Density functional theory studies of MTSL nitroxide side chain conformations attached to an activation loop

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Abstract A quantum mechanical (QM) method rooted on density functional theory (DFT) has been employed to determine conformations of the methane-thiosulfonate spin label (MTSL) attached to a fragment extracted from the activation loop of Aurora-A kinase. The features of the calculated energy surface revealed low energy barriers between isoenergetic minima, and the system could be described in a population of 76 rotamers that can be also considered for other systems since it was found that the χ3, χ4 and χ5 do not depend on the previous two dihedral angles. Conformational states obtained were seen to be comparable to those obtained in the α-helix systems studied previously, indicating that the protein backbone does not affect the torsional profiles significantly and suggesting the possibility to use determined conformations for other protein systems for further modelling studies.

Keywords DFT · EPR · Computational chemistry

1 Introduction

The most widely employed spin label for studies of the structure and dynamics of biomolecules [1, 2], through electron paramagnetic resonance (EPR) spectroscopy, is the methane-thiosulfonate spin label (MTSL). Figure 1 shows a model of MTSL attached to a fragment extracted from the activation loop of Aurora-A kinase, a serine/threonine protein kinase that regulates many cellular pathways and is overexpressed in a number of cancers [3, 4].

The MTSL is endowed of high flexibility and the knowledge of the potential surface is required in order to model its internal dynamics [5–7]. In this work, a quantum mechanical (QM) method based on density functional theory (DFT) was employed to determine features of the energy surface of the five dihedral angles of MTSL. The DFT theory was employed since it induces the electron correlations effect, representing an appropriate method to reproduce short electrostatic interactions between sulphur atoms and backbone atoms [7].

In the previous literature, similar approaches were adopted by Tombolato et al. [5] using the Hartree–Fock (HF) theory to study conformations of the MTSL in α-helix systems of the T4 Lysozyme protein [5–7] and results obtained were used to complement subsequent MD studies [8, 9]. Considering our system, the question arose whether the conformational states determined in the α-helix are the same in the activation loop of a different protein.

We performed a conformational analysis of the potential energy surfaces of the MTSL side chain attached to a fragment extracted from activation loop of Aurora-A kinase protein with the purpose to characterize its geometrical parameters and describe the system in a limited number of rotamers. We carried out this work to establish a basis for...
more advanced modelling approaches involving molecular dynamics (MD) simulations that require different initial starting conformations of the MTSL for appropriate statistical analyses.

2 Methods

2.1 QM calculations of the conformations of the MTSL

A short unit peptide was extracted from the X-ray crystal structure of the Aurora-A kinase domain (residues 122-403 C290A C393A; PDB 4CEG [10] with a resolution of 2.10 Å and R-value of 0.202) obtained after minimization and equilibration processes performed using the AMBER 15 package [11] in conjunction with ff14SB protein force field [12] in order to clean the structure and to remove bad contacts. Subsequently, the protein was solvated using the Extended Simple Point Charge (SPC/E) water model (9721 water molecules) in a truncated octahedral box with a buffer of 12 Å between the protein atoms and the edge of the box. Afterwards, a short energy minimization was performed in two steps using the Simulated Annealing with NMR-derived Energy Restraints (SANDER) module of AMBER. In the first stage, the water molecules and counter ions were relaxed with 200 cycles of minimization. In the second step, the entire system as a whole was relaxed with 1000 cycles of minimization. Subsequently, the system was heated at constant volume for 20 ps from 10 to 300 K with 10 kcal/mol weak restraints on the protein. This process was followed by two equilibration steps: the first was performed at constant pressure (1 atm) and temperature (300 K) for 200 ps with no restraints and the second was performed in a microcanonical (NVE) ensemble for 1 ns. Relaxed scans were performed subsequently using subsystems shown in Fig. 2 where the MTSL side chain was gradually built by adding atoms to the unit peptide extracted from the crystal structure of the Aurora-A kinase.

In order to provide reliable reproductions of the equilibrium geometries of the rotamers, the energy torsional profiles \( V_i(\chi_i) \) were calculated by performing relaxed scans in thirty-seven steps of 10° with the B3LYP hybrid functional [13, 14] and the 6-31G(d) basis set [15] in gas phase around each dihedral angle \( \chi_1, \chi_2, \chi_3, \chi_4 \) and \( \chi_5 \) of the MTSL side chain. The same model system was used in previous work [5–9]. The ab initio relaxed scans were performed in 37 steps of 10° using the \( \text{opt} = \text{mod redundant} \) keyword in the Gaussian 09 software [16] that fixes coordinates but optimizes or relaxes the other atoms.

These scans helped to identify the minima of the torsional energy profiles of all five dihedral angles. The Gibbs free energies were determined with the B3LYP hybrid functional and the 6-31G(d) basis set using the \( \text{Freq} = \text{hin-drot} \) keyword.

3 Results and discussion

3.1 Conformational analysis of the MTSL side chain from QM calculations

The MTSL side chain was gradually built in by adding atoms to the C\(_{\alpha}\) atom of the CH\(_3\)–CO–NH–C\(_{\alpha}\)–CO–NH–CH\(_3\) fragment extracted from the minimized and

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**Fig. 1** Structure of the Aurora-A kinase domain with the MTSL side chain (black circle) attached at position 288 within the activation loop which comprises residues 274–299 (grey). The unit peptide extracted for the DFT analysis is indicated by a blue arrow. All five dihedral angles of the MTSL side chain are enlarged and shown, with the pyrroline ring containing the nitroxide group highlighted in a red circle.

**Fig. 2** Subsystems considered for the calculations of the torsional energy profiles about the \( \chi_1 \) (S1), \( \chi_2 \) (S2), \( \chi_3 \) (S3), \( \chi_4 \) (S4) and \( \chi_5 \) (S5) dihedral angles.
The first relaxed scan was performed around $\chi_1$, which was obtained by performing relaxed scans in the range from $-180^\circ$ to $+180^\circ$, considering the rotation of the C$_\alpha$–C$_\beta$H$_2$ group attached to the CH$_3$–CO–NH–C$_\alpha$H–CO–NH–CH$_3$ fragment. The torsional energy profile showed three minima at the values of $-160^\circ$ and $+60^\circ$ (Fig. 4a).

The high energy barriers corresponded to eclipsed configurations, while the lower energy barriers correlated to configurations in which the S$_\gamma$ and S$_\delta$ atoms were interacting with the protein backbone. The first torsional profile (Fig. 4a) $V^{(1)}(\chi_1)$ showed three broad minima and two transitions between $-160^\circ$ $\leftrightarrow$ $-60^\circ$ and $-60^\circ$ $\leftrightarrow$ $+60^\circ$ that were separated by an energy barrier of $\sim 3$ and $\sim 5$ kcal/mol, respectively. These energy barriers are somewhat small, and transitions between them would be expected to occur frequently. Subsequently, the $V^{(2)}(\chi_2)$ torsional profiles were calculated considering the rotation around the C$_9$H$_2$S$_\gamma$ group attached to the C$_\alpha$ atom of the CH$_3$–NH–C$_\alpha$H–NH–CO–CH$_3$ fragment. The $\chi_1$ dihedral angle was set at $-160^\circ$ and $+60^\circ$ (minima in the previous scan). In the calculated $V^{(2)}(\chi_2)$ torsional profiles (Fig. 4b), energy barriers of $\sim 1$–2 kcal/mol were found between $-120^\circ$ $\leftrightarrow$ $+80^\circ$ for $\chi_1$ equal to $-160^\circ$ and $+60^\circ$, and $-160^\circ$ $\leftrightarrow$ $-70^\circ$ and $-160^\circ$ $\leftrightarrow$ $+80^\circ$ for $\chi_1$ equal to $-60^\circ$ were observed. These transitions would be expected to occur very frequently since they are separated by very small energy barriers, but are unlikely to do so considering the electrostatic interactions between the S$_\gamma$ sulphur atom and the protein backbone. The torsional profiles about $\chi_3$ were calculated at all these minima keeping the $\chi_1$ and $\chi_2$ dihedral angles fixed at the selected values shown in Fig. 4c. For the calculation of the $V^{(3)}(\chi_3)$ torsional profiles, the rotation around the C$_9$H$_2$S$_\gamma$–S$_\gamma$CH$_2$ group attached to the C$_\gamma$ atom of the CH$_3$–NH–C$_\alpha$H–NH–CO–CH$_3$ fragment was considered. Similar energy torsional profiles and two minima at $\pm 90^\circ$ separated by a higher energy barrier of $\sim 14$ kcal/mol were found for $V^{(3)}(\chi_3)$ for all seven possible combinations of the minima found for $\chi_1$ and $\chi_2$ (Fig. 4c). The energy minima corresponded to the structures stabilized by short electrostatic interactions between atoms of the MTSL side chain and the unit peptide model as shown in Fig. 5.

The S$_\gamma$ atom interacts with the NH group for $\chi_1 = -160^\circ$ at $\chi_2 = -120^\circ$ (Fig. 5Q1), $+80^\circ$ (Fig. 5Q2) and for $\chi_1 = -60^\circ$ at $\chi_2 = -160^\circ$ (Fig. 5Q3). The conformation at $\chi_1 = -60^\circ$, $\chi_2 = -70^\circ$ and $\chi_3 = +90^\circ$ was stabilized by two short electrostatic interactions between the S$_\delta$ atom of the MTSL chain and the C$_\gamma$H group in the fragment, and the CH$_2$ group and the CO group (Fig. 5Q4). The S$_\delta$ atom interacts with the NH group for $\chi_1 = -60^\circ$ and $\chi_2 = +80^\circ$ (Fig. 5Q5). The conformation at $\chi_1 = +60^\circ$, $\chi_2 = -120^\circ$ and $+80^\circ$ and $\chi_3 = \pm 90^\circ$ (Fig. 5Q6 and Q7) was stabilized by interactions between the S$_\gamma$ atom of the MTSL and the NH group of the protein backbone. The remaining $V^{(4)}(\chi_4)$

![Diagram](image-url)

**Fig. 3** Strategy used to find minima of the torsional energy profile around $\chi_1$, $\chi_2$, $\chi_3$, $\chi_4$ and $\chi_5$ dihedral angles from the relaxed scans. The first relaxed scan was performed around $\chi_1$, and three minima at $-160^\circ$, $-60^\circ$ and $+60^\circ$ were found. Subsequently, the energy torsional profile was calculated around $\chi_2$ at the minima of $\chi_1$, and two minima, one broad between $-160^\circ$ and $-120^\circ$ and another at $+80^\circ$, were found for $\chi_1 = -160^\circ$ and $+60^\circ$. Three minima (−160°, −70°, and +80°) were found for $\chi_1 = -60^\circ$. The relaxed scan around $\chi_3$ was performed at the minima of $\chi_2$, and two minima $\pm 90^\circ$ for all the possible combination of $\chi_1$ and $\chi_2$ were found. Three minima $\pm 70^\circ$ and $180^\circ$ were found in the torsional profiles of $\chi_4$. The torsional profile of $\chi_5$ showed minima depending on the values of $\chi_4$. The equilibrated structure of the Aurora-A kinase domain in which the Cartesian coordinates of the peptide atoms were kept fixed in the configuration obtained after the previous scan. The relaxed scans were performed following the strategy shown in Fig. 3. The energy minima of the $\chi_{i+1}$ dihedral angle were determined at the minima of the $\chi_i$ dihedral angle.
Fig. 4 Energy torsional profiles about the five dihedral angles $\chi_1$, $\chi_2$, $\chi_3$, $\chi_4$ and $\chi_5$ of the MTSL side chain. a Energy torsional profile about the dihedral angle, $\chi_1$. b Energy torsional profiles about the dihedral angle, $\chi_2$, at the minima of $\chi_1$. c Energy torsional profiles about the dihedral angle, $\chi_3$, at the minima of $\chi_2$. d-f Energy torsional profiles about the dihedral angle, $\chi_4$, at the minima of $\chi_3$. g Energy torsional profile about the dihedral angle, $\chi_5$ when $\chi_4$ is equal to $180^\circ$. h Energy torsional profile about the dihedral angle $\chi_5$ when $\chi_4$ is equal to $-70^\circ$. i Energy torsional profile about the dihedral angle $\chi_5$ when $\chi_4$ is equal to $+70^\circ$.

and $V^{(5)}(\chi_5)$ torsional profiles (Fig. 4d–i) were calculated considering the rotations around the $S_\delta$–CH$_2$ group (for $\chi_4$) and $S_\gamma$–CH$_2$ pyrroline nitroxide ring (for $\chi_5$) with the previous dihedral angles kept fixed at the selected values. Like for $V^{(1)}(\chi_3)$, the $V^{(4)}(\chi_4)$ torsional profiles (Fig. 4d–f) were found to be independent from the values of the previous dihedral angles, probably due to the distance from the protein backbone. Similar torsional profiles for $V^{(4)}(\chi_4)$ and three minima (±70° and 180°) were observed for all possible combinations of $\chi_1$, $\chi_2$ and $\chi_3$. Two low energy barriers of 1–2 kcal/mol were found between the minima at ±70° and 180°, and one high energy barrier between −70° and +70° was found. The $V^{(5)}(\chi_5)$ torsional profile was measured at ±70° and 180°, and three different profiles were
observed (Fig. 4g–i). The $V^{(5)}(\chi_5)$ profile at $\chi_4 = +70^\circ$ was calculated between $-90^\circ$ and $+90^\circ$ due to a clash between one of the methyl groups of the pyrroline nitroxide ring and the $C_\beta$ carbon of the MTSL chain for angles over this range. The shape of the torsional profiles obtained in this work is similar to those seen in $\alpha$-helices and values of the minima were found to be only slightly different [5–9], indicating the protein backbone does not significantly influence the torsional profiles. No relevant changes in the torsional profile of $\chi_1$ and $\chi_2$ were observed on extension of the atoms in the unit peptide (data not shown) but longer computational times. This was also observed for $\chi_3, \chi_4$ and $\chi_5$.

After calculation of the torsional energy profiles for all five dihedral angles in the unit peptide model, a population of 76 conformations was found at the minima of the torsional energy profiles. The torsional profiles showed low energy barriers from 1 to $\sim$20 kcal/mol and isoenergetic minima in the potential energy surface, indicating that the rotamer population would be fully sampled at room temperature and conformational states are expected to be obtained in the same amount. Also, the Gibbs free energy of the different conformations were seen to be comparable and equal to $-6,556,624$, $-6,556,624$ and $-6,556,623$ kcal mol$^{-1}$ for $Q_2$, $Q_5$ and $Q_7$ (characterized by different $\chi_1$), respectively. Similarly, it was observed for $Q_1$ and $Q_2$ (characterized by different $\chi_2$) with Gibbs free energy equal to $-6,556,624$ and $-6,556,622$ kcal mol$^{-1}$, and for conformations with $\chi_3$ equal to $+90^\circ$ and $+90^\circ$ that have a free energy equal to $-655,624$ and $-655,623$ kcal mol$^{-1}$. After having determined conformers of the MTSL side chain, we tested the effect of the geometric variation of the side chain on the magnetic parameters in order to exclude the contribution of the side chain to the EPR spectrum. Previous literature showed variations of $A_{zz}$ and $g_{zz}$ components upon geometrical variations (NO bond length and the CNOC out-of-plane dihedral angle) in the proxyl radical [17] and in aromatic radical rings [18]. Hence, we tested the effect of the geometry of the MTSL side chain on the magnetic properties in order to exclude any contribution of the chain on the spectrum. Six conformations were selected at the minima of the torsional profiles (Fig. 4) with different values of the dihedral angles (Table S1 ESI), and the magnetic parameters were computed at DFT level. Conformers showed comparable magnetic parameters, and minor changes were observed in the 94 GHz EPR spectra (Fig. S1 ESI). This indicated that the addition of the side chain and variations of its geometry do not alter significantly the spin density and shape of the molecular orbital that remained well localized on the NO moiety, like observed in the case of the proxyl radical ring described in previous work [17].

4 Conclusions and future work

The conformational analysis of the MTSL side chain on a short fragment of the Aurora-A kinase activation loop revealed torsional profiles comparable to those observed in...
fragments of α-helix studied in previous work [5–9]. This indicated that the backbone structure does not influence the torsional profiles significantly. A population of 76 conformers was found at the minima of the torsional profiles, and in addition, it was observed that the $\chi_3$, $\chi_4$ and $\chi_5$ do not depend on the previous two dihedral angles, suggesting that the determined set of rotamers can be considered also for other systems. This analysis can be used to determine some starting conformations for MD simulations of the MTSL spin-labelled Aurora-A kinase or other systems. On the basis of our own experience and previous work performed using MD of MTSL spin-labelled proteins [8, 9], it was observed that transitions of $\chi_4$ and $\chi_5$ are much faster than transitions of $\chi_1$, $\chi_2$ and $\chi_3$, so the starting structures can be established fixing $\chi_1$, $\chi_2$ and $\chi_3$. It was observed that conformers have comparable potential and Gibbs free energy allowing the fixing of values $\chi_1 = \pm 60^\circ$, $-160^\circ$, $\chi_2 = +80^\circ$ and $\chi_3 = \pm 90^\circ$ in order to perform more advanced modelling studies.

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