In Silico Analysis of Bacteriocins from Lactic Acid Bacteria Against SARS-CoV-2

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Accepted: 11 November 2021 / Published online: 27 November 2021 © The Author(s), under exclusive licence to Springer Science+Business Media, LLC, part of Springer Nature 2021

Abstract
The COVID-19 pandemic caused by a novel coronavirus (SARS-CoV-2) is a serious health concern in the twenty-first century for scientists, health workers, and all humans. The absence of specific biotherapeutics requires new strategies to prevent the spread and prophylaxis of the novel virus and its variants. The SARS-CoV-2 virus shows pathogenesis by entering the host cells via spike protein and Angiotensin-Converting Enzyme 2 receptor protein. Thus, the present study aims to compute the binding energies between a wide range of bacteriocins with receptor-binding domain (RBD) on spike proteins of wild type (WT) and beta variant (lineage B.1.351). Molecular docking analyses were performed to evaluate binding energies. Upon achieving the best bio-peptides with the highest docking scores, further molecular dynamics (MD) simulations were performed to validate the structure and interaction stability. Protein–protein docking of the chosen 22 biopeptides with WT-RBD showed docking scores lower than $-7.9$ kcal/mol. Pediocin PA-1 and salivaricin P showed the lowest (best) docking scores of $-12$ kcal/mol. Pediocin PA-1, salivaricin B, and salivaricin P showed a remarkable increase in the double mutant’s predicted binding affinity with $-13.8$ kcal/mol, $-13.0$ kcal/mol, and $-12.5$ kcal/mol, respectively. Also, a better predicted binding affinity of pediocin PA-1 and salivaricin B against triple mutant was observed compared to the WT. Thus, pediocin PA-1 binds stronger to mutants of the RBD, particularly to double and triple mutants. Salivaricin B showed a better predicted binding affinity towards triple mutant compared to WT, showing that it might be another bacteriocin with potential activity against the SARS-CoV-2 beta variant. Overall, pediocin PA-1, salivaricin P, and salivaricin B are the most promising candidates for inhibiting SARS-CoV-2 (including lineage B.1.351) entrance into the human cells. These bacteriocins derived from lactic acid bacteria hold promising potential for paving an alternative way for treatment and prophylaxis of WT and beta variants.

Keywords Beta variant · Bacteriocins · COVID-19 · Protein–protein docking · SARS-CoV-2 · MD simulations

Introduction

COVID-19 is caused by a coronavirus named severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) [1–3]. SARS-CoV-2 has spread globally and devastated public health and economies, with more than 179 million people infected and more than 3.8 million deceased to date (www.worldometers.info). The spike (S) protein, which is responsible for binding to the host receptor and viral entry, consists of S1 and S2 subunits. SARS-CoV-2 infects host cells via binding to angiotensin-converting enzyme II (ACE 2) [4], which is dominantly present on the epithelial cells [5]. Thus, blocking the spike can be considered as an effective approach to prevent SARS-CoV-2 infection. ACE2 is a regulator of gut inflammation which is more expressed in the ileum and colon than in lung [6, 7]. More than 60% of patients diagnosed with COVID-19 reported showing diarrhea, nausea,
and vomiting gastrointestinal symptoms. More importantly, patients having gastrointestinal complaints had overall experienced severe disease outcome [8–10]. Thus, a potential role of gut microbiota and COVID-19 progression and severity could be linked [11–13].

According to International Scientific Association of Probiotics and Prebiotics (ISAPP) the term postbiotics can be described as “preparation of inanimate microorganisms and/or their components that confers a health benefit on the host” [14]. These could include non-viable bacterial or complex mixture of healthy metabolic products or secreted components like short-chain fatty acids, vitamins, amino acids, organic acid enzymes, secreted proteins, or peptides such as bacteriocins [15–17]. As biotherapeutic agents, bacteriocins or individual cellular components can be directly used to target a particular disease, which may help to provide clear evidence for the effect of the biomolecules. Instead, to study the same with probiotics might yield unclear cause and effect relationship due to dynamic and complex morphology of bacteria. Thus, several bacteriocin molecules draw attention because of their known biochemical structure, storage stability, and the capability to induce mechanisms to control inflammation, adhesion of pathogens to GIT, obesity, hypertension, coronary artery diseases (CVD), cancer, and oxidative stress. And several bacteriocin preparations were patented as bio-therapeutics for a specific health claim, “immune-modulation” [16, 18].

Bacteriocins have recently drawn attention due to the COVID-19 pandemic, which is widespread around the world. Bacteriocins are ribosomally produced peptides with bactericidal effects, known to be produced by several bacteria, including lactic acid bacteria [11]. The antibacterial activities of bacteriocins have been well understood, but the basis of their antiviral activities has been embraced with a lot less attention. Various research groups worldwide probed on their antiviral activities; for example, Todorov et al. (2005) reported that the enterocin CRL35 peptide produced by Enterococcus mundtii ST4V inhibited the replication of Herpes simplex virus (HSV) type 1 and 2 [19]. Ferr et al. (2010) reported that Labyrinthopeptin A1, a carbacyclic lantibiotic of Actinomadura namibiensis DSM 6313, showed antiviral activity against HIV-1 and HSV-1 [20]. Furthermore, Serkedjieva et al. (2000) showed that the replication of H7N7 and H7N1 strains of influenza virus was highly specifically inhibited by 5.0-kDa bacteriocin of Lactococcus delbrueckii subsp. bulgaricus in chicken embryo fibroblast [21]. Quintana et al. (2014) reported that bacteriocins of Lactococcus lactis GLe03 and GLe05 showed virucidal effect on HSV-1 and poliovirus (PV-1) [22]. Recently, Anwar et al. (2020) studied the molecular dynamics (MD) simulations of plantaricin peptides (W, JLA-9, D, and BN) against spike proteins (S), human-ACE-2, and RNA-dependent RNA polymerase (RdRp). The results of their study revealed that plantaricin peptides (W, JLA-9, D, and BN) blocked the entry into the cell by binding to S protein, ACE-2, and RdRp. In another study, Manna et al. (2020) demonstrated a higher binding tendency of subtilisin, curvacin A, sakacin P, lactococcin Gb to S-protein of SARS-CoV-2, and human ACE-2 by using in silico approach [24]. To our knowledge, no bacteriocin activity against the SARS-CoV-2 beta variant (lineage B.1.351) has been reported so far. Therefore, elucidation of the bacteriocin–S protein interactions at the atomic level is critical to fill this gap in the literature. Thus, the present study focused on the potential therapeutic functions of several bacteriocins against SARS-CoV-2. We utilized an in silico protein–protein docking approach to elucidate the binding affinity of bacteriocin peptides to the RBD of SARS-CoV-2. Ultimately, the role of lactic acid bacteria-derived bacteriocins on strengthening the immune defenses against fighting coronaviruses of wild type and beta variant could be uncovered.

Materials and Methods

In Silico Preparation of Antimicrobial Peptides and Receptors

We collected 22 bacteriocin sequences from PubChem. MarvinSketch was used to draw and convert to 3D structures of the peptides (MarvinSketch 19.13, ChemAxon; https://www.chemaxon.com, Budapest, Hungary). We considered uncharged N- and C-terminal ends in our docking calculations. (In our dataset, most of the C-terminals have −NH2 and −H ending; this option led us to consider only carbonyl group the C-term). Folded structures of the peptides were predicted using the PEP-FOLD3 algorithm [25]. Then obtained folded structures were incorporated into Schrodinger’s Maestro suite (Schrodinger LLC, New York, NY) [26, 27]. PHE14 of Enterocin EJ97 (139586139) and CYS1 of Gassericin B2 (139586609) were methylated in the original structure, and we deleted methyl moiety from both sequences. Coronavirus RBD-ACE2 (PDB ID: 6LZG) complex was downloaded from Protein Data Bank. All solvent molecules and ACE2 were deleted. Protein preparation wizard was used to predict protonation states of the side chains at pH 7.4 of both peptides and RBD [28, 29]. All beta variants (K417N, K417N-E484K, and K417N-E484K-N501Y mutants) RBDs were generated using Maestro (Schrodinger LLC).

Protein–Protein Docking Simulations

HADDOCK v2.4 was used in protein–protein docking simulations [30]. Active residues at the binding pocket of RBD were obtained from the literature [31, 32]. Following residues were defined as active: LYS417, GLY446, TYR449, TYR453, LEU455, PHE456, ALA475, GLY476, GLU484,
Molecular Dynamics Simulation

Among all docking results, the best performing ligand was pediocin PA-1 (56842033) against WT and mutant RBDs. We simulated complexes of WT, K417N, K417N_E484K, and K417N_E484K_N501Y with pediocin PA-1 for 500 ns and as three independent runs. In total, we have simulated 6 µs for all simulation systems (such as WT- pediocin PA-1 500 ns × 3 times, K417N- pediocin PA-1 500 ns × 3 times). Desmond MD code (D. E. Shaw Research, New York, NY) was used to simulate systems with Martyna Tobias Klein (MTK) [35] barostat to control pressure at 1.01 bar, and with Nosé–Hoover [36, 37] thermostat to control the temperature at 310 K. OPLS3 force field was used in all simulations [38].

Reversible reference system propagator algorithms (RESPA) were used to calculate bonded and non-bonded interactions. Protein–Ligand systems were solvated in the TIP3P water model [39] and in an orthorhombic box with a 10-Å buffer distance in all x, y, and z dimensions. The 0.15 M NaCl ionic salt concentration was used to neutralize the medium. Each system was simulated for 500 ns, and a total of 5000 frames was collected and saved. Two-hundred frames out of 5000 trajectory frames were used in Molecular Mechanics Generalized Born Surface Area (MM-GBSA) calculations. The Prime module of Schrodinger’s Maestro package (Schrodinger LLC) [40, 41] was used to calculate relative binding free energies throughout the simulation along with the VSGB 2.0 [42] implicit water model.

Hydrogen Bond Analysis and Trajectory Clustering

The number of hydrogen bonds between pediocin PA-1 and RDBs was calculated using the Maestro package (Schrodinger LLC). The resultant hydrogen bond numbers for each system were converted to a probability distribution. For smoothing, “cubic splines” with a smoothing parameter of 0.99449 was used in MATLAB 2019b curve fitting toolbox (The MathWorks, Inc., Natick, MA). Before concatenation of the trajectories, all ions and water molecules were stripped out. All frames opened and aligned to starting structure in Visual Molecular Dynamics (VMD) [43]. Aligned trajectories were saved in GROMACS trr format, and gmx trjcat was used to concatenate three independent runs [44]. After concatenation, we obtained a single 15,000 frame long single trajectory for each system. Gromacs clustering algorithm was applied with the 3-Å cut-off using the gmx cluster tool [45]. The following clusters were obtained, 23, 19, 30, and 18, for the WT, K417N-RBD, K417N-E484K-RBD, and K417N-E484K-N501Y-RBD, respectively. The first member of each cluster was used to evaluate interactions at the interface and plotting figures.

Statistical Analysis

MM/GBSA scores of WT and mutants are treated as independent samples (3 replicates of 200 frames). Each frame is separated by 2.5 ns. Since distributions are not necessarily normally distributed, we used the Wilcoxon signed-rank test on datasets. MATLAB 2019b (The MathWorks, Inc., Natick, MA) statistical toolbox was used.

Results

Molecular Docking Simulations Reveal Pediocin PA-1 as the Tightest Binder

In all WT and mutants, pediocin PA-1 was the tightest binder. Salivaricin P comes second for the WT and K417N. For the K417N-E484K double and K417N-E484K-N501Y triple mutants, salivaricin B achieved second best score, where salivaricin P took third place. Among all used peptides, pediocin PA-1, salivaricin P, and salivaricin B were the best three according to docking scores. Planataricin JLA-9 was found the weakest binder among all. Interestingly, pediocin PA-1 binding energies get even lower upon the mutations, and tighter complexes are observed. In K417N-E484K double mutant, pediocin PA-1 binding was found the lowest energy complex compared to all other mutant-peptide complexes (Table 1). Pediocin PA-1 binding poses at the interface of WT and mutants represented in Fig. 1 and Figure S1. Table 2 shows the interactions between pediocin PA-1 and RBD. A3-B492, A12-B416, and A39-B484 interactions were only observed in WT docking poses. A2-B446 interaction was the only interaction that was observed solely in the K417N mutant. A2-B444, A3-B449, A3-B493, A4-B498, and A21-B505 were only observed in K417N-E484K double mutant. A5-B449, A6-B449, A6-B501, A7-B505, and A20-B505 interactions were merely observed in the...
K417N-E484K-N501Y triple mutant. A8-B453, A11-B473, A15-B403, and A43-B484 interactions were observed in all docking poses. Interestingly, A43 of the pediocin PA-1 interacted with residue number 484 of RBDs in all studied WT and mutant systems.

In salivaricin P docking poses, the following interactions are given in Table S1. Salivaricin P binding poses at the interface of WT and mutants given in Figure S2. A2-B449, A10-B498, A10-B501, A15-B487, and A18-B489 interactions were observed for only the WT-salivaricin P complex. The K417N mutant, A3-B493, A6-B492, A6-B494, A9-B494, A10-B493, A13-B489, and A17-B489 interactions were not observed in all other systems, and we can conclude that these interactions were K417N specific interactions. A1-B416, A1-B421, A1-B420, A4-B417, A7-B421, A9-B487, A10-B489, A10-B487, A11-B493, A11-B490, A14-B496, A16-B498, and A18-B505 interactions were only observed in K417N-E484K double mutant. For the K417N-E484K-N501Y triple mutant, A4-B489 and A11-B501 interactions were detected and found to be case-specific interactions.

Simulation Trajectories Identify K417N-E484K-N501Y Triple-Mutant RBD as the Most Stable System

Herein, WT-pediocin PA-1 simulations, we observed RMSD averages as 2.20 ± 0.35, 1.97 ± 0.26, and 2.21 ± 0.24 Å. In all simulations, RMSD values of RBDs converged just after 50 ns. Figure S4 shows the WT-RBD root mean square deviation (RMSD) graph. The average RMSD of the backbone atoms of the WT-RBD simulations was found as 2.13 Å (the average of the three runs). Very similar RMSD graphs were observed in the case of K417N-pediocin PA-1 complexes compared to WT.
RMSD averages were calculated 2.15 ± 0.24, 2.23 ± 0.26, and 1.97 ± 0.13 Å, respectively (see Figure S5). The average of the RMSD for the K417N mutant backbone atoms was calculated as 2.11 Å (the average of the three runs). For the K417N-E484K double-mutant-pediocin PA-1 simulations, we found backbone RMSD averages 2.38 ± 0.41, 2.15 ± 0.33, and 2.44 ± 0.40 Å, respectively (Figure S6). The average of the three runs was 2.32 Å (the average of the three runs). K417N-E484K-N501Y triple mutant has the lowest average backbone RMSD value of 2.05 Å compared to other systems (the average of the three runs). For the three replicates, the following averages were found: 1.91 ± 0.28, 2.04 ± 0.19, and 2.21 ± 0.50 Å, respectively (see Figure S7). When the backbone RMSD values were compared in all simulations, very similar behavior was observed. RBDS reached the plateau a very short time (around ~50 ns) after the start of simulation in both the WT and mutant states. The lowest RMSD average was observed in the K417N-E484K-N501Y triple mutant, while the highest RMSD average was observed for the K417N-E484K double mutant.

The Most Stable Conformation of Pediocin PA-1 Observed in K417N

K417N mutant has the lowest RMSD average for the pediocin PA-1 compared to others and is found to be 4.07 Å (the average of the three runs). The second lowest average RMSD value was found in the K417N-E484K-N501Y triple mutant as 4.44 Å (the average of the three runs). For the K417N-E484K double mutant and WT, RMSD values of pediocin PA-1 backbone atoms were found at 4.49 and 5.00 Å, respectively (the average of the three runs). Figure S13 shows the backbone RMSD values of all studied systems. Figure S14 depicts backbone RMSD values for the pediocin PA-1 at the WT-RBD complexes. RMSD averages 4.19 ± 0.55 Å, 6.13 ± 1.10 Å, and 4.68 ± 0.69 Å were obtained for the repeats, respectively. In the second run, we observed a different profile for the backbone atom RMSD of the pediocin PA-1. The backbone RMSD of the pediocin PA-1 at the K417N-RBD complexes was given in Figure S15. We observed 3.30 ± 0.93 Å, 4.12 ± 0.48 Å, and 4.80 ± 0.96 Å for the repeats, respectively. Run-2 and run-3 were not changed too much after 100 ns (see Figure S15). In K417N-E484K double-mutant RBD and pediocin PA-1 complexes, for the run-1, run-2, and run-3, we observed 5.61 ± 0.79 Å, 3.38 ± 0.38 Å, and 4.47 ± 0.92 Å, respectively (see Figure S16). For the K417N-E484K-N501Y triple-mutant RBD-pediocin PA-1 simulation repeats, we observed a very similar RMSD profile for the run-1 and run-2, especially in the last 250 ns. The averages are given in Figure S17, and for the simulation, repeats were...
found as 5.13 ± 0.62 Å, 4.80 ± 0.73 Å, and 3.38 ± 0.52 Å, respectively.

The Lowest MM/GBSA Binding Energy Observed in K417N-Pediocin PA-1 System

All MM/GBSA binding free energies for all studied systems are given in Figure S8. Figures S9–S12 depict the system-wise MM/GBSA energies for the WT, K417N, K417N-E484K, and K417N-E484K-N501Y mutants. The averages for the studied systems calculated were as follows: -101.50, -132.68, -109.63, and -129.26 kcal/mol, respectively. The lowest MM/GBSA binding energy was observed in the K417N system (see Figure S10). However, the highest score corresponds to WT (see Figure S9).

Similar to docking results, in MM/GBSA energies, WT was the highest one compared to all others. Based on the MM/GBSA binding energies, we conclude that the predicted affinity of pediocin PA-1 increases upon mutation.

Box plots of the MM/GBSA binding energy distributions are given in Fig. 3A. All mutants are significantly different from the WT. Two distributions that did not reject the null hypothesis were K417N and K417N-E484K-N501Y.

Simulation Cluster Structures Show Different Interaction Patterns

When compared the starting structure and cluster structure interactions at the interface, we observed that the A1-B484 interaction was conserved in both WT and K417N. However, the A3-B493 interaction was conserved in K417N-E484K double mutant and newly established for the WT and K417N. Although A5-B498 interaction was preserved for the WT, it was lost in the K417N mutation. The same interaction occurred in the K417N-E484K double mutant. A43-B484 interaction was conserved only in K417N and K417N-E484K mutations observed across all simulation starting structures. A17-B505 and A20-B505 interactions were conserved only in K417N and K417N-E484K mutations, respectively (see Table 2 and 3).

In Fig. 2 and Figure S22, we plotted the first member of each cluster. First members of each cluster cover 30.19%, 30.37%, 28.32%, and 28.00% of total frames for the WT, K417N, K417N-E484K, and K417N-E484K-N501Y, respectively. Polar interactions between RBD and pediocin PA-1 are given as dashes (see Fig. 2). A1-B484, A3-B484, A3-B493, A4-B494, A5-B493, A5-B498, A6-B453, A27-B498, A35-B484, and A39-B493 were observed between WT-RBD and pediocin PA-1 (Fig. 2A). For the K417N-RBD-pediocin PA-1 complex, the following polar interactions were detected in the cluster structure: A1-B484, A3-B484, A3-B493, A4-B446, A5-B501, A6-B453, A27-B498, A35-B484, and A39-B493 were observed between WT-RBD and pediocin PA-1 (Fig. 2A). For the K417N-RBD-pediocin PA-1 complex, the following polar interactions were detected in the cluster structure: A1-B484, A3-B484, A3-B493, A4-B446, A5-B501, A6-B453, A17-B505, A41-B487, and A42-B485 (Fig. 2B). The A3-B493, A4-B494, A5-B403, A5-B453, A5-B496, A5-B498, A11-B415, A11-B420, A20-B405, and A20-B406 polar interactions were found in K417N-E484K-RBD and pediocin PA-1 cluster structure (Fig. 2C). The A3-B494, A5-B494, A9-B417, A11-B415, A11-B417, A11-B420, A11-B460, A20-B505, A40-B486, and A43-B484 polar interactions were found in the K417N-E484K-N501Y triple-mutant and pediocin PA-1 cluster structure (Fig. 2D).

Table 2 Pediocin PA-1-Spike Protein RBD domain interactions. Pediocin PA-1 presented as chain A and the RBD as chain B. In A:1, A represents chain A and number (in this case, 1) after the colon depicts the residue number. 1 (bold) means interaction observed, and 0 (italics) means no interaction. Data created using simulation starting structures

| Pediocin PA-1 | RBD | WT | K417N | K417N_E484K | K417N_E484K_N501Y |
|---------------|-----|----|-------|-------------|------------------|
| A:1           | B:484 | 1  | 1     | 0           | 0                |
| A:2           | B:444 | 0  | 0     | 1           | 0                |
| A:2           | B:446 | 0  | 1     | 0           | 0                |
| A:3           | B:449 | 0  | 0     | 1           | 0                |
| A:3           | B:493 | 0  | 0     | 1           | 0                |
| A:3           | B:492 | 1  | 0     | 0           | 0                |
| A:4           | B:498 | 0  | 0     | 1           | 0                |
| A:4           | B:449 | 0  | 1     | 1           | 1                |
| A:5           | B:449 | 0  | 0     | 0           | 1                |
| A:5           | B:501 | 0  | 0     | 1           | 1                |
| A:5           | B:498 | 1  | 1     | 0           | 0                |
| A:6           | B:449 | 0  | 0     | 0           | 1                |
| A:6           | B:501 | 0  | 0     | 0           | 1                |
| A:6           | B:496 | 1  | 0     | 1           | 1                |
| A:7           | B:505 | 0  | 0     | 0           | 1                |
| A:7           | B:453 | 1  | 0     | 1           | 0                |
| A:8           | B:453 | 1  | 1     | 1           | 1                |
| A:9           | B:473 | 1  | 1     | 1           | 1                |
| A:10          | B:457 | 1  | 1     | 0           | 0                |
| A:12          | B:421 | 0  | 1     | 0           | 1                |
| A:12          | B:417 | 0  | 1     | 0           | 0                |
| A:12          | B:420 | 1  | 1     | 0           | 1                |
| A:12          | B:416 | 1  | 1     | 0           | 0                |
| A:13          | B:417 | 1  | 1     | 0           | 1                |
| A:14          | B:405 | 0  | 1     | 1           | 1                |
| A:15          | B:403 | 1  | 1     | 1           | 1                |
| A:16          | B:505 | 1  | 0     | 0           | 0                |
| A:17          | B:505 | 1  | 1     | 1           | 0                |
| A:20          | B:505 | 0  | 0     | 0           | 1                |
| A:21          | B:505 | 0  | 0     | 1           | 0                |
| A:39          | B:484 | 1  | 0     | 0           | 0                |
| A:42          | B:489 | 0  | 0     | 1           | 1                |
| A:43          | B:490 | 0  | 0     | 1           | 1                |
| A:43          | B:484 | 1  | 1     | 1           | 1                |
TYR2, TYR3, THR8, LYS20, HIS38, and HIS42 of Pediocin PA-1 Identified as Important Residues

A3-B492 interaction was observed in docking pose; however, this contact was lost in MD simulations. Interestingly, this interaction was shifted to A3-B493 and conserved as 84%, 15%, and 30% in WT simulations (Table S3). A12-B416 interaction was detected in docking pose; in MD simulations, it was completely lost. The A39-B484 interaction was WT specific, which was conserved at 12%, 9%, and 40% throughout the MD trajectories (Table S3). A2-B446 interaction was the only interaction observed in K417N mutant compared to WT, double, and triple mutants. However, during MD simulations, this interaction was conserved at 63% and 15% in K417N-1 and K447N-3 replicas (Table S3). In the second run, this contact was completely lost. On the other hand, a new interaction was formed between A2-B449 and conserved at 38%, 6%, and 61% during MD simulations (Table S3). In the case of the K417N-484 K double-mutant system, A2-B444, A3-B449, and A21-B505 interactions were determined as specific contacts. However, none of these contacts were conserved throughout the MD simulations. A3-B493 interaction was observed as 91%, 57%, and 6% during triplicate simulations (Table S3). However, the A4-B498 interaction was again disrupted, and the percentage of this interaction decreased to 6% and 7% in the second and third simulations, respectively (Table S3). The highest number of H-bonds was observed in K417N mutant.

The Highest Number of Hydrogen Bonds Observed in K417N Mutant

The H-bond probability distributions are plotted in Fig. 3B. Observed average number of hydrogen bonds were 8 ± 2, 6 ± 3, and 10 ± 3 for the WT-pediocin PA-1 complexes in 3 repeats, respectively. Figure S18 depicts the H-bonding pattern throughout the MD repeats. We observed similar patterns in the run-1 and run-2 in the last 200 ns of the simulations. The highest number of H-bonds was observed in run-3 for the WT-pediocin PA-1 complexes (see Figure S18). K417N-pediocin PA-1 H bonding patterns are given in Figure S19. After the first 100 ns, we observed very
Fig. 2 In silico analysis of the binding interactions between predicted pediocin PA-1 peptide and different mutants of the spike protein complex. (A) RBD_WT-pediocin PA-1, (B) RBD_K417N-pediocin PA-1, (C) RBD_K417N_E484K-pediocin PA-1, and (D) RBD_K417N_E484K_N501Y-pediocin PA-1. Figures were prepared using cluster structures. Polar interactions were plotted, and the interacting residues were labeled. RBD residues labeled in red, pediocin PA-1 residues colored in black.

Fig. 3 (A) Box plot representation of the MM/GBSA scores. All averages are significantly different from each other. However, the averages of K417N-RBD and K417N-E484K-N501Y-RBD averages are not significantly different. (B) Probability distributions of H-bond numbers in each system. K417N-RBD system differs from others slightly.
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they might be an alternative source for developing therapeutic species. That is why, being deeply investigated for their potent antiviral activities against SARS-CoV-2, and one of the most desired drug targets [43]. We only focused on the RBD domain of S protein because it is the key domain of S protein that binds tightly to ACE2 [31].

Protein–protein interface interactions revealed that Tyr2, Tyr3, Thr8, Lys20, His38, and His42 are essential residues for pediocin PA-1, while Tyr449, Leu455, Phe486, Tyr489, Gln493, Ser494, and Asn-Tyr505 were key residues in the RBD of S protein. Mainly, Leu455, Phe486, and Gln493 are among the key residues for the binding affinity of S protein to ACE2 receptor. Yi et al. [46] mutated 9 amino acid residues of S protein, including those three amino acids mentioned above, through single amino acid substitution mutagenesis. The binding affinity for ACE2 to the RBD was abolished upon the corresponding amino acid substitutions, indicating that they are key amino acid residues in the binding of RBD to ACE2 receptor. They also identified Ala475 and Phe486 in the SARS-CoV-2 RBD as the key amino acids for recognizing both ACE2 receptor and neutralizing antibodies, representing new binding sites for neutralizing antibodies [46]. Therefore, pediocin PA-1 could be a promising therapeutic candidate against SARS-CoV-2.

Pediocin PA-1, produced by lactic acid bacteria such as Lactococcus lactis, Pediococcus acidilactici, and Pediococcus pentosaceous, is a broad-spectrum antimicrobial agent and classified as class IIa non-lantibiotic bacteriocin [53, 54]. Mainly, it exhibits strong biological activity against Listeria monocytogenes, a foodborne pathogen and a causative agent of listeriosis [47, 48]. The mode of action for class IIa bacteriocins is based on the formation of pores in the membrane of the susceptible bacteria resulting in the dissipation of protein-motive force and the depletion of ATP [49]. In this mechanism, the bacteriocins are attracted to the target membrane receptors by electrostatic interaction due to their cationic properties [50]. On the other hand, there is no study for antiviral activity of pediocin PA-1, yet. Todorov et al. (2010) reported the antiviral activity of pediocin-like bacteriocin (99% of DNA homology to pediocin PA-1) against herpes simplex virus [51]. This study indicated the potential antiviral activity of pediocin PA-1. Moreover, salivaricin P and B were produced by Lactobacillus salivarius, Lactococcus lactis, and Streptococcus salivarius, respectively [52, 53].

Salivaricins possess selective antimicrobial activities against oral and upper respiratory tract bacteria [54]. In particular, salivaricin B is a potent antimicrobial agent with a broad-spectrum activity [55]. Salivaricins have a similar mode of action to pediocins, with an exception (salivaricin B). For instance, salivaricin 9 and salivaricin mmaye1 form pores on the target membranes of pathogenic bacteria [56, 25].

Discussion

The current pandemic of SARS-CoV-2 has been globally threatening public health [18]. Millions of people have been infected by the coronavirus, and many of them have lost their lives. Therefore, there is an urgent need for a cure for the prevention and treatment of COVID-19 [43]. Even though recently developed vaccines pave the way for prevention, there are still no specific drugs to treat COVID-19. For this urgent need for new drugs, the scientific community has been spending enormous effort to find drug molecules that inhibit the entry of SARS-CoV-2. This could be through the inhibition of key enzymes involved in its life cycle (antiviral agents) or regulation of the cytokine storm in the host (immunomodulators) [44, 45]. Many synthetic and natural compounds have been screened [46–48] to find effective drug candidates. However, bacteriocins have not been deeply investigated for their potent antiviral activities against SARS-CoV-2. Their pleiotropic biological effects, including the immunomodulatory effect on the human body, are known [49]. Considering their health-promoting properties, they might be an alternative source for developing therapeutics against SARS-CoV-2 [12, 50]. Indeed, bacteriocins were recently reported in few studies to be a potential therapeutic candidate for the treatment and prophylaxis of COVID-19 [4, 24, 51].

Very few bacteriocins from probiotic bacteria were tested using in silico approaches against SARS-CoV-2 in the literature. In silico evaluation of 22 probiotic-derived bacteriocins were conducted using molecular docking and MD simulations. The RBD of S protein of SARS-CoV-2 was selected as the drug target protein because SARS-CoV-2 enters the human cells through its attachment to the ACE-2 receptor, which is a key step in infectivity with SARS-CoV-2 and one of the most desired drug targets [43].

For the K417N-E484K double-mutant and pediocin PA-1 complexes, we observed 10 ± 2, 7 ± 3, and 8 ± 2 number of H-bonds, respectively. In the last 200 ns, the bonding pattern across the repeats changed (see Figure S20). Run-1 was the simulation with the highest average H-bond numbers compared to other repeats in double-mutant simulations. The H-bonding patterns for the K417N-E484K-N501Y triple-mutant RBD and pediocin PA-1 complexes were depicted in Figure S21. The average and the standard deviations for the runs were found 7 ± 2, 13 ± 2, and 8 ± 2, respectively. In the first 150 ns of the trajectories, all repeats had a different H bonding pattern. However, after 200 ns, run-1 and run-3 converged to a similar binding pattern, and the run-2 differentiated from the other two having the highest number of H-bonds (see Figure S21). Based on H-bond analysis, the most increased number bonding was observed in K417N system, which agrees with MM/GBSA binding energies.

very few bacterial infections from probiotic bacteria were tested using in silico approaches against SARS-CoV-2 in the literature. In silico evaluation of 22 probiotic-derived bacteriocins were conducted using molecular docking and MD similar binding patterns in all repeats. The averages for the repeats were 11 ± 3, 10 ± 3, and 11 ± 2, respectively. Also, in the last 80 ns, a decrease was observed in run-1, which differentiated slightly from run-2 and run-3 (see Figure S19). For the K417N-E484K double-mutant and pediocin PA-1 complexes, we observed 10 ± 2, 7 ± 3, and 8 ± 2 number of H-bonds, respectively. In the last 200 ns, the bonding pattern across the repeats changed (see Figure S20). Run-1 was the simulation with the highest average H-bond numbers compared to other repeats in double-mutant simulations. The H-bonding patterns for the K417N-E484K-N501Y triple-mutant RBD and pediocin PA-1 complexes were depicted in Figure S21. The average and the standard deviations for the runs were found 7 ± 2, 13 ± 2, and 8 ± 2, respectively. In the first 150 ns of the trajectories, all repeats had a different H bonding pattern. However, after 200 ns, run-1 and run-3 converged to a similar binding pattern, and the run-2 differentiated from the other two having the highest number of H-bonds (see Figure S21). Based on H-bond analysis, the most increased number bonding was observed in K417N system, which agrees with MM/GBSA binding energies.
However, salivaricin B does not induce pore formation in susceptible bacterial cells. Its mode of action is based on the interference with cell wall biosynthesis [52]. Additionally, the potential antiviral activity of salivaricin B for viral pharyngitis was also reported [58]. Due to the antiviral activities of bacteriocins such as peptide ST4V, enterocin CRL35, and enterocin ST5Ha, they were suggested to be potential antiviral agents in the prevention and treatment of COVID-19 [59]. Based on the current literature and our in silico findings, the most promising three bacteriocins found in the present study (pediocin PA-1, salivaricin P, and salivaricin B) can be considered as an alternative solution for the treatment and prophylaxis of SARS-CoV-2.

Biological stability and the survival of bacteriocins in the gastrointestinal tract (GIT) are also substantially crucial for considering their use as a therapeutic agent against SARS-CoV-2. Pediococcus acidilactici UL5, a producer of pediocin PA-1, indicated high resistance to GIT conditions and could produce pediocin PA-1 under GIT conditions but purified pediocin PA-1 lost the biological activity under GIT conditions. One potential solution for preventing pediocin PA-1 from GIT conditions is to be delivered in the microcapsules [60, 61]. Another avenue could be developing bioactive analogs of pediocin PA-1, which can provide better biological activities and be less susceptible to GIT conditions [62]. A similar study was conducted for salivaricin P, in which 11 analogs of salivaricin P were generated to prevent trypsin digestion. Eight of them were resistant to trypsin digestion while still maintaining their antimicrobial activity against L. monocytogenes, indicating the potential of engineered bacteriocin analogs for their biological activities and survival in GIT [63]. The biological stability studies of salivaricin B showed that it was resistant to proteolytic enzymes, such as pepsin, trypsin, and chymotrypsin, while it was partially sensitive to proteinase K. The biological activity was stable over a range of pH from 3 to 9.

Temperature stability was also tested for salivaricin B, which was highly stable during heating and repeated freezing and thawing cycles, and was resistant to 100 °C. An alternative approach for reducing the susceptibility of these promising bacteriocins against proteolytic enzymes is to engineer them to generate their analogs [62]. Another approach is to utilize those bacteriocin biosynthesizing probiotics since they might not be affected by the proteolytic enzymes in GIT, i.e., pediocin PA-1 producing probiotic [61]. Moreover, some probiotics are known to boost the immune system. For example, L. acidophilus upregulated the natural killer T cells in mice and played a preventive role in the influenza virus infection. In the same study, it was also found that L. acidophilus triggers the expression of viral defense genes in mice [65]. Probiotics can also bind to a virus and thus inhibit the binding of viral protein to the host receptor, leading to the prevention of entry of the virus [66]. In this sense, probiotics and their bacteriocins can provide a preventive role in entering viruses into the host cells. For example, Manna et al. conducted an in silico study for few bacteriocins (subtilisin, curvacin A, sakacin P, and lactococcin Gb). They found that bacteriocins had strong affinities to S protein or RBD of S protein [24]. Furthermore, we performed molecular docking simulations to test how the binding energies will be affected upon bacteriocins binding to delta variant (L452R-T478K double mutant). Similar to beta variant docking results, pediocin PA-1 was found as the tightest binder compared to all studied bacteriocins (see Table S4). Therefore, it is evident that both probiotic microorganisms and their bacteriocins can be considered potential preventive therapeutics or dietary supplements to enhance the immune system for the prophylaxis of viral diseases, including COVID-19.

Conclusions

In the present study, molecular docking studies of 22 biopeptide bacteriocins that were performed against RBD of S protein for WT and beta variant revealed that pediocin PA-1 and salivaricin P had the highest binding energies to WT as well as mutants. Interestingly, salivaricin B also showed a strong affinity to the mutant RBDs of S protein in spite of its relatively low binding score towards WT RBD. The MD simulations confirmed the stability of bacteriocin-RBD complexes and indicated the key interactions between bacteriocins and RBDs. Mainly, pediocin PA-1 interacted with three important amino acid residues of RBD, which are among the key residues in the binding of RBD with ACE2. It is therefore clear that three bacteriocins (pediocin PA-1, salivaricin P, and salivaricin B) are the most promising therapeutic candidates with significant in silico antiviral activity against SARS-CoV-2, including beta variant. Our findings can provide a lead in further exploring bacteriocins in the treatment and prophylaxis of COVID-19. The mutations in the spike protein are evolutionarily selected for their affinity to ACE-2 and to circumvent vaccination efforts. The bacteriocins are perhaps not part of the evolutionary forces. Therefore, it is more likely that the mutation will not impair bacteriocin-S protein interaction. Although only the beta variant was studied in this work, the in silico methodology is also suitable to delta and other possible new variants of the SARS-CoV-2. By changing the amino acid sequence of the new variants on the spike protein, new structures can be prepared and used in docking and molecular dynamics simulations. By doing so, the affinity predictions of bacteriocin-S protein profiles can be predicted.
Data Availability

The numerical calculations reported in this paper were par-tially material available at https://

Acknowledgements

Supplementary Information

Data Availability

All the results are in sole responsibility of the authors. The numerical calculations reported in this paper were par-
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Acknowledgements

This work was supported by TUBITAK, 2232 — International Fellowship for Outstanding Researchers, Project number 11C244.

Declarations

Conflict of Interest

The authors declare no competing interests.

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1007/s12602-021-09879-0.

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