
	-0.462486D-02 0.359910D-02 -
0.113561D-03	
	-0.503289D-03 0.264746D-01 -
0.602929D-02	0.390565D-03 0.594366D-03 -
0.505103D-03	0.3903030-03 0.3943000-03 =
35 -0.418335D-02 0.119983D-02	-0.213784D-02 -0.479347D-02
0.154357D-02	
36 0.105926D-02 0.292567D-03	-0.154095D-03 -0.894066D-03
0.237080D-03	
37 -0.296826D-03 -0.115595D-02	0.451618D-03 -0.147123D-02
0.199414D-03	
38 0.34469/D-02 0.169/89D-02 0.790725D-03	-0.379491D-02 0.510468D-02 -
39 0.133040D-03 -0.761011D-03	0.665258D-03 -0 141656D-02
0.143717D-03	5.555250D 05 0.141050D 02
11 12	13 14
15	
11 0.767915D-01	
12 0.801262D-01 0.307310D+00	
13 -0.590871D-02 0.573665D-02	
14 0.722121D-02 -0.709795D-02	
15 0.217836D-01 -0.189961D-01 0.171711D+00	0.766496D-01 -0.127959D+00
16 0.566443D-02 0.507231D-03	0 612635D-02 0 129413D-01
0.459892D-02	
	-0.973633D-02 -0.155654D-01 -
0.631078D-02	
18 -0.292925D-01 -0.208300D-02	0.890501D-02 0.156378D-01
0.483121D-02	
	-0.219925D-03 0.153276D-03 -
0.242028D-03	

20 0.221667D-02 0.613354D-04	0.312738D-03	-0.104758D-02	-0.140019D-02	
21 0.110237D-04	0.106507D-02	0.229087D-03	-0.265743D-04	
0.140257D-03	0 1240500 04	0 7407400 03	0 0000110 00	
22 -0.384838D-03 0.205000D-03	0.134050D-04	0./40/48D-03	0.266241D-03	
23 -0.534917D-02	-0.891875D-03	0.104549D-02	0.205104D-02	_
0.994566D-04	0.00000000000000	012010122 02	012002012 02	
24 -0.519136D-03	0.115096D-02	0.242193D-03	0.150236D-03	
0.111765D-02 25 -0.152929D-02	-0.670501D-03	-0.444032D-02	-0.455045D-02	-
0.276386D-03				
26 -0.390719D-03	-0.996987D-04	-0.127996D-02	-0.365047D-02	-
0.230085D-02				
27 -0.136409D-02	-0.345696D-03	-0.895183D-03	-0.846489D-03	
0.129802D-03 28 0.444757D-03	0 2820170-03	0 1697690-02	0 2113280-02	
0.577973D-03	J.202017D-03	0.107000-02	V.ZIIJZ0D-0Z	
29 -0.542521D-03	-0.115433D-03	-0.906777D-03	-0.183235D-02	-
0.716781D-03				
30 0.763825D-03	-0.152931D-05	0.488896D-03	0.785801D-03	
0.296360D-03				
31 0.447406D-03 0.100180D-02	0.407948D-03	-0.410375D-02	0.278751D-03	
32 0.113373D-03	0 9427880-04	0 414532D-03	0.401626D-03	_
0.502780D-03	0.9427000 04	0.4145520 05	0.4010200 05	
33 0.872590D-03	-0.643198D-03	0.394115D-02	0.464986D-03	-
0.225226D-03				
34 0.219860D-03	0.868758D-04	0.156310D-04	0.229723D-03	
0.129800D-03				
35 -0.189482D-03 0.790170D-04	0.131292D-03	-0.326911D-04	0.00000D+00	-
36 -0.914851D-04	0 453243D-04	-0.118127D-03	-0 256257D-04	
0.869752D-05	0.1552152 01	0.11012/2 00	0.2302372 01	
37 0.652878D-04	0.163669D-03	-0.458526D-03	0.478122D-04	
0.186132D-03				
38 0.935180D-04	-0.215301D-03	0.141261D-02	0.312884D-04	-
0.203120D-03	0 4165000 04	0 4105050 00	A 1151ACD AA	
39 -0.175969D-04 0.428694D-04	0.416793D-04	-0.412586D-03	0.115106D-03	
0.428694D-04 16	17	18	19	
20	± /	±0	÷ 2	
16 0.571894D+00				
17 0.724673D-01	0.465043D+00			
18 -0.378625D-01	-0.560860D-02	0.572176D+00		
19 -0.219683D+00			0.237014D+00	
20 -0.611944D-01	-0.716679D-01	0.426428D-01	0.676141D-01	
0.746130D-01 21 0.127735D+00	0 4227000 01		-0 1240120.00	_
0.458592D-01	0.432/090-01	-0.138040D+00	-0.1340120+00	-
22 -0.641312D-01	-0.111544D-01	-0.680704D-01	0.823900D-02	
0.343068D-02				
23 -0.853247D-02	-0.486087D-01	-0.308201D-01	0.250816D-02	
0.125947D-02				
24 -0.594504D-01	-0.368429D-01	-0.300155D+00	-0.608619D-02	-

	34D-02				
		0.390823D-01	-0.107669D-01	-0.232956D-01	
	70D-01	0 0270700 01	0 (015110 00	0 0002020 02	
	53D-02	-0.937078D-01	0.621511D-02	-0.908322D-02	
		0 540509D-02	-0 911532D-01	0.159606D-01	_
	89D-01	0.5405050 02	0.9113320 01	0.1390000 01	
		0.330693D-01	-0.704746D-02	0.476299D-02	_
	35D-02				
29	0.130206D-01	-0.381940D-01	0.139718D-01	-0.162979D-02	
0.1019	16D-02				
30 -	0.196653D-02	0.182470D-01	0.152532D-01	-0.183814D-02	
	90D-02				
_		-0.744970D-02	-0.678705D-03	-0.235078D-02	
	55D-02	0 4001505 00	0 0004055 00		
_		0.489158D-02	0.298485D-02	0.186597D-02	-
	67D-03	0.643724D-03	0 2609240-01	-0 2140050-02	
	84D-02	0.0437240-03	0.2000540-01	-0.5149990-02	
		-0.796603D-03	0.404955D-02	-0.984783D-03	
0.7205	24D-03				
35 -	0.247510D-02	-0.169196D-02	0.295920D-02	-0.623685D-03	
0.5480	14D-03				
		0.00000D+00	-0.990941D-03	0.312354D-03	-
	.42D-04				
		-0.373228D-03	-0.346943D-02	-0.759583D-04	-
	97D-03	-0.116480D-02	0 2555110 02	0 0701420 04	
	41D-04	-0.110400D-02	-0.355511D-02	0.9/91420-04	
		0.234187D-04	-0.117001D-02	0.663159D-04	_
	25D-03				
	21	22	23	24	
25					
	0.146622D+00				
		0.653862D-01			
		0.109985D-01			
		0.686030D-01 -0.612777D-02			
_	49D+00	-0.612///D-02	0.289630D-02	-0.1/98/5D-02	
		-0.303973D-02	0.146542D-02	0.132960D-03	
	39D-01	2.22227.22 02		3.2029002 00	
27	0.433836D-02	-0.288046D-01	0.232948D-01	-0.654212D-02	
0.1691	47D-01				
		0.533248D-04	0.109572D-02	-0.284565D-03	-
	11D+00				
	0.129614D-02	U.143053D-03	-0.107556D-02	-0.140643D-03	-
	25D-01	0.299086D-02			
	61D-02	0.299080D-02	-0.3030370-02	0.1/01400-02	
		-0.165446D-02	0.790787D-03	-0.402583D-03	_
	93D+00				
		0.120388D-02	-0.437436D-03	0.441979D-03	-
0.3523	90D-01				
	0.784960D-03	0.518451D-02	-0.287046D-02	-0.353566D-04	-
0.1703	34D-02				

34 -0.404923D-03	0.544985D-03	-0.393999D-03	-0.193777D-03	-
0.159136D-01				
35 -0.134071D-03	0.476723D-03	-0.216469D-03	-0.145765D-04	-
0.913023D-02				
36 0.525107D-04	-0.889925D-04	0.404284D-04	0.606303D-05	
0.341239D-02				
37 -0.154557D-03	-0.120062D-02	0.561504D-03	-0.140894D-03	-
0.148733D-01				
38 0.221867D-04	-0.772191D-03	0.551067D-03	0.588108D-04	-
0.979896D-02				
39 0.319495D-05	-0.348196D-03	0.111184D-03	0.00000D+00	-
0.140187D-02				
26	27	28	29	
30				
26 0.963939D+00				
	0.274839D+00			
28 -0.250020D-01				
29 -0.712441D+00				
30 0.139093D+00	-0.958204D-01	-0.128779D-02	-0.166675D+00	
0.597674D-01	0 0010655 00	0 10000000 01	0 100000 01	
31 -0.229564D-01	-0.9/1065D-03	0.128/33D-01	-0.1/6033D-01	
0.230323D-02	0 0100010 00	0 4110600 01		
32 -0.955278D-01	-0.213081D-02	-0.411860D-01	-0.404/55D-01	
0.983449D-02	0 01 00000 01	0 7220000 00	0 1000010 01	
33 -0.625441D-03	-0.916082D-01	0.733890D-02	0.102881D-01	
0.165987D-01	0 000000 01	0 0707000 00	0 1000710 00	
34 0.375541D-02	-0.23622/D-01	0.2/9/08D-02	0.1898/1D-02	
0.253504D-02	0 1067000 01	0 1010600 00		
35 0.291052D-02 0.269836D-02	-0.196720D-01	0.121268D-02	0.645698D-03	
36 0.241069D-04	0 2770100 02	0 6774270 04	-0.153527D-03	
0.298439D-03	0.3770100-02	0.0774270-04	-0.1333270-03	
37 0.755519D-02	0 2101/30-01	0 1984560-02	0.731084D-03	_
0.270887D-02	0.2101450-01	0.1904900-02	0.751004D-05	
	0 193964D-01	0 6888270-03	-0.803797D-03	_
0.303236D-02	0.1999010 01	0.0000270 03	0.003/9/2 03	
39 0.698107D-03	0.6371520-03	-0.646684D-04	0.105105D-03	
0.868158D-03	0.00,1020 00	J. JI J J J J J J J J J J J J J J J J J	J. 1051050 05	
31	32	33	34	
35			<u> </u>	
31 0.566030D+00				
32 -0.433064D-01	0.452808D+00			
33 0.428433D-02	0.632470D-02	0.543893D+00		
34 -0.122989D+00		-0.133908D+00	0.135731D+00	
	-0.490156D-01	0.269710D-01		
0.476775D-01				
36 -0.128140D+00	0.231541D-01	-0.237214D+00	0.138408D+00	-
0.254654D-01				
37 -0.135300D+00	0.438656D-01	0.132225D+00	0.115263D-01	_
0.518961D-02				
	-0.670738D-01	-0.610064D-01	-0.164998D-02	
0.205644D-02				
	-0.603077D-01	-0.206842D+00	0.177404D-01	-
0.848745D-02				
36	37	38	39	
ſ				

```
36 0.255941D+00

37 -0.156444D-01 0.147488D+00

38 0.291891D-02 -0.490771D-01 0.679189D-01

39 -0.212271D-01 -0.135405D+00 0.647460D-01 0.223940D+00

SCALE

0.89

END
```

```
      Table 34: Parameterization script (QM): Bond, angle and torsion based on QM calculated force constants for CPD
```

```
* From tutorial on A. MacKerell's website
* Modified by Cassian
* Quick measurements and molvib
*
ioformat extended
set pdbdir .
!### Set Variables ###
set residue cpd
set zeed 1 C1 1 C2 1 C3 ! seed
set frist none ! first capping residue
set lsat none ! last capping residue
set bombover 0 !needs to be set to -1 for 3-membered rings
!### Things that usually should not be changed. ####
set dontmini 0 ! Do not minimize main conformation. Should normally
not be used.
open unit 10 read card name top all36 cgenff @residue.rtf
read rtf card unit 10
close unit 10
open unit 10 read card name par all36 cgenff @residue.prm
read para card flex unit 10
close unit 10
! Read in CPD parameter file in command line
if @?str eq 1 stream @str
read sequence card
* @residue
*
1
@residue
bomlev @bombover
generate @residue first @frist last @lsat setup warn
bomlev 0
```

```
ic param
   ic seed @zeed
   ic build
   ic purge
   ic print
! Write out initial structure for testing and debugging purposes
   open unit 30 write form name @residue init.pdb
   write coor pdb unit 30
   close unit 30
!minimize structure and calculate dipole moment
if @dontmini eq 1 goto skip min
   MINI CONJ nstep 200 nprint 20 inbfrq 1000 cutnb 999.
   MINI NRAP nstep 50 tolgrd 0.00001
   prnlev 3
   coor dipole oxyz select segid @residue end
!save minimized geometry
   coor print
   open unit 30 write form name @residue min.crd
   write coor card unit 30
   open unit 30 write form name @pdbdir/@residue min.pdb
   write coor pdb unit 30
   close unit 30
label skip min
!mp2/6-31g* data
!
!C1 -O1 1.2215
quick 1 2
!C4 -O4 1.2215
quick 9 10
!C1 -C2 1.5284
quick 1 3
!C3 -C4 1.5284
quick 6 9
!C1 -C5 1.5217
quick 1 11
!C4 -C5 1.5217
quick 9 11
!C2 -C3 1.5309
quick 3 6
!C2 -H21 1.0978
quick 3 4
!C3 -H32 1.0978
quick 6 8
!C2 -H22 1.0928
quick 3 5
!C3 -H31 1.0928
quick 6 7
!C5 -H51 1.0964
guick 11 12
!C5 -H52 1.0964
```

quick 11 13 !A1 A(2,1,3) 01 - C1 - C2 125.0815 quick 2 1 3 01 - C1 - C5 125.7255 !A2 A(2,1,11) quick 2 1 11 C2 - C1 - C5 109.193 !A3 A(3,1,11) quick 3 1 11 !A4 A(1,3,4) C1 - C2 - H21 107.5191 quick 1 3 4 C1 - C2 - H22 110.1684 !A5 A(1,3,5) quick 1 3 5 C1 - C2 - C3 105.4448 !A6 A(1,3,6) quick 1 3 6 H21 - C2 - H22 107.2716 !A7 A(4,3,5) quick 4 3 5 H21 - C2 - C3 111.1477 !A8 A(4,3,6) quick 4 3 6 H22 - C2 - C3 115.0749 !A9 A(5,3,6) quick 5 3 6 C2 - C3 - H31 115.075 !A10 A(3,6,7) quick 3 6 7 !A11 A(3,6,8) C2 - C3 - H32 111.1476 quick 3 6 8 C2 - C3 - C4 105.4446 !A12 A(3,6,9) quick 3 6 9 !A13 A(7,6,8) H31 - C3 - H32 107.2717 quick 7 6 8 !A14 A(7,6,9) H31 - C3 - C4 110.1685 quick 7 6 9 !A15 A(8,6,9) H32 - C3 - C4 107.5191 quick 8 6 9 !A16 A(6,9,10) C3 - C4 - O4 125.0816 quick 6 9 10 !A17 A(6,9,11) C3 - C4 - C5 109.1928 quick 6 9 11 !A18 A(10,9,11) 04 - C4 - C5 125.7256 quick 10 9 11 !A19 A(1,11,9) C1 - C5 - C4 105.3362 quick 1 11 9 !A20 A(1,11,12) C1 - C5 - H51 111.1703 quick 1 11 12 !A21 A(1,11,13) C1 - C5 - H52 111.2602 quick 1 11 13 !A22 A(9,11,12) C4 - C5 - H51 111.2609 quick 9 11 12 !A23 A(9,11,13) C4 - C5 - H52 111.1697 quick 9 11 13 !A24 A(12,11,13) H51 - C5 - H52 106.7286 quick 12 11 13 !must be easily greppable (grep EXTRA) in output file !echu !D3 D(3,6,9,11) C2-C3-C4-C5 19.5145 qv quick 3 6 9 11

```
echo extra: C2-C3-C4-C5 19.5145 gv
echo extra: 3 6 9 11 ?phi
!D4
     D(6,9,11,1) C3-C4-C5-C1 -7.5311 gv
quick 6 9 11 1
echo extra: C3-C4-C5-C1 -7.5311 qv
echo extra: 6 9 11 1 ?phi
     D(3,1,11,9) C4-C5-C1-C2 -7.5267 qv
!D5
quick 3 1 11 9
echo extra: C4-C5-C1-C2 -7.5267 qv
echo extra: 3 1 11 9 ?phi
!D1
     D(11,1,3,6) C5-C1-C2-C3 19.5117 gv
quick 11 1 3 6
echo extra: C5-C1-C2-C3 19.5117 qv
echo extra: 11 1 3 6 ?phi
!D2
     D(1,3,6,9) C1-C2-C3-C4 -23.4181 gv
quick 1 3 6 9
echo extra: C1-C2-C3-C4 -23.4181 gv
echo extra: 1 3 6 9 ?phi
     D(1,3,11,2) C1-C2-C5-O1 -0.036 qv
!D2
quick 1 3 11 2
echo extra: C1-C2-C5-O1 -0.036 qv
echo extra: 1 3 11 2 ?phi
vibran
diag
end
!MOLVIB analysis by CHARMm
!Although the minimum is slightly asymmetric, the parameters are
!perfectly symmetric, so we use a symmetric molvib analysis.
! NOTE: The MOLVIB module is very sensitive to the format
MOLVIB NDI1 33 NDI2 33 NDI3 44 SECO
GFX
PRNT
        1
DIM
       33
            33
                 44
IC
 1
      2 0 0 !1 R1
                        R(1,2)
                                    C1 = 01
   1
                                    C1 - C2
 1
   1
      3 0 0 !2 R2
                        R(1,3)
   1 11
 1
         0 0 !3 R3
                        R(1, 11)
                                    C1 - C5
                                    C2 - H21
 1
   3
      4
         0 0 !4 R4
                        R(3,4)
                                    C2 - H22
            0 !5 R5
 1
    3
      5
          0
                        R(3,5)
 1
    3
       6
          0
            0 !6 R6
                        R(3,6)
                                    C2 - C3
 1
    6
       7
          0
            0 !7 R7
                        R(6,7)
                                    C3 - H31
    6
       8
          0
                            C3 - H32
 1
            0 !8 R8 R(6,8)
 1
    6
      9
          0
            0 !9 R9
                        R(6,9)
                                   C3 - C4
 1
    9 10
         0
            0 !10 R10 R(9,10)
                                    C4 = 04
 1
   9 11
          0 0 !11 R11 R(9,11)
                                    C4 - C5
                                    C5 - H51
 1 11 12
          0 0 !12 R12 R(11,12)
 1 11 13
          0
            0 !13 R13 R(11,13)
                                    C5 - H52
                                    01 - C1 - C2
 2
    2
      1
         3
            0 !14 A1
                        A(2,1,3)
                                                    1b2
 2
    2
      1 11 0 !15 A2
                        A(2, 1, 11)
                                    01 - C1 - C5
                                                   1b1
 2
   3
                                    C2 - C1 - C5
      1 11
            0 !16 A3
                        A(3,1,11)
                                                    rla4
      3 4 0 !17 A4
                        A(1,3,4)
                                    C1 - C2 - H21
                                                   2b3
 2
   1
 2
   1 3 5 0 !18 A5
                        A(1,3,5)
                                 C1 - C2 - H22
                                                   2b4
```

2 1 3 6 0 !19		C1 - C2 - C3	rla5 gamma
2 4 3 5 0 ! 20		H21 - C2 - H22	2 alpha
2 4 3 6 0 !21		H21 - C2 - C3	2b1
2 5 3 6 0 !22		H22 - C2 - C3	2b2
2 3 6 7 0 !23	A10 A(3,6,7)	C2 - C3 - H31	3b3
2 3 6 8 0 !24	A11 A(3,6,8)	C2 - C3 - H32	3b4
2 3 6 9 0 !25	A12 A(3,6,9)	C2 - C3 - C4	rlal gamma
2 7 6 8 0 ! 26	A13 A(7,6,8)	H31 - C3 - H32	3 alpha
2 7 6 9 0 !27	A14 A(7,6,9)	H31 - C3 - C4	3b1
2 8 6 9 0 !28	A15 A(8,6,9)	H32 - C3 - C4	3b2
2 6 9 10 0 !29	A16 A(6,9,10)	C3 - C4 - O4	4b1
2 6 9 11 0 !30		C3 - C4 - C5	rla2
2 10 9 11 0 !31	A18 A(10,9,11)	04 - C4 - C5	4b2
2 1 11 9 0 !32		C1 - C5 - C4	rla3 gamma
2 1 11 12 0 !33		C1 - C5 - H51	5b1
2 1 11 13 0 !34		C1 - C5 - H52	5b2
2 9 11 12 0 !35		C4 - C5 - H51	5b3
	A23 A(9,11,13)	C4 - C5 - H52	5b4
		H51 - C5 - H52	5 alpha
3 1 3 11 2 !38		C1-C2-C5-O1	C1
3 9 11 6 10 !39) C5-C1-C2-O4	C4
4 11 1 3 6 !40		C5-C1-C2-C3	r1t4
4 1 3 6 9 !41		C1-C2-C3-C4	r1t5
4 3 6 9 11 !42		C2-C3-C4-C5	r1t1
4 6 9 11 1 !43		C3-C4-C5-C1	r1t2
4 9 11 1 3 !44		C4-C5-C1-C2	r1t3
$\begin{array}{ccc} $		w normalization	1105
	2 2 1.		
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	4 5 1.	3 3 1.	
5 4 1.	5 4 -1.		
6 6 1.	5 4 1.		
7 7 1.	781.		
8 7 1.	8 8 -1.		
991.	10 10 1.		
	10 10 1.		
	12 13 1.		
12 12 1.			
13 12 1.	13 13 -1.		
14 19 1.	14 20 5.	15 01 1	1 5 0 0 1
15 17 1.	15 18 -1.	15 21 1.	15 22 -1.
16 17 -1.	16 18 -1.	$16\ 21\ 1.$	$16\ 22\ 1.$
17 17 -1.	17 18 1.	17 21 1.	17 22 -1.
18 25 1.	18 26 5.	10 07 1	10 00 1
19 23 1.	19 24 -1.	19 27 1.	19 28 -1.
20 23 -1.	20 24 -1.	20 27 1.	20 28 1.
21 23 -1.	21 24 1.	21 27 1.	21 28 -1.
22 32 1.	22 37 5.	00 0 5 -	
23 33 1.	23 34 -1.	23 35 1.	23 36 -1.
24 33 1.	24 34 1.	24 35 -1.	24 36 -1.
25 33 1.	25 34 -1.	25 35 -1.	25 36 1.
26 38 1.			
27 15 1.	27 14 -1.		
28 39 1.			
	29 31 -1.		
30 25 1.	30 30 -0.81	30 32 0.31	30 16 0.31
30 19 -0.81			
JU IJ 0.01			

. <u> </u>						
	30 -1.12	31 32	1.81		16 -1.81	31 19 1.12
	42 0.31	32 43	-0.81	32	44 1.	32 40 -0.81
	41 0.31					
	42 -1.81	33 43	1.12	33	40 -1.12	33 41 1.81
-1						
PED	0 15					
1	sC1=01					
2	sC1-C2					
3	sC1-C5					
4	sC2-H2sy					
5	sC2-H2as					
6	sC2-C3					
7	sC3-H3sy					
8	sC3-H3as					
9	sC3-C4					
10	sC4=04					
11	sC4-C5					
12	sC5-H5sy					
13	sC5-H5as					
14	sc-C2H2					
15	r-C2H2					
16	w-C2H2					
17	t-C2H2					
18	sc-C3H2					
19	r-C3H2					
20	w-C3H2					
21	t-C3H2					
22	sc-C5H2					
23	r-C5H2					
24	w-C5H2					
25	t-C5H2					
26	w-C1=01					
27	d-C1=01					
28	w-C4=O4					
29	d-C4=04					
30	d-5RNGsy					
31	d-5RNGas					
32	t-5RNGsy					
33	t-5RNGas					
-1						
END						

Table 35: Parameterization script (MM): Bond, angle and torsion parameter optimization for CPD

```
* From tutorial on A. MacKerell's website
* Modified by Cassian
* Potential energy scan for dihedrals
*
ioformat extended
```

```
set residue cpd
set basnam scan cpd
set zeed 1 C1 1 C2 1 C3 ! seed
set frist none ! first capping residue
set lsat none ! last capping residue
set pesbas ..
set TOPPAR ..
set peses 4
set pes1 gausst1
set pes2 gausst2
set pes3 gausst3
set pes4 gausst4
set cons 7
set con1 1 C1 1 C2 1 C3 1 C4
set con2 1 C2 1 C3 1 C4 1 C5
set con3 1 C3 1 C4 1 C5 1 C1
set con4 1 C2 1 C1 1 C5 1 C4
set con5 1 C5 1 C1 1 C2 1 C3
set con6 1 C1 1 C2 1 C5 1 O1
set con7 1 C4 1 C5 1 C3 1 O4
set meas 11
set meal @con1
set mea2 @con2
set mea3 @con3
set mea4 @con4
set mea5 @con5
set mea6 1 C3 1 C2 1 C1 1 O1
set mea7 1 C4 1 C5 1 C1 1 O1
set mea8 1 C2 1 C3 1 C4 1 O4
set mea9 1 C1 1 C5 1 C4 1 O4
set meal0 1 01 1 C1 1 C2 1 H21
set meall 1 C5 1 C1 1 C2 1 H21
set offset 31
open unit 31 append form name @basnam.clccc4
open unit 32 append form name @basnam.ccc4c
open unit 33 append form name @basnam.cc4cc1
open unit 34 append form name @basnam.c4cc1c
open unit 35 append form name @basnam.cc1cc
open unit 36 append form name @basnam.ccclo
open unit 37 append form name @basnam.c4cc10
open unit 38 append form name @basnam.ccc4o
open unit 39 append form name @basnam.clcc4o
open unit 40 append form name @basnam.oc1ch
open unit 41 append form name @basnam.cc1ch
set eun 29
open unit 29 append form name @basnam.mme
open unit 10 read card name @TOPPAR/top all36 cgenff cpd.rtf
read rtf card unit 10
close unit 10
open unit 10 read card name @TOPPAR/par all36 cgenff cpd.prm
read para card flex unit 10
close unit 10
```

```
if @?str eq 1 stream @str
open unit 10 write form name ini geom.pdb
open unit 11 write form name fin geom.pdb
read sequence card
* @residue
*
1
@residue
generate @residue first @frist last @lsat setup warn
! patch @residue @residue 1 setup warn
! autogenerate angles dihedral
set scan 1
label loop scan
set pesdir @pesbas/@pes@@scan
open unit 25 read form name @pesdir/s file.txt
label loop_pes
get pdbnam unit 25
if "@PDBNAM" eq "" goto nextscan
open unit 27 read form name @pdbnam
read coor pdb unit 27
close unit 27
write coor pdb unit 10
set cond 1
label loop cond
cons dihe @con@@cond FORCE 9999. MAIN PERIOD 0
incr cond
if @cond le @cons goto loop cond
MINI CONJ nstep 50 nprint 20 inbfrg 1000 cutnb 999.
MINI NRAP nstep 100 tolgrd 0.00001
write coor pdb unit 11
echu @eun
cons cldh
energy
echo ?ener
if @meas eq 0 goto loop_pes
set mead 1
set mun @offset
label loop mead
echu @mun
quick @mea@@mead
echo ?phi
```

```
incr mun
incr mead
if @mead le @meas goto loop_mead
goto loop_pes
label nextscan
close unit 25
incr scan
if @scan le @peses goto loop_scan
```

Table 36: Parameterization script (MM): Potential energy scans of dihedrals and impropers for CPD

A3 Scripts used for setup and run of molecular dynamic simulations

```
* Structure Optimization
ioformat extended
set pdbdir .
!### Set variables ###
set residue cpd-hocl
set frist none ! first capping residue
set lsat none ! last capping residue
set bombover 0 !needs to be set to -1 for 3-membered rings
open unit 10 read form name top_all22_prot.rtf
read rtf card unit 10
close unit 10
open unit 10 read form name par all22 prot.prm
read param card flex unit 10
close unit 10
open unit 10 read card name top all36 cgenff remas.rtf
read rtf card unit 10 append
close unit 10
open unit 10 read card name par all36 cgenff remas.prm
read para card flex unit 10 append
close unit 10
! May need dummy atom to generate HOCl
stream toppar_dum_nobel_gases.str
if @?str eq 1 stream @str
read sequence card
* CPD
*
1
CPD
bomlev @bombover
generate CPD first @frist last @lsat setup warn
read sequence card
* HOCL
*
1
HOCL
bomlev @bombover
generate HOCL first @frist last @lsat setup warn
open unit 4 card read name cpd-hocl.crd
read coor card unit 4
```

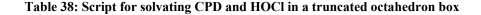
```
coor print
  open unit 30 write form name @residue init.pdb
  write coor pdb unit 30
  close unit 30
!minimize structure and calculate dipole moment
  MINI CONJ nstep 200 nprint 20 inbfrg 1000 cutnb 999.
  MINI NRAP nstep 50 tolgrd 0.00001
!save minimized geometry
  coor print
  open unit 30 write form name @residue_min.psf
  write psf card unit 30
  close unit 30
  open unit 30 write form name @residue min.crd
  write coor card unit 30
  open unit 30 write form name @residue min.pdb
  write coor pdb unit 30
  close unit 30
stop
```

Table 37: Initial minimization script for CPD and HOCI

```
*NAME: Solvate.inp
*PURPOSE: build energetically-adjusted ion-water environment around
protein
*AUTHOR: Yeo H.C.Modified by Cassian
ioformat extended
set pdbdir .
!### Set variables ###
set residue cpd-hocl
set frist none ! first capping residue
set lsat none ! last capping residue
set bombover 0 !needs to be set to -1 for 3-membered rings
open unit 10 read form name top all22 prot.rtf
read rtf card unit 10
close unit 10
open unit 10 read form name par_all22_prot.prm
read param card flex unit 10
close unit 10
open unit 10 read card name top_all36_cgenff_remas.rtf
```

```
read rtf card unit 10 append
close unit 10
open unit 10 read card name par all36 cgenff remas.prm
read para card flex unit 10 append
close unit 10
! May need dummy atom to generate HOCl
stream toppar dum nobel gases.str
if @?str eq 1 stream @str
open read formatted unit 3 name cpd-hocl min.psf
read psf card unit 3
open read formatted unit 4 name cpd-hocl min.crd
read coor card unit 4
ener
! find out where protein is and put it with center at origin
coor stat sele segi cpd .or. segid hocl end
! orient molecule with moments of inertia along x,y,z-axes
! everything will be translated/rotated
coor orie sele segid cpd .or. segid hocl end
coor stat sele segi cpd .or. segid hocl end
! Add water; here no ions are added, this should perhaps also be
done
! Box size could be autmatically determined here by adding say 8A on
each side to the
! dimensions of the protein, we also make the sizes integer numbers
(not necessary)
calc XSIZ = INT( ?XMAX - ?XMIN + 16)
calc YSIZ = INT( ?YMAX - ?YMIN + 16)
calc ZSIZ = INT( ?ZMAX - ?ZMIN + 16)
! Set size based on previous calculation
set XSIZ 24
set YSIZ 24
set ZSIZ 24
define SOLUTE sele segid cpd .or. segid hocl end
!Read the default water box and modify it
stream mod-solvent-box.str
! Variable A is used in the stream file
set A @XSIZ
! slice box into truncated-octahedral
stream setup-truncated-octahedron.str
open write card unit 1 name temp-oct.pdb
write coor pdb unit 1
* octahedron
```

```
coor conv align symm @ax @ax @ax @b @b @b !do not put this in
dynamics !!
! save the PSF and coordinates for later use
open write formatted unit 27 name cpd-hocl-solv.psf
write psf card unit 27
*
  with crystal waters solvated with TIP3 waters
open write formatted unit 27 name cpd-hocl-solv.crd
write coor card unit 27
* Energy Minimized Coordinate Set for @FILE
! For viewing in VMD
open unit 1 write card name cpd-hocl-solv.pdb
write coor pdb unit 1
* @FILE for viewing in VMD
* in rectangular box: @XSIZ x @YSIZ x @ZSIZ
stop
```



```
* CHARMM script for initial minimization
! Read in pre-defined variables
stream datadir.def
! Read initial data files
open unit 10 read form name @Otop all22 prot.rtf
read rtf card unit 10
close unit 10
open unit 10 read form name @0par_all22_prot.prm
read param card flex unit 10
close unit 10
open unit 10 read card name @Otop all36 cgenff remas.rtf
read rtf card unit 10 append
close unit 10
open unit 10 read card name @Opar all36 cgenff remas.prm
read para card flex unit 10 append
close unit 10
if @?str eq 1 stream @str
open read card unit 1 name @0cpd-hocl-solv.psf
read psf card unit 1
```

```
open read card unit 1 name @0cpd-hocl-solv.crd
read coor card unit 1
calc xsiz 24. * sqrt(0.75)
calc ysiz 24. * sqrt(0.75)
calc zsiz 24. * sqrt(0.75)
set angle 109.4712206344907
crystal define octahedral @XSIZ @YSIZ @ZSIZ @angle @angle @angle
crystal build cutoff 14.0
image byresidue sele resn tip3 end xcen 0.0 ycen 0.0 zcen 0.0
image bysegment sele segi cpd .or. segid hocl end xcen 0.0 ycen 0.0
zcen 0.0
! Specify nonbond parameters and evaluate energy once
nbonds cdie eps 1.0 atom vdw vswitch cutnb 12.0 ctofnb 10.0 ctonnb
7.0 -
 cutim 14.0 wmin 1.5 -
 ewald pmewald kappa 0.34 fftx 24 ffty 24 fftz 24 order 6
ener
mini sd nstep 200 nprint 10
mini abnr nstep 200 nprint 10
open write card unit 1 name @1/init mini.crd
write coor card unit 1
* init mini
stop
```

Table 39: Initial minimization script for solvated CPD and HOCl

```
* CHARMM script for initialization of NVE dynamics
*
! Read in pre-defined variables
stream datadir.def
! Read initial data files
open unit 10 read form name @Otop_all22_prot.rtf
read rtf card unit 10
close unit 10
open unit 10 read form name @Opar_all22_prot.prm
read param card flex unit 10
close unit 10
open unit 10 read card name @Otop_all36_cgenff_remas.rtf
read rtf card unit 10 append
close unit 10
```

```
open unit 10 read card name @0par all36 cgenff remas.prm
read para card flex unit 10 append
close unit 10
if @?str eq 1 stream @str
open read card unit 1 name @0cpd-hocl-solv.psf
read psf card unit 1
open read card unit 1 name @0cpd-hocl-solv.crd
read coor card unit 1
calc xsiz 24. * sqrt(0.75)
calc ysiz 24. * sqrt(0.75)
calc zsiz 24. * sqrt(0.75)
set angle 109.4712206344907
crystal define octahedral @XSIZ @YSIZ @ZSIZ @angle @angle @angle
crystal build cutoff 14.0
image byresidue sele resn tip3 end xcen 0.0 ycen 0.0 zcen 0.0
image bysegment sele segi cpd .or. segid hocl end xcen 0.0 ycen 0.0
zcen 0.0
! Specify nonbond parameters and evaluate energy once
nbonds cdie eps 1.0 atom vdw vswitch cutnb 12.0 ctofnb 10.0 ctonnb
7.0 -
  cutim 14.0 wmin 1.5 -
  ewald pmewald kappa 0.34 fftx 24 ffty 24 fftz 24 order 6
ener
open read card unit 1 name @1/init_mini.crd
read coor card unit 1
! Open files for writing dynamics data.
open write card unit 31 name dyn.rst
open write file unit 41 name dyn.trj
open write file unit 51 name dyn.vel
! Dynamics command with options described
DYNAMICS -
- ! algorithm
 LEAP -
  VERLET - !
 - ! start/restart/initialization info
  START - ! START for initial run; RESTART to pick up from previous
run
  IASORS 1 - ! specifies scaling (0) or assigning (other value) of
velocities
  IASVEL 1 - ! species method of assigning velocities during
heating
  ISCVEL 0 - ! specifies method of scaling velocities (0=same
factor)
 - ! time step info
                - ! number of steps
  NSTEP 50000
  TIMESTEP 0.001 - ! timestep in picoseconds
```

```
IPRFRQ 1000
                  - ! frequency for calculating averages and
fluctuations
                  - ! (must be at least as large as NTRFRQ unless
NTRFRO=0)
 - ! heating info
  IHTFRQ 2000 - ! heating frequency
  FIRSTT 98.0 - ! initial temperature
  FINALT 298.0 - ! final temperature
  TEMINC 100.0 - !temperature increment
 - ! equilibration info
  IEQFRQ 5000 - ! frequency for assigning or scaling velocities
(during equil.)
               - ! every IEQFREQ steps, always change velocities (0)
  ICHECW 1
               - ! or change only if temperature is outside window
(other value)
  TWINDH 10.0 - ! upper temperature window limit
  TWINDL -10.0 - ! lower temperature window limit
  NTRFRQ 5000 - ! frequency for removing overall translation and
rotation
 -! io units
  IUNREA -1 - ! unit number for reading restart info (-1 means no
restart)
  IUNWRI 31 - ! unit number for writing restart info
  IUNCRD 41 - ! unit number for writing coordinates (coordinate
trajectory)
  IUNVEL 51 - ! unit number for writing velocities (velocity
trajectory)
  IUNIT -1 - ! unit number to which to write miscellaneous
information
 - ! printing & writing directives
  NPRINT 100 - ! frequency for printing output info
  NSAVC 20 - ! frequency for saving coordinates
 NSAVV 20 - ! frequency for saving velociti4es
 - ! energy
 NBONDS cdie eps 1.0 atom vdw vswitch cutnb 12.0 ctofnb 10.0 ctonnb
7.0 -
         cutim 14.0 wmin 1.5 -
  EWALD pmewald kappa 0.34 fftx 24 ffty 24 fftz 24 order 6 -
             ! frequency for undating nonbond list
  INBFRQ 25
open write card unit 1 name init nve.crd
write coor card unit 1
* Coordinates for simulation after
* initial NVE heating of 50,000 steps (0.002 ps time step)
* from 98 K to 298 K. This is preparation
* for a subsequent NPT simulation.
  stop
```

Table 40: Initial NVE dynamic simulation script (slowly heating the system to 298 K)

```
* CHARMM script for initialization of NPT dynamics
! Read in pre-defined variables
stream datadir.def
! Read initial data files
open unit 10 read form name @Otop all22 prot.rtf
read rtf card unit 10
close unit 10
open unit 10 read form name @Opar all22 prot.prm
read param card flex unit 10
close unit 10
open unit 10 read card name @Otop all36 cgenff remas.rtf
read rtf card unit 10 append
close unit 10
open unit 10 read card name @Opar all36 cgenff remas.prm
read para card flex unit 10 append
close unit 10
if @?str eq 1 stream @str
open read card unit 1 name @0cpd-hocl-solv.psf
read psf card unit 1
open read card unit 1 name @0cpd-hocl-solv.crd
read coor card unit 1
calc xsiz 24. * sqrt(0.75)
calc ysiz 24. * sqrt(0.75)
calc zsiz 24. * sqrt(0.75)
set angle 109.4712206344907
crystal define octahedral @XSIZ @YSIZ @ZSIZ @angle @angle @angle
crystal build cutoff 14.0
image byresidue sele resn tip3 end xcen 0.0 ycen 0.0 zcen 0.0
image by segment sele seqi cpd .or. seqid hocl end xcen 0.0 ycen 0.0
zcen 0.0
! Specify nonbond parameters and evaluate energy once
nbonds cdie eps 1.0 atom vdw vswitch cutnb 12.0 ctofnb 10.0 ctonnb
7.0 -
  cutim 14.0 wmin 1.5 -
  ewald pmewald kappa 0.34 fftx 24 ffty 24 fftz 24 order 6
ener
open read card unit 1 name @1/init mini.crd
read coor card unit 1
open read card unit 1 name init nve.crd
read coor card unit 1
ener
! Open files for writing dynamics data.
open write card unit 31 name dyn.rst
```

```
open write file unit 41 name dyn.trj
open write file unit 51 name dyn.vel
! Dynamics command with options described
DYNAMICS -
 - ! algorithm
 CPT - !
 - ! start/restart/initialization info
  START - ! START for initial run; RESTART to pick up from previous
run
  IASORS 1 - ! specifies scaling (0) or assigning (other value) of
velocities
  IASVEL 1 - ! species method of assigning velocities during
heating
  ISCVEL 0 - ! specifies method of scaling velocities (0=same
factor)
 - ! time step info
  NSTEP 500000 - ! number of steps
  TIMESTEP 0.001 - ! timestep in picoseconds
                - ! frequency for calculating averages and
  IPRFRQ 1000
fluctuations
                  - ! (must be at least as large as NTRFRQ unless
NTRFRO=0)
 - ! heating info
  IHTFRQ 2000 - ! heating frequency
  FIRSTT 298.0 - ! initial temperature
 FINALT 298.0 - ! final temperature
  TEMINC 0.0 - !temperature increment
 - ! cpt parameters
  PCONst pmass 400.0 pgamma 20.0 tbath 298.0 -
 - ! equilibration info
  IEQFRQ 5000 - ! frequency for assigning or scaling velocities
(during equil.)
  ICHECW 1
              - ! every IEQFREQ steps, always change velocities (0)
               - ! or change only if temperature is outside window
(other value)
  TWINDH 10.0 - ! upper temperature window limit
  TWINDL -10.0 - ! lower temperature window limit
  NTRFRQ 5000 - ! frequency for removing overall translation and
rotation
 -! io units
  IUNREA -1 - ! unit number for reading restart info (-1 means no
restart)
  IUNWRI 31 - ! unit number for writing restart info
  IUNCRD 41 - ! unit number for writing coordinates (coordinate
trajectory)
  IUNVEL 51 - ! unit number for writing velocities (velocity
trajectory)
  IUNIT -1 - ! unit number to which to write miscellaneous
information
 - ! printing & writing directives
  NPRINT 100 - ! frequency for printing output info
  NSAVC 20 - ! frequency for saving coordinates
  NSAVV 20 - ! frequency for saving velociti4es
 - ! energy
```

Table 41: Initial NPT dynamic simulation script

```
* CHARMM script for NPT dynamics
! Read in pre-defined variables
stream datadir.def
! Read initial data files
open unit 10 read form name @Otop all22 prot.rtf
read rtf card unit 10
close unit 10
open unit 10 read form name @0par_all22_prot.prm
read param card flex unit 10
close unit 10
open unit 10 read card name @Otop all36 cgenff remas.rtf
read rtf card unit 10 append
close unit 10
open unit 10 read card name @Opar all36 cgenff remas.prm
read para card flex unit 10 append
close unit 10
if @?str eq 1 stream @str
open read card unit 1 name @0cpd-hocl-solv.psf
read psf card unit 1
open read card unit 1 name @0cpd-hocl-solv.crd
read coor card unit 1
calc xsiz 24. * sqrt(0.75)
calc ysiz 24. * sqrt(0.75)
calc zsiz 24. * sqrt(0.75)
set angle 109.4712206344907
crystal define octahedral @XSIZ @YSIZ @ZSIZ @angle @angle @angle
crystal build cutoff 14.0
image byresidue sele resn tip3 end xcen 0.0 ycen 0.0 zcen 0.0
image bysegment sele segi cpd .or. segid hocl end xcen 0.0 ycen 0.0
zcen 0.0
! Specify nonbond parameters and evaluate energy once
nbonds cdie eps 1.0 atom vdw vswitch cutnb 12.0 ctofnb 10.0 ctonnb
```

```
7.0 -
  cutim 14.0 wmin 1.5 -
  ewald pmewald kappa 0.34 fftx 24 ffty 24 fftz 24 order 6
ener
open read card unit 1 name @1/init mini.crd
read coor card unit 1
! Open files for writing dynamics data.
open read card unit 30 name dyn.rea
open write card unit 31 name dyn.rst
open write file unit 41 name dyn.trj
open write file unit 51 name dyn.vel
! Dynamics command with options described
DYNAMICS -
- ! algorithm
 CPT - !
 - ! start/restart/initialization info
 REST - ! START for initial run; RESTART to pick up from previous
run
  IASORS 0 - ! specifies scaling (0) or assigning (other value) of
velocities
  IASVEL 1 - ! species method of assigning velocities during
heating
  ISCVEL 0 - ! specifies method of scaling velocities (0=same
factor)
 - ! time step info
 NSTEP 500000 - ! number of steps
  TIMESTEP 0.001 - ! timestep in picoseconds
  IPRFRQ 1000 - ! frequency for calculating averages and
fluctuations
                  - ! (must be at least as large as NTRFRQ unless
NTRFRQ=0)
 - ! heating info
 IHTFRQ 2000 - ! heating frequency
 FIRSTT 298.0 - ! initial temperature
 FINALT 298.0 - ! final temperature
 TEMINC 0.0 - !temperature increment
- ! cpt parameters
 PCONst pmass 400.0 pgamma 20.0 tbath 298.0 -
 - ! equilibration info
  IEQFRQ 5000 - ! frequency for assigning or scaling velocities
(during equil.)
  ICHECW 1
               - ! every IEQFREQ steps, always change velocities (0)
               - ! or change only if temperature is outside window
(other value)
  TWINDH 10.0 - ! upper temperature window limit
  TWINDL -10.0 - ! lower temperature window limit
 NTRFRQ 5000 - ! frequency for removing overall translation and
rotation
 -! io units
  IUNREA 30 - ! unit number for readiing restart info (-1 means no
restart)
```

```
IUNWRI 31 - ! unit number for writing restart info
  IUNCRD 41 - ! unit number for writing coordinates (coordinate
trajectory)
  IUNVEL 51 - ! unit number for writing velocities (velocity
trajectory)
  IUNIT -1 - ! unit number to which to write miscellaneous
information
 - ! printing & writing directives
 NPRINT 100 - ! frequency for printing output info
 NSAVC 20 - ! frequency for saving coordinates
 NSAVV 20 - ! frequency for saving velociti4es
 - ! energy
 NBONDS cdie eps 1.0 atom vdw vswitch cutnb 12.0 ctofnb 10.0 ctonnb
7.0 -
         cutim 14.0 wmin 1.5 -
  EWALD pmewald kappa 0.34 fftx 24 ffty 24 fftz 24 order 6 -
  INBFRQ 25
             ! frequency for undating nonbond list
 stop
```

Table 42: Script for NPT dynamics simulation used for analysis

```
#!/bin/bash
#! example of job file to submit parallel MPI applications
#! lines starting with #PBS are options for the gsub command
#! Name of job
#PBS -N CPD-HOCL MPI
#! Number of nodes (in this case I require 1 nodes with 8 CPU each)
#! The total number of nodes passed to mpirun will be nodes*ppn
#PBS -1 nodes=1:ppn=8
#! Specific the shell types
#PBS -S /bin/bash
#! Name of output files for std output and error;
#! if non specified defaults are <job-name>.o<job number> and <job-
name>.e<jobnumber>
#PBS -e submit.err
#PBS -o submit.out
 #!change the working directory (default is home directory)
 cd $PBS O WORKDIR
 echo Running on host `hostname`
 echo Time is `date`
echo Directory is $PBS O WORKDIR
echo This jobs runs on the following processors:
 echo `cat $PBS NODEFILE`
```

```
#! Run the parallel MPI executable with 8 processor (nodes*ppn)
/opt/mpich2/osu ch3 mrail gen2-gnu41/bin/mpiexec -np 8
/opt/c34b1/exec/qnu/charmmqff str:toppar cpd-hocl.str -i
init mini.inp -o init mini.out
/opt/mpich2/osu ch3 mrail gen2-gnu41/bin/mpiexec -np 8
/opt/c34b1/exec/gnu/charmmgff str:toppar cpd-hocl.str -i
init nve.inp -o init_nve.out
mv dyn.rst dyn.rst.nve
mv dyn.trj dyn.trj.nve
mv dyn.vel dyn.vel.nve
/opt/mpich2/osu ch3 mrail gen2-gnu41/bin/mpiexec -np 8
/opt/c34b1/exec/gnu/charmmgff str:toppar cpd-hocl.str -i
init npt.inp -o dyn.out.1
cp dyn.rst dyn.rea
mv dyn.rst dyn.rst.1
mv dyn.trj dyn.trj.1
mv dyn.vel dyn.vel.1
for i in 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20
   do
        /opt/mpich2/osu ch3 mrail gen2-gnu41/bin/mpiexec -np 8
/opt/c34b1/exec/gnu/charmmgff str:toppar cpd-hocl.str -i
equil npt.inp -o dyn.out.$i
  cp -u dyn.rst dyn.rst.$i
  cp -u dyn.rst dyn.rea
 mv dyn.trj dyn.trj.$i
 mv dyn.vel dyn.vel.$i
      done
```

 Table 43: Shell script to run various molecular dynamics scripts via queue on computer cluster and appropriately archive the output files

```
* read in traj and write out a grp of frames - cassian
*
! Read in pre-defined variables
stream datadir.def
! Read initial data files
open unit 10 read form name @0top_all22_prot.rtf
read rtf card unit 10
close unit 10
open unit 10 read form name @0par_all22_prot.prm
read param card flex unit 10
close unit 10
open unit 10 read card name @0top_all36_cgenff_remas.rtf
read rtf card unit 10 append
close unit 10
open unit 10 read card name @0par_all36_cgenff_remas.prm
```

```
read para card flex unit 10 append
close unit 10
! Read in CPD + HOCl parameter file in command line
if @?str eq 1 stream @str
open read card unit 1 name @0cpd-hocl-solv.psf
read psf card unit 1
open read card unit 1 name @0cpd-hocl-solv.crd
read coor card unit 1
set a 9999
set b 9999
set c 9999
set a2 9999
set b2 9999
set c2 9999
set d O
set m O
! Selection criteria
set cutoff1 2.4
set cutoff2 3.2
set cutoff3 3.2
set trajfirst 1
set trajlast 20
calc diff = @trajlast - @trajfirst + 1
calc termination = 500000 * @diff / 20
set i @trajfirst
set j @trajlast
label trajread
open read unit @i file name @ldyn.trj.@i
incr i
if @i le @j goto trajread
open unit 99 write form name hbond3.tmp
open unit 98 write form name hbond3.pdb
!save the hbond distance
open unit 97 write form name hbond3.dat
echu 97
echo FRAME HOCL...O
                   OH2...H (RESID) (O type) O...H20
TRAJECTORY FIRSTU @trajfirst NUNIT @diff
set 1 0
set q 1
label loop
     traj read
     incr 1 by 1
! 24 = size of box of water
calc xsiz 24. * sqrt(0.75)
calc ysiz 24. * sqrt(0.75)
```

```
calc zsiz 24. * sqrt(0.75)
set angle 109.4712206344907
crystal define octahedral @XSIZ @YSIZ @ZSIZ @angle @angle @angle
crystal build cutoff 14.0
! Recenter CPD in box of water
coor stat select type C5 end
set xmin ?XMIN
set ymin ?YMIN
set zmin ?ZMIN
image byresidue sele resn tip3 end xcen @xmin ycen @ymin zcen @zmin
image bysegment sele segi cpd .or. segid hocl end xcen @xmin ycen
@ymin zcen @zmin
UPDATE IMGFRQ 1
COOR MINDistance SELECT type HG1 END select type O4 .OR. type
O1 END
     set keto1 ?MINDA2
     COOR DISTANCE CUT @cutoff1 SELECT type HG1 END select (type
04 .OR. type 01) END
     IF ?MIND .lt. @cutoff1 THEN
      set atemp ?MIND
     define wat select (type C5 .around. 4.0) .and. type OH2 end
     set nwat ?NSEL
     IF @nwat .gt. 0 THEN
     set var1 1
     label loop1
           COOR MINDistance select (type H51 .OR. type H52) END
SELECT wat .subset. @var1 END
           IF ?MIND .lt. @cutoff2 THEN
                 set btemp ?MIND
                 IF @btemp .lt. @b THEN
                       set waterid ?SELRESI
                       IF ketol .eq. 2 THEN
                             COOR MINDistance SELECT (resi @waterid
.and. (type H1 .or. type H2)) END select (type O4) END
                       else
                             COOR MINDistance SELECT (resi @waterid
.and. (type H1 .or. type H2)) END select (type O1) END
                       ENDIF
                       IF ?MIND .lt. @cutoff3 THEN
                       set ctemp ?MIND
                       IF @ctemp .lt. @c THEN
                       set a @atemp
                            set b @btemp
                            set c @ctemp
                 ENDIF
                 ENDIF
           ENDIF
        ENDIF
          incr var1 by 1
```

```
if @var1 .le. @nwat goto loop1
 !open unit 98 write form name hbond@g.pdb
   !coor write pdb unit 98
   !incr q
   echo @1 @a @b @c
  calc interval = @1 - @d
  IF @interval .gt. 10000 THEN
  IF @b .lt. 9999 THEN
  IF @c .lt. 9999 THEN
  set d @1
   incr m
   !set count 1
  open write formatted unit 96 name struc@m.psf
    write psf card unit 96
   !open unit 96 write form name struc@m.pdb
   !coor write pdb unit 96
    open unit 96 write form name struc@m.crd
    coor write card unit 96
   set a2 @a
   set b2 @b
   set c2 @c
   set a 9999
   set b 9999
   set c 9999
ENDIF
ENDIF
 ELSE
  !incr count
  IF @a .le. @a2 THEN
  IF @b .le. @b2 THEN
  IF @c .lt. @c2 THEN
  open write formatted unit 96 name struc@m.psf
    write psf card unit 96
    !open unit 96 write form name struc@m.pdb
    !coor write pdb unit 96
    open unit 96 write form name struc@m.crd
    coor write card unit 96
   set a2 @a
   set b2 @b
   set c2 @c
   set a 9999
  set b 9999
   set c 9999
  ELSE
        set a 9999
        set b 9999
        set c 9999
ENDIF
 ELSE
        set a 9999
        set b 9999
        set c 9999
ENDIF
 ELSE
        set a 9999
```

```
set b 9999
            set c 9999
    ENDIF
    ENDIF
    ENDIF
    ENDIF
      if @1 .lt. @termination goto loop
! protein - solvent:
! Note: assume that the trajectory is RECENTERED (protein near the
box center)
! do not use verbose option, since we are not interested in
individual waters here.
! Note that the trajectory is not closed between these commands, and
can be reused
coor contact CUTHB @cutoff1 verbose select type HG1 end select type
01 .OR. type 04 end -
     firstu @trajfirst nunit @diff iunit 99
stop
```


 Table 44: Script for reading trajectory and saving coordinates of reactant precursors based on predetermined criteria

 A4 Scripts used for calculation energy barrier in fully-solvated system.

```
* CPD
* Run minimization for replica along reaction coordinate
bomlev -2
stream datadir.def
open unit 10 read form name @Otop all22 prot.rtf
read rtf card unit 10
close unit 10
open unit 10 read form name @Opar all22 prot.prm
read param card flex unit 10
close unit 10
open unit 10 read card name @Otop all36 cgenff remas.rtf
read rtf card unit 10 append
close unit 10
open unit 10 read card name @Opar all36 cgenff remas.prm
read para card flex unit 10 append
close unit 10
! Topology and param file of CPD
if @?str eq 1 stream @str
set struc struc1
! Identify key atoms of catalytic water and CPD
set hrid 429
set hsid WT4
set alphah1 H52
set alphah2 H51
set keto 01
! Read PSF file of solvated CPD
read psf card name ../@struc.psf
! Read coordinates of product
open coor card unit 12 name p@struc.crd
read coor card unit 12
coor copy COMP
! Read coordinates of reactant
open coor card unit 12 name r@struc.crd
read coor card unit 12
!----- Needed to define Q-Chem env. vars. ------
 envi qchemcnt
               "qchem-replica.inp"
 envi qcheminp "q1.inp"
 envi qchemexe "qchem"
```

```
envi qchemout "qchem.out"
 ENVI QCHEMPWD "@9"
1------
! Define OM atoms
define qm sele segid CPD .or. segid hocl .or. (resid @hrid .and.
segid @hsid) end
define buffer select .not. qm end
REPLicate A NREPlica 16 SELECT qm END
! Need to redefine atom selection after relicate command
define qm sele segid CPD .or. segid hocl .or. (resid @hrid .and.
segid @hsid) end
define buffer select .not. qm .and. .not. segid A* end
COOR orient rms mass select .not. (qm .or. seqid A*) end
! Fix 'buffer' region:
cons fix select buffer end
1-----
! { setup initial guess coordinates for all intermediates }
set 1 1
set a 0.0
label loop
coor duplicate sele qm end sele segid A@1 end
coor duplicate sele qm end sele segid A@1 end comp
coor average fact @a sele segid A@1 end
incr a by 0.066667 ! 1/15 (since a starts at 0 and ends at 15)
incr 1 by 1
if @1 .lt. 16.5 goto loop
!-----
_ _ _ _ _ _ _ _ _ _ _ _ _ _ _
DELEte ATOM SELEct qm END
DEFIne replicas SELEct SEGId A* END
! Fix 'buffer' region:
define buffer select .not. segid A* end
cons fix select none end
cons fix select buffer end
! { average the non active reactant and product atoms }
COOR average sele .not. replicas end
COOR copy comp
! { set up an appropriate interaction matrix }
BLOCK 2
CALL 2 SELEct replicas END
 COEF 1 1 1.0
```

```
COEF 2 2 0.0625 ! 16
COEF 2 1 0.0625 ! 16
END
scalar wmain set 1.0
scalar wmain set 0.0 sele type h* end
SCALAR wmain set 4 sele (resid @hrid .and. (segid @hsid .or. segid
A*) .and. type h*) .or. (resn cpd .and. type @alphah1) end
SCALAR wmain set 2 sele (resid @hrid .and. (segid @hsid .or. segid
A*) .and. type o*) end
SCALAR wmain set 1.5 sele (resn cpd .and. type @keto) .or. (resn
hocl .and. type o*) .or. (resn cpd .and. type c5) end
SCALAR wmain set 1.5 sele (resn hocl .and. type h*) .or. (resn cpd
.and. type @alphah2) end
coor copy comp
coor copy comp weight
! write psf for next script/step
write psf card name replica.psf
! Run the Conventional Replica/Path Method
 rpath krms 600.0 rmax 0.25 kangle 100.0 kmax 600.0 cosmax 0.90
rota -
      tran weight
! { fix the endpoints }
cons fix select none end
cons fix sele segid al .or. segid al6 .or. buffer end
qchem RPATh parallel 1 remove sele replicas end
!------
open write formatted unit 10 name replica ini.crd
write coor card unit 10
MINI SD NSTEP 50 NPRINT 50
open write formatted unit 10 name replicaSD50.crd
write coor card unit 10
set 1 100
label loop22
MINI ABNR NSTEP 100 NPRINT 10
open write formatted unit 10 name replicaABNR@1.crd
write coor card unit 10
incr 1 by 100
if @1 .le. 500 goto loop22
```

```
! { plot energy as a function of the path }
open write card unit 20 name energy.dat
set 1 1
label eloop
BLOCK 2
 CALL 1 sele all end
CALL 2 sele replicas .and. .not. segid A@1 end
COEF 1 1 1.0
COEF 2 1 0.0
COEF 2 2 0.0
END
ENERGY
write title unit 20
* @1 ?ener
incr 1 by 1
if @1 .lt. 16.5 goto eloop
STOP
```

Table 45: Script for mapping out minimum energy path using Replica Path method with all MM waters fixed

```
* CPD
* Run minimization for replica along reaction coordinate
* AND locate Transition State using CI-NEB
bomlev -2
stream datadir.def
open unit 10 read form name @Otop all22 prot.rtf
read rtf card unit 10
close unit 10
open unit 10 read form name @Opar all22 prot.prm
read param card flex unit 10
close unit 10
open unit 10 read card name @Otop all36 cgenff remas.rtf
read rtf card unit 10 append
close unit 10
open unit 10 read card name @Opar all36 cgenff remas.prm
read para card flex unit 10 append
close unit 10
! Topology and param file of CPD
if @?str eq 1 stream @str
```

```
set struc struc1
! Identify key atoms of catalytic water and CPD
set hrid 429
set hsid WT4
set alphah1 H52
set alphah2 H51
set keto 01
! Read PSF file of solvated CPD
read psf card name ../../@struc.psf
! Read coordinates of product
open coor card unit 12 name ../p@struc.crd
read coor card unit 12
coor copy COMP
! Read coordinates of reactant
open coor card unit 12 name ../r@struc.crd
read coor card unit 12
!----- Needed to define Q-Chem env. vars. -----
envi qchemcnt "qchem-replica.inp"
envi qcheminp "q1.inp"
envi qchemexe "qchem"
envi qchemout "qchem.out"
ENVI OCHEMPWD "@9"
!-----
! Define QM atoms
define qm sele segid CPD .or. segid hocl .or. (resid @hrid .and.
segid @hsid) end
define buffer select .not. qm end
REPLicate A NREPlica 16 SELECT gm END
! Need to redefine atom selection after relicate command
define qm sele segid CPD .or. segid hocl .or. (resid @hrid .and.
seqid @hsid) end
define buffer select .not. qm .and. .not. segid A* end
COOR orient rms mass select .not. (qm .or. segid A*) end
! Fix 'buffer' region:
cons fix select buffer end
1-----
! { setup initial guess coordinates for all intermediates }
set 1 1
set a 0.0
label loop
```

```
coor duplicate sele qm end sele segid A@1 end
coor duplicate sele qm end sele seqid A@1 end comp
coor average fact @a sele segid A@1 end
incr a by 0.066667 ! 1/15 (since a starts at 0 and ends at 15)
incr 1 by 1
if @1 .lt. 16.5 goto loop
1-----
_ _ _ _ _ _ _ _ _ _ _ _ _ _ _
DELEte ATOM SELEct gm END
! Read PSF file of solvated CPD
read psf card name replica.psf
! Read coordinates of previous replicas
open coor card unit 12 name ci-nebsd50.crd
!read coor card comp unit 12
read coor card unit 12
!-----
-----
DEFINe replicas SELEct SEGID A* END
! Fix 'buffer' region:
define buffer select .not. seqid A* end
cons fix select none end
cons fix select buffer end
! { average the non active reactant and product atoms }
COOR average sele .not. replicas end
COOR copy comp
! { set up an appropriate interaction matrix }
BLOCK 2
CALL 2 SELEct replicas END
COEF 1 1 1.0
COEF 2 2 0.0625 ! 16
COEF 2 1 0.0625 ! 16
END
scalar wmain set 1.0
scalar wmain set 0.0 sele type h* end
SCALAR wmain set 4 sele (resid @hrid .and. (segid @hsid .or. segid
A*) .and. type h*) .or. (resn cpd .and. type @alphah1) end
SCALAR wmain set 2 sele (resid @hrid .and. (segid @hsid .or. segid
A*) .and. type o*) end
SCALAR wmain set 1.5 sele (resn cpd .and. type @keto) .or. (resn
hocl .and. type o*) .or. (resn cpd .and. type c5) end
SCALAR wmain set 1.5 sele (resn hocl .and. type h*) .or. (resn cpd
.and. type @alphah2) end
coor copy comp
coor copy comp weight
```

```
! Run the Conventional Replica/Path Method
 rpath krms 600.0 rmax 0.25 kangle 100.0 kmax 600.0 cosmax 0.90
rota -
       tran weight nebf etan cimg ppmf
! { fix the endpoints }
cons fix select none end
cons fix sele segid al .or. segid al6 .or. buffer end
qchem RPATh parallel 1 remove sele replicas end
1_____
MINI SD NSTEP 50 NPRINT 50
open write formatted unit 10 name ci-nebSD100.crd
write coor card unit 10
set 1 200
label loop22
MINI SD NSTEP 100 NPRINT 10
open write formatted unit 10 name ci-nebsd@1.crd
write coor card unit 10
incr 1 by 100
if @1 .le. 500 goto loop22
1-----
! { plot energy as a function of the path }
open write card unit 20 name energy.dat
set 1 1
label eloop
BLOCK 2
CALL 1 sele all end
CALL 2 sele replicas .and. .not. segid A@1 end
COEF 1 1 1.0
COEF 2 1 0.0
COEF 2 2 0.0
END
ENERGY
write title unit 20
* @1 ?ener
*
incr 1 by 1
if @1 .lt. 16.5 goto eloop
STOP
```

Table 46: Script for locating transition state of system using CI-NEB with all MM waters fixed

```
* CPD
* Run minimization for replica along reaction coordinate
* keeping a hydration shell flexible arounf the QM region
bomlev -2
stream datadir.def
open unit 10 read form name @Otop all22 prot.rtf
read rtf card unit 10
close unit 10
open unit 10 read form name @Opar all22 prot.prm
read param card flex unit 10
close unit 10
open unit 10 read card name @Otop all36 cgenff remas.rtf
read rtf card unit 10 append
close unit 10
open unit 10 read card name @Opar all36 cgenff remas.prm
read para card flex unit 10 append
close unit 10
! Topology and param file of CPD
if @?str eq 1 stream @str
set struc struc1
! Identify key atoms of catalytic water and CPD
set hrid 429
set hsid WT4
set alphah1 H52
set alphah2 H51
set keto 01
! Read PSF file of solvated CPD
read psf card name ../min-struc-shell/@struc.psf
! Read coordinates of product
open coor card unit 12 name ../min-struc-shell/t@struc.crd
read coor card unit 12
coor copy COMP
! Read coordinates of reactant
open coor card unit 12 name ../min-struc-shell/r@struc.crd
read coor card unit 12
!----- Needed to define Q-Chem env. vars. -----
envi qchemcnt "qchem-replica.inp"
envi qcheminp "q1.inp"
envi qchemexe "qchem"
envi qchemout "qchem.out"
ENVI OCHEMPWD "@9"
!-----
```

```
! Define QM atoms
define qm sele seqid CPD .or. seqid hocl .or. (resid @hrid .and.
seqid @hsid) end
define buffer1 select .not. qm end
! Define flexible hydration shell
define water sele ((segid WT4 .and. (resid 11 .or. resid 13 .or.
resid 62 .or. resid 147 .or. resid 216 .or. resid 241 .or. resid
422)) .or. (segid WT6 .and. resid 131)) -
show end
REPLicate A NREPlica 16 SELECT (qm .or. water) END
! Need to redefine atom selection after relicate command
define qm sele seqid CPD .or. seqid hocl .or. (resid @hrid .and.
seqid @hsid) end
define buffer1 select .not. gm .and. .not. seqid A* end
define water sele ((segid WT4 .and. (resid 11 .or. resid 13 .or.
resid 62 .or. resid 147 .or. resid 216 .or. resid 241 .or. resid
422)) .or. (segid WT6 .and. resid 131)) -
show end
COOR orient rms mass select .not. (qm .or. seqid A* .or. water) end
define buffer select buffer1 .and. .not. water end
! Fix 'buffer' region:
cons fix select buffer end
1_____
! { setup initial guess coordinates for all intermediates }
set 1 1
set a 0.0
label loop1
coor duplicate sele (qm .or. water) end sele seqid A@1 end
coor duplicate sele (qm .or. water) end sele segid A@1 end comp
coor average fact @a sele segid A@1 end
incr a by 0.1428571 ! 1/7 (since a starts at 0 and ends at 15)
incr 1 by 1
if @1 .lt. 8.5 goto loop1
open coor card unit 12 name ../min-struc-shell/p@struc.crd
read coor card unit 12
coor copy COMP
! Read coordinates of reactant
open coor card unit 12 name ../min-struc-shell/t@struc.crd
read coor card unit 12
```

```
set a 0.0
label loop2
incr a by 0.125 ! 1/8 (since a starts at 9 and ends at 16)
coor duplicate sele (qm .or. water) end sele segid A@1 end
coor duplicate sele (qm .or. water) end sele seqid A@1 end comp
coor average fact @a sele segid A@1 end
incr 1 by 1
if @1 .lt. 16.5 goto loop2
-
!-----
                          -----
DELEte ATOM SELEct (qm .or. water) END
! Read PSF file of solvated CPD
read psf card name replica.psf
! Read coordinates of previous replicas
open coor card unit 12 name replicaabnr600.crd
!read coor card comp unit 12
read coor card unit 12
DEFIne replicas SELEct SEGID A* END
DEFIne qm SELEct (SEGId A* .and. (resid @hrid .or. resname cpd .or.
resname hocl)) show END
! Fix 'buffer' region:
define buffer select .not. segid A* end
cons fix select none end
cons fix select buffer end
! { average the non active reactant and product atoms }
COOR average sele .not. replicas end
COOR copy comp
! { set up an appropriate interaction matrix }
BLOCK 2
CALL 2 SELEct replicas END
COEF 1 1 1.0
COEF 2 2 0.0625 ! 16
COEF 2 1 0.0625 ! 16
END
scalar wmain set 1.0
scalar wmain set 0.0 sele type h* end
SCALAR wmain set 4 sele (resid @hrid .and. (segid @hsid .or. segid
A*) .and. type h*) .or. (resn cpd .and. type @alphah1) show end
SCALAR wmain set 2 sele (resid @hrid .and. (segid @hsid .or. segid
A*) .and. type o*) show end
SCALAR wmain set 1.5 sele (resn cpd .and. type @keto) .or. (resn
hocl .and. type o*) .or. (resn cpd .and. type c5) show end
SCALAR wmain set 1.5 sele (resn hocl .and. type h*) .or. (resn cpd
.and. type @alphah2) show end
```

```
coor copy comp
```

```
coor copy comp weight
! Run the Conventional Replica/Path Method
rpath krms 600.0 rmax 0.25 kangle 100.0 kmax 600.0 cosmax 0.90 rota
      tran weight nebf etan cimg ppmf
! { fix the endpoints }
cons fix select none end
cons fix sele segid al .or. segid al6 .or. buffer end
qchem RPATh QCLJ parallel 1 remove sele qm end
!-----
set 1 100
label loop22
MINI SD NSTEP 100 NPRINT 10
open write formatted unit 10 name ci-nebSD@1.crd
write coor card unit 10
incr 1 by 100
if @1 .le. 500 goto loop22
1_____
! { plot energy as a function of the path }
open write card unit 20 name energy/energy-ci_500.dat
set 1 1
label eloop
BLOCK 2
CALL 1 sele all end
CALL 2 sele replicas .and. .not. segid A@1 end
COEF 1 1 1.0
COEF 2 1 0.0
COEF 2 2 0.0
END
ENERGY
write title unit 20
* @1 ?ener
incr 1 by 1
if @1 .lt. 16.5 goto eloop
stop
```

Table 47: Script for locating transition state using CI-NEB for system with flexible hydration shell

\$comment			
qchem control file needed for the			
qchem-replica/ci-neb.inp			
\$end			
\$rem			
exchange	B3LYP		
basis	6-31+G**		
qm_mm	true		
jobtype	force		
symmetry	off		
sym_ignore	true		
print_input	false		
qmmm_print	true		
SCF_ALGORITHM	DIIS_GDM		
SCF_GUESS_ALWAYS	true		
THRESH	14		
SCF_CONVERGENCE	9		
MAX_SCF_CYCLES	200		
\$end			
\$molecule			
0 1			
\$end			

 Table 48: QM script read by CHARMM to create actual script to be run by QChem. Used in Replica

 Path method and CI-NEB method

\$external_charges		
-12.66269371 -3.72031255	-0.82295700	-0.83400000
0.00024239 3.15057423		
-12.46711514 -4.08620449	-1.68400236	0.41700000
0.00007331 0.40001352		
-13.03480408 -2.85588642	-1.16791530	0.41700000
0.00007331 0.40001352		
-12.82328583 -0.88125814	-6.57305158	-0.83400000
0.00024239 3.15057423		
-12.43442879 -0.05557931	-6.15039695	0.41700000
0.00007331 0.40001352		
-12.32563613 -0.88686097	-7.40489782	0.41700000
0.00007331 0.40001352		
-8.21232033 -2.74258280	5.04212284	-0.83400000
0.00024239 3.15057423		
-8.40833473 -1.77126777	4.86145020	0.41700000
0.00007331 0.40001352		
-8.27885723 -2.76478148	6.04030132	0.41700000
0.00007331 0.40001352		
-11.67862480 8.36726152	-9.01817243	-0.83400000
0.00024239 3.15057423		
-11.98195331 9.23895475	-8.73214214	0.41700000
0.00007331 0.40001352		

-12.57279938 7.96145526	-9.25817983	0.41700000	
0.00007331 0.40001352 3.44952856 -0.70802351	11 04027074	0 02400000	
0.00024239 3.15057423	-11.0403/9/4	-0.83400000	
4.19502234 -0.41123434	-10 54358408	0 41700000	
0.00007331 0.40001352	10.51550100	0.11/00000	
2.76111905 -1.03030439	-10.34711382	0.41700000	
0.00007331 0.40001352			
1.74631441 -7.57735586	-7.82149363	-0.83400000	
0.00024239 3.15057423			
1.88608599 -7.97766495	-6.95973730	0.41700000	
0.00007331 0.40001352			
0.95238292 -8.00864410	-8.19403648	0.41700000	
0.00007331 0.40001352 -8.90489964 8.30332777	4 22062510	0 02400000	
0.00024239 3.15057423	-4.33863510	-0.83400000	
-9.47780470 9.03232691	-4 47938116	0 41700000	
0.00007331 0.40001352	1,1,000110	0112700000	
-8.57478909 8.57199309	-3.45566349	0.41700000	
0.00007331 0.40001352			
-7.58822541 9.37925360	-1.99477844	-0.83400000	
0.00024239 3.15057423			
-7.33173518 9.14004347	-1.09726529	0.41700000	
0.00007331 0.40001352 -8.25068621 10.13018725	_1 00125027	0 41700000	
0.00007331 0.40001352	-1.00133027	0.41700000	
-8.04161835 -1.21567106	-4.70295620	-0.83400000	
0.00024239 3.15057423			
-8.63120365 -1.92100966	-5.04181099	0.41700000	
0.00007331 0.40001352			
-8.11833286 -1.30447316	-3.73433805	0.41700000	
0.00007331 0.40001352 -4.79387236 2.43332100	7 42520464	0 02400000	
0.00024239 3.15057423	-7.43530464	-0.83400000	
-4.88113356 1.47559214	-7.60975504	0.41700000	
0.00007331 0.40001352			
-5.71967506 2.66631365	-7.72223759	0.41700000	
0.00007331 0.40001352			
-0.78158152 3.00790405	-9.29691982	-0.83400000	
0.00024239 3.15057423	0.00418664	0 41 5 0 0 0 0	
-0.92151082 2.10098171 0.00007331 0.40001352	-8.98417664	0.41700000	
-0.55826366 3.04506445	-10 22206688	0.41700000	
0.00007331 0.40001352	10.22200000	0.41/00000	
-15.28397678 2.52064809	-9.35715883	-0.83400000	
0.00024239 3.15057423			
-15.33644496 3.37625131	-8.87684839	0.41700000	
0.00007331 0.40001352			
-15.90883388 2.12835606	-8.72084635	0.41700000	
0.00007331 0.40001352 2.06923771 7.52523041	0.79183286	-0.83400000	
0.00024239 3.15057423	0.17103200	0.00100000	
	0.25203398	0.41700000	
0.00007331 0.40001352			
2.15169120 6.52610636	0.92274112	0.41700000	

0.00007331 0.40001352		
-4.75245047 3.31913185	-10.09435272	-0.83400000
0.00024239 3.15057423	0 10400400	0 41 5 0 0 0 0
-5.01067591 3.26426482	-9.13492489	0.41700000
0.00007331 0.40001352 -4.87542582 4.27081680	10 000000	0 41700000
-4.87542582 4.27081880 0.00007331 0.40001352	-10.23380089	0.41700000
-11.10755942 6.90023927	-1 96551757	-0.83400000
0.00024239 3.15057423	-1.00001101	-0.83400000
-11.67873881 6.29674725	-2 37477489	0 41700000
0.00007331 0.40001352	2.37177105	0.11,00000
-11.42030928 7.74367647	-2.26106115	0.41700000
0.00007331 0.40001352		
-9.95538367 -7.17085660	-1.88822378	-0.83400000
0.00024239 3.15057423		
-10.81655134 -6.88799012	-2.17057241	0.41700000
0.00007331 0.40001352		
-9.58955277 -6.44168390	-2.42013372	0.41700000
0.00007331 0.40001352		
-8.20581011 11.30084059	1.41081138	-0.83400000
0.00024239 3.15057423		
-7.73376469 10.50504038	1.12831684	0.41700000
0.00007331 0.40001352	1 0/5051/0	
-8.92558913 10.99959776	1.94595142	0.41700000
0.00007331 0.40001352		0.02400000
0.16168584 -2.01792860	-7.50541592	-0.83400000
0.00024239 3.15057423 0.34712616 -2.90599823	-7 07946990	0 41700000
0.00007331 0.40001352	-7.07040000	0.41700000
-0.60477573 -1.85593629	-6 95776033	0.41700000
0.00007331 0.40001352	0.99770099	0.11,00000
-2.12588978 3.02616072	-6.71686792	-0.83400000
0.00024239 3.15057423		
-3.03456879 2.88080072	-6.92847157	0.41700000
0.00007331 0.40001352		
-1.76671541 3.39728117	-7.53661919	0.41700000
0.00007331 0.40001352		
-6.14285946 7.88858700	-5.11608028	-0.83400000
0.00024239 3.15057423		
-7.08336496 7.83981323	-4.87488365	0.41700000
0.00007331 0.40001352		
-5.97302198 6.92667627	-5.08902025	0.41700000
0.00007331 0.40001352		
1.81689727 5.69858503	-7.30674553	-0.83400000
0.00024239 3.15057423		0 41700000
1.59058917 4.86141920	-/.684366/0	0.41700000
0.00007331 0.40001352 2.70646715 5.43788433	6 02526426	0 41700000
0.00007331 0.40001352	-0.92320430	0.41/00000
-6.23954790 5.53375944	-9 18444555	-0 83400000
0.00024239 3.15057423	2.10111000	0.03100000
-6.95070474 6.09495386	-8.81293361	0.41700000
0.00007331 0.40001352	0.012/0001	
-6.49491709 4.66287192	-8.81128567	0.41700000
0.00007331 0.40001352		

-9.15484623 11.43463251	-1.47783427	-0.83400000
0.00024239 3.15057423	0 77041570	0 41700000
-8.91523747 12.08645365	-0.//8415/2	0.41700000
0.00007331 0.40001352		
-10.12292866 11.60295698	-1.57418231	0.41700000
0.00007331 0.40001352		
-4.96940088 7.32975769	5.59079790	-0.83400000
0.00024239 3.15057423		
-4.97037077 8.25142002	5.25582504	0.41700000
0.00007331 0.40001352		
-4.40525436 6.91930676	4.96578026	0.41700000
0.00007331 0.40001352		
-0.26684002 10.00807020	-10.06604914	-0.83400000
0.00024239 3.15057423		
0.03129144 9.58866331	-9.17263989	0.41700000
0.00007331 0.40001352		
-0.95131163 10.56023714	-9 62617354	0 41700000
0.00007331 0.40001352	5.0201,551	0.11,00000
-11.15907404 5.46717755	0 61492972	-0 83400000
	0.01493973	-0.83400000
0.00024239 3.15057423	0 04100514	0 41700000
-11.00866053 5.88342504	-0.24188514	0.41700000
0.00007331 0.40001352		
-10.59244559 5.96438555	1.28520636	0.41700000
0.00007331 0.40001352		
-4.83281708 -6.92203426	-1.25783682	-0.83400000
0.00024239 3.15057423		
-4.91476154 -6.10188293	-0.64053756	0.41700000
0.00007331 0.40001352		
-3.88136363 -6.92576265	-1.46580851	0.41700000
0.00007331 0.40001352		
-1.64866905 9.96153948	-6.94652442	-0.83400000
0.00024239 3.15057423		
-1.67398457 10.90520594	-7.14385967	0.41700000
0.00007331 0.40001352		
-1.66991572 10.06001780	-5.95639495	0.41700000
0.00007331 0.40001352		
-0.23070198 5.22515583	-2.17702746	-0.83400000
0.00024239 3.15057423	212//02/10	
-0.68355227 5.50536346	-3 00527048	0 41700000
0.00007331 0.40001352	5.0052/010	0.11,00000
0.65467203 5.28325462	-2.50035000	0.41700000
0.00007331 0.40001352	-2.50055000	0.41/00000
	1 (0993009	0 02400000
	-4.60773897	-0.83400000
0.00024239 3.15057423	4 20140544	0 41 5 0 0 0 0 0
	-4.32142544	0.41700000
0.00007331 0.40001352		
0.97568953 0.66868222	-5.61498117	0.41700000
0.00007331 0.40001352		
-10.85094322 -1.28269304	-8.74101127	-0.83400000
0.00024239 3.15057423		
-11.16956712 -0.75009549	-9.47861684	0.41700000
0.00007331 0.40001352		
-10.24647237 -0.64405835	-8.33378614	0.41700000
0.00007331 0.40001352		
-3.79904056 -0.90402341	-9.47147846	-0.83400000
	-	

0.00024239 3.15057423		
-4.06642818 -1.54297495	-8.79240990	0.41700000
0.00007331 0.40001352		
-4.66850042 -0.62892956	-9.77049541	0.41700000
0.00007331 0.40001352	0 70000010	0 02400000
-6.40939454 12.73173353	-2./0289318	-0.83400000
0.00024239 3.15057423 -6.89332251 11.97487089	2 2 2 2 7 4 5 0 2	0 41700000
	-2.362/4592	0.41/00000
0.00007331 0.40001352 -5.60239176 12.65092108	-2 20562956	0 41700000
0.00007331 0.40001352	-2.20002000	0.41/00000
0.18268676 12.24280665	-2 97719757	-0 83400000
0.00024239 3.15057423	2.51115151	0.03400000
-0.32673602 12.72819779	-2 30513470	0 41700000
0.00007331 0.40001352	2.30313170	0.11,00000
-0.16427783 12.64990208	-3.74899320	0.41700000
0.00007331 0.40001352		
1.70999646 2.95553684	-8.22914314	-0.83400000
0.00024239 3.15057423		
1.62235367 2.07860875	-7.79374266	0.41700000
0.00007331 0.40001352		
0.96318227 2.97343135	-8.90891552	0.41700000
0.00007331 0.40001352		
-10.94326019 -5.29858589	0.67193198	-0.83400000
0.00024239 3.15057423		
-11.55381584 -4.77557135	0.18947107	0.41700000
0.00007331 0.40001352		
-11.55175591 -5.99943066	1.06350756	0.41700000
0.00007331 0.40001352		
-14.95601198 -0.71559520	-5.01344677	-0.83400000
0.00024239 3.15057423	E E10(E011	0 11 0 0 0 0 0
-14.32886336 -0.70913930	-5./196/311	0.41700000
0.00007331 0.40001352		0 11 000000
-15.76153776 -0.45086856 0.00007331 0.40001352	-5.52604957	0.41700000
-5.94910002 9.58284092	2 54445242	-0.83400000
0.00024239 3.15057423	5.54445245	-0.83400000
-5.84677458 8.84972477	2 95315051	0 41700000
0.00007331 0.40001352	2.)))1001	0.41/00000
-6.83834124 9.41940403	3 83400202	0 41700000
0.00007331 0.40001352	5.05100202	0.11,00000
-1.56935549 0.35127306	-9.18744278	-0.83400000
0.00024239 3.15057423		
-2.36397362 -0.17355590	-9.39029121	0.41700000
0.00007331 0.40001352		
-1.75811040 0.50599879	-8.16943741	0.41700000
0.00007331 0.40001352		
-14.07500622 -3.92943330	1.44328885	-0.83400000
0.00024239 3.15057423		
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-8.08180046 -5.60862160	-4.11813736	0.41700000
0.00007331 0.40001352		
-8.06923866 -4.93459272	-2.73378325	0.41700000
0.00007331 0.40001352		
-4.98947447 6.34384319	-11.47277468	-0.83400000
0.00024239 3.15057423		
-4.38051813 6.99816831	-11.06174009	0.41700000
0.00007331 0.40001352		
-5.62951296 6.29497203	-10.76720922	0.41700000
0.00007331 0.40001352		
-8.27862644 -6.22921848	0.11456603	-0.83400000
0.00024239 3.15057423		
-9.20361233 -5.98716402	0.10752918	0.41700000
0.00007331 0.40001352		
-8.23515320 -6.65712786	-0.77678341	0.41700000
0.00007331 0.40001352		
	4.53264741	-0.83400000
0.00024239 3.15057423		
-11.38798830 6.89944844	4.40146951	0.41700000
0.00007331 0.40001352		
-12.55618403 7.59442238	3.63910709	0.41700000
0.00007331 0.40001352		
-6.23802818 -2.05695356	-12.25023545	-0.83400000

0.00024239 3.15057423		
-6.13116277 -2.92388261	-11.77491231	0.41700000
0.00007331 0.40001352		
-5.61272014 -2.27334703	-12.91469634	0.41700000
0.00007331 0.40001352	4 80088500	0.00400000
-6.09605598 3.33998847	4.70277500	-0.83400000
0.00024239 3.15057423	E 08146016	0 11 0 0 0 0 0
-6.38206244 4.09658241	5.27146816	0.41700000
0.00007331 0.40001352	4 01005440	0 41700000
-5.20117903 3.37160611 0.00007331 0.40001352	4.91095448	0.41/00000
-5.52061605 7.17474270	2 26965106	0 83400000
0.00024239 3.15057423	2.20005190	-0.83400000
-6.20627928 6.43898106	2 35506892	0 41700000
0.00007331 0.40001352	2.33300092	0.41/00000
-4.70384932 6.67641544	2 00695515	0 41700000
0.00007331 0.40001352	2.00000010	0.11,00000
-13.65620129 7.45130690	-5.26471823	-0.83400000
0.00024239 3.15057423		
-14.23529415 7.70012941	-5.97015684	0.41700000
0.00007331 0.40001352		
-12.97290819 8.15358284	-5.38182562	0.41700000
0.00007331 0.40001352		
-0.15509634 -2.72927952	6.89051485	-0.83400000
0.00024239 3.15057423		
-0.82812691 -3.00526023	7.53619385	0.41700000
0.00007331 0.40001352		
-0.73481238 -2.54001141	6.12987900	0.41700000
0.00007331 0.40001352		
-12.08586591 9.46859130	-6.14232176	-0.83400000
0.00024239 3.15057423		
-12.37590473 10.00505051	-6.89231603	0.41700000
0.00007331 0.40001352		
-11.43954294 9.98039016	-5.61479585	0.41700000
0.00007331 0.40001352	2 01155560	0.00100000
-13.77178452 -1.77177949 0.00024239 3.15057423	3.01155/68	-0.83400000
-13.99905369 -1.46797081	2 07250072	0 41700000
0.00007331 0.40001352	3.8/2599/2	0.41700000
-13.96769497 -0.96654649	2.47607974	0 41700000
0.00007331 0.40001352	2.7/00/2/4	0.11/00000
-9.07558553 3.02668914	-12 63456457	-0 83400000
0.00024239 3.15057423	12.03130137	0.03100000
-9.34321707 3.15119704	-13.57319062	0.41700000
0.00007331 0.40001352		
-8.43977182 3.74490556	-12.58055227	0.41700000
0.00007331 0.40001352		
-12.64775536 9.15957074	-2.72061105	-0.83400000
0.00024239 3.15057423		
-12.22316954 10.00026898	-2.50208421	0.41700000
0.00007331 0.40001352		
-13.41526100 9.49654822	-3.12602325	0.41700000
0.00007331 0.40001352		
-14.14414046 6.12802414	0.60974650	-0.83400000
0.00024239 3.15057423		

-13.50532505 5.49088292	0.29932980	0.41700000	
0.00007331 0.40001352			
-14.30748865 5.77580695	1.50289587	0.41700000	
0.00007331 0.40001352			
-3.58556791 4.65147798	-13.03767273	-0.83400000	
0.00024239 3.15057423			
-2.69634958 5.00402955	-12.93018744	0.41700000	
0.00007331 0.40001352			
-4.19563766 5.29573004	-12.55840990	0.41700000	
0.00007331 0.40001352			
-12.75514815 3.79680629	-0.63400074	-0.83400000	
0.00024239 3.15057423			
-12.09488985 4.44156812	-0.37973495	0.41700000	
0.00007331 0.40001352			
-12.15647814 3.19973974	-1.14255233	0.41700000	
0.00007331 0.40001352			
-2.82846212 5.86562014	4.46032476	-0.83400000	
0.00024239 3.15057423			
-1.94455612 5.60928631	4.17307854	0.41700000	
0.00007331 0.40001352			
-2.69034266 6.77059746	4.29284573	0.41700000	
0.00007331 0.40001352			
-0.74459024 -1.47454067	-13.34422085	-0.83400000	
0.00024239 3.15057423			
-0.96498151 -0.51782318	-13.23167774	0.41700000	
0.00007331 0.40001352			
-1.36986490 -1.90669199	-12.73271248	0.41700000	
0.00007331 0.40001352			
-11.42049568 1.74410542	-6.20227831	-0.83400000	
0.00024239 3.15057423			
-11.45701950 2.53933820	-5.56972616	0.41700000	
0.00007331 0.40001352			
-11.59960716 2.22951516	-7.01957052	0.41700000	
0.00007331 0.40001352			
2.00765657 -5.96039009	-4.22972727	-0.83400000	
0.00024239 3.15057423			
1.66777205 -5.67990160	-3.34176111	0.41700000	
0.00007331 0.40001352			
2.59278393 -5.23373747	-4.40355873	0.41700000	
0.00007331 0.40001352			
-1.80957465 13.41002819	-1.08127956	-0.83400000	
0.00024239 3.15057423			
	-1.07766084	0.41700000	
0.00007331 0.40001352			
-2.11009364 14.30828163	-0.96596484	0.41700000	
0.00007331 0.40001352			
3.06266475 -2.00935292	-8.57498169	-0.83400000	
0.00024239 3.15057423			
2.29461861 -2.45595694	-8.22233963	0.41700000	
0.00007331 0.40001352			
	-8.21736526	0.41700000	
0.00007331 0.40001352			
5.41639662 6.98136330	-7.56164074	-0.83400000	
0.00024239 3.15057423			
4.51864910 7.33842087	-7.54427862	0.41700000	
			_

0.00007331 0.40001352		
5.68595123 7.35090637	-8.42472267	0.41700000
0.00007331 0.40001352		
-12.95901034 5.69611732	-3.22328277	-0.83400000
0.00024239 3.15057423		
-13.77139923 5.68432896	-2.69825788	0.41700000
0.00007331 0.40001352		
-13.34648249 6.33859949	-3.87178178	0.41700000
0.00007331 0.40001352		
2.86162949 4.83362484	1.04894245	-0.83400000
0.00024239 3.15057423		
2.94785142 4.43282461	0.15239438	0.41700000
0.00007331 0.40001352		
2.96798277 4.03178787	1.60738480	0.41700000
0.00007331 0.40001352		
-0.37394977 3.92270803	2.80023980	-0.83400000
0.00024239 3.15057423		
-0.68419307 3.90205503	1.88045883	0.41700000
0.00007331 0.40001352		
-0.54673797 3.04112530	3.16742992	0.41700000
0.00007331 0.40001352		
-1.33274293 -1.08798683	-4.14268970	-0.83400000
0.00024239 3.15057423		
-0.58509195 -0.49760401	-4.26913738	0.41700000
0.00007331 0.40001352		
-1.57219243 -0.88222152	-3.26186395	0.41700000
0.00007331 0.40001352		
0.51386356 -4.55409622	4.08526278	-0.83400000
0.00024239 3.15057423		
0.54449826 -5.05407763	4.86631298	0.41700000
0.00007331 0.40001352		
1.17675662 -3.82966375	4.26508665	0.41700000
0.00007331 0.40001352		
-9.03044796 0.01655403	-7.12107897	-0.83400000
0.00024239 3.15057423		
-8.20575523 -0.06905016	-6.59003067	0.41700000
0.00007331 0.40001352		
-9.60659695 0.42270315	-6.47083044	0.41700000
0.00007331 0.40001352		
-5.23969854 -8.46711982	-8.28344024	-0.83400000
0.00024239 3.15057423		
-5.52945674 -9.31833971	-7.88173354	0.41700000
0.00007331 0.40001352		
-4.30740798 -8.55647839	-8.23987974	0.41700000
0.00007331 0.40001352		
1.25795798 1.03646688	-12.03802130	-0.83400000
0.00024239 3.15057423		
0.44199638 0.96703128	-12.53276369	0.41700000
0.00007331 0.40001352		
1.25303463 0.28819088	-11,38923762	0.41700000
0.00007331 0.40001352		
-3.86322170 8.42595885	-10.16992955	-0.83400000
0.00024239 3.15057423		0.00100000
-4.20128625 9.35292456	-10 00118308	0 41700000
0.00007331 0.40001352	10.00110000	0.11,00000
0.10001002		

-3.28508107 8.33410094	-9.41658883	0.41700000	
0.00007331 0.40001352 -1.39618202 1.11947588	-12.98610136	-0.83400000	
0.00024239 3.15057423			
-1.43460079 2.01820723	-13.32191441	0.41700000	
0.00007331 0.40001352 -2.25057884 1.17017154	10 50040046	0 41700000	
0.00007331 0.40001352	-12.53048346	0.41/00000	
-11.98060472 -4.45980895	-5.87099279	-0.83400000	
0.00024239 3.15057423			
-12.82667083 -4.14411701 0.00007331 0.40001352	-6.23088182	0.41700000	
-11.49747241 -4.45754302	-6 66067851	0 41700000	
0.00007331 0.40001352	0.00007031	0.11,00000	
-15.00323031 2.41336863	-0.69267030	-0.83400000	
0.00024239 3.15057423			
-14.16930363 2.92504148 0.00007331 0.40001352	-0.74159809	0.41700000	
-15.41205666 2.73811428	-1.49574371	0.41700000	
0.00007331 0.40001352			
-6.21218586 5.15519667	-4.94121981	-0.83400000	
0.00024239 3.15057423		0 41700000	
-6.72395086 4.43384933 0.00007331 0.40001352	-5.30///9/9	0.41/00000	
-5.42491817 4.68438816	-4.57948351	0.41700000	
0.00007331 0.40001352			
-2.29626624 11.47527525	-9.14929299	-0.83400000	
0.00024239 3.15057423 -2.29822628 12.39635966	-8 84270148	0 /1700000	
0.00007331 0.40001352	-0.042/0140	0.41/00000	
-3.22399269 11.25381300	-9.08112816	0.41700000	
0.00007331 0.40001352			
-5.04480140 11.05734179 0.00024239 3.15057423	-9.18205981	-0.83400000	
-5.45388369 11.94699976	-9.04231505	0.41700000	
0.00007331 0.40001352			
-5.47470669 10.61978934	-8.38914208	0.41700000	
0.00007331 0.40001352	11 04500104	0.00400000	
-12.99631064 4.39289984 0.00024239 3.15057423	-11.04582184	-0.83400000	
-12.52935117 5.03446301	-11.59578340	0.41700000	
0.00007331 0.40001352			
	-11.06841629	0.41700000	
0.00007331 0.40001352 -14.18057526 -3.40689767	-7.67142881	-0.83400000	
0.00024239 3.15057423	-/.0/142001	-0.03400000	
	-8.59677041	0.41700000	
0.00007331 0.40001352			
-14.29993022 -2.60616792 0.00007331 0.40001352	-7.08094276	0.41700000	
1.48241453 -3.17238231	-13.02713272	-0.83400000	
0.00024239 3.15057423			
0.96281199 -2.33442588	-12.99476597	0.41700000	
0.00007331 0.40001352	10 5700740	0 41700000	
0.82027153 -3.64223047	-13.5/238/43	0.41/00000	

0.0000	7331 0.400	01352		
\$end				
\$lj pa:	rameters			
1	0.00003187	3.9199	95436	
2	0.00019123	3.0290	05564	
3	0.00009562	3.5992	23082	
4	0.00005578	2.3876	50856	
5	0.00005578	2.3876	50856	
6	0.00009562	3.5992	23082	
7	0.00005578	2.3876	50856	
8	0.00005578	2.3876	50856	
9	0.00003187	3.9199	95436	
10	0.00019123	3.0290	05564	
11	0.00009562	3.5992	23082	
12	0.00005578	2.3876	50856	
13	0.00005578	2.3876	50856	
14	0.00007331	0.4000	01352	
15	0.00030613	3.1448	37248	
16	0.00054661	3.4032		
17	0.00024239	3.150		
18	0.00007331	0.4000		
19	0.00007331	0.4000	01352	
\$end				
scan.i: \$end	п <u>Б</u>			
\$rem		600		
igdesp		699		
symmet		off		
sym_ig:		true		
exchan basis	ge	b3lyp 6-31+g**		
qm mm		true		
jobtyp	۵	force		
symmet		off		
sym ig:	-	true		
print		false		
qmmm p		true		
	gorithm	diis gdm		
	ess_always	true		
thresh		14		
	nvergence	9		
	f_cycles	200		
\$end				
\$molec	ule			
0 1				
C	-4.53025		0.3474980463	-3.4482620388
0	-4.28218		-0.6443653521	-4.1159852232
C	-4.25898		0.4728237024	-1.9548600563
H	-3.18531	.44153	0.4014444707	-1.7546327052

-			
Н	-4.7318365026	-0.3880481106	-1.4758713928
С	-4.8803182970	1.8151120642	-1.5142897172
Н	-4.1078092431	2.5797624642	-1.3630062200
Н	-5.4671954870	1.7724772590	-0.5921306735
С	-5.7213707362	2.2742501489	-2.6949769955
0	-6.6676775358	3.0452749043	-2.6486124046
С	-5.2295935503	1.5944700901	-3.9544916891
Н	-4.4735430443	2.2400199363	-4.4323231722
Н	-5.9951562143	1.4005382003	-4.7094519761
Н	-7.6213871300	3.9841879802	-1.5545514805
0	-8.1913255525	4.6144490141	-1.0423151450
CL	-7.3445458593	6.1039254073	-1.2278823717
0	-5.9399657407	-0.4077018344	-6.4716743463
Н	-5.1471612627	-0.5542510367	-5.9306930634
Н	-6.6122922685	-0.9632560473	-6.0326838892
\$end			
L			

 Table 49: Script generated by CHARMM to be run by QChem for each replica in Replica Path method and CI-NEB method

Appendix B: Topology and parameter files

B1 Topology and parameter files for sugar moleclues.

REST	AXYS			0.000	<u>ר</u>	lalpł	na-I	D-xylo	nvr	anos r	non	omer		
NED I	111110			0.000									nose	e(ARB)
GROU							200	aboa		2000	_ `			
ATOM	C1	CC3162	2	0.3	340									
ATOM		HCA1			090									
ATOM	01	OC311		-0.6										
ATOM	HO1	HCP1			120									
ATOM		CC3263	3		020									
ATOM	H51	HCA2			090									
ATOM	H52	HCA2		0.0	090									
ATOM	05	OC3C63	1	-0.4										
GROU														
ATOM	C2	CC3163	1	0.1	L40									
ATOM	H2	HCA1		0.0	090									
ATOM	02	OC311		-0.6	550									
ATOM	HO2	HCP1		0.4	120									
GROU														
ATOM	C3	CC3163	1	0.1	L40									
ATOM	H3	HCA1		0.0	090									
ATOM	03	OC311		-0.6	550									
ATOM	HO3	HCP1		0.4	120									
GROU														
ATOM	C4	CC316	1	0.1	L40									
ATOM	H4	HCA1		0.0	090									
ATOM	04	OC311		-0.6	550									
ATOM	HO4	HCP1		0.4	120									
BOND	C1	01		C1	H1	C	01	HO1		C1	05		C1	C2
BOND	C2	H2		C2	02)2	HO2		C2	C3		C3	H3
BOND	C3	03		03	HO3	(23	C4		C4	H4		C4	04
BOND	04	HO4		C4	C5	(25	H51		C5	H52	2		
BOND	C5	05												
DONO	BLNK	HO1												
	BLNK													
DONO	BLNK	HO3												
DONO	BLNK	HO4												
ACCE														
ACCE														
ACCE	03													
ACCE														
ACCE														
!	I		K	L		R(IK)		T(IKC	J)	PHI	r	Γ(JKL))	R(KL)
IC	01	C2 *(C1	H1		1.411	L5	105.8	32 -1	120.20	6 3	110.81	L	1.0905
IC	01		C1	C2		1.411	L5			118.92		111.32	2	1.5218
IC	02	C3 *(C2	H2		1.419	90	110.9	99 -1	116.9	7 3	108.77	7	1.0892

IC	02	C1 *C2	C3	1.4190	111.74	-123.77	110.04	1.5167
IC	03	C4 *C3	H3	1.4198	107.51	119.38	108.28	1.0957
IC	03	C2 *C3	C4	1.4198	111.19	119.03	110.08	1.5102
IC	04	C5 *C4	H4	1.4163	107.94	-120.61	109.07	1.0972
IC	04	C3 *C4		1.4163		-118.91		1.5171
IC	05	C1 C2		1.4059	111.32	53.56		1.5167
IC	C1	C1 C2 C2 C3				-54.55		
				1.5218				1.5102
IC	C2	C3 C4		1.5167	110.08			1.5171
IC	C3	C4 C5		1.5102		-58.25		1.4384
IC	C4	C5 05		1.5171	109.77		114.27	1.4059
IC	C5	05 C1	C2	1.4384	114.27	-57.30	111.32	1.5218
IC	05	C1 01	HO1	1.4059	113.00	67.18	108.04	0.9634
IC	C1	C2 O2	HO2	1.5218	111.74	-44.11	105.37	0.9665
IC	C2	C3 O3	HO3	1.5167	111.19	-51.80	106.26	0.9641
IC	C3	C4 04		1.5102	111.43	47.58	105.89	0.9645
		NONE LA						
REST	BXYS		0.000 !	heta-D-	xvlopyr;	anos monc	mer	
КЦОТ	DATO		0.000 .	beea b	луторут		MICI	
GROU								
	C 1	001100	0 240					
ATOM		CC3162	0.340					
ATOM		HCA1	0.090					
ATOM		OC311	-0.650					
ATOM		HCP1	0.420					
ATOM	C5	CC3163	0.020					
ATOM	H51	HCA2	0.090					
ATOM	H52	HCA2	0.090					
ATOM	05	OC3C61	-0.400					
GROU								
ATOM	C2	CC3161	0.140					
ATOM		HCA1	0.090					
ATOM		OC311						
			-0.650					
ATOM	HOZ	HCP1	0.420					
GROU								
ATOM		CC3161	0.140					
ATOM		HCA1	0.090					
ATOM		OC311	-0.650					
ATOM	HO3	HCP1	0.420					
GROU								
ATOM	C4	CC3161	0.140					
ATOM	H4	HCA1	0.090					
ATOM		OC311	-0.650					
ATOM		HCP1	0.420					
GROU	1101	1101 I	0.120					
01000								
BOND	C1	01	C1 H1	01	HO1	C1 C	5 C	1 C2
BOND		H2	C2 O2	02	HO2			3 H3
BOND		03	03 H03	C3	C4			4 04
BOND		HO4	C4 C5	C5	H51	C5 H	152	
BOND		05						
	BLNK							
DONO	BLNK	HO2						
DONO	BLNK	HO3						
	BLNK							
ACCE								

ACCE								
ACCE								
ACCE	04							
ACCE	05							
!	I	J K	\mathbf{L}	R(IK)	T(IKJ)	PHI	T(JKL)	R(KL)
IC	01	C2 *C	1 H1	1.3890	105.75	114.54	108.17	1.0950
IC	01	05 *C	1 C2	1.3890	111.55	117.06	110.06	1.5340
IC	02	C3 *C		1.4154		-118.21		1.0919
IC	02	C1 *C		1.4154		-125.56		1.5253
IC	03	C4 *C		1.4157		120.65		1.1068
IC	03	C2 *C		1.4157		120.03		1.5177
IC	04	C5 *C		1.4252		-120.61		1.1024
IC	04	C3 *C						
IC				1.4252		-122.08		1.5287
	05			1.4254	110.06	54.09		1.5253
IC	C1	C2 C		1.5340	111.08	-51.23		1.5177
IC	C2	C3 C		1.5253	109.86	53.25		1.5288
IC	C3	C4 C		1.5177		-57.46		1.4274
IC	C4	C5 0		1.5288		62.25		1.4254
IC	C5	05 C		1.4274	113.77	-60.97	110.06	1.5340
IC	C1	C2 0	2 HO2	1.5340	110.87	-100.51	112.13	0.9638
IC	C2	C3 0	3 HO3	1.5253	108.09	-165.88	112.08	0.9730
IC	C3	C4 0	4 HO4	1.5177	108.31	134.18	106.97	0.9713
PATC	FIR	S NONE	LAST NONE					
RESI	BNAG		0.000	! N-Acy	tyl-Gluo	cosamine		
GROU								
ATOM	C1	CC3162	0.340					
ATOM	H1	HCA1	0.090					
ATOM	01	OC311	-0.650					
ATOM	HO1	HCP1	0.420					
ATOM	C5	CC3163	0.110					
ATOM		HCA1	0.090					
ATOM		OC3C61						
GROU	00	000001	0.100					
ATOM	C2	CC3161	0.070					
ATOM		HCA1	0.090					
ATOM		NH1 u	-0.47					
ATOM		H	0.31					
GROUI		CIII 2	0 07					
ATOM			-0.27					
ATOM		HA	0.09					
ATOM		HA	0.09					
ATOM		HA	0.09					
GROUI								
ATOM	CLP	С	0.51					
ATOM	OL	0	-0.51					
GROUI	Р							
GROU								
ATOM		CC3161	0.140					
ATOM		HCA1	0.090					
ATOM		OC311	-0.650					
ATOM		HCP1	0.420					
	1100	TUCTT	0.420					
GROU								

ATOM	C4	CC31	61	0.140						
ATOM	H4	HCA1		0.090						
ATOM	04	OC31	1	-0.650						
ATOM	HO4	HCP1		0.420						
GROU										
ATOM		CC32	1	0.050						
ATOM		HCA2		0.090						
ATOM		HCA2		0.090						
	06	OC31		-0.650						
ATOM	HO6	HCP1		0.420						
BOND	C1	01	C1	H1	01	HO1	C1	05	C1	C2
BOND	C2	H2	C2	NL	NL	HL	NL	CLP	C2	C3
BOND	C3	03	03	HO3	C3	C4	C4	H4	C4	04
BOND		HO4	C4	C5	C5	Н5		C6	C6	H61
BOND		H62	C6	06	06	HO6		05	C3	Н3
BOND		HL1			CL	HL3		CLP	00	110
DOUB		LP OL		11112	CL	шэ	СЦ			
	CLP (
			I OL							
	BLNK									
	BLNK									
	BLNK									
DONO	BLNK	HO6								
ACCE	01									
ACCE	03									
ACCE	04									
ACCE	05									
ACCE										
!	I	J	K	L	R(IK)	T(IKJ)	PHI	יד,) ד	(L)	R(KL)
IC	01	C2	*C1	 H1	1.3890	105.75				1.0950
IC	01	05	*C1	C2	1.3890	111.55	117.0			1.5340
IC	03	C4	*C3	H3						
					1.4157	110.61	120.			1.1068
IC	03	C2	*C3	C4	1.4157	108.09	120.	109		1.5177
IC	04	C5	*C4	H4	1.4252	110.90		108		1.1024
IC	04	C3	*C4	C5	1.4252	108.31		111		1.5287
IC	C6	05	*C5	Н5	1.5099	108.10				1.1042
IC	C6	C4	*C5	05	1.5099	111.57			.69	1.4274
IC	06	H62	*C6	H61	1.4132	110.47	-120.3	2 107	.85	1.0945
IC	06	C5	*C6	H62	1.4132	110.45	-121.5	3 108	.99	1.0959
IC	05	C1	C2	C3	1.4254	110.06	54.0	9 111	.08	1.5253
IC	C1	C2	C3	C4	1.5340	111.08	-51.2	3 109	.86	1.5177
IC	C2	C3	C4	C5	1.5253	109.86	53.2			1.5288
IC	C3	C4	C5	05	1.5177	111.17	-57.4			1.4274
IC	C4	C5	05	C1	1.5288	108.69	62.2			1.4254
IC	C5	05	C1	C2	1.4274	113.77	-60.9			1.5340
IC	C4	C5	C1 C6	06	1.5287	111.57	180.	110		1.4132
IC		C1								
	05		01	HO1	1.4254	111.55	-60.	107		0.9684
IC	C1	C2	02	HO2	1.5340	110.87	-60.	112		0.9638
IC	C2	C3	03	HO3	1.5253	108.09	60.	112		0.9730
IC	C3	C4	04	HO4	1.5177	108.31	60.	106		0.9713
IC	C5	C6	06	HO6	1.5099	110.44	-60.	107	.72	0.9641
ic c		p *nl		0.0 0.		0.0 0.	0			
ic n	l cl	*cl	p ol	0.0 0.	0 180.0	0.0 0.	0			
ic c		p nl	-	0.0 0.						
ic o	-	p nl		0.0 0.						
		-								

ic hl1 cl clp nl 0.0 0.0 180.0 0.0 0.0 ic hl2 cl clp nl 0.0 0.0 60.0 0.0 0.0 ic hl3 cl clp ol 0.0 0.0 120.0 0.0 0.0 ic c2 clp *nl hl 0.0 0.0 180.0 0.0 0.0 ic hl1 hl2 *cl hl3 0.0 0.0 240.0 0.0 0.0 ic hl nl c2 h2 0.0 0.0 0.0 0.0 0.0 PATC FIRS NONE LAST NONE RESI PCA 0.00 ! first resi in CPO (Pyroglutamic acid) GROUP ATOM CB CT2 -0.18 ATOM HB1 0.09 HA ATOM HB2 ΗA 0.09 GROUP -0.28 ATOM CG CT2 ATOM HG1 0.09 HA ATOM HG2 HA 0.09 ATOM CD CC 0.72 ATOM OE OC -0.69 ATOM N NH1 -0.40 ATOM HN 0.31 Η 0.07 ATOM CA CT1 ATOM HA HB 0.09 GROUP ATOM C С 0.51 ATOM O 0 -0.51 BOND CD N BOND CB CA CG CB CD CG BOND N HN N CA C CA BOND C CA HA CB HB1 CB HB2 CG HG1 +NBOND CG HG2 DOUBLE O C CD OE -C CA HN C CA +N O IMPR N DONOR HN N ACCEPTOR OE CD ACCEPTOR O C IC -C CA *N HN 1.3471 124.4500 180.0000 113.9900 0.9961 IC -C Ν CA С 1.3471 124.4500 180.0000 107.2700 1.5216 IC N CA С +N1.4512 107.2700 180.0000 117.2500 1.3501 IC +N CA *C 0 1.3501 117.2500 180.0000 121.0700 1.2306 IC CA С +N+CA 1.5216 117.2500 180.0000 124.3000 1.4530 IC N С *CA CB 1.4512 107.2700 121.9000 111.7100 1.5516 С *CA HA IC N 1.4512 107.2700 -118.0600 107.2600 1.0828 IC N CA CB CG 1.4512 111.0400 180.0000 115.6900 1.5557 IC CG CA *CB HB1 1.5557 115.6900 121.2200 108.1600 1.1145 IC CG CA *CB HB2 1.5557 115.6900 -123.6500 109.8100 1.1131 IC CA CB CG CD 1.5516 115.6900 180.0000 115.7300 1.5307 IC CD CB *CG HG1 1.5307 115.7300 117.3800 109.5000 1.1053 IC CD CB *CG HG2 1.5307 115.7300 -121.9600 111.0000 1.1081

 Table 50: Topology file for sugars, for n-acytyl-glucosamine and for pyroglutamic acid, which are found in CPO, and for which parameters were created by analogy

PRES SSO4 0.000 ! patch between MAN-MAN(C1-O4) DELETE ATOM 101 DELETE ATOM 1H01 DELETE ATOM 2HO4 BOND 1C1 204 GROU ATOM 1C1 CC3162 0.240 ATOM 1H1 HCA1 0.090 ATOM 1H1 HCA1 0.090 ATOM 1C5 CC3163 0.080 ATOM 1H5 HCA1 0.090 ATOM 105 OC3C61 -0.500 GROU ATOM 2C4 CC3161 0.360 ATOM 2H4 HCA1 0.090 ATOM 2H4 HCA1 0.090 ATOM 2O4 OC311 -0.450 PRES XSO4 0.000 ! patch between xys-sugar(C1-O4) DELETE ATOM 101 DELETE ATOM 1HO1 DELETE ATOM 2HO4 BOND 1C1 204 GROU ATOM C1 CC3162 0.210 ATOM H1 HCA1 0.090 ATOM C5 CC3263 0.020 ATOM H51 HCA2 0.090 ATOM H52 HCA2 0.090 ATOM 05 0C3C61 -0.500 GROU ATOM 2C4 CC3161 0.360 ATOM 2H4 HCA1 0.090 ATOM 204 OC311 -0.450 PRES SSO3 0.000 ! patch between MAN-MAN(C1-O3) DELETE ATOM 101 DELETE ATOM 1HO1 DELETE ATOM 2HO3 BOND 1C1 2O3 GROU ATOM 1C1 CC3162 0.240 ATOM 1H1 HCA1 0.090 ATOM 1C5 CC3163 0.080 ATOM 1H5 HCA1 0.090 ATOM 105 OC3C61 -0.500 GROU ATOM 2C4 CC3161 0.360 ATOM 2H4 HCA1 0.090 ATOM 204 OC311 -0.450 PRES XSO3 0.000 ! patch between XYS-MAN(C1-O3)

DELETE ATOM 101							
DELETE ATOM 1HO1							
DELETE ATOM 2HO3							
BOND 1C1 2O3							
GROU							
ATOM C1 CC3162	0.210						
ATOM H1 HCA1	0.090						
ATOM C5 CC3263	0.020						
	0.090						
	0.090						
ATOM 05 0C3C61							
GROU	0.500						
	0 360						
ATOM 2C4 CC3161 ATOM 2H4 HCA1	0.090						
ATOM 204 0C311	0.090						
AIOM 204 0C311	-0.450						
PRES SSO6	0.000 !	patch	between	MAN-MAN (C1-06)	
DELETE ATOM 101							
DELETE ATOM 1H01							
DELETE ATOM 2HO6							
BOND 1C1 206							
GROU							
ATOM 1C1 CC3162	0.240						
ATOM 1H1 HCA1	0.090						
ATOM 1C5 CC3163	0.080						
ATOM 1H5 HCA1	0.090						
ATOM 105 0C3C61							
GROU							
ATOM 2C6 CC321	0.250						
ATOM 2H61 HCA2	0.090						
ATOM 2H62 HCA2	0.090						
	-0.430						
A10M 200 0C311	-0.430						
PRES XSO6	0.000 !	patch	between	XYS-MAN (C1-06)	
DELETE ATOM 101							
DELETE ATOM 1H01							
DELETE ATOM 2HO6							
BOND 1C1 206							
GROU							
	0 010						
ATOM C1 CC3162	0.210						
ATOM H1 HCA1	0.090						
ATOM C5 CC3263	0.020						
ATOM H51 HCA2	0.090						
ATOM H52 HCA2	0.090						
ATOM 05 OC3C61	-0.500						
GROU							
ATOM 2C6 CC321	0.250						
ATOM 2H61 HCA2	0.090						
ATOM 2H62 HCA2	0.090						
ATOM 206 OC311	-0.430						
PRES SASN	0.000 !	patch	between	MAN-ASN (C1-ND2)	
		T		(,	

DELETE ATOM 101 DELETE ATOM 1H01 DELETE ATOM 2HD22 BOND 1C1 2ND2 GROU ATOM C1 CC3162 0.240 ATOM H1 HCA1 0.090 CC3163 0.080 HCA1 0.090 ATOM C5 ATOM H5 HCA1 ATOM 05 0C3C61 -0.500 GROUP ATOM 2ND2 NH2 -0.32 ATOM 2HD21 H 0.32 PRES STHR 0.000 ! patch between MAN-THR(C1-OG1) DELETE ATOM 101 DELETE ATOM 1H01 DELETE ATOM 2HG1 BOND 1C1 20G1 GROU ATOM C1 CC3162 0.240 0.090 ATOM H1 HCA1 ATOM C5 CC3163 0.080 ATOM H5 HCA1 0.090 ATOM 05 0C3C61 -0.500 GROUP
 ATOM 2CB
 CT1
 0.17

 ATOM 2HB
 HA
 0.09

 ATOM 2OG1
 OH1
 -0.26
 PRES SSER 0.000 ! patch between MAN-SER(C1-OG) DELETE ATOM 101 DELETE ATOM 1H01 DELETE ATOM 2HG1 BOND 1C1 20G GROU ATOM C1 CC3162 0.240 ATOM H1 HCA1 0.090 ATOM C5 CC3163 0.080 ATOM H5 HCA1 0.090 ATOM 05 0C3C61 -0.500 GROUP ATOM 2CB CT2 0.08 ATOM 2HB1 HA 0.09 ATOM 2HB2 HA 0.09 ATOM 20G 0H1 -0.26 PRES XSER 0.000 ! patch between AXYS-SER(C1-OG) DELETE ATOM 101 DELETE ATOM 1HO1 DELETE ATOM 2HG1

BOND 1C1 20G GROU ATOM C1 CC3162 0.210 ATOM H1 HCA1 0.090 ATOM C5 0.020 CC3263 ATOM H51 HCA2 0.090 ATOM H52 HCA2 0.090 OC3C61 -0.500 ATOM 05 GROUP ATOM 2CB CT2 0.08 ATOM 2HB1 HA 0.09 ATOM 2HB2 HA 0.09 ATOM 20G OH1 -0.26 0.00 ! Patch for HEME to CYS link PRES PCYS ! Patch residues must be 1-CYS, 2-HEME DELETE ANGLE 2NA 2FE 2NC 2NB 2FE 2ND DELETE ATOM 1HG1 GROUP CT2 ATOM CB -0.05 0.09 ATOM HB1 HA ATOM HB2 HA 0.09 ATOM SG -0.13 S BOND 1SG 2FE ANGLE 1CB 1SG 2FE 1SG 2FE 2NA 1SG 2FE 2NB ANGLE 1SG 2FE 2NC 1SG 2FE 2ND DIHE 1CB 1SG 2FE 2NA IC 1CB 1SG 2FE 2NA 1.8180 100.60 0.0000 90.000 2.0000 IC 1CB 1SG 2FE 2NB 1.8180 100.60 90.000 90.000 2.0000 IC 1CB 1SG 2FE 2NC 1.8180 100.60 180.00 90.000 2.0000 IC 1CB 1SG 2FE 2ND 1.8180 100.60 -90.000 90.000 2.0000 IC 1CA 1CB 1SG 2FE 1.5380 112.50 0.2000 100.60 2.4500

Table 51: Patches used to attach sugar-sugar molecules, sugar-amino acids and heme-cystine

```
* $Id: par_allxx_sugar.inp,v 1.40 2008/07/08 18:06:46 ganesh Exp $
*>>>>> All-hydrogen parameters used in the <<<<<<<<
*>>>>> development of the CHARMM carbohydrate force field<<<<<<
*>>>>>> Direct comments to Alexander D. MacKerell Jr. <<<<<<<*
*>>>>>> 410-706-7442 or email:
* Initial sugar parameter file
*
! par22: par_all22_prot.inp
! alex: Alex MacKerell
! sng: Shannon Greene
```

```
Olgun Guvench
! og:
! viv: Igor Vorobyov
! erh: Elizabeth Hatcher
!
BONDS
!V(bond) = Kb(b - b0) **2
!Kb: kcal/mole/A**2
!b0: A
!
! atom types
              Kb b0
!
! monosaccharide CO
CC3162 OH1
            428.00 1.400
! monosaccharide CN
CC3161 NC2D1 320.00 1.430
CC3161 NH1
           320.00 1.430
CC3162 NH2
            320.00 1.430
!**************** ANGLES ***********
ANGLES
!V(angle) = Ktheta(Theta - Theta0)**2
!V(Urey-Bradley) = Kub(S - S0)**2
!Ktheta: kcal/mole/rad**2
!Theta0: degrees
!Kub: kcal/mole/A**2 (Urey-Bradley)
!S0: A
!
            Ktheta Theta0 Kub S0
! atom types
!
!
! monosaccharide CCN
      CC3161 NH1 70.000 113.50
CC3161
        CC3162 NH2 70.000 113.50
CC3161
CC3162 CC3161 NH1 70.000 113.50
!
! monosaccharide CNC
CC3161 NH1 C 50.000 120.00
! monosaccharide NCO
OC3C61 CC3162 NH2 110.0 112.0
!
! monosaccharide NCH
NH1 CC3161 HCA1 48.000 108.00
NH2
      CC3161 HCA1 48.000 108.00
NH2
      CC3162 HCA1 48.000 108.00
!
! monosaccharide HNC
CC3161 NH1 H 35.000 117.00
!
! monosaccharide CCO
CC3161 CC3162 OH1
                    75.700 107.00
! monosaccharide OCO
```

```
OH1 CC3162 OC3C61 45.000 116.50
!
! monosaccharide COC
CC3162 OH1 CT1 95.00 109.70
!
! monosaccharide HCO
                      55.000 108.89
HCA1 CC3162 OH1
! ****** V(dihedral) = Kchi(1 + cos(n(chi) - delta))
DIHEDRALS
! monosaccharide CCCO
! C3-C2-C1(anomeric)-O5
CC3161 CC3161 CC3162 OH1 0.31 3 180.0
! monosaccharide CCCN
NH1 CC3161 CC3162 NH2 0.20 3 0.0
     CC3161 CC3161 CC3161 0.20 3 0.0
NH1
NH2 CC3162 CC3161 CC3161 0.20 3 0.0
! monosaccharide CCOC
! C2-C1-O5-C5
CC3161 CC3162 OH1 CT1 0.20 3 0.0
                                   0.0
CC3161 CC3162 OH1 CT2 0.20 3
CC3163 CC3162 OC311 CC3162 0.20 3 0.0
CC3263 CC3161 OC311 CC3162 0.20 3
                                        0.0
CC3161 CC3161 OC311 CC3162 0.20 3
                                        0.0
CC3161 CC3162 OC311 CC3161 0.20 3
                                        0.0
! C1-O-C4-C5/C3
CC3162 OC311 OC3C61 CC3163 0.20 3
                                      0.0
CC3162 OC311 OC3C61 CC3161 0.20 3
                                        0.0
! monosaccharide OCCO
! (H)O-C(anomeric)-C-O(H)
OH1CC3162CC3161OC3112.651180.0OH1CC3162CC3161OC3110.0020.0
OH1 CC3162 CC3161 OC311 0.13 3 180.0
! monosaccharide HCCO
HCA1 CC3161 CC3162 OH1 0.14 3 0.0
! monosaccharide HCCN
                           0.20 3 0.0
HCA1 CC3162 CC3161 NH1
HCA1
       CC3161 CC3162 NH2 0.20 3 0.0
HCA1 CC3161 CC3161 NH1 0.20 3 0.0
! monosaccharide OCCN
OC311 CC3161 CC3161 NH1 0.20 3 0.0
OC3C61 CC3162 CC3161 NH1
                            0.20 3 0.0
OC311 CC3162 CC3161 NH1 0.20 3
                                      0.0
! monosaccharide OCNC

        OC3C61
        CC3162
        NH2
        CC
        0.0
        3
        0.

        OC3C61
        CC3162
        NH2
        H
        0.0
        3
        0.0

                                            0.0
```

CC3161NH1CO2.502180.0CC3162NH2CCO2.502180.0 ! monosaccharide CNCC CC3161 NH1 C CT3 1.60 1 0.0 CC CT2 1.60 1 CC3162 NH2 0.0 CC NH2 CC3162 CC3161 0.50 1 180.0 NH1CC3161CC31620.501180.0NH1CC3161CC31610.501180.0 С С ! monosaccharide HCOC CC3162 OH1 CT1 0.284 3 0.0 HCA1 HCA1 CC3162 OH1 CT2 0.284 3 0.0 CC3161 OC311 CC3162 0.284 3 0.0 HCA1 CC3162 OC311 CC3161 0.284 3 HCA1 0.0 ! monosaccharide HNCC CC3162 CC3161 NH1 H 0.00 1 0.0 CC3161 CC3162 NH2 H 0.00 1 0.0 H NH1 CC3161 CC3161 0.00 1 0.0 ! monosaccharide HNCH 0.00 1 0.0 0.00 1 0.0 HCA1 CC3161 NH1 H CC3162 NH2 HCA1 Н ! monosaccharide CNCH CC3161 NH1 C 0.00 1 HCA1 0.0 0.00 1 HCA1 CC3162 NH2 CC 0.0 ! OCOC OC3C61 CC3162 OH1 CT2 0.2 3 0.0 OC3C61 CC3162 OC311 CC3161 0.2 3 0.0 CC3162 OC3C61 CC3163 0.41 1 180.0 OH1 CC3162 OC3C61 CC3163 0.89 2 0.0 OH1 CC3162 OC3C61 CC3163 0.05 3 0.0 OH1 NH1 CC 370.000 1.3450 FE NPH 270.2000 2.0000 FE S 300.0000 2.4500 CT2 S FE 100.0 50.0 100.6 FE S NPH 90.0 CT1 NH1 CC 50.000 120.0000 Н NH1 CC 34.000 123.0000 NH1 CC CT2 80.000 116.5000 CC NH1 80.000 122.5000 OC NPH FE S CT2 0.0400 4 0.00 CT1 NH1 CC 0.2000 1 С 0.2000 1 1.6000 1 2.5000 2 180.00 CT2 CC NH1 CT1 0.00 CT2 CC NH1 CT1 180.00 CT2 CT1 NH1 CC 1.8000 1 0.00 NH1 CC CT2 2.5000 2 180.00 Н

HB	CT1	NH1	CC	0.0000	1	0.00		
OC	CC	NH1	CT1	2.5000	2	180.00		
OC	CC	NH1	H	2.5000	2	180.00		

 Table 52: Lines appended by Cassian to CHARMM parameter file in order to include parameters developed for entries in topology file in Appendix B1

VITA

CASSIAN D'CUNHA

1995 – 1999	Bachelor of Engineering: Chemical Engineering University of Mumbai Mumbai, India
2000 - 2003	Master of Science: Computer Science Florida International University Miami, Florida, USA
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H. L. Alvarez, D. Chatfield, D. A. Cox, E. Crumpler, C. D'Cunha, R. Gutierrez, J. Ibarra, E. Johnson, K. Kumar, T. Milledge, G. Narasimhan, S. M. Sadjadi, C. Zhang, "CyberBridges A Model Collaboration Infrastructure for e-Science," Proceedings of the 7th IEEE International Symposium on Cluster Computing and the Grid (CCGrid'07), Rio de Janeiro, Brazil, May 2007.

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