Molecular Dynamics Simulations of Insulin: Elucidating the Conformational Changes that Enable Its Binding

Anastasios Papaioannou¹,²*, Serdar Kuyucak², Zdenka Kuncic¹,²

¹ Charles Perkins Centre, University of Sydney, Sydney, NSW, Australia, ² School of Physics, University of Sydney, Sydney, NSW, Australia

* a.papaioannou@physics.usyd.edu.au

Abstract

A sequence of complex conformational changes is required for insulin to bind to the insulin receptor. Recent experimental evidence points to the B chain C-terminal (BC-CT) as the location of these changes in insulin. Here, we present molecular dynamics simulations of insulin that reveal new insights into the structural changes occurring in the BC-CT. We find three key results: 1) The opening of the BC-CT is inherently stochastic and progresses through an open and then a “wide-open” conformation—the wide-open conformation is essential for receptor binding, but occurs only rarely. 2) The BC-CT opens with a zipper-like mechanism, with a hinge at the Phe24 residue, and is maintained in the dominant closed/inactive state by hydrophobic interactions of the neighboring Tyr26, the critical residue where opening of the BC-CT (activation of insulin) is initiated. 3) The mutation Y26N is a potential candidate as a therapeutic insulin analogue. Overall, our results suggest that the binding of insulin to its receptor is a highly dynamic and stochastic process, where initial docking occurs in an open conformation and full binding is facilitated through interactions of insulin receptor residues with insulin in its wide-open conformation.

Introduction

Insulin is a polypeptide hormone that plays a crucial role in regulating glucose levels in higher organisms. Defects in insulin signalling can lead to insulin resistance, which is the hallmark of diabetes mellitus. The key molecular mechanisms and kinetics in the insulin-signaling pathway have been extensively studied but a complete picture is yet to emerge. In particular, the dynamics of how insulin binds to the insulin receptor (IR) and activates signal transduction needs to be better understood for pharmacological applications. Insulin comprises two polypeptide chains, a 21 amino acid A-chain made of two α-helices, and a 30 amino acid B-chain containing a central α-helix (see Fig 1A). The two chains are constrained by two inter-chain (A7–B7 and A20–B19) and one intra-chain (A6–A11) disulphide bridges (see Fig 1B). The first crystal structure of insulin was determined by Adams et al. in 1969 [1] followed by other structures of insulin and its analogs in the storage form (hexamers and dimers) [2,3]. Insulin monomer
triggers signal transduction through binding to the tyrosine kinase-type IR with high affinity, which is a disulphide-linked (αβ)_2 homodimer [3]. Two surfaces on the insulin molecule have been identified to interact with the IR [4]. The first comprises predominantly the hormone-dimerizing residues and interacts with the α_CT segment from one IR α-chain and the central β-sheet of the L1 domain of the other α-chain (primary binding site) [4–9]. The second comprises predominantly the hormone-hexamerizing residues and is suggested to interact with the IR site at the junction of FnIII-1 and 2 of the α-chain (site 2) [4,6–11].

Attempts to dock insulin to the IR using the available crystal structures have been unsuccessful to date [12,13], indicating that conformational changes are required to enable its binding. Several recent experimental studies [14–20] have further revealed that substantial conformational changes occur in both insulin and the primary site of IR during binding. These are: initial detachment (opening) of the B-chain C-terminal (BC-CT) of insulin from its hydrophobic (helical) core to expose key binding residues (see Fig 1A and 1B), followed by a rotation and re-modeling of the α_CT portion of IR upon the surface of the β-sheet of the L1 domain of IR, and lastly docking of the insulin B-chain α-helix in a parallel orientation to the α_CT segment, creating strong hydrophobic interactions in between. Extensive mutagenesis and structural studies of insulin substantiate the critical role of the BC-CT, and in particular the residues F24–Y26, in binding to IR [17,18,21–25]. Furthermore, the first crystal structure of the insulin–IR complex has revealed the extent of the structural changes and the importance of the BC-CT residues of insulin, especially F24–Y26, as well as the crucial role of H710 and F714 of the α_CT segment [14,26]. In a recent experimental study [15], a hinge in the BC-CT (F24 residue) was identified, which is critical for insulin to engage the α_CT segment of IR (see Fig 1B and 1C). In contrast to these numerous experimental studies of the insulin-IR complex, there

---

**Fig 1. Structural details of the insulin molecule.** (A) The structure of the insulin (PDB ID: 2G4M). A-Chain (orange); B-chain N-terminal (residues F1–G8) (red), B-chain α-helix (residues S9–C19) (black), B-chain C-terminal (residues G20–T30) (blue). The colored spheres represent the Cα atoms of the first and last residues of the A and B-chains. (B) The new ribbon representation shows the hydrophobic core of insulin, comprising the residues F24, L15, Y26, L11, V12, G8 (side chains shown in cyan), I2a and V3α (side chains shown in green). The hydroxyl group of Y26 and the oxygen atom of G8 are shown in red, and the disulfide bridges in yellow. (C) The hydrogen bonds between the residues neighboring Y24 and the α chain residues, G23(O)–N2α(N) and F25(N)–Y19α(O), are illustrated in blue dashed lines. Oxygen and nitrogen atoms are represented in red and blue, respectively.

doi:10.1371/journal.pone.0144058.g001
have been relatively few computational studies investigating the conformational and dynamic properties of insulin [17,24,27,28]. Furthermore, all the previous experimental and computational studies on insulin focused on the structural changes that occur in the BC-CT during activation and on mutations in the critical residues, F24 (all the residues mentioned without extra labelling belong to chain B. The A chain residues are denoted by a subscript A) and Y26 [15,17–19,21–25,27,28]. Two key questions regarding the activation of insulin remain unanswered: the frequency of the BC-CT opening, which is necessary for insulin to bind to the IR, and the mechanism that triggers this opening. We, therefore, performed MD simulations of wild-type (WT) and mutated insulin to gain new insights into the active form of insulin, i.e., the opening of the BC-CT, and the behavior of its hydrophobic core consisting of the residues G8, L11, V12, L15, F24 and Y26 from the B-chain and I2A and V3A from the A-chain (see Fig 1B). The opening of the BC-CT was studied by examining the distances between the neighboring BC-CT and \( \alpha \)-helix residues (see Table 1), and constructing the potential of mean force (PMF) from the distance distributions. Finally, the mechanism that triggers opening of the hydrophobic core was studied via mutations of the critical residues on the BC-CT and observing the behavior of water molecules in the immediate vicinity of the core.

### Methods

#### Insulin Modelling

In the MD simulations, we used a crystal structure of insulin with a resolution of 1.8 Å (PDB ID: 2G4M [29]). All simulations were constructed using the VMD software following the same protocol. The structure of insulin was solvated with approximately 3500 water molecules. Water molecules associated with the crystal structure, which help to stabilize the protein, were also included in the simulation box. Nine Na\(^+\) and seven Cl\(^-\) ions were placed randomly in the solvated system to neutralize it and to set the NaCl concentration to 0.1 M. The extra 2 Na\(^+\) ions are needed to neutralize the negative charges of the protein and thus avoid any artifacts arising from the use of periodic boundaries. The simulations were then subjected to an initial minimization of water molecules followed by equilibration. The \( x \), \( y \) and \( z \) dimensions of the water box were approximately 54, 52 and 45 Å, respectively. In the first stage of equilibration, the system was equilibrated with 1 atm pressure coupling until correct water densities were obtained. In the next stage, the restraints on the side chain and backbone atoms of the protein were gradually relaxed by reducing them from \( k = 30 \) to 0.1 kcal/mol/Å\(^2\). Finally, the protein was released, i.e., all restraints on the protein were removed, and the resulting system was then ready for MD simulations. Equilibration of the system was monitored from the RMSD data.

| \( C_\alpha \) atom of the BC-CT residue and the closest \( C_\alpha \) atom in the B-chain \( \alpha \)-helix | Distances in the crystal structure 2G4M [Å] | Average distances from MD simulations where insulin is in the closed state [Å] |
|---|---|---|
| F24(\( C_\alpha \))–L15(\( C_\alpha \)) | 6.7 | 7.0±0.4 |
| F25(\( C_\alpha \))–L15(\( C_\alpha \)) | 8.5 | 9.4±0.5 |
| Y26(\( C_\alpha \))–V12(\( C_\alpha \)) | 7.2 | 8.1±0.9 |
| T27(\( C_\alpha \))–V12(\( C_\alpha \)) | 10.2 | 10.5±1.1 |
| P28(\( C_\alpha \))–G8(\( C_\alpha \)) | 9.1 | 9.8±2.1 |
| L29(\( C_\alpha \))–G8(\( C_\alpha \)) | 12.2 | 12.3±2.4 |
| T30(\( C_\alpha \))–G8(\( C_\alpha \)) | 14.5 | 13.8±2.7 |

doi:10.1371/journal.pone.0144058.t001
MD Simulations
The MD simulations were performed using the NAMD package [30] employing the CHARMM36 force field [31,32]. The NpT ensemble was used with periodic boundary conditions in combination with the Particle Mesh Ewald (PME) method, which calculates the long-range electrostatic interactions in the periodic system. The temperature and pressure were maintained at 300 K and 1 atm, respectively, by applying the Langevin coupling with a damping coefficient of 5 ps⁻¹. The simulations were carried out with a switching cut-off within a distance of 10–13 Å for Lennard-Jones interactions. The list of non-bonded interactions was truncated at 13.5 Å applying an update frequency of 1 ps. The time step employed in all simulations was 2 fs and trajectory data were recorded every 10 ps over the course of the simulation.

WT Insulin
We performed a 690 ns MD simulation for WT insulin, which was split into six shorter ones to improve sampling (5×120 ns and 1×90 ns). Different relaxation times were used for the starting configuration in each MD simulation, i.e., 250, 300, ..., 500 ps. The Root Mean Square Deviations (RMSDs) of insulin for individual MD simulations are shown in Fig 2. Each RMSD has a distinct initial behavior, indicating the independence of the starting configurations. All the statistical analyses, discussed in the Results subsection “Characterization of the active form of insulin”, are based on the long MD simulation of WT insulin.

Water molecules within the hydrophobic core
We investigated the role of water molecules in the opening of the BC-CT by calculating the number of water molecules within a sphere that covers the whole volume of the hydrophobic core during the long MD simulations. The sphere was centred on the Cγ atom of the L15 residue, located in the centre of the hydrophobic core, and had a radius of 9 Å. While the hydrophobic core had a few residual water molecules in the closed state, the sphere included additional water molecules outside the core, which averaged to nine for closed conformations. Thus, nine water molecules were subtracted from the total number of water molecules found in the sphere.

Potential of Mean Force (PMF) calculation
The PMF describes how a system’s free energy changes as a function of a specific reaction coordinate. To characterize the dynamics of the BC-CT opening, we chose for this purpose the distance of the Cα atoms of the BC-CT residues from the closest Cα atoms in the B-chain α-helix (see Table 1). We note that these coordinates do not necessarily represent any direct coupling between the pairs of residues, but were chosen because they provide the best description for the opening of the BC-CT. As we had a sufficiently long sampling of insulin in the MD simulations, the PMF was determined directly from the density data using the Boltzmann equation

\[ W = -kT \ln \left( \frac{\rho}{\rho_0} \right) \]  

where \( T = 300 \) K is the system temperature, \( \rho \) is the density obtained from distance distributions, and \( \rho_0 \) is a reference density. With this information, we can determine how much free energy is required to change insulin’s conformation from the closed/inactive to the open/active forms.
Mutated Insulin

We also considered several mutations on the Y26 residue to examine the behavior of the active conformation of insulin and to compare the simulation results with the experimental findings. The mutant insulin structures were created using the mutator plugin in the VMD software [33] and a 100 ns MD simulation was performed in each case using the same simulation protocol as described in the "MD simulations".

Fig 2. RMSDs of six MD simulations of insulin. The first five MD simulations have a duration of 120 ns, while the sixth one 90 ns. The RMSDs are given with respect to the crystal structure 2G4M.

doi:10.1371/journal.pone.0144058.g002
Breaking of the F24 hinge

In order to break the hinge at the F24 residue and facilitate a wider opening of the BC-CT, we applied a repulsive harmonic force to the C\textsubscript{\alpha} atoms of the F24 and P28 residues relative to their partners in the B-chain \alpha-helix (see Table 1). The potential \( U \) used is given by

\[
U = \begin{cases} 
\frac{1}{2} k (r - r_0)^2, & r < r_0 \\
0, & r > r_0 
\end{cases}
\]

where \( k \) is the spring constant, \( r \) is the distance between the C\textsubscript{\alpha} atoms of the chosen pairs, and \( r_0 \) is the target distance. A force constant of 30 kcal/mol/Å\(^2\) was used and \( r_0 \) was determined by adding 9.8 Å to the average distances given in Table 1. This potential was applied to the C\textsubscript{\alpha} atoms of the pairs of residues F24–L15 and P28–G8 for 200 ps, which was sufficient to break the hinge. After that, a 30 ns MD simulation was performed without any forces applied to insulin.

Results and Discussion

Characterization of the active form of insulin

Visual inspection of the long MD simulations of WT insulin shows that it stays predominantly in the closed conformation and the BC-CT makes only occasional excursions away from the B-chain \alpha-helix. The MD simulations of insulin in the closed state were validated by comparing with the corresponding crystal structure 2G4M. As shown in Table 1, the average distances between the C\textsubscript{\alpha} atoms of BC-CT and B-chain \alpha-helix agree with the experimental values within 1 Å. We note that the fluctuations in distances increase from F24 to T30, which points to an effective interaction that diminishes with the distance from the hinge towards the C-terminal. As discussed in Fig 1B, the side chains of F24 and Y26 contribute to the formation of the hydrophobic core, which is responsible for maintaining insulin in the closed state. Table 2 lists the strongest hydrophobic interactions involving the F24 and Y26 side chains and compares the average distances obtained from the MD simulations with the experimental values. Again there is a good agreement between the simulation results and experiments, which further validates the MD simulations.

Examination of the time series of the distances listed in Table 1 shows that the F24–L15 distance does not change, consistent with a hinge behavior, while others increase to varying degrees. As an example, Fig 3 shows the distances between the C\textsubscript{\alpha} atoms of the pairs Y26–V12 and P28–G8 from the MD simulation. These two distances provide the most useful indicators

Table 2. The strong interactions in the hydrophobic core that are responsible for maintaining insulin in its closed conformation.

| Interaction          | Distance in the crystal structure [Å] | Average distance during MD simulations where insulin stays at its closed conformation [Å] |
|----------------------|---------------------------------------|-----------------------------------------------------------------------------------------|
| F24(Aromatic)–L15(C) | 5.2                                   | 4.6±0.5                                                                                 |
| Y26(Aromatic)–I2A(C) | 5.2                                   | 5.5±2.1                                                                                 |
| Y26(Aromatic)–L11(C) | 5.2                                   | 5.8±1.3                                                                                 |
| Y26(Aromatic)–V12(C) | 3.8                                   | 4.5±1.0                                                                                 |

The distances were calculated from the center of the benzene ring (aromatic) to the nearest C atom. The same trajectory data as in Table 1 were used in calculating the average distances.

doi:10.1371/journal.pone.0144058.t002
for opening of the BC-CT, and hence warrant further consideration. Their average distances are approximately 8.1 and 9.8 Å, respectively. It is evident from all the MD simulations that there is no consistent or systematic dynamic behavior characterizing the opening like a switch or quasi-periodic motion between two states. Insulin activation instead occurs stochastically at uncorrelated simulation times and frequencies, and appears to be a random event.

To analyze the frequency of the opening quantitatively, Fig 4 plot the distance histograms obtained from the 690 ns of trajectory data as probability densities, representing the probability of a distance occurring during the MD simulations. The peak values in the distance
distributions occur at approximately 8.2 Å for both Y26–V12 and P28–G8. The threshold for the open conformation was identified by the entry of water molecules into the hydrophobic core. A non zero number of water molecules inside the hydrophobic core indicates a transition from the closed to the open conformation. Based on this criterion, we found that a distance of less than 8.7 and 9.5 Å for the Y26–V12 and P28–G8 distributions, respectively, corresponds to the closed conformation of insulin. This is indicated by the black histograms in Fig 4, which clearly also indicates that the most preferable conformation of insulin is the closed/inactive form (64% from the Y26–V12 distribution). The probability of the open conformation (cyan histograms) decreases rapidly with increasing Cα atom distance. Fig 4 also suggests the existence of a third conformation, corresponding to the tail end of the distribution (blue histograms). We refer to this as the “wide-open” conformation, defined as the open conformation of insulin that enables it to fit to the IR, attributable to a ≈50° rotation (wide opening) of the F24-T30 BC-CT part about F24 [15]. The boundary between the open and wide-open conformations is found to occur when the Y26-V12 and P28-G8 distances are approximately 10.7 and 15 Å, respectively. From the distance histograms in Fig 4, the wide-open conformation has a low probability of occurrence (5% from the Y26–V12 distribution) and thus it is a relatively rare event. This suggests that initial docking of insulin to the IR may occur in an “open” conformation and the “wide-open” conformation required for full docking may be facilitated through interactions with the IR residues.

We next consider the energetics of insulin opening by converting the distance distributions for the pairs listed in Table 1 to PMFs using Eq 2. Fig 5A shows the PMFs for all the pairs.
**Fig 5. PMFs calculation.** (A) The PMFs as a function of the distance between the pairs shown in Table 1. (B) Illustration of the pairs. The spheres in blue and red represent the Cα atoms of the BC-CT and B-chain α-helix residues, respectively.

doi:10.1371/journal.pone.0144058.g005
involved in the opening of insulin, plotted as a function of distances between the C\(_{\alpha}\) atoms of the BC-CT and B-chain \(\alpha\)-helix (cf. Table 1 and Fig 5B). To facilitate comparison of the PMFs, the minima are set to zero and the curves are aligned at the same equilibrium distance (because the PMFs are effective interactions, the equilibrium distances are relative, not absolute). The C-terminal residue T30 is not included in the PMFs as it moves approximately randomly. The next BC-CT residue (L29) is seen to be the most flexible. Its conformation changes from closed to open within 1 kcal/mol. The amount of energy needed for conformational changes increases systematically from L29 to the hinge residue F24. This suggests that the BC-CT opens in a zipper-like fashion. This opening is initiated in the end residues of the C-terminal and continues through successive residues on the BC-CT until the hinge residue F24 (see Fig 6).
Our simulations further suggest that the mechanism of insulin opening is driven by the entry of water molecules into the hydrophobic core. Fig 7 shows the correlations between the number of water molecules within the hydrophobic core with the opening of the BC-CT during two randomly selected intervals from the 690 ns MD simulations. These correlations indicate that the main criterion for the opening is the Y26(Cα)–V12(Cα) distance. When this increases by $\approx 4 \, \text{Å}$, approximately half a dozen water molecules enter the hydrophobic core, thereby stabilizing the open configuration for several ns. The simulations show that the lower extremity of the hydrophobic core, in close proximity to the Y26 side chain, is never completely devoid of water. It always retains 2–3 water molecules, which randomly make H-bonds with the hydroxyl group of Y26 and/or the carbonyl oxygen of G8. Our results suggest that the interaction of these water molecules with the Y26 hydroxyl group causes its aromatic side chain to fluctuate. We observed that these fluctuations randomly drive the aromatic side chain away from the core, thereby creating more space and opportunity for water to weaken and break the hydrophobic interactions and as a result allow additional water molecules to enter the core.

Understanding the behavior of the hydrophobic core

The hydrophobic core in insulin is formed by the residues G8, L11, V12, L15, F24, Y26, I2A and V3A (see Fig 1B). While both the F24 and Y26 side chains contribute significantly to the formation of the hydrophobic core, they play very different roles in the opening process and it is important to understand the cause of this behavior. The F24 side chain is completely buried in the protein with no exposure to water and has a strong hydrophobic interaction with the L15 side chain. The position of F24 is further bolstered due to the residues flanking F24 forming H-bonds with the A-chain, i.e., G23(O)–N21A(N) and F25(N)–Y19A(O) (see Fig 1C). Thus the network of interactions around F24 result in a very stable hinge point at F24 and this explain why F24 retains its conformation during the opening of insulin, as observed in our MD simulations and also observed experimentally by Menting et al. [15]. In contrast, the side chain of Y26 is exposed to water and is more flexible due to the interaction of its hydroxyl group with the neighboring water molecules. Also T27 flanking Y26 is not involved in any stabilizing interactions with other residues. Thus the motion of Y26 is not restricted by its neighbor and the hydrophobic interactions between the Y26 side chain and the other residues in the core can be broken more easily, which triggers the opening of the BC-CT.

To examine the role of the Y26 side chain in the activation of insulin, we performed MD simulations of insulin with the mutations, Y26F, Y26N, Y26A and Y26G, for which there exists available mutagenesis data [18,22,24]. The effect of these mutations on the opening of the BC-CT is shown in Fig 8, while Table 3 lists the corresponding probability of each conformation. When the aromatic side chain of Y26 is replaced with a non-aromatic side chains, as in Y26N, Y26A and Y26G, the active form of insulin becomes the dominant conformation. In particular, the Y26G mutation causes the largest change from the WT insulin with the wide-open conformation becoming dominant (88% probability compared to 5%). The absence of an aromatic side chain weakens the hydrophobic interactions and thereby creates more space for water molecules to enter the hydrophobic core. The result in each case is a more flexible BC-CT and a much wider opening compared to that of WT insulin (cf. Fig 8). These simulation results corroborate the experimental results of Zakova et al. [24], where the mutations Y26G and Y26N were observed to result in an open conformation of insulin.

The role of the hydroxyl group in the Y26 side chain becomes evident upon closer inspection of the Y26F mutation. As seen from Fig 8 and Table 3, this mutation has negligible effect on the behavior of the opening, which is in agreement with the mutagenesis data [24]. Our simulation results show that the phenylalanine side chain occupies almost the same area as the
tyrosine side chain and interacts similarly with the other residues of the core. A closer inspection of Fig 8 and Table 3 reveals a slightly more stable closed conformation for the Y26F mutation compared to WT insulin. We attribute this to the destabilizing effect of the hydroxyl-water interactions in WT insulin, which are absent in the Y26F mutation. Overall, our results indicate that Y26 is the critical residue in the activation of insulin. We note that the mutations that lead to a more active form of insulin have been found to generally also reduce its binding affinity to the IR [24], thus indicating that Y26 also plays an important role in binding of insulin. Of these mutations, only Y26N has been found to exhibit IR selectivity [24].

We next investigate the stability of the hinge at the F24 residue. Initially, we performed 30 ns MD simulations of insulin with the mutations, F24A and F24H. In both cases, we found that the hinge is highly stable and did not break during the simulations. As a result, a more dynamic process was then used. A harmonic force is applied to the Cα atoms of the F24 and P28 residues for 200 ps to break the hinge and open the BC-CT in the WT insulin, followed by a 30 ns MD simulation of the protein with no forces applied. An identical simulation was performed for the F24E mutation to highlight the role of the aromatic side chain in stabilizing the hinge. Fig 9 shows the distances between the Cα atoms for the pairs F24–L15 and Y26–V12 for both the

![Graph showing distances between Cα atoms for F24-L15 and Y26-V12 for different mutations]

**Fig 8. Effect of Y26 mutations on the BC-CT opening.** The distance between the Cα atoms of the Y26–V12 residues, which determines the criterion for the BC-CT opening, for the Y26G, Y26A, Y26N and Y26F mutations. The closed conformation distance in WT insulin is shown for reference (black dashed line).

doi:10.1371/journal.pone.0144058.g008
WT and the mutated insulin over the course of the simulation. The hinge, after breaking, is seen to reform in less than a nanosecond in WT insulin, which returns to its closed conformation after a few nanoseconds as indicated by the F24–L15 and Y26–V12 distances in Fig 9. Thus even after the hinge is broken via an external force, the F24 and Y26 aromatic side chains quickly return to their previous conformations, restoring the strong hydrophobic interactions with the other residues in the core and maintaining the insulin in its closed conformation for the rest of the simulation. In contrast, the F24E mutation disrupts the hydrophobic core by allowing water molecules to enter the core. This leads to the breakage of the hinge at F24 leaving insulin in a wide-open conformation during the whole simulation. We note that the F24E mutation results in a much wider opening of the BC-CT compared to the mutations of Y26, which clearly can be attributed to the breaking of the hinge.

Conclusions

Our simulation results have revealed for the first time that the activation of insulin is stochastic in nature, with no obvious deterministic behavior. The BC-CT opens randomly and its dominant conformation is the closed/inactive one. The open conformation of the BC-CT does not match the receptor-bound interface. We have found an additional “wide-open” conformation, which uniquely enables insulin to bind to the IR. However, our simulations indicate that this wide-open conformation is a rare event (<5% probability).

Our PMF results also reveal that the energy needed for the BC-CT residues to change their conformation from the closed to the open form is lowest for the residues near the C-terminal and increases towards the residues closer to the hinge. From this we deduce that the BC-CT opens like a zipper, i.e., the opening is initiated in the last residue and the other residues open progressively until the hinge is reached. We found that although the last residues of the BC-CT (P28–T30) are the most flexible, they are not capable of initiating the opening of insulin necessary for its activation. The hydrophobic interactions in the core maintain insulin in its closed conformation and the opening is feasible only when the core breaks at its outer limit (Y26). We also found that the opening process is triggered by the fluctuations in hydrophobic interactions, which allow entry of water molecules into the hydrophobic core, thereby disrupting the hydrophobic interactions between the nonpolar side chains of Y26 and the other residues in the core. The hinge at F24 was found to be very stable. We attribute this to bolstering of the hydrophobic interactions of F24 with the hydrogen bonds of the flanking residues.

Finally, our MD simulations have revealed that the Y26N, Y26A and Y26G mutations yield a higher probability of the wide-open conformation needed for IR binding, owing to more space being available in the core for water molecules to enter and weaken the hydrophobic interactions therein. Of these mutations, only the Y26N mutation was found to exhibit significant IR selectivity in previous experimental studies [24]. Thus, this particular mutation
Fig 9. Stability of the hinge. Distances between the C\alpha atoms of the F24–L15 and Y26–V12 pairs, which represent the hinge and the BC-CT opening, respectively, for the WT (blue) and the mutated (red) insulin, after the hinge is broken and the protein is in its wide-open conformation. The trajectories are compared to the closed conformation distance of WT insulin (black dashed line).

doi:10.1371/journal.pone.0144058.g009
presents potential as a therapeutic insulin analogue. We plan to investigate this further with insulin-IR docking simulations.

In conclusion, our MD simulations provide new insights on the conformational dynamics of insulin that are necessary to enable its binding to the IR. In particular, we propose that binding of insulin to the IR most likely follows a two-stage process: the first is the stochastic opening of the BC-CT when insulin is in the vicinity of the αCT segment of the IR; in the second stage, conformational changes occur in both insulin (to the wide-open conformation) and the αCT segment of the IR, thereby facilitating switching of insulin from its free to bound conformation.

Acknowledgments
AP acknowledges support from a University of Sydney International Scholarship (USydIS). MD simulations were carried out using the Raijin cluster at the National Computational Infrastructure (NCI, Canberra, Australia) and the Avoca cluster at the Victorian Life Sciences Computation Initiative (VLSCI, Melbourne, Australia).

Author Contributions
Conceived and designed the experiments: AP SK ZK. Performed the experiments: AP. Analyzed the data: AP. Wrote the paper: AP SK ZK.

References
1. Adams MJ, Blundell TL, Dodson EJ, Dodson GG, Vijayan M, Baker EN, et al. Structure of rhombohedral 2 Zinc insulin crystals. Nature. 1969; 224: 491–495.
2. Baker EN, Blundell TL, Cutfield JF, Cutfield SM, Dodson EJ, Dodson GG, et al. The structure of 2Zn pig insulin crystals at 1.5 A resolution. Philos. Trans. R. Soc. Lond. B. Biol. Sci. 1988; 319: 369–456. PMID: 2905485
3. De Meyts P, Whittaker J. Structural biology of insulin and IGF1 receptors: implications for drug design. Nat. Rev. Drug Discov. 2002; 1: 769–783. PMID: 12360255
4. De Meyts P. Insulin and its receptor: structure, function and evolution. Bioessays. 2004; 26: 1351–1362. PMID: 15551269
5. Smith BJ, Huang K, Kong G, Chan SJ, Nakagawa S, Menting JG, et al. Structural resolution of a tandem hormone-binding element in the insulin receptor and its implications for design of peptide agonists. Proc. Natl. Acad. Sci. U. S. A. 2010; 107: 6771–6776. doi: 10.1073/pnas.1001813107 PMID: 20348418
6. Ward CW, Lawrence MC. Ligand-induced activation of the insulin receptor: a multi-step process involving structural changes in both the ligand and the receptor. Bioessays. 2009; 31: 422–434. doi: 10.1002/bies.200800210 PMID: 19274663
7. Ward CW, Lawrence MC. Similar but different: ligand-induced activation of the insulin and epidermal growth factor receptor families. Curr. Opin. Struct. Biol. 2012; 22: 360–366. doi: 10.1016/j.sbi.2012.03.014 PMID: 22521506
8. De Meyts P, Sajid W, Paisgaard J, Theede AM, Gauguin L, Aladdin H, et al. Insulin and IGF-I receptor structure and binding mechanism. In: Saltiel AR, Pessin JE, editors. Mechanisms of Insulin Action. Springer, New York; 2007. pp. 1–32.
9. De Meyts P. Insulin/receptor binding: the last piece of the puzzle? Bioessays. 2015; 37: 389–397 doi: 10.1002/bies.201400190 PMID: 25630923
10. McKern NM, Lawrence MC, Streltsov VA, Lou MZ, Adams TE, Lovrecz GO, et al. Structure of the insulin receptor ectodomain reveals a folded-over conformation. Nature. 2006; 443: 218–221. PMID: 16957736
11. Whittaker L, Hao C, Fu W, Whittaker J. High-affinity insulin binding: insulin interacts with two receptor ligand binding sites. Biochemistry. 2008; 47: 12900–12909. doi: 10.1021/bi801693h PMID: 18991400
12. Vashishth H, Abrams CF. Docking of insulin to a structurally equilibrated insulin receptor ectodomain. Proteins. 2010; 78: 1531–1543. doi: 10.1002/prot.22670 PMID: 20112420
13. Vashisth H, Abrams CF. All-atom structural models of insulin binding to the insulin receptor in the presence of a tandem hormone-binding element. Proteins. 2013; 81: 1017–1030. doi: 10.1002/pro.24255 PMID: 23348915

14. Menting JG, Whittaker J, Margetts MB, Whittaker LJ, Kong GK, Smith BJ, et al. How insulin engages its primary binding site on the insulin receptor. Nature. 2013; 493: 241–245. doi: 10.1038/nature11781 PMID: 23302862

15. Menting JG, Yang Y, Chan SJ, Phillips NB, Smith BJ, Whittaker J, et al. Protective hinge in insulin opens to enable its receptor engagement. Proc. Natl. Acad. Sci. U. S. A. 2014; 111: E3395–E3404. doi: 10.1073/pnas.1412897111 PMID: 25092300

16. Ward CW, Menting JG, Lawrence MC. The insulin receptor changes conformation in unforeseen ways on ligand binding: sharpening the picture of insulin receptor activation. Bioessays. 2013; 35: 945–954. doi: 10.1002/bies.201300065 PMID: 24037759

17. Zakova L, Kletvikova E, Veverka V, Lepsik M, Watson CJ, Turkenburg JP, et al. Structural integrity of the B24 site in human insulin is important for hormone functionality. J. Biol. Chem. 2013; 288: 10230–10240. doi: 10.1074/jbc.M112.448050 PMID: 23447530

18. Jirasek J, Zakova L, Antolikova E, Watson CJ, Turkenburg JP, Dodson GG, et al. Implications for the active form of human insulin based on the structural convergence of highly active hormone analogues. Proc. Natl. Acad. Sci. U. S. A. 2010; 107: 1966–1970 doi: 10.1073/pnas.0911785107 PMID: 20133841

19. Glendorf T, Sorensen AR, Nishimura E, Petterson I, Kjeldsen T. Importance of the solvent-exposed residues of the insulin B-chain alpha-helix for receptor binding. Biochemistry. 2008; 47: 4743–4751 doi: 10.1021/bi800054z PMID: 18376848

20. Whittaker J, Whittaker LJ, Roberts CT, Phillips NB, Ismail-Beigi F, Lawrence MC, et al. Helical element at the hormone-binding surface of the insulin receptor functions as a signaling element to activate its tyrosine kinase. Proc. Natl. Acad. Sci. U. S. A. 2012; 109: 11166–11171. doi: 10.1073/pnas.1205681109 PMID: 22736795

21. Pandyarajan V, Smith BJ, Phillips NB, Whittaker L, Cox GP, Wickramasinghe N, et al. Aromatic anchor at an invariant hormone-receptor interface: function of insulin residue B24 with application to protein design. J. Biol. Chem. 2014; 289: 34709–34727. doi: 10.1074/jbc.M114.608562 PMID: 25305014

22. Zakova L, Barth T, Jiracek J, Barbova J, Zorad S. Shortened insulin analogues: marked changes in biological activity resulting from replacement of TyrB26 and N-methylation of peptide bonds in the C-terminus of the B-chain. Biochemistry. 2004; 43: 2323–2331. PMID: 14979729

23. Zakova L, Kazdova L, Hanclova I, Protivinska E, Sanda M, Budesinsky M, et al. Insulin analogues with modifications at position B26. Divergence of binding affinity and biological activity. Biochemistry, 2008; 47: 5858–5868. doi: 10.1021/bi800054z PMID: 18452310

24. Zakova L, Kletvikova E, Lepsik M, Collins M, Watson CJ, Turkenburg JP, et al. Human insulin analogues modified at the B26 site reveal a hormone conformation that is undetected in the receptor complex. Acta Crystallogr. D. Biol. Crystallogr. 2014; 70: 2765–2774. doi: 10.1107/S1399004714017775 PMID: 25286859

25. Kosinova L, Veverka V, Novotna P, Collins M, Urbanova M, Moody NR, et al. Insight into the structural and biological relevance of the T/R transition of the N-terminus of the B-chain in human insulin. Biochemistry. 2014; 53: 3392–3402. doi: 10.1021/bi500073z PMID: 24819248

26. Menting JG, Lawrence CF, Kong GK, Margetts MB, Ward CW, Lawrence MC. Structural congruency of ligand binding to the insulin and Insulin/Type 1 insulin-like growth factor hybrid receptors. Structure. 2015; 23: 1–12.

27. Zoete V, Meuwly M, Karplus M. A comparison of the dynamic behavior of monomeric and dimeric insulin shows structural rearrangements in the active monomer. J Mol Biol. 2004; 342: 913–929. PMID: 15342246

28. Legge FS, Budi A, Treutlein H, Yarovsky I. Protein flexibility: multiple molecular dynamics simulations of insulin chain B. Biophys. Chem. 2006; 119: 146–157. PMID: 16129550

29. Mueller-Dieckmann C, Panjikar S, Schmidt A, Mueller S, Kuper J, Geerlof A, et al. On the routine use of soft X-rays in macromolecular crystallography. Part IV. Efficient determination of anomalous substructures in biomacromolecules using longer X-ray wavelengths. Acta Crystallogr. D. Biol. Crystallogr. 2007; 63: 366–380. PMID: 17327674

30. Phillips JC, Braun R, Wang W, Gumbart J, Tajkhorshid E, Villa E, et al. Scalable molecular dynamics with NAMD. J. Comput. Chem. 2005; 26: 1781–1802. PMID: 16222654

31. MacKerell AD, Brooks B, Brooks CL, Nilsson L, Roux B, Won Y, et al. CHARMm: The energy function and its parameterization. In: Schleyer PvR, editor-in-chief. The Encyclopedia of Computational Chemistry. John Wiley and Sons, Chichester; 1998. pp. 271–277.
32. Brooks BR, Brooks CL III, Mackerell AD Jr, Nilsson L, Petrella RJ, Roux B, et al. CHARMM: the biomolecular simulation program. J. Comput. Chem. 2009; 30: 1545–614. doi:10.1002/jcc.21287 PMID: 19444816

33. Humphrey W, Dalke A, Schulten K. VMD: visual molecular dynamics. J. Mol. Graph. 1996; 14: 33–38. PMID: 8744570