Comparative Simulation Studies of Native and Single-Site Mutant Human Beta-Defensin-1 Peptides

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ABSTRACT

Human defensins play important roles in a broad range of biological functions, such as defense and immunity. Yet, little is known about their molecular properties, i.e. secondary structure stability, structural variability, important side chain interactions, surface charge distribution, and resistance to thermal fluctuations, and how these properties are related to their functions. To assess these factors we studied the native human β-defensin-1 monomer and dimer as well as several single-site mutants using Molecular Dynamics simulations. The results showed that disulfide bonds are important determinants in maintaining the defensins’ structural integrity, as no structural transitions were observed at 300 K and only minor structural unfolding was detected upon heating to 500 K. The α-helix was less thermally stable than the core β-sheet structure held together by hydrogen bonds and hydrophobic interactions. The monomer α-helix stability was directly correlated, whereas the end-to-end distance was inversely correlated to the experimentally measured β-defensin-1 chemotactic activity, in the order: Mutant 2 (Gln24Glu) > Mutant 3 (Lys31Ala) = Wild Type > Mutant 1 (Asn4Ala). The structural stability of the β-defensin-1 dimer species exhibited an inverse correlation to their chemotactic activity. In dimers formed by Mutants 2 and 3, we observed sliding of one monomer upon the surface of the other in the absence of unbinding. This dynamic sliding feature may enhance the molecular oligomerization of β-defensin-1 peptides contributing to their antibacterial activity. It could also help these peptides orient correctly in the CC Chemokine Receptor 6 binding site, thereby initiating their chemotactic activity. In agreement with this notion, the remarkable sliding behavior was observed only for the mutants with the highest chemotactic activity.
INTRODUCTION

Human defensins are a large family of related structured polypeptides comprised of members of two large classes of α-defensins (1,2) and β-defensins (3–6). The first member of the human β-defensins, HBD-1, was isolated from human plasma where it occurs at nanomolar concentration (4). It is expressed in skin keratinocytes and various epithelial cells (7–13). Human β-defensins exhibit a number of physiological functions (14), including: anti-HIV activity (15), innate and adaptive immunity (16), and chemotactic activity for macrophages and monocytes (17–19). There is a body of evidence that human β-defensins also exert chemotactic effects on T-cells and immature dendritic cells, mediated via the CC chemokine receptor 6 (CCR6) similar to that of the larger chemokine selective CCR6 agonist - macrophage inhibitory protein-3α (MIP-3α) (20–23). Human β-defensins also have direct antimicrobial effects (24–26). For example, HBD-1 and some of its mutants are active against E. coli when tested in vitro (27). It has been hypothesized that defensins exert their antibacterial effect due to their positively charged surface (19, 28–30), which enables them to initially attach to the bacterial membrane (31). In this picture, the hydrophobic part of the molecule inserts into the lipid bilayer, leading to destabilization and disruption of the cell membrane, formation of multiple pores (32), and leakage of the cell contents (33).

The basic β-defensin structure is composed of an N-terminal α-helix and three antiparallel β-strands, the latter held together by three intra-molecular disulfide bonds. The most important conserved motif is the GXC in the second β-strand, where G is Gly, C is the fourth Cys in the peptide, and X is any amino acid (34). In the case of HBD-1, this motif is represented by Gly25-Thr26-Cys27. The proline residue in the β1-β2 turn of β-defensins, most of the Gly residues (35), and the β-bulge at the conserved Gly found in the β2-strand, are also conserved (34). Based upon X-ray studies, the reported HBD-1 structure is comprised of 36 amino acids having six cysteines forming three conserved disulfide bonds, namely Cys5/Cys34, Cys12/Cys27, and Cys17/Cys35 (34). HBD-1 is mostly reported to be a monomer, but a tetrameric structure of HBD-1, has also been resolved (34). The X-ray structures of single-site mutants of HBD-1 have been reported as well (27). NMR diffusion measurements of the radius of hydration (36) and Nuclear Overhauser effect experiments showed that HBD-1 exists in the monomeric form in aqueous solution (37).

Despite the established physiological importance of the human β-defensins, only a few computer-based modeling studies for human β-defensins have been reported to date (38–40). In one paper (40), Molecular Dynamics (MD) simulation study using GROMACS package (41–43) has been carried out to study HBD-1 conformational behavior with and without disulfide bridges. The authors reported the unfolding behavior of HBD-1 and the distances between sulfur atoms during the course of the MD simulations (40). Although HBDs are good candidates for computational studies due to their small size, the literature is lacking MD simulation studies for HBDs, and there are no simulation studies addressing HBDs chemotactic properties. Only one study (27) reported single-site mutants and compared their biological activities to the wild type (WT). Here, we carried out comprehensive simulation studies of the structural and dynamic properties of the native HBD-1 peptide in its monomeric and dimeric forms as well as three single-site mutants of HBD-1 using all-atom MD simulations. We have
chosen these specific mutants because they exhibit important functional differences. They vary in their level of chemotactic and antibacterial activities compared to the WT.

Here, we report the main results of our studies, in which we resolved and contrasted the structural and dynamic properties of the native HBD-1 peptide with those of the single-site mutants. For each system, we analyzed the secondary structure dynamics using secondary structure propensity and Ramachandran plots. We examined dynamics of binary contacts between side chains utilizing contact maps and maps of stable contacts, molecular flexibility using root-mean-square deviations (RMSD) and root-mean-square fluctuations (RMSF), surface charge distributions employing solvent accessible surface areas (SASA), and the end-to-end distances. In this comparison, we relate the observed structural and dynamic changes to the altered side chains of the mutants, and discuss their implications for the experimentally measured functional differences in chemotactic activity reported for these mutants (27). The most significant findings are the following. We found correlations between the measured chemotactic activity and some of the molecular properties, such as the α-helix stability, end-to-end distance, and stability of the dimeric form (Table 1). The dimers formed by mutants Mut2 (Gln24Glu) and Mut3 (Lys31Ala) exhibited sliding and rolling behavior of the two monomer chains over each other. The basic residues were found to be exposed to solvent, which points to their importance in antibacterial and chemotactic activity of HBD-1.

METHODS

**Structural Models:** The molecular models have been constructed using the X-ray structures from the Protein Data Bank (PDB) (44) (see Table SM1 in Supplementary Material (SM)). The initial structures for the HBD-1 mutants in their dimeric form were obtained by substituting the appropriate residue in the HBD-1 wild type (WT) dimer structure using Visualization Molecular Dynamics (VMD) mutator (45). These substitutions are Asn4Ala (Mut1), Gln24Glu (Mut2), and Lys31Ala (Mut3). The rationale for choosing these three mutants was that their chemotactic activity was significantly different when compared to the WT; Mut1 (Mut2) showed lower (higher) chemotactic activity, whereas Mut3 was almost as active as the WT (27) (see Table 1). The mutation sites were also diverse; point mutations were in the α-helix (Mut1), in the β2-strand (Mut2), and in the β2-β3 turn (Mut3) (27).

**Computer Simulations:** *Equilibrium MD simulations:* All-atom MD simulations in explicit water were performed using the NAMD package (46) with CHARMM22 force field (47) and TIP3 model for water molecules (48). For each peptide, a water box has been created so that in each direction x, y, and z the distance between the molecule and the box boundaries was ≥12 Å. For example, the box size for the WT monomer was 48.5 Å × 50.4 Å × 45.8 Å. All the systems have been neutralized using chloride counter-ions. The density of each system was maintained at 1 g/cm³ to mimic ambient water conditions. After initial minimization, all the systems were heated to $T = 300$ K at constant volume and equilibrated at $T = 300$ K for 4 ns. Long 30 ns equilibrium simulation runs have been performed for each system, using the NPT-ensemble and a 1 fs integration step. The van-der-Waals (VdW) interactions were gradually turned off at a
distance between 10 Å and 12 Å. The list of non-bonded interactions was truncated at 13.5 Å. The short-range non-bonded interactions and the long-range electrostatic forces were computed for every step. The Particle Mesh Ewald (PME) summation was employed to describe the long-range electrostatics (49, 50). Heating Simulations: We carried out 10 ns heating simulations, in which the obtained structures equilibrated at \( T = 300 \) K have been heated from \( T_{in} = 300 \) K to \( T_{fin} = 500 \) K. GROMACS Simulations: Additional 25 ns simulation runs for the native WT monomer (Table SI1) have been performed using the GROMACS package (41–43) (GROMOS96 45a3 (51) and OPLS (52) force fields) to compare our findings with the results reported in Ref. (40) and to exclude possible correlations between the results obtained and the force fields used. In these runs, we employed the same simulation setup used in Ref. (40), albeit with a larger water box (49.4 Å × 51.3 Å × 46.7 Å). In simulations using GROMOS96 and OPLS force fields, we also utilized PME method (49, 50). A summarized description of all MD simulation runs is given in Table SM1 in the Supplementary Material (SM).

Data Analyses: We analyzed the secondary structure dynamics using time evolution of secondary structure propensity, i.e., propensities (probabilities) of amino acid residues to form different secondary structure motifs (e.g. α-helices, β-strands, etc.), and the Ramachandran plots (distribution of dihedral angles \( \Phi \) and \( \Psi \)). We used the structure overlap function \( \langle \chi \rangle \) (53), \( 0 \leq \chi(t) \leq 1 \), to estimate the extent of similarity of the transient conformation observed at time \( t \) to the reference structure (initial state) at time \( t = 0 \). When \( \chi = 0 \), the two structures are completely dissimilar, and when \( \chi = 1 \) they are exactly the same. The structure overlap function is given by \( \chi(t) = \left[ 2/N(N-1) \right] \sum_{i,j=1}^{N} \theta \left[ \delta r_{ij}^0 - \left| r_{ij}^t - r_{ij}^0 \right| \right] \), where \( N \) is the number of atoms in the system, \( \theta \) is the Heaviside step-function (\( \theta(x) = 1 \) when \( x > 0 \) and zero otherwise), \( \delta = 0.2 \) is the tolerance to the thermal fluctuations (at \( T = 300 \) K). Here, \( r_{ij}^t \) is the distance between the \( i \)-th and \( j \)-th atoms at time \( t \), and \( r_{ij}^0 \) is the same distance in the reference state \( (t = 0) \). The contact maps of interacting amino acids represented the trajectory averaged picture of binary contacts between side chains (both inter- and/or intra-monomer interactions). A contact between \( C_\alpha \)-atoms of a pair of residues, for which the \( C_\alpha-C_\alpha \) distance is < 6.5 Å cut-off, is counted as a contact. Dynamic maps of stable contacts is a powerful tool to follow the pattern of formation and rupture of stable binary contacts between coupled residues (both native and non-native) in a dimer. This tool has been used to study protein-protein interactions in the HBD-1 dimers. All the stable contacts, which persisted for at least 1000 ps in a simulation run, have been displayed as a function of time. The contact maps and plots of the total number of inter-monomer contacts have been used to analyze the transient structures and kinetics of protein-protein interactions. The root-mean-square deviations (RMSD), which quantify the average displacements of \( C_\alpha \)-particle relative to their positions in the reference state, were used to probe the global peptide fluctuations. The root-mean-square fluctuations (RMSF) allow one to access local conformational fluctuations. We used a “running average” structure as a reference state in each 100 ps time interval to eliminate rotational and translational contributions (54, 55). The solvent accessible surface area (SASA) per residue provides insights into solvent accessibility of the side chains on the surface of the molecule (56). This measure has been used to describe potential surface interactions and receptor binding properties. The plots of the
total SASA versus time have been used to measure these properties of the peptide molecule as a whole. We monitored the end-to-end distance as a function of time as a suitable measure of the global conformational transitions in the HBD-1 peptide and its derivatives, such as unfolding, thermal denaturation, and dissociation (in HBD-1 dimers).

RESULTS

Structural Features: The compact structure of the 36 amino acid native HBD-1 wild type (WT) (Fig. 1a) extracted from the dimer (Fig. 1b) is stabilized by the β-sheet and three conserved disulfide bonds between Cys residues 5 and 34, 12 and 27, and 17 and 35 (34). In the native WT peptide, residues His2-Ser7 form an α-helix, while residues Gln11-Leu13, Ile23-Cys27, and Ala32-Cys35 form three β-strands connected by two turns localized to residues Pro18-Thr21 and Tyr28-Lys31 (34). The native HBD-1 dimer structure, characterized by an asymmetric arrangement of chains A and B (Fig. 1b), is stabilized by the inter-monomer interactions among amino acids in the β2-strand of chain A and in the β2-β3 turn of chain B. All the conserved residues and structure elements (34, 35) discussed earlier were retained in all defensins we simulated in this study. Comparing the WT monomer equilibrated for 4 ns (Fig. 1c) to its initial structure (Fig. 1a), we found that the α-helix and β-sheet showed some structural differences in terms of their orientation with respect to the bulk of the molecule (Fig. 1). Two surface representations of different orientations of the equilibrated WT monomer (Fig. 1d) showed extensive patches of residues with similar charge and polarity. Specifically, we observed basic and hydrophobic patches that are known to be crucial for binding and inserting into bacterial membranes (31–33). These findings correlate well with the known antibacterial activity of defensins (19, 28–30), and possibly their role in chemotaxis (27). The results of heating simulations for the native peptide WT and mutants Mut1, Mut2 and Mut3 showed that, for all the systems, the number of hydrogen bonds decreased and the radius of gyration increased with increasing temperature from $T_{in} = 300$ K to $T_{fin} = 500$ K (data not shown). This should be expected as proteins tend to expand with increasing temperature while losing some or all of their hydrogen bonds that stabilize their native fold. However, this effect was minimal due to the constraints posed by the three disulfide bonds.

Secondary Structure Dynamics: We examined the stability of the secondary structure elements by analyzing the time evolution of secondary structure propensity of amino acid residues and the Ramachandran plots. The β-sheet structure formed from the three anti-parallel β-strands showed essentially similar behavior in the WT, Mut1, Mut2, and Mut3. This is hardly surprising since the three conserved disulfide bond topology exists in all the peptides, giving rise to a highly stable β-sheet structure. The time evolution of secondary structure propensity for the α-helical region in WT, Mut1, Mut2, and Mut3 showed some minor differences (Fig. 2). No pronounced distortion of the secondary structure was observed in equilibrium MD simulation runs, although His2 and/or Tyr3 in WT and α-helix in Mut1 were found to be partially unstructured (Fig. 2). All simulation runs for all the peptides showed the presence of some π-helix character except for Mut2, which had the most stable α-helix. Mut1 had the least stable α-helix (Fig. 2), and the order of decreasing stability for the α-helix was Mut2 > Mut3 ≈ WT > Mut1. This
correlates well with the experimentally measured chemotactic activity of HBD-1: Mut2 (130 ± 8%) > Mut3 (109 ± 14%) = WT (100 ± 11%) > Mut1 (40 ± 8%) (27) (Table 1).

We compared the structural integrity of the native HBD-1 peptide and its mutants in the monomeric and dimeric forms (Fig. 3). Mut1 (Mut2) dimer was the most (least) stable in terms of secondary structure, thus, giving an inverse correlation with the reported HBD-1 chemotactic activity (27) (Table 1). This implies that HBD-1 monomers are more active CCR6 ligands than their dimeric counterparts, similar to the MIP-3α monomer that is reported to be more adept (than the dimer) at binding to the CCR6 receptor (57). The α-helix in Mut2 monomer was more stable than the same structural elements in both chains comprising the Mut2 dimer, showing less π-helical character (Fig. 3). The structurally asymmetric dimer chains A and B exhibited different dynamic behavior. For example, the dimer β1-strand of chain A was the least stable secondary structure element, while β3-strand of chain A was the most stable.

We carried out heating simulations for WT and the three mutants to probe the thermal stability of these peptides, and to test the strength of the disulfide bonds. As shown in Figure 2, at $T = 460 \text{ K}$ the α-helix in the WT molecule started to unravel from Asp1 through Cys5 (i.e. up to the first disulfide bond position), and remained unstructured up to $T = 500 \text{ K}$. Mut1 was relatively stable during heating; at $T = 500 \text{ K}$ residues 2-4 formed a 310-helix (Fig. 2). Hence, introducing an Asn4Ala-substitution (Mut1) in WT thermally stabilized the α-helix. Upon heating Mut1, transient β1-strand deformation was observed in the 430-450 K temperature range (data not shown). During heating of Mut2, no significant changes to the α-helix were observed (Fig. 2); it was only transiently deformed. In heating simulations for Mut3, at $T = 475 \text{ K}$ the α-helix formed a turn and at $T = 460 \text{ K}$ the β1-strand completely lost structure (Fig. 2).

The results of analysis of secondary structure propensities and Ramachandran plots for all HBD-1 monomers and dimers showed that no significant structural changes occurred. At higher temperature, residues 2-7 in WT and Mut3 showed slightly lower α-helix propensities and some loss of structure (see also Fig. 2). Ramachandran plots for Mut2 monomer, obtained from equilibrium and heating simulations, exhibited similar distributions of amino acids in the β-sheet region but different distribution for the α-helical region (Fig. 4a). As expected for Mut2 monomer, residues 3-7 showed lower α-helical character at elevated temperature (Fig. 4c) compared to room temperature (Fig. 4b).

**Tertiary Structure Analysis:** For WT and the three mutants, we compared the final structures obtained at the end of 30 ns equilibrium simulation runs with the corresponding initial structures using the structure overlap function $\chi$ (see section Methods). We found that the former and the latter were not very dissimilar. We ranked HBD-1 peptides in decreasing order of their $\chi$ value, averaged over three independent trajectories, as follows: Mut3 (0.96) > Mut2 (0.94) > WT (0.92) > Mut1 (0.9). This was only slightly different from the ranked chemotactic activity measured experimentally (27), Mut2 > Mut3 = WT > Mut1 (Table 1), indicating that a direct correlation exists between the extent of conformational fluctuations in HBD-1 monomers and their chemotactic activity. We also carried out a similar analysis for the dimers of the WT and the mutants and observed a greater variation in $\chi$ as compared to their monomeric forms. The highest $\chi$ was observed for the WT dimer (0.96) and Mut1 dimer (0.95), and the
lowest $\chi$ was observed for the Mut2 dimer (0.83). These results are opposite to those of the monomers and agree with the differences in secondary structure stability discussed above, where Mut1 (Mut2) dimer was the most (least) stable (Table 1).

We next analyzed transient and final structures obtained in the course of heating simulations. The WT showed a large decrease in $\chi$ (Fig. 5 solid black), which corresponds to partial unraveling of the $\alpha$-helix starting at $T = 460$ K (Fig. 2). The curve of $\chi$ for Mut1 showed a transient drop at $T = 430-450$ K due to deformation of the $\beta_1$-strand mentioned above. For Mut2 and Mut3, $\chi$ varied considerably and after reaching $T = 460$ K decreased sharply to $\chi \approx 0.8$ and 0.7, respectively (Fig. 5). For Mut3, the decrease in $\chi$ at $T = 460$ K was due to the $\beta_1$-strand becoming more unstructured, which was concomitant with unraveling of the $\alpha$-helix discussed previously (see also Fig. 2). We used the final $\chi$ values observed in 10 ns heating simulations (Fig. 5) to rank structural stability of the HBD-1 monomers against the thermal perturbation. We obtained the following order: Mut1 (0.96) > WT (0.88) > Mut2 (0.8) > Mut3 (0.7) (Fig. 5). Using the average $\chi$ values resulted in the same order. These quantities are positively correlated with the experimental chemotactic activities reported for these molecules (27) (Table 1).

**Dynamics of Binary Contacts:** An examination of the dynamics of formation and rupture of intra- and inter-monomer contacts allowed us to gather residue level information to help us explore a temporal evolution of protein-protein interactions in the HBD-1 peptides. Indeed, contact maps, extracted from equilibrium and heating simulations, clearly illustrate the specific interacting residue sites and their stability at equilibrium ($T = 300$ K) and at elevated temperature (from $T_{in} = 300$ K to $T_{fin} = 500$ K). We analyzed the dynamics of intra-monomer contacts for monomers of the WT and the mutants, and the dynamics of inter-monomer contacts for dimers.

For the Mut1 (Asn4Ala) monomer, Ala4 formed stronger, longer lived contacts than Asn4 of the WT monomer. The polar contact of Asn4 with residue Gln24 was present in the WT monomer but was absent in Mut2 (Gln24Glu). The contacts formed in Mut3 (Lys31Ala) by Ala31 with both Tyr14 and Ser15 were weaker than those formed by Lys31 in the WT monomer. The weaker interactions are due to the shorter methyl group side chain of Ala compared to the three methylene bridge in the Lys side chain. All the contacts formed by the basic residues were monitored due to their importance in the antibacterial and chemotactic activities (19, 27–30). In all monomer equilibrium simulations, we observed Lys33 and Lys36 establishing binary contacts, while Lys22, Arg29, and Lys31 formed no contacts. No significant differences were detected in the distributions of contacts obtained from equilibrium simulations and heating simulations. However, in heating simulations a few minor changes were detected in the $\alpha$-helix in WT and Mut3 (Fig. 2; see, also, Fig. 6), and in the $\beta_2$-strand of Mut2 (Fig. 3a). In Mut3, the contacts between residues in the $\alpha$-helix (residues 2-7) and $\beta_2$-strand (residues 23-27) became progressively weaker as shown in the contact maps and structural snapshots for $T = 320$ K, 400 K, and 480 K (Fig. 6). At 320 K, only Tyr3 and Ser7 in the $\alpha$-helix are lacking contacts; the other 4 residues all form strong contacts. At $T = 400$ K, the $\alpha$-helix has moved away from the $\beta_2$-strand (Fig. 6; see, also, Fig. 2); Asn4, Cys5, and Val6 are still forming strong contacts. At 480 K, the $\alpha$-helix has unfolded (Fig. 2) and has moved away from the $\beta_2$-strand resulting in significantly weaker contacts (Fig. 6). As expected, the exception to this trend was observed for Cys5 (disulfide bond).
Equilibrium simulations for the Mut1 dimer revealed stable contacts in the \( \alpha \)-helical region, especially at Asn4 position (data not shown). These contacts were not detected in the WT dimer, which could be due to the Asn4Ala mutation creating new contacts at position 4. This might be one of the reasons why the Mut1 dimer was more stable than the WT dimer. The curve of \( \chi \) for the Mut2 dimer showed a decrease in \( \chi \) values at 30 ns (~0.60); the decreases in \( \chi \) values at ~6, 18, 22 and 24 ns were accompanied by decreases in the number of stable contacts (Fig. 7a and 7b). At the end of the simulation run, the Mut2 dimer had different coupled residues compared to the initial structure (Fig. 7b), strongly suggesting that the two chains had been sliding over each other. For the Mut2 dimer, we observed formation of the stable contacts Glu24A/Thr26B, Glu24A/Arg29B, Glu24A/Tyr3B and Glu24A/Asn4B, which were not detected for the WT dimer. This was due to the different physicochemical properties of the Gln and Glu residues. The Mut3 mutation (Lys31Ala) influenced stable contacts in the Mut3 dimer, where residue Ala31 in chain B formed weaker stable contacts than residue Lys31 in chain B in the other HBD-1 dimers. In all runs for the Mut3 dimer, we observed formation of contacts Thr26A/Ala31B, Ile23A/Ala31B and Thr26A/Ala31B (Fig. 7d). For Mut3, some transition regions in the time-dependent map of stable contacts are obvious. However, we see that some contacts might suddenly appear or disappear, or might exist only for a short period of time (Fig. 7d). The two successive snapshots (Fig. 7c) correspond to the structures observed at 15 ns and 18 ns on either side of the abrupt 17 ns drop in \( \chi \) to ~0.65. In these two representations, the two monomer chains are clearly in different relative orientations. This conformational transition is reflected in the changes in the map of stable contacts (Fig. 7d), which can be interpreted in terms of the monomers sliding to their new positions without any attempt to dissociate.

The dynamics of formation-rupture of non-covalent interactions can be monitored using the total number of inter-monomer contacts \( Q(t) \) between coupled residues as “a reaction coordinate” (Fig. 8). We see from independent diverging trajectories of \( Q(t) \) for the Mut2 dimer that formation-rupture of binding contacts is a stochastic process. For example, in the second run at \( T = 300 \) K for Mut2 dimer, we observed a reversible dissociation transition at ~22 ns followed by a redimerization transition at ~25 ns. In the third trajectory, the monomers in the Mut2 dimer formed more contacts compared to their native state at the end of a 30 ns run (Fig. 8 solid green). The WT dimer was stable maintaining ~15-20 contacts in the course of a 30 ns run. The Mut2 dimer was the least stable in terms of the total number of inter-monomer contacts. We also examined the \( Q \) versus \( t \) pattern for the WT and all mutant dimers, and arranged them in the following order of their decreasing stability: Mut1 > Mut3 > WT > Mut2. We found that the higher dimer stability corresponded to the weaker measured chemotactic activity (27) (with a slight difference in WT and Mut3 ranking).

**Root-Mean-Square Fluctuations and Deviations:** The root-mean-square fluctuations (RMSF) for each \( i \)-th amino acid residue, \( RMSF = \sqrt{\frac{1}{t_{tot}} \sum_{j=1}^{t_{tot}} (x_i(t_j) - \bar{x}_i)^2} \), where \( x_i(t_j) \) and \( \bar{x}_i \) are the position for \( C_\alpha \) - atom \( i \) at time \( t_j \) and average value of the reference position of \( C_\alpha \) atom \( i \), respectively, and \( t_{tot} \) is the length of the simulation run (in ns), provides a measure of molecular fluctuations at the local scale. We calculated RMSF using the results of
equilibrium simulations at $T = 300$ K and the PDB structures for the monomers and dimers of the WT and the three mutants. For all monomeric systems, the RMSF profiles were similar and the RMSF values varied in the 1.8-2.3 Å range (data not shown). The highest RMSF values were observed for the residues forming the turns Ser8-Gly10, Tyr14-Lys22, and Tyr28-Lys31, the N-terminus (Tyr3), and the C-terminus (Lys36). By comparing the results of equilibrium and heating simulations, we found that at elevated temperatures the WT monomer had lower RMSF values (Fig. SM1a). This may be due to the hydrophobic effect, where solvent exclusion provides increasing stability with rising temperatures. This effect was observed for all the HBD-1 peptides except for Mut3 (Fig. SM1b).

The RMSF analysis for dimers revealed that the WT dimer was the most stable compared to Mut1, Mut2 or Mut3 (smallest RMSF; data not shown). Only the first ten residues of Mut1 dimer showed lower RMSF values than those of WT, including the mutation site Asn4Ala. This was due to the shorter side chain of Ala versus the Asn replacement, thereby showing a smaller fluctuation at that position. In the Mut2 dimer, the Gln24Glu mutation did not result in a larger RMSF value at that position compared to the WT dimer (data not shown), perhaps because both Glu and Gln have similar side chain size. The two chains in a dimer interact thereby affecting RMSF for each chain, unlike two standalone monomers. Accordingly, the RMSF profile for the dimer in question is not identical to the RMSF profile for the monomers, and the two dimer chains are also different especially at the dimerization site – the region of functional asymmetry – where fluctuations are expected to be minimal. This, together with the time evolution of secondary structure propensity (Fig. 3b), corroborate the theory of dimer asymmetry. In the equilibrium simulations of the WT dimer, the dimerization site was identified to be the β2-strand of chain A interacting with the residues in β2- and β3-strands, and β2-β3 turn in chain B (residues 23-27; see Fig. 9). This site had low fluctuations (Fig. 9b) and, consequently, higher interaction levels on the side chain contact map (Fig. 9a). Aside from the dimerization site, all dimers showed the minimum RMSF value at all Cys residues due to the disulfide bond constraints, and at the β-strands due to the formation of the β-sheet architecture.

We also analyzed the time-dependence of the root-mean-square deviations (RMSD) given by $RMSD(t) = \sqrt{\frac{\sum_{i=1}^{N} \left( x_i(t) - x_i(t_0) \right)^2}{N}}$, as a molecular-scale measure of dynamic transitions in the HBD-1 peptides, where $x_i(t)$ is the position of the $i$-th $C_\alpha$ atom at time $t$, in the average structure; $x_i(t_0)$ is the reference position for the same atom in the initial structure ($t = 0$), and $N$ is the total number of $C_\alpha$ atoms. In Figure 10, we display representative trajectories of RMSD for the dimers of WT, Mut2, and Mut3 (one of the three simulation runs for Mut2 and Mut3 are same as in Fig. 7). The RMSD profile for the WT dimer remained flat for the entire 30 ns simulation run at $T = 300$ K, indicating that the native WT dimer was stable (Fig. 10a). We see that the onset of the conformational transition in Mut2 and Mut3 coincides with sudden changes in their RMSD values (Fig. 7 and 10). For example, there is a noticeable increase in the RMSD value for Mut3 dimer after 16 ns (Fig. 10c dotted red), which is negatively correlated with the sharp drop in the $\chi$ value at ~17 ns (Fig. 7c), due to the redistribution of stable bonding contacts in the dimer (Fig. 7d). Here, the interaction pattern, in which the β2-
strand of chain A formed contacts with the β2-strand and β2-β3 turn of chain B, has changed to the new pattern, in which the β2-β3 turns in both chains interact together strongly through the Gly30A/Arg29B and Arg29A/Tyr28B contacts, and β2-strand couples to β2-β3 turns through the Thr26A/Arg29B contact (see also Fig. 7c for relevant structures). Hence, chain A appears to be sliding over chain B without any dissociation attempt, as discussed previously. Similarly, the RMSD value for the Mut2 dimer showed stepwise increases (Fig. 10b solid black), which correlates well with the previously described sudden drops in $\chi$ value at ~6, 18, 22 and 24 ns (Fig. 7a), and weakening of the dimerization site (see Fig. 7b).

**Solvent Accessible Surface Area:** We analyzed the solvent accessible surface area (SASA) per residue averaged over three equilibrium simulation runs for the monomer WT and mutants. This measure provides information about the accessibility of the side chains of hydrophilic amino acids exposed to water, and, hence, it can be used to single out and monitor the residues participating in associations with other biomolecules. Surprisingly, we observed roughly similar SASA values for all residues and for all monomers during equilibrium simulations at $T = 300$ K except for the mutation sites (data not shown). Specifically, substitution Asn4Ala (Mut1) and Lys31Ala (Mut3) showed lower while Gln24Glu substitution (Mut2) showed higher SASA values compared to the WT. We attributed this to the length of the side chain; the shorter the side chain, the less it would be exposed to the solvent. The results for SASA calculations for heating simulations (Fig. 11) revealed the same qualitative pattern. All the mutation sites were less solvent exposed than their WT counterparts. The relative similarity between the results of equilibrium and heating simulations imply that, although at high temperature the HBD-1 peptides loose some of their secondary structure, the overall tertiary structure, which determines the surface arrangement of polar residues, is preserved.

Equilibrium simulations at $T = 300$ K for dimers revealed that the SASA patterns for each monomer chain in dimeric and monomeric forms are very similar (data not shown). This stands in contrast to the results obtained from RMSF analysis (Fig. 9b). Some residues, e.g. Ala16, Ile19, Lys33, Lys36 in the case of WT dimer, showed noticeable differences between the two monomeric chains in the dimer, which also corroborates our previous findings for the secondary structure propensity (Fig. 3b) and RMSF (Fig. 9b) regarding asymmetrical dimers. We also analyzed the total (molecular) SASA as a function of time. A representative example, obtained from heating simulations for the WT monomer (Fig. SM2a) showed higher exposure to water at the end of the heating simulations ($\approx 4,200$ Å$^2$) compared to the equilibrium simulations ($\approx 4,000$ Å$^2$). This coincided with the $\alpha$-helix unraveling in the WT monomer at high temperature (Fig. 2), making the N-terminal more water exposed. A similar SASA pattern was observed for Mut3 monomer (data not shown). For the Mut1 monomer, equilibrium and heating simulations showed the same SASA values because, as we described previously, the molecule remained the most stable. During the heating simulation for the Mut2 monomer (Fig. SM2b green), the SASA value increased at $T = 470$ K due to the β2-strand deformations, and at $T = 475$ K due to the transiently deformed $\alpha$-helix (Fig. 2).

**End-to-end Distance:** The global transitions in proteins, such as unfolding, can be accessed by analyzing their end-to-end distance. We examined this quantity for the WT
and the mutants monomers using the results of equilibrium and heating simulations. At higher temperature, WT and Mut3 showed larger end-to-end distance values than their average end-to-end distance obtained from three equilibrium simulation runs at $T = 300$ K, mostly due to $\alpha$-helix unfolding at $T = 460$ K for WT and $T = 475$ K for Mut3 displayed in Fig. 2 (see results for WT in Fig. SM2c). By contrast, the end-to-end distance for Mut1 remained constant at $\sim 14$ Å all through the course of heating simulations (data not shown), which was due to the relative stability already noted for that monomer (Fig. 2). Upon heating, the end-to-end distance for Mut2 initially increased gradually to $\sim 17$ Å (Fig. SM2d green), following the changes in the secondary structure, i.e. partial deformations of the $\beta_2$-strand and $\alpha$-helix at around $T = 476$ K, and then dropped to $\sim 12$ Å due to restoration of the $\alpha$-helix (Fig. SM2d green; see, also, Fig. 2). We ranked the monomer WT and mutants in the order of their increasing end-to-end distances obtained at the end of heating simulations (at $T = 500$ K). Interestingly, the end-to-end distances, Mut2 (12 Å) < Mut3 (13 Å) $\approx$ WT (13.5 Å) < Mut1 (14.5 Å), were inversely correlated with the magnitude of the experimental chemotactic activities (27) (Table 1).

**DISCUSSION**

In this study, we examined the dynamics and stability of HBD-1 WT and three single-site mutants. This enabled us to provide new insights into the structural differences between these species in order to understand their biological functions as antibacterial and chemotactic agents. Understanding the molecular basis for the antibacterial structure-activity relationships of HBD-1 WT and mutants might help in designing and developing novel antibiotics, as well as help in preventing the development of bacterial resistance to current antibiotics. This initiative has already been taken by Polymedix to develop non-peptide antibiotics that resemble $\beta$-defensins (58). Understanding the molecular basis for HBD-1 WT and mutants’ chemotactic activity might also aid in discovering potential treatments for pathologies involving the immune system.

When overlaid with the WT monomer, the X-ray structures of all mutants studied had the same overall structure and secondary structures as the native WT; the only structural differences were in the turns (27). However, structural dynamic changes could result from the specific mutations that lead to the measured differences in biological activity. The investigators who created and reported the mutants’ structures (27) purposefully left unaltered the conserved residues such as the cysteines (disulfide positions), GXC motif (34), and glycine and proline residues (35), due to their critical roles in determining the defensins’ 3D structure and function. The $\beta$-bulge (34) was conserved among WT and all mutants, where its first residue (number 24) adopted $\alpha$-helical parameters, e.g. dihedral angles $\Phi$ and $\Psi$ in the Ramachandran plots as reported for classical $\beta$-bulges (59). Our Ramachandran plot showed that for residue 24 of all monomers, $\Phi$ ranged from -94 to -85 and $\Psi$ from -57 to -44, i.e. in the range for classical $\alpha$-helices (data not shown).

The dimers studied were all asymmetric, as reported for the PDB structure for the WT peptide (34) (Fig. 1b), where interaction occurs through the $\beta_2$-strand of chain A and the $\beta_2$-$\beta_3$ turn of chain B. This asymmetry is also reflected in the time evolution of the
dimer’s secondary structure propensity (Fig. 3b), side chain contact maps (Fig. 9a), RMSF (Fig. 9b), and SASA (data not shown). A result of this asymmetry is that all the dimers’ individual monomer chains behave differently when compared to the standalone monomers. This is because the two chains in a dimer interact with a different portion of their sequence and each monomer affects the other’s dynamics. An example of this is clearly seen when comparing the time evolution of secondary structure propensity for Mut2 monomer (Fig. 3a) to its dimer (Fig. 3b). Here, in order of α-helix stability, Mut2 monomer > chain A of Mut2 dimer > chain B of Mut2 dimer; Mut2 monomer shows less π-helix character within the α-helix region (Fig. 3).

The initial monomer and dimer structures together with those observed in equilibrium simulations at \( T = 300 \) K had their positively charged side chains surface exposed and grouped into separated patches of cationic and hydrophobic residues. This was clearly observed, e.g. in the SASA per residue plots and could be visually observed in VMD, looking similar to the WT equilibrated structure illustrated in Figure 1d. The positively charged groups on the surface of the molecule have been established to be an important feature for maintaining chemotactic activity (27). It is thought that the spatially separated cationic and hydrophobic regions allow defensins to insert into the bacterial membranes. The basic residue patches are thought to interact with the anionic phospholipids allowing an oligomeric defensin structure to rearrange, insert hydrophobically into the membrane and create a lethal hole in the bacterial membrane, causing antibacterial activity (35, 37). Comparing the equilibrated WT monomer (Fig. 1c) with its surface representation (Fig. 1d), the five basic residues (Arg29, and Lys22, 31, 33, and 36) are observed to be more concentrated towards the C-terminus. These residues are present at the β3-strand, and the end of β1-β2 and β2-β3 turns. On the other hand, the hydrophobic cluster is more concentrated at the β1-β2 turn. One hydrophobic residue is present per secondary structure element: Val6, Leu13, Ile23, and Ala32 in the α-helix, β1-, β2-, and β3-strands, respectively. These general features of 3D structure described for the WT monomer were also true for all the mutants used in our study due to their similar 3D structures (27).

The three conserved disulfide bonds play an important role in maintaining the tertiary structure motif of defensins. Any effect of the specific mutations on the disulfide bond dependent dynamic properties therefore is a legitimate concern. During all simulations, the three disulfide bond lengths showed minimum change at ambient and elevated temperatures. This is not surprising and highlights the crucial role of disulfide bonds in maintaining a correct constrained tertiary structure backbone. Disulfide bond based tertiary structure is thought to exist in order to provide defensins with resistance to proteolytic degradation in the environments where they are secreted and active (60). Despite the presence of the disulfide bonds, heating the molecules resulted in their swelling. This was reflected in the decrease in the number of intra-monomer hydrogen bond and an increase in the radius of gyration, as expected for heating simulations (61, 62). Whenever the α-helix lost structure, as in the case of WT and Mut3 monomers, this occurred towards the end of the trajectory and always in regions away from the disulfide bonds. For example, the WT α-helix was partially unfolded from Asp1 through Cys5 at \( T = 458 \) K (Fig. 2).

The single-site mutations were found to affect the molecular properties. During the equilibrium simulations of Mut1 (Asn4Ala) monomer, residue 4 had on average a
lower SASA than the WT, which is due to the shorter side chain of Ala compared to Asn. In Mut1, residue 4 also formed a strong intra-monomeric contact with Gln24, that was absent in the WT. This could be attributed to the difference in side chain polarity; Asn is polar while Ala is not. The Asn4Ala mutation also caused Mut1 dimer to form more stable contacts, as evidenced from equilibrium simulation at \( T = 300 \) K, compared to the WT, giving it a greater overall stability. During all equilibrium simulations for Mut1 dimer, the first ten residues exhibited lower RMSF than those of the WT. This is due to the shorter side chain of Ala compared to Asn causing less fluctuation. Heating the Mut1 (Asn4Ala) monomer from \( T_m = 300 \) K to \( T_{fin} = 500 \) K resulted in a more stable α-helix than that found in the WT (Fig. 2), suggesting that at higher temperature this mutation stabilized the α-helix. This stability is likely due to either the hydrophobic nature of the Ala substitution, or because Ala is known to overstabilize the helix (63).

In the case of equilibrium simulations for Mut2 (Gln24Glu) monomer, because Gln and Glu amino acids have almost the same chain length, there was no significant difference compared to the WT regarding their average RMSF or SASA at residue 24 throughout the simulation. Upon heating, the Gln24Glu mutation did not only affect Mut2 β2-strand structure stability (wherein residue 24 lies), but also affected the α-helix. This is because these two secondary structure elements interact with each other. All equilibrium simulations for Mut2 (Gln24Glu) dimer showed the Glu24A/Thr26B stable contact (Fig. 7b) that was absent in the WT dimer. Other new stable contacts, absent in all other HBD-1 dimers tested due to Gln24A mutation, included Glu24A/Tyr3B, Glu24A/Asn4B, and Glu24A/Arg29B. The Glu24A/Arg29B contact involved the negatively charged side chain of Glu interacting with the cationic Arg. The rest of the new contacts were established due to hydrogen bond formation involving the terminal carboxylate group of Glu or due to the resulting difference in contacts between the WT and Mut2 species.

Equilibrium simulations for Mut3 (Lys31Ala) monomer demonstrated lower SASA at the mutation site compared to the WT, due to the shorter side chain of Ala compared to Lys. Heating Mut3 resulted in distortions in both β1- and β3-strands, because they interact directly with each other in a manner similar to the Mut2 discussed earlier. The Mut3 contacts between Ala31 and both Tyr14 and Ser15 were weaker than those contacts formed by Lys31 in the case of the WT. This was due to the side chain of Ala being too short to maintain the contact during the simulation run. It could also be due to the polar nature of Tyr, Ser, and Lys, making these residues incapable of interacting with the non-polar Ala. Ala31 contacts in Mut3 dimer were also relatively weak compared to Lys31 in the other tested mutant dimers. Some of the Mut3 trajectories showed a Thr26A/Ala31B contact (Fig. 7d), while one had a transiently stable Ile23A/Ala31B contact and a strong Thr26A/Ala31B contact. This again could be attributed to the short Ala side chain that cannot reach more residues in the other chain.

Analysis of the time evolution of secondary structure propensity agreed well with the results from the structure overlap function, propensities, and Ramachandran plots, where the monomers showed almost equally stable α-helices throughout the course of their equilibrium simulations at \( T = 300 \) K (Fig. 2). Heating the molecules from \( T_m = 300 \) K to \( T_{fin} = 500 \) K caused some noticeable changes in the secondary structure (Fig. 2). The α-helix was unstructured at \( T = 460 \) K in the WT, at \( T = 500 \) K in the Mut1, transiently in the 476-488 K range in the Mut2, and at \( T = 475 \) K in the Mut3 (Fig. 2). On the other
hand, β-strands showed only minor changes compared to the extent of alterations in the α-helices. Specifically, the β1-strand was transiently unstructured between $T = 430-450$ K in Mut1, and at $T = 460$ K in Mut3, while the β2-strand was unstructured at $T = 470$ K in case of Mut2.

During heating simulations, the loss of secondary structure elements was reflected in various analysis methods. Temporal changes in the end-to-end distance were directly correlated with the secondary structure dynamic changes. When the molecules started to unravel, their two ends moved away from each other, thereby increasing the end-to-end distance, accompanied by an increase in SASA (see, e.g., Fig. SM2). The gradual loss of Mut3 monomer side chain contacts during heating from $T_{in}=300$ K to $T_{fin}=500$ K (Fig. 6) is also a result of the loss of secondary structure elements. Ramachandran and propensity plots (see Fig. 4) reflected secondary structure changes throughout the simulation runs. All these secondary structure changes also coincided with decreases in the structure overlap function (Fig. 5), and showed that the order of monomer stability was Mut1 > WT > Mut2 > Mut3. This order was only slightly different from the α-helix dynamics results (Fig. 2 right panel) having a decreasing order Mut2 > Mut1 > Mut3 > WT. This minor difference likely results from the fact that the $\chi$ results involved differences between the entire structure, involving all secondary structure elements’ dynamics, rather than just the α-helix dynamics. The heating simulation for Mut1 was the most stable trajectory, possessing an almost intact α-helix (only deformed at $T = 500$ K) and a transient β1-strand deformation between 430 K and 450 K. This was followed by WT that exhibited a unstructured α-helix at $T = 460$ K, followed by Mut2 having β2-strand and transient α-helix destructuring at $T = 470$ K, and $T = 475-490$ K, respectively. Then, Mut3 was the least stable due to its unstructured β1-strand and α-helix at $T = 460$ K and $T = 475$ K, respectively, that both persisted until the end of the trajectory.

In a previously reported MD simulation study (40), the authors observed an early α-helix unfolding with total loss of hydrogen bonds at $T = 300$ K. We did not observe α-helix unfolding until ~22 ns of the WT monomer equilibrium simulation at the same temperature (Fig. SM3a), and no total loss of hydrogen bonds was detected. Our MD simulations also suggest that different force fields used in simulations could affect the output results and thus should be carefully chosen. The outcome of using different force fields in our simulations suggested that the difference between the α- and π-helix structure is so minor that it could be neglected. This makes sense when one considers the minor difference between the dihedral angles observed for the two helix types.

Because we aimed at investigating HBD-1 antibacterial and chemotactic activities, all the analyzed terms were related to these biological functions whenever possible. All the contacts formed by the basic residues were monitored as we expected them to have minimal intra- and/or inter-molecular interactions. This feature is consistent with the low number of acidic residues found in the sequence capable of favorable charge interactions. Free patches of basic residues are likely a requirement for the defensins performing their antibacterial and chemotactic activities (19, 27–30). For all monomer equilibrium simulations at $T = 300$ K, only Lys33 and Lys36 exhibited cumulative side chain contacts throughout the simulation runs, while Lys22, Arg29, and Lys31 formed almost none. This would potentially make them free to bind to sites on the CCR6 receptor, allowing HBD-1 to exert its chemotactic effect, and also make these positive charges available to carry out the antibacterial effect, whose effectiveness depends on the
molecule's total surface positive charge (19, 28–30).

Increasing the molecule’s surface positive charge density via oligomerization is believed to increase the defensins’ antibacterial effect at the bacterial membrane (38). Although the WT has mostly been reported experimentally as a monomer (27, 34, 36), it was also crystallized as a dimer (34). The single-site mutants used in our study were only reported crystallographically as monomers (27). However, we were able to demonstrate the potential multiple monomer interaction sites for the Mut2 and Mut3 species. This hypothesis was based on the observed sliding behavior without any dimer dissociation. This sliding phenomenon could help promote the oligomerization of the tested mutants. One of the equilibrium simulation runs for Mut2 dimer showed a stepwise decrease in $\chi$ at 6 ns, 18 ns, 22 and 24 ns (Fig. 7a) indicating loss of similarity with the initial PDB structure. This was accompanied by a decrease in the total number of inter-monomer contacts (Fig. 8 solid black), and an increase in RMSD (Fig. 10b solid black). Its dynamic maps of stable contacts (Fig. 7b) revealed weaker contacts together with a variation in the interaction site - switching from interaction between chain A $\alpha$-helix with chain B $\beta$2-strand to a new interaction between chain A $\beta$2-strand and chain B $\beta$2-$\beta$3 turn. At 6 ns some contacts between $\alpha$-helix and $\beta$2-strand were lost, then at 22 ns the interaction site weakened where only Arg29B formed three stable contacts with Lys22-Ile23-Gln24 (Fig. 7b). The weakest Mut2 dimer trajectory (Fig. 8 dotted red) lost contacts at 6 ns, totally unbound at ~22 ns, then redimerized via the formation of a few contacts at ~25 ns. This change in interaction sites for the HBD-1 Mut2 and Mut3 dimer species (intermolecular sliding behavior) was manifested in the $\chi$ (Fig. 7a and 7c), dynamic maps of stable contacts (Fig. 7b and 7d), and in the inter-monomer contacts plot (Fig. 8).

This behavior suggests that HBD-1 molecules have multiple potential inter-monomer interacting sites in vivo, and this could be a biological strategy that evolution has devised to aid the molecules in performing their functions as antibacterial and chemotactic agents. Perhaps the sliding behavior enables HBD-1 to oligomerize at the bacterial membrane, increasing their overall surface positive charge density, thus enhancing their antibacterial effect, as hypothesized in an earlier modeling study (38). It could also facilitate HBD-1 binding to the CCR6 receptor site via sliding dynamics that ‘finds’ the correct dimer conformation in order to bind, thereby initiating chemotactic activity. If the active species initiating chemotaxis is a monomer, and dimer vs monomer as the active species is not a settled issue in the literature (57), then its ability to exhibit sliding behavior with CCR6 within the binding region may be the mechanism whereby receptor binding occurs. The observed sliding behavior of Mut2 and Mut3, but not WT and Mut1, could explain their higher chemotactic activity compared to WT and Mut1, where the reported measured chemotactic activities were as follows: Mut2 > Mut3 = WT > Mut1 (27) (Table 1). Longer trajectories need to be generated to see whether WT and Mut1 possess this sliding behavior; being more stable than Mut2 and Mut3 dimers does not mean that they lack this trait because this could be attributed to insufficient sampling time. If WT and Mut1 dimers were proved to lack this behavior, this could mean that not only the mutation possibly caused this sliding phenomenon, but also that the site of mutation is crucial for whether sliding occurs in the dimer. Mut1 has the mutation site at the $\alpha$-helix, while Mut2 has the mutation at the $\beta$2-strand and does not seem to be as stable as Mut3, whose mutation occurs at the end of the $\beta$2-$\beta$3 turn.

Comparing the HBD-1 monomers reported chemotactic activity (27) to the $\alpha$-
helix stability observed in equilibrium simulations at $T = 300$ K, we concluded that the chemotactic activity is directly proportional to their $\alpha$-helix stability (Table 1). The $\alpha$-helix has been reported to be crucial for binding to CCR6 and therefore important for chemotaxis (64, 65). This is undoubtedly the reason for the low chemotactic activity of Mut1, since it is mutated within the $\alpha$-helix leading to a destabilized structure. We also observed that the dimer stability was inversely proportional to the chemotactic activity (Table 1). In terms of the total number of inter-monomer contacts, Mut1 dimer was the most stable molecule followed by Mut3, then the WT, with Mut2 the least stable. This decrease in dimer stability was observed to correlate with a greater chemotactic activity with a slight difference between the WT and Mut3 ranking. Together, these results present an interesting model that implies HBD-1 monomers are more active as CCR6 ligands than the dimers in binding to CCR6, as is the case for MIP-3α (57).

The order of the calculated average $\chi$ values for monomer equilibrium simulations, Mut3 (0.96) > Mut2 (0.94) > WT (0.92) > Mut1 (0.9), was directly correlated to the experimental chemotactic activities with a slight difference in Mut2 and Mut3 rankings (Table 1). On the other hand, end-to-end distances for the same equilibrium simulations showed an ordered relationship opposite to that of the magnitude of the experimental chemotactic activities: Mut2 (12 Å) < Mut3 (13 Å) ≈ WT (13.5 Å) < Mut1 (14.5 Å) (Table 1). These data suggest that the more compact the structure, the better the surface exposure/orientation of the side chains needed to bind to the CCR6 receptor. This correlation also supports the theory that the monomer, rather than the dimer, could be the active in vivo binding ligand for CCR6.

CONCLUSION

In this study, we examined the dynamics and stability of HBD-1 WT and three single-site mutants (Mut1, Mut2 and Mut3) using all-atom MD simulations of their monomers and dimers and theoretical analysis. Our results shed some light on the underlying basis of the antibacterial and chemotactic properties of HBD-1 that is rarely studied in the literature. During all the equilibrium simulations at room temperature, the reported conserved structural features thought important for HBD-1 to perform their functions were maintained with minor or no changes. Among these are the GXC motif, glycine and proline residues, and the $\beta$-bulge. The results also confirm the conservation of disulfide bonds and their role in maintaining the defensins’ structural integrity where only minor length changes were observed even upon heating the molecules.

The single-site mutation types and positions affected the molecules’ side chain contacts and stability, and accordingly their biological functions. HBD-1 monomers’ $\alpha$-helix dynamic stability was directly related to their reported experimental chemotactic activity (27), whereas the dimers’ $\alpha$-helix dynamic stability was inversely related (Table 1; see, also, Fig. 2). There was an observed sliding behavior in the case of Mut2 and Mut3 that is thought to give them the higher chemotactic activity when compared to Mut1 and WT (Table 1). A possible explanation is that sliding alters the dimer contacts and tertiary structure, and could aid in the HBD-1 oligomerization that potentially enhances their chemotactic activity. The same mechanism is thought to promote the HBD-1 antibacterial activity by enhancing the surface positive charge density. The
importance of the surface positive charge to the molecule’s biological activity was also mirrored by their effects on the structure and dynamic properties. All the basic amino acids were exposed to the surface as observed in the 3D structure representations (Fig. 1d), SASA, and minimal side chain contacts for basic residues. The potential dependence of chemotactic activity on the monomer and dimer calculated average $\chi$, and end-to-end distances (Table 1) suggests that closer resemblance to the native structure and a more compact 3D structure results in better surface exposure and orientation to bind to the CCR6 receptor. Our results also suggest that the monomer, rather than the dimer, is the active in vivo chemotactic agent. Other structural observations proved the $\alpha$-helix to be less thermally stable than the $\beta$-sheet. The dimer structure asymmetry was seen in 3D representations (Fig. 9a), time evolution of the secondary structure propensity (Fig. 3b), side chain contact maps (Fig. 9a), RMSF (Fig. 9b), and SASA (data not shown).

Understanding the molecular basis for the SAR of HBD-1 could help in the process of drug discovery of novel drugs for targeting the immune and pathogen defense systems, while preventing any potential development of drug resistance.

**Supplementary Material** dealing with details of simulations and some of the results of structural analysis is available at no charge from the authors directly; the supplementary data can also be purchased from Adenine Press for US $50.00. It can also be downloaded free of charge from the author's server at the URL.

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**REFERENCES**

Table 1: A summary comparison of experimental chemotactic activity with the simulation results for the native HBD-1 peptide (WT) and the three single-site mutants (Mut1, Mut2 and Mut3). For each attribute, the four molecules are ranked in descending order using Roman numerals with I being the highest and IV being the lowest. A slight ranking difference between our findings and the reported data is shown in *italics*.

<table>
<thead>
<tr>
<th></th>
<th>WT</th>
<th>Mut1 (Asn4Ala)</th>
<th>Mut2 (Gln24Glu)</th>
<th>Mut3 (Lys31Ala)</th>
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<td>IV (40 ± 8%)</td>
<td>I (130 ± 8%)</td>
<td>II (109 ± 14%)</td>
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<td>Stability of α-helix in monomeric form (Fig. 2)</td>
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<td>IV</td>
<td>I</td>
<td>II</td>
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<td>Average χ – value from monomer simulations (Fig. 5)</td>
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<td>IV (0.90)</td>
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<td>Absent</td>
<td>Present</td>
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<td>II (0.95)</td>
<td>IV (0.83)</td>
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Figure 1: The native HBD-1 peptide (WT) in the monomeric and dimeric forms. Panel (a): The WT HBD-1 peptide extracted from the HBD-1 dimer. Panel (b): The WT HBD-1 dimer (PDB entry: 1IJV). Panel (c): The WT HBD-1 peptide at the end of 4 ns equilibration shows some structural changes compared with the initial structure (a). Panel (d): Surface representations of the equilibrated WT HBD-1 peptide oriented as panel (c), then rotated 180 degrees along the vertical axis showing distinct patches of basic and hydrophobic residues on the peptide surface. In panels (a), (b), and (c), the WT monomer is shown in cyan, and chains A and B are shown in blue and red, respectively; the cysteine residues and disulfide bonds (Cys5/Cys34, Cys12/Cys27, and Cys17/Cys35) are represented as yellow lines and bonds respectively. In panel (d), the hydrophobic, basic, acidic, and polar residues are shown, respectively, in white, blue, red, and green color.
Figure 2: Time evolution of the secondary structure propensity for the α-helical portion (residues 2-7) of the monomers of native HBD-1 peptide (WT) and mutated HBD-1 peptides (Mut1, Mut2 and Mut3) from equilibrium simulations at constant temperature $T = 300$ K (left panel), and from thermal unfolding simulations at increasing temperature (from $T_{in} = 300$ K to $T_{fin} = 500$ K; right panel). The α-, π-, and $3_{10}$-helices are shown in purple, red, and blue color, respectively; unstructured turn and coil regions are presented, respectively, in cyan and white color. The unstructured regions, observed at the end of the simulations of thermal unfolding for WT and Mut3 are encircled.
Figure 3: Time evolution of the secondary structure propensity for the mutated HBD-1 peptide Mut2 monomer (panel (a)) and dimer (panel (b)) obtained from equilibrium simulations at room temperature $T = 300$ K. The β-sheets are shown in yellow. The color denotation for the α-, π-, and $3_{10}$-helices, and for unstructured turn and coil regions is the same as in Figure 2.
Figure 4: Panel (a): The Ramachandran plot (i.e. distribution of dihedral angles $\Phi$ and $\Psi$) for the mutated HBD-1 peptide Mut2 obtained from equilibrium simulations at room temperature ($T = 300$ K; black dots), and from heating simulation at increasing temperature (from $T_{in} = 300$ K to $T_{fin} = 500$ K; green squares). Each data point represents one residue whose dihedral angles were averaged over the simulation run. Panel (b): The propensity of residues 2-7 in the mutated HBD-1 peptide Mut2 to form an $\alpha$-helix obtained from equilibrium simulations. Panel (c): Same as in panel (b) but obtained from heating simulation at increasing temperature from $T_{in} = 300$ K to $T_{fin} = 500$ K.
Figure 5: The structure overlap function $\chi$ versus time for the native HBD-1 peptide (WT) and mutated peptides (Mut1, Mut2 and Mut3) obtained from heating simulations at increasing temperature from $T_{in} = 300$ K to $T_{fin} = 500$ K. Color and plot denotations are presented in the graph.
Figure 6: Side chain contact maps for mutated HBD-1 peptide Mut3 obtained from heating simulation at \( T = 320 \) K, 400 K, and 480 K. The strength of intra-molecular binary contacts between interacting amino acids varied from the weakest (shown in white color) to the strongest (black); the yellow squares with the letter “C” point out the Cys5/Cys34 disulfide bonds in the \( \alpha \)-helix. Contacts formed between residues Asp1-Val6 in the \( \alpha \)-helix and residues Gln24-Tyr28 in the \( \beta_2 \)-strand are circled. In the rightmost panel, the residues forming the secondary structure elements are boxed; \( \alpha \)-helix: residues 2-7, and \( \beta \)-sheet: residues 11-13, 23-27, and 32-35. The amino acid residues mediating the \( \alpha \)-helix interaction with the \( \beta_2 \)-strand are shown explicitly.
Figure 7: Panels (a) and (c): The structure overlap function $\chi$ versus time from equilibrium simulations (at $T = 300$ K) for the dimers of the mutated HBD-1 peptide Mut2 first run (panel (a)), and Mut3 second run (panel (c)). In panel (c), transient structures of Mut3 dimer, recorded at 15 ns and 18 ns, show some sliding and rolling motion, which results in the disruption of contacts between residues Ile23A, Gly25A and Thr26A with Arg29B at 15 ns, and residues Thr26A and Arg29B at 18 ns (structures are color coded as in Figure 3). Panels (b) and (d): Equilibrium dynamics of side chain contact formation/rupture for stable binary interactions between amino acid residues stabilizing the dimeric form of Mut2 (panel (b)) and Mut3 (panel (d)). For Mut2 dimer, these contacts are listed from bottom to top in panel (b): Asp1/Ile23, Asp1/Thr26, Asn4/Ile23, Asn4/Gly25, Asn4/Thr26, Asn4/Lys33, Phe20/Gly30, Lys22/Asp1, Lys22/Arg29, Ile23/Arg29, Ile23/Gly30, Ile23/Lys31, Glu24/Asp1, Glu24/Tyr3, Glu24/Asn4, Glu24/Thr26, Glu24/Arg29, Gly25/Thr26, Gly25/Gly30, Thr26/Gly30, Thr26/Lys31, and Gly30/Lys31. For Mut3 dimer, these contacts are listed from bottom to top in panel (d): Asp1/Gly30, Asp1/Lys33, Tyr3/Lys22, Tyr3/Ile23, Tyr3/Gly25, Asn4/Ile23, Asn4/Gly25, Asn4/Thr26, Lys22/Asp1, Lys22/Arg29, Ile23/Thr26, Ile23/Arg29, Ile23/Ala31, Glu24/Asp1, Glu24/Lys31, Glu24/Arg29, Glu25/Thr26, Gly25/Gly30, Thr26/Arg29, Thr26/Gly30, Thr26/Ala31, Arg29/Asp1, Arg29/Tyr28, Gly30/Arg29, and Gly30/Gly30. All red vertical lines connecting upper and lower panels represent the specific times at which large changes to the overall structure and the dynamics of binary contacts have occurred.
Figure 8: The total number of inter-monomer contacts between interacting amino acid residues in chains A and B (see Figure 1) obtained from equilibrium simulations at $T = 300$ K for the dimers of native HBD-1 peptide (WT; average of two trajectories) and mutated peptide (Mut2). The three representative simulation runs, displayed by solid black, dotted red, and solid green curves, show stochastic variability in the number of inter-monomer association interactions. The first trajectory for the Mut2 dimer (solid black curve), also discussed in Figure 6, had noticeable structural changes that occurred at $\sim 6$, $18$, $22$, and $24$ ns (marked by vertical arrows). In the second run for the Mut2 dimer (dotted red curve), a complete dissociation transition was observed at $t \approx 22$ ns, which was followed by redimerization at $t \approx 25$ ns.
Figure 9: Panel (a): The time-averaged map of interacting residues (side chain contact map) in chain A (shown in blue color) and chain B (red) for the dimer of WT HBD-1 peptide obtained from equilibrium simulation at $T = 300$ K. The side chain contact map shows structural evidence for the asymmetric dimerization site between chains A and B, encircled in the structure and on the map. The contacts’ representations ranged from black (strongest coupling) to white (weakest interaction) depending on the contact strength throughout the simulation run; the yellow squares point out the position of the disulfide bonds (Cys 5/34, 12/27, and 17/35). Panel (b): The root-mean-square fluctuations (RMSF) for amino acid residues in chains A and B for the same run displayed in panel (a), showing similar yet non-identical pattern for the two dimer chains. Higher RMSF values were observed at the C- and N-termini and in the turn regions, while the minimum RMSF values were observed for the $\alpha$-helical portion, disulfide bonds and the dimerization site localized to residues 23-26 and 26-33 of chain A and B, respectively (encircled in red).
Figure 10: The root-mean-square deviation (RMSD) for the dimers of native (WT) HBD-1 peptide (panel (a)), and mutants Mut2 (panel (b)) and Mut3 (panel (c)) obtained from equilibrium simulations at $T = 300$ K. The first, second, and third simulation runs are represented by the solid black, dotted red, and solid green curves, respectively. The solid black (dotted red) trajectory for Mut2 (Mut3) dimer are discussed in Figure 6. The WT dimer was found to be the most stable (RMSD ≈ 2 Å), while the curves for Mut2 and Mut3 showed the highest RMSD (≈ 7-10 Å). In panels (b) and (c), the vertical arrows mark the times at which noticeable structural changes had occurred as discussed in Figure 6.
Figure 11: The solvent accessible surface area (SASA) per residue obtained from heating simulations of the native HBD-1 peptide (WT, shown in solid black color) and mutants Mut1 (dotted red), Mut2 (solid green), and Mut3 (dashed blue) at increasing temperature (from $T_{in} = 300$ K to $T_{fin} = 500$ K). Each mutant had smaller SASA value at its mutation site (encircled) as compared to the WT peptide with the smallest SASA difference for Mut2 (substitution Gln24Glu).