Molecular dynamics approach to the I431V mutational impact on thyroid hormone receptor-beta

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Abstract

The point mutations like I431V on thyroid hormone receptor-beta (THR-β) gene cause resistance to thyroid hormones (RTH) with the clinical diagnosis of elevated free triiodothyronine (T3) and free thyroxin (T4) but not suppressed thyroid stimulating hormone (TSH) on the blood serum. Some ultrasonographic (USG) reports of the patients with RTH show thyroid gland disorder with goiter or nodule(s) or cyst(s) and some USG reports even with RTH are normal. I431V-mutant causes more steric hindrance while binding T3 into THR-β than the native wild type THRT3. The residue on the 431-codon is dynamic in nature showing its flexibility over the course of entry and release of T3-hormone into/from the ligand binding pocket. The more increased solvent accessible surface area of I431V-mutant than that of native I431-residue makes the partial unfolding of the globular THR-β protein. The smaller height of radial distribution function between I431-mutant and T3 shows the decrease in probability of finding the atomic particles nearby T3-hormone in THRT3-MT than in THRT3-WT. The electrostatic interaction energy between native I431 and T3 is negative, but it is positive between I431V-mutant and T3. Moreover, the internal energy of I431V-mutant has been found smaller than that of native I431-residue in THRT3 systems.

Keywords: Thyroid hormone receptor; resistance to thyroid hormones; point mutation; interaction energy; internal energy.

1. Introduction

Thyroid hormone receptors (THRs) are the nuclear receptor proteins that actively bind triiodothyronine (T3: hormone secreted by thyroid gland) to mediate the biological activities associated with gene transcription. THRs regulate gene expressions upon binding to the DNA sequences called as thyroid response elements (TREs). THRs have two isoforms THR-α and THR-β that are in hetero-diversity in the different organs. T3 has the higher affinity to bind into both isoforms than thyroxin (T4). While T3
binds to the ligand binding domain (LBD) of hydrophobicity, THR shows its structural change [1-4]. The carboxy-terminal activation domain forms an amphi-pathic helix[2, 3] as a part of cavity or LBD. The dynamic nature of helix-12 (H12 in Fig. 2-c) of THR is important for the hormone binding, ligand dissociation, activation mechanism and regulation of transcription activity [5].

The mutational impact on THR reduces the hormone binding affinity and gene expression activity. Ultimately, the mutations leading to the resistance to thyroid hormone (RTH) cause the endocrine disorder. The inactivated THR-β gene by means of homozygous mutant (Thrβ⁻⁻) in mice results in a goiter and elevated levels of thyroid hormones [6]. The mutant T327I found in THR-β gene shows the paucity of symptoms and signs of thyroid dysfunction suggesting RTH despite the increased serum levels of T3, T4 and thyroid stimulating hormone (TSH) [7]. A rare mutation A268G found in axon 9 of THR-β gene results RTH with elevated serum T3 and T4, TSH in the normal range and the episodic palpitations associated with mild dizziness [8]. A heterogeneous mutation G344R is identified in axon 10 of THR-β gene of 10-month-old-female resulting in paucity of symptoms of RTH with normal T3, T4 and TSH levels [9]. Also, R383H mutation is found in THR-β gene of a woman with inappropriate central secretion of TSH despite of elevated T3 and T4 [10]. The RTH patients are clinically euthyroid or hypothyroid depending on severity of mutation. A point mutation R429Q found in mice has the elevated serum thyroid hormones but the inappropriate TSH consistent with hypothalamic-pituitary RTH [11]. The active sites of these point mutations along with the site of I431V-mutant [12] in the THR-β gene are indicated in Fig. (2-c). The helix-12 serves as a selective gatekeeper which is responsible for corepressor specificity by the wild type receptor (THR-WT). The point mutations in THR are all positioned to impact the gatekeeper functions and then to change corepressor specificity by the mutated receptor (THR-MT) [13, 14]. THR-WTs regulate normal physiological and developmental pathways, but THR-MTs result neoplastic and endocrine diseases.

![Chemical structures of (a) triiodothyronine (T3): C₁₅H₁₂I₃NO₄ with molar mass = 650.98 g/mol, (b) isoleucine (ILE): C₆H₁₃NO₂ with molar mass = 131.17 g/mol and (c) valine (VAL): C₅H₁₁NO₂ with molar mass = 117.15 g/mol[prepared from PyMOL molecular graphics system].](attachment:image)

Fig. 1: Chemical structures of (a) triiodothyronine (T3): C₁₅H₁₂I₃NO₄ with molar mass = 650.98 g/mol, (b) isoleucine (ILE): C₆H₁₃NO₂ with molar mass = 131.17 g/mol and (c) valine (VAL): C₅H₁₁NO₂ with molar mass = 117.15 g/mol[prepared from PyMOL molecular graphics system].

The I431V point mutation is detected experimentally in THR-β gene of a 14-year-old Brazilian girl with elevated T3 and T4 but unsuppressed TSH showing RTH with goiter [12]. This mutation occurs
by replacing isoleucine (ILE) amino acid residue by valine (VAL) in 431 codon of THR-β gene. I431V mutation exerts impairment of ligand-dependent release of corepressor, silencing mediator for retinoid and thyroid receptors (SMRT) with dominant negative effects on THR-WT and reducing potency. The chemical structures of T3, ILE and VAL are shown in Fig. 1. ILE is heavier and larger in size than VAL. T3 binding assay [12] shows that the hormone binding affinity of I431V-mutant THR-β is 2.6 times lower than that in wild type THR-β. THR-WT, THRT3-WT and THRT3-MT show the distinct features reflecting in terms of root mean square deviation (RMSD), radius of gyration (R_g), root mean square fluctuation (RMSF), IR spectra, radial distribution functions, solvent accessible surface area (SASA), interaction energy, internal energy, etc. The structural and physical properties of the wild type and the mutated protein-hormone systems can be studied by using molecular dynamics simulations (MDS).

2. Methodology

The structure of wild type THRT3 was taken from the x-ray crystallographic data of protein data bank code-3GwS [15]. The CHARMM force field topologies and parameters [16, 17, 18, 19] were used for MDS of the protein-hormone systems. The hetero-atomic T3-hormone was parameterized first to describe its molecular geometry, atomic masses, charges and torsion barriers [19]. The generation of the complete systems of THR-WT, THRT3-WT and THRT3-MT starting from their protein structure files to the solvation in neutral water-ion environment, equilibration, production runs and analysis of the results were performed by using the interfaces of visual molecular dynamics (VMD) [20] and nanoscale molecular dynamics (NAMD) [21] computational packages.

From the wild type native structure of THRT3, the mutated THRT3 was made after the point mutation on 431-codon replacing ILE by VAL with the help of VMD-Mutagenesis. Unliganded THR-WT, T3-liganded THRT3-WT and I431V-mutant THRT3-MT were solvated separately in the TIP3P water-box.
which was neutralized with Na\(^+\) & Cl\(^-\) ions in the concentration of 0.15 mol/L. Each water-box has the periodic cell basis vectors 50.35, 61.92 and 74.78 Å along x, y and z-axes, respectively.

Configuration files were prepared by writing TCL-scripts to simulate the systems for which Langevin piston temperature, pressure and damping coefficient are 310 K, 1 bar and 1 ps\(^{-1}\), respectively. Oscillation period of the piston was 100 fs with decay time of 50 fs and periodic cell fluctuation was isotropic. The Langevin dynamics [22, 23] was applied for all atoms except hydrogens of the systems. Particle mesh Ewald (PME) was active with the grid dimensions of (54, 64, 80 Å) and the PME coefficient of 0.258 for the electrostatic force evaluation [24]. In adjusting the force field parameters used in MDS, 1-4 scaling was 1.0; cut-off, switching and pair-list distances for the electrostatic and van der Waals interactions were 12, 10 and 14 Å, respectively. Finally, each system was geometrically optimized with 3000 conjugate gradient steps and then it was equilibrated up to 2ns at 310 K. To calculate the trajectory of atomic particles, the velocity Verlet algorithm [25] was used during MDS of the systems. All the simulations were performed adjusting integrator parameters of 2 fs/step constrained with rigid bonds of H-atoms.

Conformational stability of the protein-hormone systems was tested by plotting RMSD, RG, RMSF and SASA of protein backbone, T3-hormone and residues of interest: I431 and I431V of THR-WT, TRH3-WT and TRH3-MT. The dihedral angle distributions were analyzed from Ramachandran plots to test the steric hindrance among \(\alpha\)-helices, \(\beta\)-sheets and side chains and residues of the systems. The electrostatic and van der Waals interactions were observed between the native or mutated residue on 431-codon and T3-hormone by using NAMD-Energy-GUI. Also, number of H-bonding, probability density from radial distribution functions, IR spectral densities, internal energies of the native wild type and mutated systems were calculated, analyzed and compared by plotting the corresponding results over the simulation time.

3. Result and Discussion

THR-\(\beta\) nuclear receptor isoform consists of a chain length of 259 amino acid residues. The protein structure of wild type THR-\(\beta\) that we take for the simulation has total mass of 27640 amu and 3895 atoms. The active ligand bound to the LBD of THR-\(\beta\) is the thyroid hormone (T3) which has mass of 651 amu and 35 atoms. The plots of dihedral angles (\(\phi,\psi\)), i.e. Ramachandran plots for THR-WT, TRH3-WT and TRH3-MT are shown in Fig. 3. In these plots, most of the amino acid residues of parallel, anti-parallel and right twisted \(\beta\)-sheets lie in the blue region of I-quadrant; that of right handed \(\alpha\)-helix lie in the blue region of II-quadrant and left handed \(\alpha\)-helix lie in the green region of III-quadrant [26, 27]. The residue GLY has no side chain so that it has no steric hindrance and it is allowed in the white region. The white region is sterically disallowed for all other amino acids except GLY. The blue regions are the most allowed regions for \(\alpha\)-helices and \(\beta\)-sheets. The green area is allowed for the shorter van der Waal radii of atoms and the related amino acids. By comparing three plots of Fig. 2, we observe that smaller number of amino acid residues lies in the disallowed region for TRH3-WT than for THR-WT and TRH3-I431V. Thus, the larger steric hindrances occur for the unliganded and mutated THR-\(\beta\). The clinical analysis shows that TRH3-WT is related to euthyroid or normal thyroid functions, THR-WT is related to overt thyroidism and TRH3-MT is related to RTH [6-14].
The constancy of root mean square deviation (RMSD) and radius of gyration (RG) of the molecules THR-β protein and T3-hormone over the time frames of equilibration run (Fig. 4, 5) show the conformational stability and validity of force field topology and parameters used for MDS of wild type and mutated nuclear receptors. The more fluctuating RMSD and RG of residue I431 show its dynamic property. This dynamic property of the residues is significantly important while trapping T3 in and releasing T3 from the LBD of thyroid hormone receptors.

The values of RMSD and RG of the molecules with the standard deviations of the data are listed in the Table 1. During their 2 ns equilibration run in neutral water-ion environment, RG of 431-residue of THR-WT is being smaller with larger RMSD, i.e. it is bending more towards the center of mass of the system than that of THRT3-WT and THRT3-MT. In other words, the nature of RMSD of 431-residue change from unliganded to liganded as well as wild type to mutated THR-β. The RMSD or RG of the protein is almost same with small fluctuations for these three cases. The protein RMSD is about 2 Å and the protein RG is about 19 Å for the wild type and the mutated THR-β systems. The RMSD
and RG of T3-hormone while bounded in the LBD of nuclear receptors are 0.7 Å and 4.3 Å respectively.

**Fig. 5:** RG-plots of 431 residue, THR-protein and T3-hormone during equilibration of (a) wild type THR, (b) wild type THRT3 and (c) I431V-mutant THRT3.

**Table 1:** Average values of root mean square deviation (RMSD) and radius of gyration (RG) of unliganded wild type and T3-liganded wild type and I431V-mutant THR-β including with the standard deviation of the data obtained after MDS by their equilibration runs.

<table>
<thead>
<tr>
<th>Molecule</th>
<th>THR-WT</th>
<th>THRT3-WT</th>
<th>THRT3-MT</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>I431</td>
<td>protein</td>
<td>I431</td>
</tr>
<tr>
<td>RMSD(Å)</td>
<td>4.6 ± 2.4</td>
<td>2.0 ± 0.2</td>
<td>3.6 ± 1.6</td>
</tr>
<tr>
<td>RG(Å)</td>
<td>10.3 ± 3.7</td>
<td>18.8 ± 0.1</td>
<td>12.1 ± 3.1</td>
</tr>
</tbody>
</table>

In this study, the root mean square fluctuations (RMSF) of most of the residues of THRT3-WT are observed to be a little bit larger than that of THR-WT and THRT3-MT as shown in Fig. 6. In the I431V mutational site, the values of RMSF are 3.2 Å for ILE of THR-WT, 4.1 Å for ILE of THRT3-WT and 3.0 Å for VAL of THRT3-MT. The residues in helix-1 have the highest RMSF so that the helix-1 in THRT3-WT is more flexible.

**Fig. 6:** RMSF of amino acid residues of THR-WT, THRT3-WT and I431V-mutant THRT3-MT during their 2 ns equilibration run.
Solvent accessible surface area (SASA) of THR-WT, THRT3-WT and THRT3-MT are observed to be slightly different in the extension radius of 1.4 Å. After 500 frames of energy minimization and equilibration, the SASA with the standard deviation of the data from frame to frame simulation up to 2 ns is 13620.1 ± 253.07 Å² for THR-WT, 13952.4 ± 130.9 Å² for THRT3-WT and 13721.0 ± 147.9 Å² for THRT3-MT. In this way, the SASA of 431-residue is 14.8 ± 9.0 Å² for THR-WT, 12.6 ± 10.1 Å² for THRT3-WT and 17.4 ± 8.1 Å² for THRT3-MT. The lower SASA indicates the higher thermodynamic stability of the protein. The SASA of T3-hormone is observed to be zero indicating the hydrophobicity of LBD of THR-β. The mutation on the solvent accessible residue causes the conformational change by partial unfolding of the protein with increasing SASA [28]. Such effect is more intensive if the mutation occurs on the surface residue than on the interior of the protein. The I431V mutation is the interior one so that no significant change in SASA of wild type and mutated THR-β has been observed after their 2 ns equilibration. However, the total SASA of THR-WT and THRT3-MT are more fluctuating than that of THRT3-WT up to 3000 frames of the equilibration runs (Fig. 7-a). Furthermore, SASA of I431V-mutant is higher and less fluctuating than that of native and wild type I431-residue of THR-β (Fig.7-b).

The distributions of number of H-bonds along with the frame order of the simulations up to 2 ns equilibrations are shown in Fig. 8. Here, the average values of H-bond number with the standard deviation of the data after 500 frames with the donor-acceptor distance 4.0 Å and cut-off angle 40° among N and O atoms are 5 ± 2 for I431-THR(WT), 16 ± 4 for T3-THR(WT), 4 ± 2 for I431-THR(MT) and 15 ± 4 for T3-THR(MT).

During MDS of the wild type and mutated THR-β systems, the distance between I431 and T3 in THRT3-WT varies as 10.2 ± 2.6 Å and the distance between I431V and T3 in THRT3-MT varies as 13.1 ± 2.0 Å as shown in Fig. 9-b. Such distance is smaller and more fluctuating in THRT3-WT than in THRT3-MT. This is one of the indicators of T3 resistant features of mutated nuclear receptor proteins that cause RTH. Radial distribution function [29] or pair auto-correlation function g(r) in a system of particles measures probability of finding a particle at r distance away from the reference particle. The distribution function g(r) between I431-residue and T3-hormone has the higher value than that between I431V and T3.
and it lies in the range of 5 Å and 18 Å as shown in Fig. 9-c. In other words, the number density of mutated residue is less than that of the native wild type residue to respond for T3-hormone and TREs.

![Graph](image)

**Fig. 8:** Variation of number of H-bonds between 431-residue and protein vs. simulation time in case of (a) wild type and (b) I431V-mutant THRT3 systems.

![Graph](image)

**Fig. 9:** (a) Positions of I431V-mutant and T3-hormone in THRT3-MT, (b) changing distance between 431-codon and T3 while equilibrating THRT3-WT and THRT3-MT systems and (c) radial distribution functions between 431-residue and T3-hormone in case of wild type and mutated THRT3.

IR spectral densities using total dipole moment of the atoms in 431-residue and T3 are calculated in the frequency range of 0 to 2000 cm$^{-1}$ and their plots are different for THRT3-WT and THRT3-MT as shown in Fig. 10. The maximum IR-intensity of T3 in the wild type receptor is 0.041 at 1837.42 cm$^{-1}$ and that of I431 (wild type) is 0.027 at 1637.88 cm$^{-1}$. Again, the maximum IR-intensity of T3 in the mutated receptor is 0.037 at 1313.63 cm$^{-1}$ and that for I431V-mutant is 0.029 at 1571.37 cm$^{-1}$. The distribution has larger number of high intensity peaks for T3 and I431V in THRT3-MT than in THRT3-WT.

Total non-bonding interaction energy is the sum of electrostatic and van der Waal’s interaction energy. The changing values of interaction energy between 431-residue and T3-hormone in wild type and mutated THR-β systems are plotted with respect to the frame order in Fig. 11 and these are averaged.
over the time frames from 500 to 4013. The average values of electrostatic and van der Waals interaction energy (from NAMD-Energy-GUI using switching and cut-off distances of 10 Å and 12 Å, respectively) with their standard deviations are -0.25 ± 0.17 kcal/mol and -0.18 ± 0.07 kcal/mol between I431 and T3 of THRT3-WT and 0.21 ± 0.08 kcal/mol and -0.07 ± 0.05 kcal/mol between I431V and T3 of THRT3-MT, respectively. Thus, the electrostatic energy between I431 and T3 in THRT3-WT is negative and more fluctuating and that between I431V and T3 in THRT3-MT is positive and less fluctuating over the simulation time.

![Graph](a)

**Fig. 10:** IR spectra of (a) I431 residue and T3-hormone in wild type THRT3 and (b) I431V-mutant and T3-hormone in mutated THRT3.

![Graph](a)

**Fig. 11:** Electrostatic and van der Waals interaction energies (a) between I431 and T3 in wild type THRT3 and (b) between I431V and T3 in mutated THRT3.

Internal energy of the biomolecular system is the sum of bonding and non-bonding potentials. Coulomb energy is the largest and its fluctuation during SMD makes the total internal energy fluctuate about its mean value. Using NAMD-Energy-GUI with switching and cut-off distances of 10 Å and 12 Å, the internal energy of 431-residue, bounded T3-hormone and protein of THR-WT, THRT3-WT and THRT3-MT are calculated and the average values for the corresponding systems are listed in the Table 2. The variations of the related internal energy with respect to the simulation time are plotted in the Fig. 12. The plots for the individual potential terms show that the major contributor of fluctuations in the internal energy is the electrostatic energy of the related system.
Table 2: Internal energy of the components of unliganded wild type, T3-liganded wild type and I431V-mutant THR-β systems averaged over the values from the last 0.25 ns of their equilibrations where the including error represents standard deviation of the data.

<table>
<thead>
<tr>
<th>Molecule</th>
<th>THR-WT</th>
<th>THRT3-WT</th>
<th>THRT3-MT</th>
</tr>
</thead>
<tbody>
<tr>
<td>I431 protein</td>
<td>42.82 ± 3.21</td>
<td>42.88 ± 3.18</td>
<td>36.13 ± 3.07</td>
</tr>
<tr>
<td>T3</td>
<td>-822.38 ± 67.67</td>
<td>128.73 ± 4.52</td>
<td>124.41 ± 4.63</td>
</tr>
<tr>
<td>I431V protein</td>
<td>-644.21 ± 109.87</td>
<td>-109.87 ± 3.07</td>
<td>-756.54 ± 104.22</td>
</tr>
</tbody>
</table>

Fig. 12: Fluctuating total internal energy along the frame order of (a) I431-residue and T3, and (b) THR-proteins for unliganded as well as T3-liganded wild type and I431V-mutant THR-β systems in the last 0.25 ns of their equilibrations.

4. Conclusion

Conformational properties and interactions of mutational residues and T3-hormone of unliganded and T3-liganded thyroid hormone receptor-beta (THR-β) can be studied by means of molecular dynamics simulations (MDS). The almost constant values of root mean square deviation (RMSD ≈ 2 Å) and radius of gyration (RG ≈ 19 Å) of native and I431-mutant THR-β proteins show their conformational stability during MDS. The native I431-residue has largely fluctuating RMSD and RG showing its dynamic property in adapting and releasing T3-hormone. The experimentally verified I431V mutation on THR-β gene (Ferreira et al., 2010) is the subject of this computational study. This mutation causes resistance to thyroid hormones [12]. Solvent accessible surface area (SASA) is larger for the I431V-mutant indicating partial unfolding of the globular protein. However, the root mean square fluctuation of I431V-mutant is smaller in comparison with that of native wild type I431. The SASA of T3 is zero verifying the hydrophobicity upon binding to the ligand binding domain (LBD) of THR-β.

Comparing the radial distribution functions g(r) for the atomic particles between I431 and T3 as well as I431V and T3, the probability density is higher in the former wild type than the latter mutated type for which g(r) covers the distance of 5 Å to 18 Å. IR spectral density calculations for I431-residue and T3-hormone in the frequency range of 0-2000 cm⁻¹ imply that greater number of intensity maxima is found in THRT3-MT than in THRT3-WT. This signifies higher dipole moment distributions by supporting less globular stability of the mutated THR-β than that of the wild type THR-β. Furthermore, electrostatic interaction energy between I431V-mutant and T3 is positive whereas it is negative between
native I431-residue and T3. In this study, internal energy of the native I431-residue is 42.88 ± 3.18 kcal/mol and that of wild type I431V-mutant is 36.13 ± 3.07 kcal/mol in THRT3 systems. The major contributor of such change in the internal energy is the electrostatic energy.

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