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# The Identification of a Novel Peptide Derived from Lactoferrin Isolated from Camel Milk with Potential Antimicrobial Activity

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## ABSTRACT

Background and Aim: Antimicrobial peptides have attracted significant attention in recent decades because of their properties, such as rapid bactericidal effects, having a wide spectrum of activity, and a rare development of drug resistance. The purpose of this study was to examine the antibacterial activity of a peptide derived from the lactoferrin isolated from camel milk against *Staphylococcus aureus*, *Pseudomonas aeruginosa*, and *Acinetobacter baumannii*.

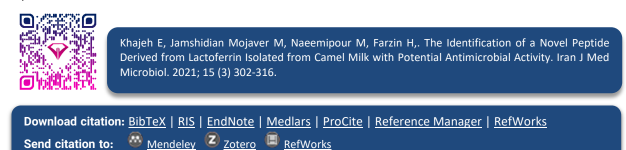
Materials and Methods: In the present study, by means of bioinformatics, an antibacterial peptide was potentially identified as candidates in lactoferrin of camel milk, and an appropriate peptide was selected based on defined criteria. The Pepsin-Camel-Lac1 peptide was synthesized. The methyl thiazolyl diphenyl-tetrazolium bromide assay was conducted to examine the toxicity of Pepsin-Camel-Lac1 against a cell line. Three pathogenic bacteria, namely *S. aureus*, *P. aeruginosa*, and *A. baumannii* were analyzed to assess the antibacterial activity of Pepsin-Camel-Lac1 peptide.

**Results:** The results showed that the newly-identified peptide had no toxicity against the cell line. The minimum inhibitory concentration values of Pepsin-Camel-Lac1 against *S. aureus, P. aeruginosa,* and *A. baumannii* were 31.25  $\mu$ g/mL, 31.25  $\mu$ g/mL, and 62.5  $\mu$ g/mL, respectively.

Conclusion: It seems that the growth of *S. aureus, P. aeruginosa,* and *A. baumannii* was not affected by Pepsin-Camel-Lac1 treatment in the bacterial culture medium.

Keywords: baumannii	Milk proteir	ns; Pepsin; Antibact	erial activity; Staphylococcu	s aureus; Pseudomonas aeruginosa; Acinetoba	cter		
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#### Introduction

Milk contains a large number of proteins, some of which have been well characterized, such as lactoferrin, which exhibits antibacterial activity (1). Two iron-binding proteins, namely lactoferrin, and transferrin, have been shown to have biological activity in the milk of many species (2). The two proteins almost possess similar properties, such as molecular weight (80 kDa) and the ability to bind to iron (3). Lactoferrin is a monomeric glycoprotein with a high affinity to iron (4). The presence of lactoferrin has been confirmed in a large number of mammals, and the amino acid sequence of this protein has been identified in human, pig, horse, camel, cattle, buffalo, goat, and mouse (5). The third structure of lactoferrin has been characterized in the five species of human, cow, buffalo, camel, and horse, all of which have the same structures (6-10). All nutrients found in human milk are also present in camel milk. A number of investigations have recently reported the therapeutic properties of camel milk, such as anti-cancer, antidiabetic, anti-microbial potential (11-14). Most secretions fluids, such as milk, saliva, tears, and neutrophil granules, contain this glycoprotein (15). In the immune system, lactoferrin is introduced as a part of the innate immune system because of its sensitive position on the mucosal surface as one of the agents existing at the first-line of defense to prevent the entry of microbial agents (16). Camel milk lactoferrin has been identified as a factor against Gram-positive and Gram-negative bacteria, as well as fungi and parasites (17). Proteins and peptides of the milk, such as lactoferrin, lactoperoxidase, and lysozyme, play an influential role in overall antimicrobial activity in milk (18). The milk proteins and peptides can be natural substitutes for antibiotics due to their antibacterial properties (19). Antimicrobial peptides (AMPs) are oligopeptides with varied numbers of amino acids (from 5 to more than 100) (20, 21). Some studies have analyzed the antimicrobial activity of peptides derived from lactoferrin (22-26). Camel milk lactoferrin has antibacterial activity against Escherichia coli, Salmonella typhimurium, Listeria monocytogenes, Staphylococcus aureus and Pseudomonas aeruginosa (11, 27-29). Thanks to bioinformatics tools and knowing the cleavage sites of enzymes, the identification of AMPs in natural sources, such as milk, has been feasible (30-32). Different methods are used to characterize and produce active AMPs (33). These methods are based on proteolysis using bioinformatics software and statistical analyses to assess the role of peptides released from cleaved proteins. In this study, in order to find a novel AMP, different AMPs, derived from camel milk lactoferrin, were designed by bioinformatics approaches, and then, the most appropriate AMP was chosen according to the required criteria. The designed peptide was synthesized and

analyzed in terms of the antibacterial activity against *P. aeruginosa, A. baumannii,* and *S. aureus*.

#### **Materials and Methods**

#### **Bioinformatics Analysis**

#### Selection of Effective Antimicrobial Peptide Derived from Lactoferrin

Camel milk lactoferrin is a glycoprotein that contains 708 amino acids and possesses two domains and two iron-binding sites (34). The strategy used for AMP synthesis from camel milk lactoferrin was in accordance with bioinformatics approaches of a study conducted by Dziuba et al. (32). Based on the cleavage sites identified for various proteolytic enzymes, insilico studies indicated that pepsin is capable of releasing numerous active peptides from lactoferrin (32, 35, 36). The initial structure of camel milk lactoferrin was obtained from the Uniport database with an ID of Q9TUM0. Next, pepsin was used to hydrolyze the protein, as this enzyme (pH>2) has proteolytic activity and is commonly applied to obtain bioactive peptides (37-39). Four online software tools, namely BIOPEP database, Collection of Antimicrobial Peptides Server, Antimicrobial Peptide Calculator and Predictor and Peptide cutter database were utilized to evaluate the antimicrobial properties of cleaved peptides derived from lactoferrin (40-43). The online server BIOPEP-UWM (www.uwm.edu.pl/biochemia) was utilized since it contains a library of bioactive peptides and comprises of the two essential coordinated parts, namely grouping databases and apparatuses for the assessment of proteins as the antecedents of bioactive peptides, which includes the proteolytic procedure plans. Another online database, CAMP (http://www.camp.bicnirrh. res.in), contains information about the conserved sequences, representing some examples and Hidden Markov Models (HMMs) of 1386 AMPs that are categorized into 45 families. On the other hand, the online database APD2 (http:-//aps.unmc.edu/AP/expectation/forecastmain.php) contains 2684 antimicrobial peptides from six kingdoms (266 from microbes, 4 from archaea, 8 from protists, 13 from parasites, 329 from plants, and 2018 from creatures) (44).

#### **Experimental Procedures**

#### Material

*S. aureus* (1074, ATCC: 6538), *P. aeruginosa* (1060, ATTCC: 27853), and *A. baumannii* (1003, ATCC: BAA-747) were purchased from the Pasteur Institute of Iran.

#### Peptide Synthesis

The chosen peptide was named Pepsin-Camel-Lac1 (PCL1) synthesized by MIMOTOPES Pty Ltd, Australia.

In order to prepare the designed peptide, 1 mg of the designed peptide was dissolved in 1 mL of sterile water and used for the experiments.

#### **Toxicity Assay**

The Methyl thiazolyl diphenyl-tetrazolium bromide (MTT) assay was utilized to assess the toxicity of PCL1 against a human cell line. The assay is based on the decrease of the tetrazolium salt into blue formazan crystals in viable cells (45, 46). The cell line was cultured in the RPMI 1640 medium supplemented with 10% Fetal Bovine Serum (FBS), 100 ml penicillin, and 100 mg/mL streptomycin. During the cell culture period, cells were incubated at 37°C in a 95% atmosphere and 5% CO<sub>2</sub>. When the cells reached 80-90% confluence, the viability of cells was examined and passaged. In this experiment, 100 µL of the cell suspension was added to each well of a 96-well plate, each containing 100 mL of the culture medium. The cells were treated with the pre-defined concentrations of PCL-1 and then incubated for 24 h. After dilution of PCL-1 with sterile water, cells were treated with PCL-1, and then 20 µL of the MTT solution (5 mg/mL) was added to the wells and incubated for 24 hours. After the incubation period, 50 µL of dimethyl sulfoxide (DMSO) was added to the wells to dissolve the purple formazan crystals. Afterward, according to the obtained results, the inhibitory concentration (IC50) of PCL-1 was calculated, in which 50% of the cells underwent cell death. The experiments have been carried out in triplicates, and the values were represented as the percentage of cell viability in comparison with the control cells.

#### Preparation of Microbial Suspension Solution

The lyophilized batches containing bacteria mentioned earlier were transferred to the liquid Mueller Hinton Broth (MHB) and incubated at 37°C for 24 h. In order to determine the antimicrobial effects, a fresh 24-hour culture medium was prepared, then 1 ml of the 24 h microbial suspension was transferred into a tube containing the MHB medium, and the turbidity of the prepared microbial suspension was assessed by McFarland standards using a spectrophotometer at a wavelength of 625 nm. The final suspension was used for the evaluation of drug susceptibility testing, MIC, and cell culture (47).

#### Determination of MIC and MBC Values by Micro-Dilution Method

The MHB medium was used to evaluate the antimicrobial effects of PCL-1 by minimum inhibitory concentration (MIC) and minimum bactericidal

concentration (MBC) methods. In general, a sterile flat 96-well plate was used after determining the half-McFarland turbidity. In this method, 50mL of MHB medium was added to well 2 to well 11. Then,  $50\mu$ L of the PCL-1 peptide was added to well 2 to well 9. Next,  $50 \mu$ L of the microbial suspension was added to well 1 well 10. Afterward, the plate was incubated for 24 h. Well 1 that contains bacteria were considered as the negative control, while and well 11 containing the MHB medium was considered the positive controls. In order to assess the MBC values, the contents of the wells were cultured on the Mueller Hinton Agar medium. Lack of bacterial growth is indicative of MBC determination (47, 48).

#### Agar Well Diffusion Assay

The antimicrobial activity test was performed using the Agar well diffusion assay (49). This technique is broadly used to assess antimicrobial activity. In this method, the microbial suspension is distributed on the plate surface. Then, a hole is created with a diameter of 6-8 mm in sterile conditions, and then specific concentrations of the PCL-1 peptide were added to the hole and left to release into the pores of Agar. Different concentrations of PCL-1 were prepared as follows; PCL-1 was transferred into 5 sterile microtubes, then 100 mL of sterile water was added into microtube 2 to 5. Next, 100 mL of PCL-1 was added to microtube 1 and was serially transferred into other microtubes until the last microtube. After that, 100µL of the contents of each microtube was isolated and added to the plate agar. After incubating the plates at 37°C for 24h, the results were examined based on the diffused area around the created hole (47).

#### Results

#### **In-Silico Proteolysis of Lactoferrin**

The sequence of amino acids of lactoferrin was exposed in in-silico proteolysis performed by the BIOPEP database. In-silico proteolysis can be carried out by means of some bioinformatics tools, as we employed the peptide cutter database (https://-web.expasy.org/peptide cutter/.) (42, 50). For in-silico protein hydrolysis, a single enzyme was used and the PCL-1 peptide cleavage site was bolded (Table 1). The pepsin enzyme (pH>2) was utilized to create different peptides from lactoferrin, resulted in the production of 50 peptides with the length of 5-21 amino acids. For the statistical prediction of antimicrobial properties, the produced peptides were submitted (Table S1). A short linear peptide, PCL1, met the required criteria, as shown in Figure 1.

 Table 1. Results of peptide released from BIOEP database camel milk lactoferrin.

Enzyme	No. of cleavages	Positions of cleavage sites
<u>Pepsin (pH&gt;2)</u>	188	2 3 5 8 9 10 11 13 14 15 16 17 18 27 40 41 62 77 78 81 82 83 84 87 90 92 100 101 110 111 122 123 124 125 127 128 130 131 138 143 144 149 150 153 155 156 157 163 170 171 188 190 191 207 209 210 211 214 215 217 226 227 233 234 237 246 247 248 265 266 285 286 287 288 289 296 297 304 305 306 307 308 316 318 319 323 324 325 326 327 336 337 338 339 342 343 349 358 359 365 366 374 375 379 380 401 402 403 404 410 411 412 413 416 417 418 419 430 441 442 451 453 467 469 470 485 486 492 494 502 504 505 506 520 522 523 542 544 545 548 549 551 560 561 567 568 578 579 582 584 585 587 588 589 590 591 592 593 608 625 626 629 630 631 636 637 647 648 650 651 657 659 660 666 667 669 670 676 678 679 684 690 691 698 700 704 705 706

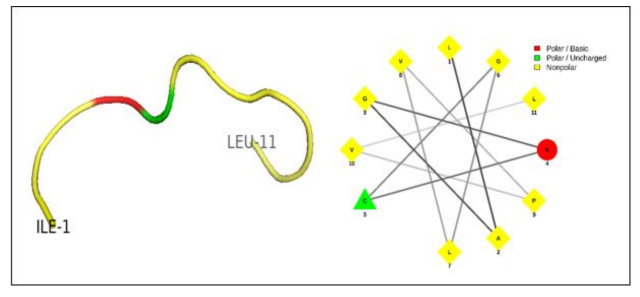


Figure 1. The PCL1 peptide picture consisted of 11 amino acids and has a liner structure.

#### **Physicochemical Properties of AMPs**

The physicochemical properties of antimicrobial peptides designed by the BIOPEP database were characterized using free web tools and the algorithms for the chosen AMP are shown (<u>Table S1</u>). The molecular weights of created peptides were in a range of 446 to 2493 Da (<u>Table S1</u>). Among 50 peptides created, 26 peptides had a pl value of less than 7, while the rest has above 7. It is now known that the molecular charge and hydrophobicity play a significant role in the functionality of AMPs (**51-53**). Among 50 peptides derived from lactoferrin, 14 peptides had a negative charge, 25 possess a positive charge, and 11 were neutral. In the present study, only positive AMPs were selected for further analyses. Because cationic peptides constitute a significant percentage of AMPs,

as a result of their higher affinity to bind to negatively charged cell membranes of bacteria and eventually disrupt the bilayer lipid structure (54, 55). Amphipathic property is another marked parameter to choose efficient AMPs since these structures allow them to bind to hydrophilic regions; hence AMPs with both hydrophilic and hydrophobic domains have a higher priority to be used for antibacterial tests (56). The amino acid composition also plays a role in designing the antibacterial peptides. Gram-positive and Gram-negative antimicrobial peptides have a similar ratio of cysteine and lysine residues (57). As shown in Table 1, the PCL1 peptide composed of 63% hydrophobic region, 27% hydrophilic region, and 9% charged amino acids; Also, it contains 18% lysine and 9% cysteine.

 Table 2. Physical and chemical characteristics of the Pepsin-Camel-Lac1 peptide.

Position of cleavage site	Enzyme	Peptide sequence	Peptide length [aa]	Peptide mass [Da]	Seq. ID	Charge	pl
430	Pepsin (pH>2)	>30 IAGKCGLVPVL	11	1069.371	30	8.22	+1

Sequence ID	Name of cleaving enzyme(s)	Resulting peptide sequence	Peptide length [aa]	Peptide mass [Da]	Position of cleavage site	pl	Charge
1	Pepsin (pH>2)	>1 AASKKSVRW	9	1032.211	27	11.12	+3
2	Pepsin (pH>2)	>2 CTTSPAESSKCAQ	13	1312.432	40	6.25	0
3	Pepsin (pH>2)	>3 QRRMKKVRGPSVTCVKKTSRF	21	2493.033	62	12.2	+8
4	Pepsin (pH>2)	>4 ECIQAISTEKADAVT	15	1578.755	77	4.14	-2
5	Pepsin (pH>2)	>5 LRPIAAEV	8	868.044	100	6.25	0
6	Pepsin (pH>2)	>6 GTENNPQTH	9	996.989	110	5.5	-1
7	Pepsin (pH>2)	>7 YAVAIAKKGTN	11	1135.329	122	9.72	+2
8	Pepsin (pH>2)	>8 KSCHTGL	7	744.864	138	8.22	+1
9	Pepsin (pH>2)	>9 GRSAG	5	446.464	143	10	+1
10	Pepsin (pH>2)	>10 NIPMG	5	530.640	149	5.5	0
11	Pepsin (pH>2)	>11 TGPPEP	6	596.638	163	3.81	-1
12	Pepsin (pH>2)	>12 LQKAVAK	7	756.944	170	10	+2
13	Pepsin (pH>2)	>13 FSASCVPCVDGKEYPNL	17	1829.073	188	4.47	-1
14	Pepsin (pH>2)	>14 CAGTGENKCACSSQEP	16	1584.711	207	10	-1
15	Pepsin (pH>2)	>15 LQDGAGDVA	9	844.877	226	3.53	-2
16	Pepsin (pH>2)	>16 VKDSTV	6	647.726	233	6.25	0
17	Pepsin (pH>2)	>17 PAKADRDQY	9	1063.135	246	6.25	0
18	Pepsin (pH>2)	>18 LCPNNTRKPVDASQECH	17	1912.125	265	7.09	0
19	Pepsin (pH>2)	>19 ARVPSHAVVARSVNGKEDL	19	2005.265	285	9.06	+1
20	Pepsin (pH>2)	>20 LVKAQEK	7	814.980	296	8.88	+1
21	Pepsin (pH>2)	>21 GRGKPSA	7	671.754	304	11.12	+2
22	Pepsin (pH>2)	>22 GSPAGQKD	8	758.786	316	6.25	0
23	Pepsin (pH>2)	>23 RIPSKIDSG	9	972.109	336	8.88	+1
24	Pepsin (pH>2)	>24 ITAIRG	6	629.757	349	10	+1
25	Pepsin (pH>2)	>25 LRETAAEVE	9	1017.104	358	4.14	-2
26	Pepsin (pH>2)	>26 RRAQVV	6	727.865	365	11.97	+2
27	Pepsin (pH>2)	>27 RRAQVV	8	807.830	374	11.97	+2
28	Pepsin (pH>2)	>28 SRQSNQSVVCATASTTEDCIA	21	2171.339	401	4.47	-1
29	Pepsin (pH>2)	>29 KGEADA	6	589.603	410	4.47	-1
30	Pepsin (pH>2)	>30 IAGKCGLVPVL	11	1069.371	430	8.22	+1
31	Pepsin (pH>2)	>31 AESQQSPESSG	11	1106.068	441	3.53	-2
32	Pepsin (pH>2)	>32 DCVHRPVKG	9	1010.179	451	8.22	+1

#### Table S 1. Physical and chemical characteristics of the synthesized peptides.

Sequence ID	Name of cleaving enzyme(s)	Resulting peptide sequence	Peptide length [aa]	Peptide mass [Da]	Position of cleavage site	pl	Charge
33	Pepsin (pH>2)	>33 AVAVVRKANDKITW	14	1570.855	467	10	+2
34	Pepsin (pH>2)	>34 RGKKSCHTAVDRTAG	15	1586.789	485	10	+3
35	Pepsin (pH>2)	>35 NIPMGP	6	627.756	492	5.5	0
36	Pepsin (pH>2)	>36 KNTDSCRF	8	970.068	502	8.22	+1
37	Pepsin (pH>2)	>37 SQSCAPGSDPRSKL	14	1432.571	520	8.22	+1
38	Pepsin (pH>2)	>38 CAGNEEGQNKCVPNSSERY	19	2085.207	542	4.7	-1
39	Pepsin (pH>2)	>39 LAENVGDVA	9	886.957	560	3.53	-2
40	Pepsin (pH>2)	>40 VKDVTV	6	659.781	567	6.25	0
41	Pepsin (pH>2)	>41 DNTDGKNTEQ	10	1121.082	578	4.14	-2
42	Pepsin (pH>2)	>42 NGTRKPVTEAESCHL	15	1641.819	608	7.19	0
43	Pepsin (pH>2)	>43 PVAPNHAVVSRIDKVAH	17	1810.089	625	9.06	+1
44	Pepsin (pH>2)	>44 GRNGQDCPGK	10	1031.111	647	8.22	+1
45	Pepsin (pH>2)	>45 QSKTKN	6	704.781	657	10	+2
46	Pepsin (pH>2)	>46 NDNTEC	6	694.671	666	3.53	-2
47	Pepsin (pH>2)	>47 QGKTTY	6	696.758	676	8.88	+1
48	Pepsin (pH>2)	>48 LGPQY	5	576.650	684	5.5	0
49	Pepsin (pH>2)	>49 VTAIAK	6	601.744	690	8.88	+1
50	Pepsin (pH>2)	>50 RRCSTSP	7	805.907	698	10	+2

#### Table S 2. Predict Antimicrobial Peptides.

	Results with Support Vector Machine (SVM) classifier				Results with Artificial Neural Network (ANN) classifier	Results with Discriminant Analysis classifier	
Seq. ID.	Class	AMP Probability	Class	AMP Probability	Class	Class	AMP Probability
1	AMP	0.941	AMP	0.584	AMP	NAMP	0.094
2	NAMP	0.016	NAMP	0.3475	NAMP	NAMP	0.029
3	NAMP	0.038	AMP	0.6	NAMP	NAMP	0.421
4	NAMP	0.095	NAMP	0.2235	NAMP	NAMP	0.004
5	NAMP	0.004	NAMP	0.331	NAMP	NAMP	0.003
6	NAMP	0.138	AMP	0.603	NAMP	NAMP	0.004
7	AMP	0.595	NAMP	0.215	AMP	AMP	0.522
8	AMP	0.714	NAMP	0.332	NAMP	NAMP	0.024
9	NAMP	0.000	NAMP	0.289	NAMP	NAMP	0.000
10	NAMP	0.002	NAMP	0.347	AMP	NAMP	0.001
11	AMP	0.996	AMP	0.691	NAMP	NAMP	0.000
12	NAMP	0.239	NAMP	0.374	AMP	NAMP	0.147
13	NAMP	0.144	NAMP	0.145	NAMP	NAMP	0.033
14	NAMP	0.230	NAMP	0.298	NAMP	NAMP	0.056

	Results with Support Vector Machine (SVM) classifier				Results with Artificial Neural Network (ANN) classifier	Results with Discriminant Analysis classifier	
15	AMP	0.929	NAMP	0.354	NAMP	NAMP	0.001
16	NAMP	0.008	NAMP	0.347	NAMP	NAMP	0.000
17	NAMP	0.011	AMP	0.5275	NAMP	NAMP	0.001
18	NAMP	0.063	NAMP	0.083	NAMP	NAMP	0.004
19	NAMP	0.345	NAMP	0.17	NAMP	NAMP	0.036
20	NAMP	0.002	NAMP	0.297	NAMP	NAMP	0.004
21	NAMP	0.001	NAMP	0.3515	NAMP	NAMP	0.001
22	NAMP	0.000	NAMP	0.3385	NAMP	NAMP	0.000
23	NAMP	0.093	NAMP	0.265	NAMP	NAMP	0.009
24	NAMP	0.000	NAMP	0.4965	AMP	NAMP	0.263
25	NAMP	0.001	NAMP	0.2205	NAMP	NAMP	0.000
26	NAMP	0.008	NAMP	0.455	NAMP	NAMP	0.019
27	NAMP	0.025	NAMP	0.378	NAMP	NAMP	0.000
28	NAMP	0.033	NAMP	0.1255	NAMP	NAMP	0.002
29	NAMP	0.216	NAMP	0.455	NAMP	NAMP	0.000
30	NAMP	0.146	AMP	0.5805	AMP	AMP	0.949
31	NAMP	0.387	NAMP	0.364	NAMP	NAMP	0.000
32	NAMP	0.003	NAMP	0.3025	AMP	NAMP	0.000
33	NAMP	0.399	AMP	0.567	AMP	NAMP	0.326
34	NAMP	0.188	NAMP	0.186	NAMP	NAMP	0.003
35	NAMP	0.005	NAMP	0.4345	AMP	NAMP	0.003
36	NAMP	0.004	NAMP	0.3385	NAMP	NAMP	0.187
37	NAMP	0.008	NAMP	0.1685	NAMP	NAMP	0.003
38	NAMP	0.178	NAMP	0.2445	NAMP	NAMP	0.039
39	AMP	0.593	NAMP	0.3495	NAMP	NAMP	0.001
40	NAMP	0.007	NAMP	0.443	NAMP	NAMP	0.000
41	NAMP	0.026	AMP	0.627	NAMP	NAMP	0.000
42	NAMP	0.060	NAMP	0.0225	NAMP	NAMP	0.001
43	NAMP	0.256	NAMP	0.0655	NAMP	NAMP	0.014
44	NAMP	0.002	NAMP	0.2105	NAMP	NAMP	0.001
45	AMP	0.999	AMP	0.501	AMP	NAMP	0.063
46	AMP	0.984	NAMP	0.336	NAMP	NAMP	0.017
47	AMP	0.760	NAMP	0.3345	NAMP	NAMP	0.002
48	NAMP	0.052	NAMP	0.294	AMP	NAMP	0.041
49	NAMP	0.000	AMP	0.503	AMP	NAMP	0.169
50	AMP	0.999	NAMP	0.4025	NAMP	NAMP	0.086

#### Prediction of the Antimicrobial Activity of Produced Peptides During In-Silico Proteolysis from Lactoferrin

The in-silico proteolysis of lactoferrin leads to the generation of short peptides. Among the created peptides, the PCL-1 peptide shows anti-bacterial, antiviral, and anti-tumor activity due to its high positive charge and ability to interact with negatively charged substrates, such as glycosaminoglycan, lipopolysaccharide, heparan, phosphatidylserine, and nucleic acids (58, 59). The design of peptides varies depending on the biological activity of these macromolecules (22). Many peptides have a wide range of activity, such as the antibacterial and anti-cancer properties (60). For example, the design of AMPs derived from the lactoferrin protein has been conducted using the CAMP tools. To this aim, Four software tools have been employed, namely Random Forest, Support Vector Machines, Artificial Neural Network, and Discriminant Analysis (41). In this analysis, the resultant score was in excess of 0.45; therefore, the PCL1 peptide was listed as AMPs, and a positive classification was obtained for at least two statistical methods (Table 3).

 Table 3. The characteristics of PCL1 peptide released during in-silico proteolysis and prediction.

	SVM classifier		RF C	Classifier	ANN classifier	DA classifier	
Sequence	Class	AMP Probability	Class	AMP Probability	Class	Class	AMP Probability
IAGKCGLVPVL	NAMP	0.146	AMP	0.5805	AMP	AMP	0.949

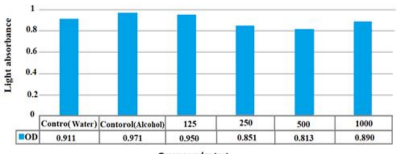
Four statistical models in the CAMP database: SVM, RF, ANN and DA classiers that used for proteolysis and prediction peptides.

#### **Toxicity Assay**

In order to achieve a linear regression equation of the concentrations of PCL-1 versus the growth inhibition, Microsoft Excel software was used, and the values were expressed as the means and standard deviation of the means of three separate experiments. The below equation was used to measure the growth curve and draw a plot:

Growth inhibition= (control OD-sample OD)/control OD\*100 (61)

Finally, the IC50 values of specimens, representing a concentration of the peptide by which 50% of bacteria are eliminated, were obtained from the growth curve. The SPSS software version 16 was used for the analysis of the obtained data, and the difference between various treatments was analyzed by one-way analysis of variance (one-way ANOVA) followed by Tukey's post hoc test (Table 4). The MTT results showed that PCL-1 has no significant effect on the viability of the cell line used (Figure 2).



Groups under test

Figure 2. MTT cell line with PCL1 peptide with sig= 0.00

Table 4. Results of analysis using SPSS technology of mixed PCL1 peptide levels on cell lines.

	Sum of Squares	df	Mean Square	F	Sig
Between Groups	1.500	5	0.3	8.498	0.001
Within Groups	0.424	12	0.035		
Total	1.924	17			

#### Determination of MIC and MBC Values by Micro-Dilution Method

The MIC values of bacterial strains were calculated by the micro-dilution method. After two hours of incubation, the turbidity of samples was read at wavelengths of 490 and 630nm using an ELISA reader. The well in which bacterial growth inhibition occurred was regarded as the MIC values and used to determine the MBC value. Following the incubation period, the first dilution in which bacteria did not grow on the agar plate was regarded as the MBC value. According to Figure 3, the MIC values for *S. aureus* in response to PCL-1 was reported to be 31.25µg/mL, while the MBC value was 500µg/mL (Figure 3A).

The MIC and MBC values for *P. aeruginosa* were  $31.25 \ \mu\text{g/mL}$  and  $125 \ \mu\text{g/mL}$ , respectively (Figure 3B). In addition, the values of MIC and MBC for *A. baumannii* were  $62.5 \ \mu\text{g/mL}$  and  $500 \ \mu\text{g/mL}$ , respectively (Figure 3C).

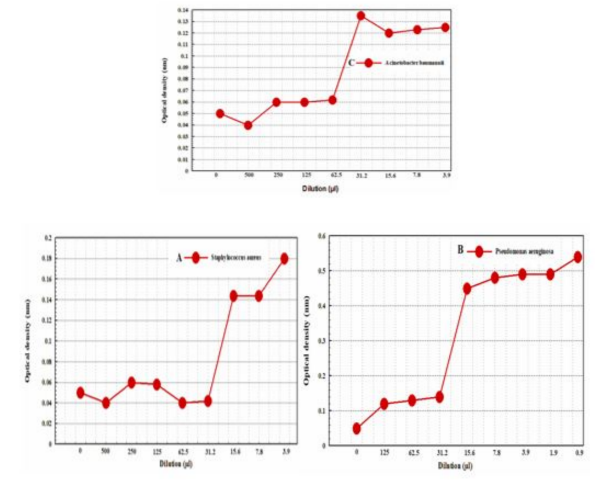


Figure 3. MIC and MBC analyze outcomes for the three bacterial strains that were tested in A, B, and C for S.Aureus, P.Aeruginosa, and A.Baumannii.

#### Agar Well Diffusion Assay

The diameter created around the holes in agar indicates the effect of PCL-1 on the inhibition growth

of bacterial strains. The results demonstrated that PCL-1 had no significant effect on the growth inhibition of the three bacterial strains when compared with the control group (Figure 4).

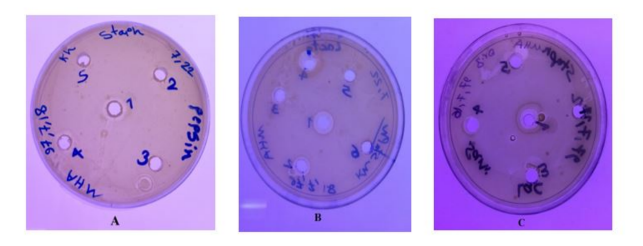


Figure 4. Results of agar well diffusion assay for Staphylococcus Aureus, Pseudomonas Aeruginosa, and Acinetobacter Baumannii are respectively in A, B, and C.

## Discussion

In an in-silico proteolysis conducted by Dziuba et al., 11 new peptides with potential antimicrobial activity were picked from all the digested peptides in the milk protein (32). Jahani et al. demonstrated that lactoferrin to be effective as an antibacterial against Gram-positive (Staphylococcus epidermidis, Bacillus cereus) and Gram-negative (Campylobacter jejuni, Salmonella spp.) bacteria. However, it was more effective on the Gram-positive instead of the Gramnegative bacteria (62). In line with the investigation of peptides derived from milk lactoferrin through the milk fermentation and assessment of their antibacterial effects. Korheonan et al. reported that milk proteins contain a rich source of peptides that could be potentially released and activated. For example, when these peptides are activated during gastricintestinal digestion or milk fermentation, they could be involved in modulatory processes in living systems (33). Bushra Niaz et al. reported that due to the association of lactoferrin with enterobacterial lipopolysaccharide (LPS), it acts as a permeabilizing agent of Gram-negative bacteria and destabilizes the bacterial membrane and thus enhance bacterial permeability. Lactoferrin is not an effective antibacterial agent of its own, but allows those peptides that have been derived from it, whose hydrophobicity impose a limit on their effectiveness, to enter the bacterial membrane as an antibacterial agent.(63). As a result of direct interaction between protein or lactoferrin-derived peptides, Farnaud and Evans identified that it possesses bactericidal activities (25). Cameleers believe that camel milk is more resistant to spoilage than the milk of other animals, and it has a longer shelf life. The effect of lactoperoxidase can be one of the significant factors leading to camel milk resistance to spoilage. In this work, antimicrobial peptides were identified from camel milk lactoferrin using bioinformatics; then, the PCL-1 peptide was

selected, and the activity of this peptide was evaluated in-vitro. In the first step, the hydrolysis of lactoferrin was performed using the pepsin enzyme using the BIOPEP database by which 50 peptides were produced containing 5-21 amino acids. The physicochemical properties of the synthesized peptides were investigated using online tools, such as the APD2 and Peptide cutter databases, to select appropriate AMPs. The general properties of the generated peptides, including molecular weight, charge, and PI values, were analyzed to select the appropriate antimicrobial peptides (Table S1). Among the created peptides, two peptides met the initial criteria, but since the presence of lysine and cysteine amino acids in AMPs is an advantage, the peptide with an ID of 30 (PCL-1 peptide) was selected for further analyses (Table 1). Four tools of the CAMP database (ANN, DA, RF, SVM) were employed to determine the antibacterial activity of the created peptides in this study. The PCL-1 peptide was classified as AMPs since it had parameters with scores of higher than 0.45, including RF: 0.5805 and DA: 0.949 (Table 3). After synthesizing the PCL-1 peptide, the viability of bacteria in response to PCL-1, MBC, MIC, agar well diffusion was assessed in order to determine the antimicrobial activity, as well as toxicity against the human cell line. The MTT assay showed that this peptide had no toxicity against the human cell line. Our findings also showed the MIC and MBC values for S. aureus, P. aeruginosa, and A. baumannii were 31.25/500, 21.25/125, and 62.5/500, respectively. The agar well diffusion assay showed that the analyzed bacteria were resistant to the PCL-1 peptide. The experimental results of this study showed that the PCL-1 peptide had no antimicrobial properties.

## Conclusion

There are naturally occurring bioactive peptides in fermented dairy products, such as yogurt, butter, milk, and cheese, which are inactivated in primary sources, and they could be activated by digestion of milk, milk fermentation, exposure to proteolytic starter media, or hydrolysis using proteolytic enzymes. Milk-derived peptides exhibit super-beneficial activity, even through oral consumption. In this study, pepsindigested peptides in milk lactoferrin were identified by bioinformatics tools. After the synthesis of PCL-1, the MTT assay was applied to assess the toxicity of the selected peptide against the human cell line. The results showed that the chosen peptide had no toxicity against the human cell line or bacterial strains.

## **Abbreviations**

Antimicrobial peptides: AMPs, Collection of Antimicrobial Peptides: CAMP, Antimicrobial Peptide Calculator and Predictor: APD2, Pepsin-Camel-Lac1: PCL1, Methyl thiazolyl diphenyl-tetrazolium bromide: MTT, Mueller Hinton Broth: MHB, Minimum inhibitory

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concentration: MIC, Minimum bactericidal concentration: MBC, Agar well diffusion assay: AWDA, Random Forest: RF, Support Vector Machines: SVM, Artificial Neural Network: ANN, Discriminant Analysis: DA.

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## **Ethics Approval and Consent to Participate**

This article does not contain any studies with human participants or animals performed by any of the authors.

## **Consent for Publication**

All authors gave their consent for publication.

## **Conflict of Interest**

The authors declared no conflict of interest.

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# شىناسايى يك پپتيد جديد بەدستآمدە از لاكتوفرين جداشدە از شير شتر با فعاليت بالقوه

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# چکيده

**زمینه و اهداف**: پپتیدهای ضد میکروبی در دهههای اخیر به دلیل برخی خاصیتها از جمله اثرات سریع ضد باکتری، داشتن طیف وسیعی از فعالیت و توسعه نادر مقاومت دارویی، توجه قابل توجهی را به خود جلب کردهاند. بررسی فعالیت ضد باکتریایی یک پپتید مشتقشده از لاکتوفرین از شیر شتر علیه *استافیلوکوکوس اورئوس، سودوموناس آئروژینوزا و آسینتوباکتر* بومانی هدف این مطالعه بود.

مواد و روش کار: در مطالعه حاضر، با استفاده از بیوانفورماتیک، چندین پپتید ضد باکتری به طور بالقوه بهعنوان کاندیداهای لاکتوفرین شیر شتر شناسایی شدند؛ سپس یک پپتید مناسب بر اساس معیارهای بیوانفورماتیک تعریف شده، انتخاب شد. پپتید انتخابی Pepsin-Camel-Lacl سنتز شد. سپس با روش MTT خاصیت سمی پپتید بر ردهٔ سلولی بررسی گردید. سه باکتری بیماریزا، یعنی استافیلوکوکوس اورئوس، سودوموناس آئروژینوزا و اسینتوباکتر آسینتوباکتر بومانی، برای ارزیابی فعالیت ضد باکتریایی پپتید Pepsin-Camel-Lacl تجزیه و تحلیل شدند.

یافتهها: نتایج نشان داد که پپتید تازهشناسایی شده فاقد خاصیت سمیت در برابر ردهٔ سلولی است. نتایج MIC مربوط به پپتید پپسین برای باکتری *استافیلو کو کوس اورئوس، سودوموناس آئروژینوزا* و آسینتوب*اکتر بومانی* به ترتیب ۳۱.۲۵، ۳۱.۲۵ و ۶۲/۵ بود.

**نتیجهگیری**: به نظر م<sub>یار</sub>سد که رشد *استافیلوکوکوس اورئوس، سودوموناس آئروژینوزا* و *اسینتوباکتر بومانی* تحت تأثیر درمان Pepsin-Camel-Lac1 در محیط کشت باکتریها قرار نگرفت.

کلید واژهها: پروتئینهای شیر، پپسین، فعالیت ضد باکتریایی، *استافیلوکوکوس اورئوس، سودوموناس آئروژینوزا،* آسینتوباکتر بومانی

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