The Role of Water in the Stability of Wild Type and Mutant Insulin Dimers

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Abstract

Insulin dimerization and aggregation play important roles in the endogenous delivery of the hormone. One of the important residues at the insulin dimer interface is PheB24 which is an invariant aromatic anchor that packs towards its own monomer inside a hydrophobic cavity formed by ValB12, LeuB15, TyrB16, CysB19 and TyrB26. Using molecular dynamics and free energy simulations in explicit solvent, the structural and dynamical consequences of mutations of Phe at position B24 to Gly, Ala, and d-Ala and the des-PheB25 variant are quantified. Consistent with experiments it is found that the Gly and Ala modifications lead to insulin dimers with reduced stability by 4 and 5 kcal/mol from thermodynamic integration and 4 and 8 kcal/mol from results
using MM-GBSA, respectively. Given the experimental difficulties to quantify the thermodynamic stability of modified insulin dimers, such computations provide a valuable complement. Interestingly, the Gly-mutant exists as a strongly and a weakly interacting dimer. Analysis of the molecular dynamics simulations shows that this can be explained by water molecules that replace direct monomer-monomer H-bonding contacts at the dimerization interface involving residues B24 to B26. It is concluded that such solvent molecules play an essential role and must be included in future insulin dimerization studies.

**Introduction**

Insulin is a small, aggregating protein that plays an eminent role in regulating glucose uptake in cells. In its crystal form, the WT hormone is a hexamer consisting of three dimers with either two or four Zn atoms bound to it. Each dimer consists of two monomers (chain A with 21 amino acids and chain B with 30 amino acids), connected by two inter-chain (Cys\(^A7\)–Cys\(^B7\), Cys\(^A20\)–Cys\(^B19\)) and one intra-chain (Cys\(^A6\)–Cys\(^A11\)) disulfide bonds (see Figure 1). Under physiological conditions insulin monomers readily aggregate to form dimers. Experimental and computational studies have found that the main stabilizing contributions to self-association are nonpolar interactions through directionality provided by hydrogen bonds. Dissociation of the dimer to form two monomeric insulins is of great physiological importance as the monomer is the functionally relevant state of the hormone. One suggested possibility to suppress aggregation is to modify the dimerization interface through suitable substitutions or chemical modifications.

The current view of the insulin structure-function relationship is derived primarily from insulin hexamer and dimer crystal structures, as well as from studies of the structure-activity correlations of chemically modified and/or naturally occurring mutant insulins in solution. Mutagenesis of the dimer-forming surface of insulin can yield analogues with a
reduced tendency to aggregate and pronounced differences in the pharmacokinetic properties with potentially promising therapeutic applications.\textsuperscript{7,10} Typical experimental methods for quantitative studies of insulin dimerization are isothermal titration calorimetry (ITC)\textsuperscript{3} or NMR spectroscopy. ITC requires relatively high protein concentrations, while NMR spectroscopy can be slow for such purposes. Several NMR studies of active monomeric insulin mutants show a rearrangement of the C-terminal end of chain B.\textsuperscript{8,14} The removal of residues B26–B30 in despentapeptide prevents dimerization without any significant changes in the rest of the molecule.\textsuperscript{11,13,17} This truncated insulin is at least 50 % as potent as human insulin.\textsuperscript{8,14} A combined Raman spectroscopy and microscopy study of insulin in different aggregation states (monomer, dimer, hexamer and fibril) shows that dimerization damps fluctuations at an intermolecular $\beta$-sheet.\textsuperscript{18} Experimental alanine scanning finds substitution of alanine at various positions to reduce insulin affinity for the receptor by more than 20-fold.\textsuperscript{19} While the residues that are most likely to be directly involved in binding are A1, A2, A3, A19, B12, B23 and B24, any substitution of residues A1–A3 has been shown to impair function.\textsuperscript{15}

Phenylalanine at position B24 is invariant among insulin sequences and is located at the dimerization interface maintaining the orientation of the B-chain of the monomer.\textsuperscript{5,12,22} These observations together with studies of low-potency B24 analogues suggest that the Phe$^{B24}$ amino acid residue plays an important role\textsuperscript{23} in the activity of insulin, while Ser$^{B24}$ (Ref.\textsuperscript{24,25}), Leu$^{B24}$ (Ref.\textsuperscript{26}), and His$^{B24}$ (Ref.\textsuperscript{27}) analogues show reduced binding potency. However, it was also found that certain B24 substitutions, such as Gly,\textsuperscript{16,28} D-Ala,\textsuperscript{11,29} D-His,\textsuperscript{27} Met and Cha\textsuperscript{30} are well tolerated in view of the affinity of insulin to its receptor. The bioactivity of such insulins has also been described as “anomalous” because it can not be readily explained by crystal models.

Experimental data for dimerization free energies of insulin analogues is scarce due to several experimental challenges. The role of Phe$^{B24}$ in stabilizing the insulin dimer has also
Figure 1: Structure of insulin dimer (PDB Code: 4INS). Chain A1 (blue), Chain B1 (red), Chain A2 (grey), Chain B2 (orange). Residue 24 in both monomers is shown in stick representation.

been studied to some extent. Unlike WT insulin, the Gly\textsubscript{B24} mutant does not dimerize in aqueous solution at pH 1.9.\textsuperscript{8} Furthermore, alanine scanning of the dimerization interface revealed that the Ala\textsubscript{B24} analogue is monomeric and does not readily aggregate.\textsuperscript{31,32} This suggests that the Ala\textsubscript{B24} and Gly\textsubscript{B24} analogue dimers are less stable than the WT dimer. Furthermore, ITC measurements of N-methylated insulin dimer analogues at positions B24, B25, and B26\textsuperscript{33,34} revealed considerably reduced (by a factor of 5) dimerization capabilities compared with human insulin.

Because the dimer↔monomer equilibrium is one of the essential steps in forming the receptor binding-competent monomeric form of insulin and the process is difficult to study quantitatively by experiments, MD simulations are an attractive alternative to characterize the stability of insulin dimers. In the present work the stability and dynamics of the insulin dimer analogues (Gly\textsubscript{B24}, Ala\textsubscript{B24}, D-Ala\textsubscript{B24} and des-Phe\textsubscript{B25}) are investigated using atomistic...
simulations in explicit solvent. The relative stabilities of the analogues are compared with the native WT dimer and with qualitative results from experiments. Both, protein-protein and protein-water contacts are analyzed to characterize the role of water and hydrogen bonding at the dimer-forming surface. Previous computational studies of the WT dimer and of alanine scans have provided important complementary information to experimental characterizations and are a rational basis to extend such an approach to modified insulins using explicit free energy simulations. Hence, the present work lays the foundation to extend insulin dimerization studies to arbitrary modifications at the interface.

This article is organized as follows: Section 2 introduces the methods. In Section 3 the dynamics and stability of insulin dimers and the inter-molecular hydrogen bonding between monomer-monomer and water-protein (water-bridged) is presented and discussed. Finally conclusions are drawn in Section 4.

Methods

Molecular Dynamics Simulations

Molecular dynamics (MD) simulations were carried out using CHARMM together with the “all-atom” CHARMM force field including the CMAP correction and periodic boundary conditions (PBC). Additional validation simulations for the insulin dimer were carried out using Gromacs and the CHARMM36 force field as described in the supporting information. The starting coordinates for the MD simulations were the X-ray structure of the WT porcine insulin dimer resolved at 1.5 Å (Protein Data Bank (PDB), Code: 4INS). The structure contains the coordinates of the insulin dimer and two aggregated Zn atoms. Zn atoms are removed as they been shown to be relevant only for hexamer formation. Hydrogen atoms were added to the X-ray structure. The resulting structure was used to gen-
erate mutants computationally. For this, the Phe residue at position B24 was mutated into Glycine (Gly), Alanine (Ala), and D-Alanine (D-Ala) yielding mutants Gly$^{B24}$, Ala$^{B24}$, and D-Ala$^{B24}$, respectively. Furthermore, the wildtype (WT) insulin dimer without the Phe$^{B25}$ amino acid on both monomers, a des-Phe$^{B25}$ mutant was also studied.

The wildtype dimer and mutants were solvated in a 77.6 × 62.8 × 55.8 Å box of TIP3P water molecules. All MD simulations were carried out in explicit water. Water molecules overlapping the protein were removed which leads to a system with approximately 1550 protein atoms and 8495 water molecules. The solvent was equilibrated at 300 K for 30 ps with the insulin frozen. Then 2000 steps of steepest descent (SD) minimization were carried out. The entire system was heated to 300 K during 15 ps using harmonic constraints with a force constant of 5 kcal/mol Å$^2$ on the position of the backbone atoms. The system was further equilibrated for 120 ps by gradually decreasing harmonic constraints on the backbone atoms. For all simulations the Verlet leapfrog integrator was used for time propagation with a time step of 1 fs. A 12 Å cutoff was applied to the shifted electrostatic and switched van der Waals interactions and images for periodic boundary conditions were updated every 10 time steps. All distances to hydrogen atoms were constrained by using SHAKE. For the WT dimer and mutants (Gly$^{B24}$, Ala$^{B24}$, D-Ala$^{B24}$ and des-Phe$^{B25}$) multiple individual trajectories were run starting from different structures taken from the equilibration run (Table S1). Simulations in the NPT and NVT ensembles were run using the extended system constant pressure and temperature (CPT) algorithms with a Hoover thermostat. In addition, microcanonical NVE simulations were run as well.

**Calculation of Binding Free Energy**

In the present work, the stability of the WT and mutant insulin dimers is determined from two complementary approaches. One of them is the molecular mechanics-generalized Born
surface area (MM-GBSA) approach\cite{50,51} and the other one is thermodynamic integration.\cite{52,53}

The thermodynamic cycles used for computing the binding free energy $\Delta G_{\text{bind}}$ using MM-GBSA and thermodynamic integration (TI) are shown in Figures 2A and B, respectively. Computational details for MM-GBSA are provided in the Supporting Information.

\begin{equation}
\Delta G_{\text{bind}} = \Delta G_{\text{bind}}^0 - \Delta G_{\text{solv}}^{\text{Monomer I + Monomer II}} + \Delta G_{\text{solv}}^{\text{Dimer vac}}
\end{equation}

\begin{equation}
\Delta G_{\text{bind}}^1 = <\Delta G_{\text{bind}}^0 - \Delta G_{\text{solv}}^{\text{Monomer I + Monomer II}} + \Delta G_{\text{solv}}^{\text{Dimer vac}}> + \Delta T \Delta S
\end{equation}

\begin{equation}
\Delta\Delta G_{\text{stab}} = \Delta G_{\text{Dimer}} - 2x\Delta G_{\text{Monomer}}
\end{equation}

\begin{equation}
\Delta\Delta G_{\text{stab}} = \Delta G_{2} - \Delta G_{1}
\end{equation}

Figure 2: Thermodynamic cycles used to determine stabilization free energies from MM-GBSA simulations (A) and from thermodynamic integration (B). In (B, left panel) the thermodynamic cycle used to compute the protein dimerisation stability differences, and the right panel shows the thermodynamic cycle used to compute the stability differences by mutating the B24 side chain only.

To corroborate the results from the MM-GBSA simulations and to investigate the role of
water molecules on the thermodynamic stability, dimer stabilization free energies were also
determined from thermodynamic integration (TI) in the presence of explicit water molecules.
TI applies a scaling parameter $\lambda$ to switch between an initial (state A, $\lambda = 0$) and the final
(state B, $\lambda = 1$) state by gradually damping all nonbonded interactions. The $\lambda = 0$ and
$\lambda = 1$ states correspond to the grown and annihilated nonbonded interactions on either
the protein dimer or on the monomer, respectively. Initial coordinates were taken from the
equilibrated simulations. TI simulations were performed using CHARMM’s PERT module
using soft-core potentials$^{54–56}$ for the LJ interactions that applied only on the repulsive part
of the LJ potential (it was used when the electrostatic interactions were turned off). For this
computational approach, restraining potentials$^{57}$ affecting the translational, rotational and
conformational freedom of the protein may be activated and released during the simulations
to aid convergence and improve the sampling.

Free energy simulations at each $\lambda$–value were carried out in the $NPT$ ensemble, using the
Hoover heat-bath method$^{49}$ with pressure coupling at $T = 298$ K, $p = 1$ atm, and with the
masses of the temperature and pressure piston set to roughly 20 % and 2 % of the system’s
mass, respectively. A friction coefficient of 50 ps$^{-1}$ was used. The interval $0 < \lambda < 1$
was divided into 40 equidistant steps to ensure accuracy. For each $\lambda$–value the system was
re-equilibrated for 60 ps followed by 100 ps of dynamics during which information was accu-
mulated. $\lambda$ was changed from initial to final value using the slow-growth protocol,$^{58}$ which
allowed the system to re-equilibrate between steps. Contrary to the MM-GBSA simulations
(see SI), TI run in this fashion includes all enthalpic and entropic contributions. Assuming
that the change in the entropic contribution $\Delta\Delta S_{WT-Mutant}$ remains approximately constant
for the various mutants it is expected that the stability ranking from the two methods re-
mains the same which is a testable hypothesis.

The protein dimer stability difference $\Delta\Delta G_{\text{stab}} = \Delta G_{\text{dimer}} - 2 \times \Delta G_{\text{monomer}}$ was computed
within the "same trajectory method" \textsuperscript{11} such as to close the thermodynamic cycle, see Figure 2B (left panel).

The free energy of mutating Phe\textsubscript{B24} (WT) into Gly and Ala was also calculated directly using the dual topology approach \textsuperscript{54,59} for mutations F24G and F24A in the dimer (Figure 2, Panel B right). The effect of mutating the B24 side chain on the stability of the protein $\Delta \Delta G$ was calculated according to $\Delta G_2 - \Delta G_1$, where $\Delta G_1$ and $\Delta G_2$ are the free energies of mutating the side chain of a WT residue into another residue (mutant), in the aqueous phase, as a sole residue and in the protein, respectively. For these simulations, no restraints were used and the interval $0 < \lambda < 1$ was divided into 34 steps. Windows at the two ends of the $\lambda$ interval were more finely spaced. For each of these steps the system was re-equilibrated for 30 ps followed by 60 ps of dynamics during which information was accumulated. The results are averages and standard deviations of five runs.

\section*{Results and Discussion}

In the following, first, the stabilities from MM-GBSA and thermodynamics integration simulations of the dimers are discussed. Then, the findings are discussed in the context of the dynamics and interactions along the dimerization interface. Finally, the role of water molecules is considered.

\section*{Dimerization Free Energies}

One of the main objectives of the present study is to determine the relative stabilities of mutated insulin dimer analogues relative to the WT dimer. Experimentally, a dimerization energy of $-7.2 \pm 0.8$ kcal/mol \textsuperscript{60} in favour of the dimer was determined for WT insulin
which compares with $\Delta G_{\text{bind}} = -11.9 \pm 6.7$ kcal/mol (absolute binding free energy) and $\langle \Delta G_{\text{bind}}^0 \rangle + \langle \Delta G_{\text{desolv}} \rangle = -38.7 \pm 5.8$ kcal/mol (enthalpic contribution to $\Delta G_{\text{bind}}$) from previous molecular dynamics simulations.

Table 1 reports the different contributions to the calculated absolute binding free energies of dimerization for the different insulin analogues including the WT. The enthalpic contribution $G_{\text{enthalpic}}$ is the sum of $E_{\text{vdW}}, E_{\text{elec}}, G_{\text{solv,elec}},$ and $G_{\text{solv,nb}}$ which is $\sim -45$ kcal/mol for the WT dimer. Compared to this, Gly$^{B24}$, Ala$^{B24}$, and D-Ala$^{B24}$ (approximately $-38, -38, -33$ kcal/mol, respectively) are enthalpically less stable. The des-Phe$^{B25}$ analogue is least stable ($\sim -16$ kcal/mol). The destabilization energy of about 7–12 kcal/mol for the different energetically low-lying analogues of the insulin dimer is related to replacing the two Phe$^{B24}$ residues (in both monomers) which contribute an average stabilization of $-3.96$ and $-3.36$ kcal/mol, to the total stabilization of the WT protein (see Figure S1). These contributions are consistent with results reported in an earlier MM-GBSA study which found $-3.92$ and $-2.68$ kcal/mol, respectively. Experimentally, the des-Phe$^{B25}$ insulin dimer was also found to be unstable as it reported exclusively monomeric insulin for this variant. The individual contributions (see Table 1) to the binding free energy suggest that stabilization is predominantly due to nonpolar terms $E_{\text{vdW}}$ and $G_{\text{solv,nb}}$. The favorable $E_{\text{elec}}$ contribution from the two monomers is canceled by the desolvation energy $G_{\text{solv,elec}}$ upon dimerization. This is found for all modified insulins investigated here and supports previous investigations of the WT.

Including the entropic contribution $T(\Delta S_{\text{trans}} + \Delta S_{\text{rot}} + \Delta S_{\text{vis}})$ to $\Delta G_{\text{bind}}$, allows more direct comparison with experimentally determined values. For the WT the stabilization free energy is $\Delta G_{\text{bind}} = -16.0 \pm 6.9$ kcal/mol when accounting for entropic contributions, in qualitative agreement with the experimental binding free energy of $-7.2 \pm 0.8$ kcal/mol. As a comparison, previous simulation work based on one trajectory found an enthalpic stabilization
Table 1: Binding free energy (kcal/mol) decomposition for the insulin dimerization of various insulin analogues: vDW, electrostatic, and solvation (elec. and nonpolar) contributions, using the same trajectory method by MD simulations.

|                      | WT         | Gly<sup>B24</sup> | Ala<sup>B24</sup> | D-Ala<sup>B24</sup> | des-Phe<sup>B25</sup> |
|----------------------|------------|-------------------|-------------------|---------------------|-----------------------|
| ⟨Δ<sub>vdw</sub>⟩   | -66.59(4.4) | -53.39(4.3)       | -71.35(4.0)       | -60.39(5.1)         | -44.47(5.2)           |
| ⟨Δ<sub>ele</sub>⟩   | -114.38(34.3) | -107.84(40.7)    | -111.55(26.5)     | -104.85(33.9)       | -56.15(32.2)          |
| ⟨Δ<sub>ele,desolv</sub>⟩ | 146.23(32.6) | 132.83(38.4)   | 155.99(23.8)      | 142.77(30.8)        | 92.43(29.6)           |
| Δ<sub>np,desolv</sub>⟩ | -10.28(0.6)  | -9.76(0.5)       | -11.49(0.4)       | -10.54(0.5)         | 7.99(0.6)             |
| ⟨Δ<sub>bind</sub>⟩ + | -45.02(6.5)  | -38.16(7.3)      | -38.40(7.5)       | -33.00(9.5)         | -16.19(6.5)           |
| −T⟨Δ<sub>trans</sub>⟩ | 13.02<sup>a</sup> | 13.02<sup>a</sup> | 13.02<sup>a</sup> | 13.02<sup>a</sup>   | 13.00<sup>a</sup>     |
| −T⟨Δ<sub>rot</sub>⟩ | 14.36(0.02)  | 14.31(0.02)      | 14.35(0.01)       | 14.33(0.01)         | 14.32(0.02)           |
| −T⟨Δ<sub>vib</sub>⟩ | 1.61(3.7)    | -1.43(4.0)       | 3.50(3.5)         | -1.87(4.1)          | -4.52(3.6)            |
| −T⟨Δ<sub>S</sub>⟩  | 28.99(3.7)   | 25.90(4.0)       | 30.86(3.5)        | 25.47(4.1)          | 22.80(3.6)            |
| ⟨Δ<sub>bind</sub>⟩  | -16.03(6.9)  | -12.26(8.0)      | -7.54(8.0)        | -7.53(10.9)         | 6.61(7.2)             |

<sup>a</sup> The standard deviation of −T⟨Δ<sub>trans</sub>⟩ is not defined, because it is a function of mass, which has constant value.

<sup>†</sup> The standard state is taken to be 1 M as was used by Tidor <i>et al.</i><sup>61</sup></table>

of −38.6±5.8 kcal/mol for the WT dimer which decreased to −11.9±6.7 kcal/mol when including entropic contributions.<sup>11</sup> This is consistent with earlier studies<sup>61</sup> on the dimerization of WT insulin which found an unfavourable entropic contribution of ≈ 30 kcal/mol (depending on the size and shape of the protein). On the other hand, for relative stabilization free energies upon mutation the entropic part is less important except for the contribution due to vibrations. For the protein variants considered here, TΔ<sub>vib</sub> ranges from −2 to 3.5 kcal/mol and hence can contribute up to 6 kcal/mol to the differential stabilization of one protein variant relative to another one.

The results in Table 1 show that Ala<sup>B24</sup> is the entropically least favored substitution while des-Phe<sup>B25</sup> is most favoured. Adding the entropic and enthalpic contributions, the total binding free energies of dimerization of insulins leads to stabilisation ranging from −16 to −7 kcal/mol among the low-lying analogues, <i>i.e.</i>, WT, Gly<sup>B24</sup>, Ala<sup>B24</sup> and D-Ala<sup>B24</sup>. The
des-Phe$^{B25}$ mutant with $\Delta G_{\text{bind}}$ of $\sim 6$ kcal/mol is energetically unfavorable and is expected to be monomeric in solution.

Overall, 46 independent (10 for WT, Gly$^{B24}$, Ala$^{B24}$ and des-Phe$^{B25}$, 6 for D-Ala$^{B24}$) free energy simulations were performed, each 10 ns long, which amounts to a typical aggregate of 100 ns for each system studied. These simulations were also run in different statistical mechanical ensembles and Table S1 in the SI provides a comprehensive summary. Considering the enthalpic part of $\Delta G_{\text{bind}}$ for all the trajectories (WT and B24 insulin dimer analogues) two situations can be distinguished (see Table S1 and Figures S2 to S7 in the SI for illustrations): 1) Stable, low-lying ($\sim -45$ kcal/mol) analogues with binding energies comparable to the WT dimer, and 2) Less-stable, high-lying ($\sim -20$ kcal/mol) analogues with decreased stability. The ten different WT trajectories show similar binding energies. Even Ala$^{B24}$, and des-Phe$^{B25}$ show similar binding energies among various trajectories. However, Gly$^{B24}$ and D-Ala$^{B24}$ show appreciable differences in binding energies among various trajectories. Specifically, for the Gly$^{B24}$ mutant a strongly (SI) and a weakly interacting (WI) dimer is found which will be discussed further below.

Dimer stabilization free energies were also determined from thermodynamic integration (TI, see methods) which provide a direct validation of the MM-GBSA results. TI simulations were carried out for the WT (Phe$^{B24}$) and the 3 mutants (Gly$^{B24}$, Ala$^{B24}$ and des-Phe$^{B25}$) using the single topology method. The computed stabilization free energy of $-8.4$ kcal/mol for the WT dimer differs by about 1 kcal/mol from the experimentally measured value of $-7.2 \pm 0.8$ kcal/mol$^{60}$ which provides a direct validation of the TI simulations. Relative to WT, the Gly$^{B24}$ and Ala$^{B24}$ mutants are destabilized by $\Delta \Delta G_{\text{WT/GlyB24}}^{\text{TI}} = 3.8$ kcal/mol and $\Delta \Delta G_{\text{WT/AlaB24}}^{\text{TI}} = 5.4$ kcal/mol, respectively, compared with $\Delta \Delta G_{\text{WT/GlyB24}}^{\text{MM-GBSA}} = 3.7$ kcal/mol and $\Delta \Delta G_{\text{WT/AlaB24}}^{\text{MM-GBSA}} = 8.5$ kcal/mol from MM-GBSA simulations, see Table 1. The des-Phe$^{B25}$ variant is unstable in TI as was also found from MM-GBSA. Hence, the TI and
MM-GBSA simulations are consistent with one another and support the experimental observation that the Ala$^{B24}$ mutant is marginally stable/unstable in solution$^{31,32}$ which serves as an additional validation of the present simulations.

Table 2: Stability free energy (kcal/mol) of the various insulin dimer analogues. The error on the computed values is reported in parentheses.

|        | WT       | Gly$^{B24}$ | Ala$^{B24}$ | des-Phe$^{B25}$ |
|--------|----------|-------------|-------------|-----------------|
| $\Delta \Delta G_{\text{Stability}}$ | −8.4 (0.2) | −4.6 (0.2) | −3.0 (0.2) | 2.3 (0.2) |

The free energy of mutating Phe$^{B24}$ (WT) into Gly and Ala was also calculated directly (Figure 2, Panel B right) using the dual topology approach$^{54,59}$ for mutations F24G and F24A in the dimer. These computations yield $\Delta \Delta G_{\text{PheB24} \rightarrow \text{GlyB24}} = 3.2 \pm 0.2$ kcal/mol and $\Delta \Delta G_{\text{PheB24} \rightarrow \text{AlaB24}} = 4.2 \pm 0.2$ kcal/mol. In other words, the Gly$^{B24}$ and Ala$^{B24}$ mutants are destabilized by 3.2 and 4.2 kcal/mol relative to the WT protein, respectively, which compares and is consistent with values of 3.8 and 5.4 kcal/mol from the first set of TI simulations, see Table 2.

In summary, the MM-GBSA and TI simulations all agree in that WT (Phe$^{B24}$) is most stable, followed by Gly$^{B24}$ and Ala$^{B24}$ mutants. The des-Phe$^{B25}$ variant is unstable. Furthermore, the differential stabilization free energies of two TI simulations differ by 1 to 1.5 kcal/mol which is, however, acceptable given the very different ways in which they were carried out.

The Weakly and Strongly Interacting Gly$^{B24}$ Dimer

The MM-GBSA simulations found a strongly (SI) and a weakly (WI) interacting Gly$^{B24}$ dimer. For the SI variant ($\Delta G \approx −48$ kcal/mol) the dimer is stabilized almost as strongly as the WT dimer, and for WI, the dimer is considerably destabilized by almost 30 kcal/mol.
(ΔG ≈ −20 kcal/mol). Figure S1 shows per-residue contributions to the total binding free energies of WT, SI and WI GlyB24 dimers. This analysis indicates that the differences mainly arise from contributions to the electrostatic <E_{elec}> and solvation energy <G_{elec,desolv}> (see Figure S2). For the WT and GlyB24-SI insulin dimer the per-residue binding free energies follow a similar pattern whereas for GlyB24-WI they differ (see Figure S1). For instance, most of the residues have favorable contributions to the total binding free energies for WT and SI (orange and yellow bars in Figure S1, respectively), whereas for the WI dimer (purple bars in Figure S1) these contributions are clearly reduced or even reversed which gives rise to reduced stabilization. Four residues of WI destabilize the dimer by > 2 kcal/mol.

One residue that contributes significantly to the differences between WI and SI is GluB13 (see Figure S1) which makes unfavorable contributions of about 2.5 and 3.4 kcal/mol to the B1 and B2 chain, respectively. The differences were further analyzed and the electrostatic part was found to be primarily responsible for that, see Figure S8. The pattern of the per-residue contributions to the total dimerization free energy (see Figure S1) suggests that the two monomers in WT, SI and WI dimers are equivalent although they are not strictly symmetric as was reported, e.g., for the crystal structure of the B9 (Ser→Glu) mutant insulin dimer62 which was not symmetric.

In Figure 3B and D two H-bond distances between the side chains of residues HisB10 and GluB13 are reported for SI and WI, respectively. For GlyB24-SI only a transiently formed intramonomer hydrogen bond is found, see Figure 3B. Contrary to that the two side chains HisB10 and GluB13 form an intramonomer H-bond which makes the donor and acceptor atoms of the two side chains unavailable for dimerization contacts, see Figure 3D. This, in turn, reduces the stability as determined in the current protocol (MM-GBSA) and partially explains the difference of 20 kcal/mol between the SI and WI-dimer for the GlyB24 mutant. In addition to the loss of ≈ 6 kcal/mol, the dimer stability is also reduced due to four β-sheet
Figure 3: Superimposed (backbone) B1-chains of two snapshots for SI and WI variants of the Gly\textsubscript{B24} insulin dimer analogue. The orange and green traces in (A) and (C) correspond to one monomer of the SI and WI dimer together with a CPK representation of the His\textsuperscript{B10} and Glu\textsuperscript{B13} residues. Panels (B) and (D) highlight the absence (B) and presence (D) of the hydrogen bond between His\textsuperscript{B10} and Glu\textsuperscript{B13}.

H-bonds which are almost absent in the WI dimer (average of 0.7 H-bonds) but exist most of the time for the SI (3.7 H-bonds), see Figure 4A.

**Hydrogen Contacts at the Interface**

In the insulin dimer the Phe\textsuperscript{B24}-Phe\textsuperscript{B25}-Tyr\textsuperscript{B26} segment of monomer I (chain B1) forms an antiparallel $\beta$-sheet with the adjacent Tyr\textsuperscript{B26}-Phe\textsuperscript{B25}-Phe\textsuperscript{B24} segment of monomer II (chain
B2, see Figure 4B). Inter-chain H-bonds are formed between Phe$^{B24}$(I) and Tyr$^{B26}$(II) and between Phe$^{B24}$(II) and Tyr$^{B26}$(I). Therefore, substitutions in this region influence dimer formation. Previous work$^{60}$ has reported that insulin dimerization is enthalpically controlled and the four inter-monomer H-bonds in the apolar environment are the prime driving force for insulin assembly. However, MD simulations$^4$ have shown that insulin dimerization primarily results from nonpolar interactions, in particular B24–B26 residues make the largest favorable contributions and the role of the H-bonds is to provide the necessary directionality of the interactions.

Figure 4: (A) The number of interfacial H-bonds between protein residues 24–26 of chains B1 and B2 and their probabilities for WT, Gly$^{B24}$-SI, Gly$^{B24}$-WI, Ala$^{B24}$ and D-Ala$^{B24}$ mutants. Average number of H-bonds are: 3.6 (WT), 3.7 (Gly$^{B24}$-SI), 0.7 (Gly$^{B24}$-WI), 3.2 (Ala$^{B24}$) and 2.2 (D-Ala$^{B24}$). Here, for analysis purposes, an H-bond is defined by a donor-acceptor (H-O) distance of ≤ 2.4 Å although several other complementary characterizations and criteria exist.$^{63}$ For these directional H-bonds at the dimerization interface the definition of an angle is not mandatory as the distinction between a “formed/established” and a “broken” H-bond for the purpose of the present analysis is straightforward, see Figure 6. Depending on this definition the persistence times of the dimerization contacts change somewhat but not the conclusions that are drawn from the analysis. (B) Backbone representation of the H-bonds $d_1$ to $d_4$ and the weaker CH–S contacts $\rho_1$ and $\rho_2$ along the dimerization interface, see also Figure 6.
Figure 4A reports the population of H-bonds between the two monomers involving residues 24–26 of chains B1 and B2. These involve (see Figure 4B) intermonomer contacts $\rho_1$ ($\text{CysB19}_\text{SG} \cdot \text{CA} \cdot \text{XB24}$) in monomer I, $\rho_2$ ($\text{CysB19}_\text{SG} \cdot \text{CA} \cdot \text{XB24}$) in monomer II, and intramonomer (dimerization) H-bonds $d_1$ ($\text{XB24}_\text{N} \cdot \text{O} \cdot \text{TyrB26}$), $d_2$ ($\text{XB24}_\text{O} \cdot \text{N} \cdot \text{TyrB26}$), $d_3$ ($\text{TyrB26}_\text{N} \cdot \text{O} \cdot \text{XB24}$) and $d_4$ ($\text{TyrB26}_\text{O} \cdot \text{N} \cdot \text{XB24}$), where X is Phe for WT or Ala and Gly for the mutants considered and where the first residue belongs to monomer I and the second corresponds to monomer II.

Probability distributions of H-bonds (see Figure 4A) for WT and Gly$^{\text{B24}}$-SI mutants are quite similar and show an average of 3.6 and 3.7 H-bonds, respectively, compared to only 0.7 H-bonds for the Gly$^{\text{B24}}$-WI mutant. The H-bond distribution for the Ala$^{\text{B24}}$ mutant (yellow) resembles that of the WT dimer with an average H-bond population of 3.2. For the D-Ala$^{\text{B24}}$ analogue at most three H-bonds are found. Finally, no H-bond is present for the des-Phe$^{\text{B25}}$ mutant indicating its inability to aggregate to a dimer.

The individual H-bonds at the dimerization interface can have different populations, see Table 3. For WT and Gly$^{\text{B24}}$-SI the four inter-monomer H-bonds are present for $\sim 90\%$ of the simulation time. Specifically, for the WT occupations of 98 $\%$ were found for the $\text{TyrB26}_\text{H} \cdot \text{O} \cdot \text{PheB24}$ contacts and somewhat reduced occupancies for the $\text{PheB24}_\text{H} \cdot \text{O} \cdot \text{TyrB26}$ contacts (84 $\%$ for the B1–B2 and 78 $\%$ for the B2–B1 contact), see Table 3. Except for the B2–B1 $\text{PheB24}_\text{H} \cdot \text{O} \cdot \text{TyrB26}$ contact (47 $\%$) these occupancies are in reasonable agreement with previous work. Given the symmetric nature of the interface such a low occupancy is unexpected and the present results appear to be more realistic. Furthermore, the simulation time in the current work is considerably longer and the size of the simulation box is also larger, hence certain differences are not unreasonable.

For the weakly interacting dimer the H-bond occupancies are considerably reduced to between 0 and 34 $\%$, see Table 3. These direct, inter-monomer H-bond occupancies correspond to the average number of H-bonds found in the H-bond distributions in Figure 4A. The
CH···S contacts (which are not labelled “H-bonds” here) are weak interactions and for model systems such as C₂H₂···SH₂ they were found to be stabilized by −1.34 kcal/mol at the MP2/aug-cc-pVQZ level of theory.⁴¹

**Role of Water at the Interface**

In search for a molecular explanation why the Gly⁵²⁴ mutant is prone to destabilize, the hydration environment of the intermonomer H-bonds at the dimerization interface was further analyzed. The stability of the insulin dimer interface would decrease if monomer-monomer contacts would be replaced by protein-water interactions. Hence, the interfacial contacts, in particular the four β-sheet H-bonds (see Figure 4B) conserved and/or replaced by water-mediation in various insulin dimer analogues are explored. For this, the H-bonds around position B24 were analyzed and the results are summarized in Table 3.

For the WT dimer the water-protein H-bonds are occupied by 10 % to 30 % whereas for the strongly interacting Gly mutant almost no H-bonds to the water are found. Conversely, the weakly interacting Gly-dimer has occupations ranging from 15 % to 68 % which correlates with the reduced number of direct protein-protein H-bonds. This suggests that water molecules along the dimerization interface can replace direct protein-protein contacts. In general, low occupancy of protein-protein contacts implies a high population of water-mediated contacts.

In order to further characterize the stability, inter- and intramonomer contacts and close encounters with solvent water molecules for the different insulin dimers, and to support the findings described so far, additional 40 ns MD simulations for the Ala⁵²⁴ and Gly⁵²⁴ mutants and the WT were carried out, see Figure 6. The average RMSD compared to the 4INS reference structures are 2.0 Å, 2.5 Å, and 2.8 Å for the WT Ala⁵²⁴, and Gly⁵²⁴ mutants,
Table 3: Monomer-monomer interfacial H-bonds involving residues 24 to 26 for WT, Gly\textsuperscript{B24-SI} and Gly\textsuperscript{B24-WI}, Ala\textsuperscript{B24} and D-Ala\textsuperscript{B24} mutants. The percentage of simulation time, during which the H-bonds were formed are reported for direct inter-monomer H-bonds, and water-protein H-bonds. Here, an H-bond is defined by a donor-acceptor (H-O/H\textsubscript{w}-O/H-O\textsubscript{w}) distance $\leq$ 2.4 Å although many other possible definitions exist. For Gly\textsuperscript{B24-WI} and D-Ala\textsuperscript{B24}, H-bond occupancy in parenthesis corresponds to structures with two different water molecules.

| Chain | Donor      | Acceptor   | Inter-monomer H-bond occupancy/\% | Water-protein H-bond occupancy/\% |
|-------|------------|------------|----------------------------------|-----------------------------------|
| WT    |            |            |                                  |                                   |
| monI  | Phe\textsuperscript{B24}.H | monII | Tyr\textsuperscript{B26}.O | 84 | 14 |
| monII | Tyr\textsuperscript{B26}.H | monI  | Phe\textsuperscript{B24}.O | 98 | 10 |
| monII | Phe\textsuperscript{B24}.H | monI  | Tyr\textsuperscript{B26}.O | 78 | 28 |
| monI  | Tyr\textsuperscript{B26}.H | monII | Phe\textsuperscript{B24}.O | 98 | 29 |
| GlyB24-SI |            |            |                                  |                                   |
| monI  | Gly\textsuperscript{B24}.H | monII | Tyr\textsuperscript{B26}.O | 81 | 4 |
| monII | Tyr\textsuperscript{B26}.H | monI  | Gly\textsuperscript{B24}.O | 96 | 6 |
| monII | Gly\textsuperscript{B24}.H | monI  | Tyr\textsuperscript{B26}.O | 95 | 0 |
| monI  | Tyr\textsuperscript{B26}.H | monII | Gly\textsuperscript{B24}.O | 95 | 0 |
| GlyB24-WI |            |            |                                  |                                   |
| monI  | Gly\textsuperscript{B24}.H | monII | Tyr\textsuperscript{B26}.O | 19 | 30 (28) |
| monII | Tyr\textsuperscript{B26}.H | monI  | Gly\textsuperscript{B24}.O | 34 | 68 (21) |
| monII | Gly\textsuperscript{B24}.H | monI  | Tyr\textsuperscript{B26}.O | 0  | 15 (82) |
| monI  | Tyr\textsuperscript{B26}.H | monII | Gly\textsuperscript{B24}.O | 13 | 56 (32) |
| AlaB24 |            |            |                                  |                                   |
| monI  | Ala\textsuperscript{B24}.H | monII | Tyr\textsuperscript{B26}.O | 92 | 7 |
| monII | Tyr\textsuperscript{B26}.H | monI  | Ala\textsuperscript{B24}.O | 99 | 27 |
| monII | Ala\textsuperscript{B24}.H | monI  | Tyr\textsuperscript{B26}.O | 38 | 86 |
| monI  | Tyr\textsuperscript{B26}.H | monII | Ala\textsuperscript{B24}.O | 87 | 7 |
| D-AlaB24 |            |            |                                  |                                   |
| monI  | D-Ala\textsuperscript{B24}.H | monII | Tyr\textsuperscript{B26}.O | 90 | 4 |
| monII | Tyr\textsuperscript{B26}.H | monI  | D-Ala\textsuperscript{B24}.O | 59 | 46 |
| monII | D-Ala\textsuperscript{B24}.H | monI  | Tyr\textsuperscript{B26}.O | 0  | 22 (75) |
| monI  | Tyr\textsuperscript{B26}.H | monII | D-Ala\textsuperscript{B24}.O | 73 | 31 (1) |

† The “non-bridging type II” motif (Figure 5d) counts one inter-monomer + one water-protein H-bond. “bridging type I” (Figure 5d) and “bridging type III” (Figure 5f) count two water-protein H-bonds. Because of this the total occupancy can be > 100 \%.
Figure 5: Water molecules interacting at the insulin dimer interface involving B24–B26 inter-monomer H-bond motifs. Only donor and acceptor atoms are highlighted in ball-and-stick representation. H-bonds are depicted by dashed lines (orange for intermonomer, black for water-protein contacts). (a) “non-bridging type I” one water-protein H-bond in the presence of two inter-monomer H-bonds, (b) “non-bridging type II” one water-protein H-bond in the presence of one inter-monomer H-bond, (c) “non-bridging type III” one water-protein H-bond in the absence of inter-monomer H-bonds, (d) and (e) “bridging type I” a single water-bridged H-bond in the absence of inter-monomer H-bonds, (f) “bridging type II” two water-bridged H-bonds in the absence of inter-monomer H-bonds. An H-bond is defined by a donor-acceptor (H-O) distance of 2.4 Å.

respectively (see also Figure S10). For these simulations the H-bonds along the dimerization interface and additional geometrical determinants including the water-occupancy were analyzed. This yielded several types of water-protein H-bonds, shown and described in Figures 5a to f. “Bridging” water-protein H-bonds water have significant interactions with both monomers and thus provide stabilization of the dimer. However, in an MM-GBSA analysis where explicit water molecules do not appear, their influence on the stability is not included. The water-protein H-bond occupancy is higher in Gly^{B24}-WI mutant than in WT and Gly^{B24}-SI (see Table 3). Throughout the MD simulations of Gly^{B24}-WI, consistently one or two water molecules are involved in H-bonds between Gly^{B24} and Tyr^{B26}.
Figure 6: CA-SG ($\rho$) and intermonomer N-O distances ($d$, both in Å) as a function of time for A) the glycine mutant, B) the alanine mutant and C) the wildtype. Rows 1, 2, 4 and 5 report the intermonomer hydrogen bonds between residues 24 and Tyr$^{B26}$ of the two B-chains. Rows 3 and 6 show the Cys$^{B19}$SG–CA$^{XB24}$ separations for monomers I and II, respectively. Symbols mark times when one (red cross) or two (green plus) water oxygen atoms are within 3.5 Å of both the N- and O-atoms of the corresponding H-bond. Dashed lines are drawn to guide the eye. Generally whenever $\rho_1$ (or $\rho_2$) is below 4.1 Å at least one of the intermonomer H-bonds is broken; the magenta dashed line marks this 4.1 Å threshold. The horizontal blue dashed line denotes the minimum distance of $\rho_1$ that was sampled in the Gly simulation. Vertical dashed lines indicate key points in the simulations. For feature I $\rho_1$ is at a minimum (0.5 Å below the 4.1 Å threshold) in the Gly simulation and intermonomer bond breakage is clearly observed for all four intermonomer bonds. For feature II - where $\rho_1$ meets the 4.1 Å threshold for the Ala$^{B24}$ mutant and intermonomer bond breakage is observed.
Figure 6 reports several important distance time series to further characterize the dimerization interface. They include inter- and intramonomer distances $\rho_1$, $\rho_2$ and $d_1$ to $d_4$, see Figure 4B. In Figure 6 the symbols mark times when one (red cross) or two (green plus) water molecules are within 3.5 Å of both atoms of the intermonomer N-O pair. Individual water molecules have also been found to be relevant in simulations of the insulin monomer in water, in HIV-I protease or in controlling rebinding of NO to microperoxidase.

For Gly<sub>B24</sub> frequent and spontaneous insertion of one or even two water molecules to replace the direct protein-protein NH–O bond is found. This is sometimes but not exclusively accompanied by losing the NH–O contact and differs considerably for the Ala mutant and the WT protein for which the NH–O Hydrogen bonds are intact for most of the simulation and water molecules are considerably less frequently close to the hydrogen bond. Again, the Gly<sub>B24</sub> mutant clearly displays a two-state behaviour as already found above: with the NH–O hydrogen bond intact (as is the case for WT and most of the Ala mutant simulations) which corresponds to the SI Gly<sub>B24</sub> dimer, or with the H-bond broken and typically replaced by a solvent water which is the situation in the WI dimer.

This analysis also provides molecular-level insight into the origins of intermolecular H-bond breaking and water-insertion. The Phe→(Ala,Gly) mutations replace a bulky phenylalanine residue at position 24 by considerably smaller CH<sub>3</sub> (Ala) or H (Gly) moieties (see Figure 7). Hence, the mutations lead to increased conformational freedom of the side chains. This in turn affects the distance(s) $\rho_1$ (and/or $\rho_2$), see Figure 6. As a reference, for the WT protein this distance ranges from 4.5 Å to 5.5 Å but decreases to below 4.0 Å for the Gly mutant. For the Ala mutant the separation is closer to the situation in the WT protein. The data in Figure 6 suggests that there are two conditions which lead to breaking of the protein-protein hydrogen bond: (1) the distance $\text{CysB19SG–CA}^{X24}$ in at least one monomer, i.e. $\rho_1$ or $\rho_2$, must be below a threshold of 4.1 Å; and (2) water molecules must be within the vicinity of the intermonomer N-O pair. Furthermore, the data suggests that the smaller $\rho_1$ or $\rho_2$, the
Figure 7: Schematic illustration for the relative sizes of residue 24 in wildtype insulin (Phe\textsuperscript{B24}) and the two mutants (Ala\textsuperscript{B24} and Gly\textsuperscript{B24}). Cys\textsuperscript{B19} is also highlighted to show the variation in steric hinderance between the three cases and to indicate the impact on the key distance, $\rho_1$ or $\rho_2$.

larger the distance between the intermonomer N-O pair (e.g. ‘I’ compared to ‘II’ in Figure 6). Note also that the behavior of the protein-protein H-bonds were closely connected and the breakage of one was often correlated with the breakage of the others.

The present analysis reveals that in Gly\textsuperscript{B24}-WI extensive water-mediated H-bonds entirely replace the $\beta$-sheet H-bonds. This is also consistent with the DCCM maps (see Figure S9) where the inter monomer H-bonds are absent for the WI dimer. As a result, the Gly\textsuperscript{B24}-WI dimer has fewer stabilizing inter-monomer interactions resulting in a considerably lower $\Delta G_{bind}$ compared to WT and Gly\textsuperscript{B24}-SI within the present MM-GBSA approach. Ala\textsuperscript{B24} has inter-monomer H-bond occupancies similar to WT, whereas D-Ala\textsuperscript{B24} partly differs from WT, in particular one H-bond (D-Ala\textsuperscript{B24} (monomer II):H···Tyr\textsuperscript{B26} (monomer I):O) is missing throughout the simulation (see Table 3). Consequently, D-Ala\textsuperscript{B24} was found to have a larger number of water-protein H-bonds than Ala\textsuperscript{B24}. The inter-molecular hydrogen bonds, one
of the main factors driving WT insulin dimerization, are absent in the des-Phe$^{B25}$ mutant, thereby preventing its dimerization.

Conclusions

Residue PheB24 plays an essential role in insulin folding, assembly, stability, receptor binding and hormonal signalling. In the present work changes in protein dimer stability resulting from mutations at position B24 were studied using (1) MD simulations with explicit water and (2) free energy simulations using MM-GBSA and TI. To the best of our knowledge, the present work is the first systematic computational study of the relative stabilities and dynamics of dimeric insulin analogues at position B24. The simulations support and extend earlier findings of the importance of residue B24 for the structural integrity of the hormone.

MM-GBSA and TI provide reliable information about the stabilisation of insulin dimers (WT and mutants). Compared to the experimentally determined stabilisation of $-7.2 \pm 0.8$ kcal/mol for the WT, TI finds $-8.4 \pm 0.2$ kcal/mol which is in good quantitative agreement. MM-GBSA is useful for qualitative and comparative purposes but not for quantitative studies. On the other hand, the relative stability changes from TI and MM-GBSA agree quite favourably and suggest that MM-GBSA is useful for ranking the stabilities of WT and mutant dimers.

Substitutions at position B24 of the insulin dimer-forming surface by Gly, Ala and D-Ala amino acid residues give dimeric insulin analogues with reduced dimer stability relative to the WT dimer. The des-PheB25 is exclusively monomeric, as was found by NMR experiments and serves as an additional validation in the present work. The presence of a WI and SI variant of the Gly$^{B24}$ mutant originates from H-bonds which are direct B1$\leftrightarrow$B2 protein-
protein contacts in the SI dimer but water-bridged in the WI dimer and from changes in the orientation of residue B13. This highlights that modifications at one site (B24) can have substantial functional effects. Bridging water molecules replacing H-bonding interactions along dimerization (and oligomerization) interfaces is most likely an ubiquitous feature and should probably be included in estimating dimerization energies which is, however, not routinely done.

Simulations can thus complement existing experiments and provide molecular-level insight for observed differences between chemically related systems. Also, they provide important information for situations in which experiments are technically difficult or impossible as they may be the only direct method to probe the structure, energetics and dynamics at atomic resolution.

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Supporting Information Available

Binding free energy decomposition data, per residue energetic contribution data and dynamical cross correlation maps. This material is available free of charge via the Internet at http://pubs.acs.org/.
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