Design of a RANK-Mimetic Peptide Inhibitor of Osteoclastogenesis with Enhanced RANKL-Binding Affinity

Jeonghwan Hur, Ambarnil Ghosh, Kabsun Kim, Hai Minh Ta, Hyunju Kim, Nacksung Kim, Hye-Yeon Hwang, and Kyeong Kyu Kim

The receptor activator of nuclear factor κB (RANK) and its ligand RANKL are key regulators of osteoclastogenesis and well-recognized targets in developing treatments for bone disorders associated with excessive bone resorption, such as osteoporosis. Our previous work on the structure of the RANK-RANKL complex revealed that Loop3 of RANK, specifically the non-canonical disulfide bond at the tip, performs a crucial role in specific recognition of RANKL. It also demonstrated that peptide mimics of Loop3 were capable of interfering with the function of RANK in osteoclastogenesis. Here, we reported the structure-based design of a smaller peptide with enhanced inhibitory efficiency. The kinetic analysis and osteoclast differentiation assay showed that in addition to the sharp turn induced by the disulfide bond, two consecutive arginine residues were also important for binding to RANKL and inhibiting osteoclastogenesis. Docking and molecular dynamics simulations proposed the binding mode of the peptide to the RANKL trimer, showing that the arginine residues provide electrostatic interactions with RANKL and contribute to stabilizing the complex. These findings provided useful information for the rational design of therapeutics for bone diseases associated with RANK-RANKL function.

INTRODUCTION

Receptor activator of nuclear factor κB (RANK) and its corresponding ligand (RANKL) belong to the tumor necrosis factor (TNF)-receptor ligand family and are known as key regulators of bone resorption and bone formation. Osteoclasts are multinucleated cells involved in the process of bone resorption, which is essential for bone remodeling. The interaction of RANKL with RANK on the osteoclast precursors leads to the differentiation and fusion of these precursors into mature osteoclasts. Osteoclasts are specialized bone-resorbing cells that play a critical role in maintaining bone homeostasis.

Keywords: bone resorption, docking, molecular dynamics simulation, peptide inhibitor, receptor activator of nuclear factor-κB (RANK), structure

Received 20 October, 2015; revised 24 January, 2016; accepted 26 January, 2016; published online 26 February, 2016

Mol. Cells 2016; 39(4): 316-321
http://dx.doi.org/10.14348/molcells.2016.2286

© The Korean Society for Molecular and Cellular Biology. All rights reserved.
This is an open-access article distributed under the terms of the Creative Commons Attribution-NonCommercial-ShareAlike 3.0 Unported License. To view a copy of this license, visit http://creativecommons.org/licenses/by-nc-sa/3.0/.
Table 1. Amino acid sequence of the inhibitory peptides that mimic Loop3 of RANK.

| Peptide name       | Sequence                  |
|--------------------|---------------------------|
| Loop3 in RANK      | 118-GYH WNSDCEC CRR N-131|
| L3-3               | 1-YC WNSDCEC CVYRR-13     |
| L3-3S              | 1-YC SNsdCEC CVYRR-13     |
| L3-3n              | 1-YC WNSDCEC CRRA-12      |
| L3-3A              | 1-SDCECS-6                |
| L3-3B              | 1-SDCECSR-8               |

Important residues that participate in cyclization or phi-phi interactions are shown in bold letters. Substituted or deleted residues are also indicated.

MATERIAL AND METHODS

Protein and peptide preparation

The pVFT3s (Korean patent 10-0690230) harboring the ectodomain region (residues 157-316) of mouse RANK cDNA was transformed into *Escherichia coli* BL21 (DE3) cells (Novagen, USA). The protein was expressed overnight at 18°C following induction with 0.2 mM isopropyl-b-D-thiogalactosidase. After Ni-NTA column (GE Healthcare, USA) chromatography, purified protein was digested overnight with TEV protease at 4°C and loaded onto a Superdex-200 gel-filtration column (GE Healthcare) equilibrated with 25 mM Tris-HCl (pH 7.5) and 50 mM NaCl. The peptides were custom-synthesized (BIO-FD&C, Korea) and dissolved in water to prepare the stock.

Osteoclast differentiation assay

The maturation of mouse osteoclast-precursor cells was examined by the tartrate-resistant acid phosphatase (TRAP) assay (Oshiro et al., 2002). Briefly, osteoclast precursors were cultured in a 96-well plate (2 × 10^4 cells/well) in α-Minimal Essential Medium supplemented with 10% fetal bovine serum in the presence of 20 ng/ml of RANKL and 30 ng/ml human macrophage colony-stimulating factor. RANK or the peptide to be tested was added to a final concentration of ~10-100 μM. Activity of treated cells was assessed using naphthol AS Bi phosphate (Sigma Aldrich) and dissolved in water to prepare the stock. Multinucleated TRAP-stained cells were counted under the microscope. Each experiment was repeated three times for statistical analysis.

Cell viability measurement

The possible cytotoxicity of peptides was estimated by quantitative colorimetric MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl-tetrazolium bromide] (Sigma Aldrich) assay as previously described (Denizot and Lang, 1986). Cells were cultured in a 96-well plate (3 × 10^4 cells/well), and each peptide was added to the medium to varying final concentrations (~10-100 μM). Cell viability was represented by the reduction of MTT to formazan, which was quantitated by absorbance at 570 nm.

Statistical analysis

All data were expressed as the mean ± SD. Statistical analyses were performed using the unpaired Student t test, and p < 0.05 was considered statistically significant.

Biolayer interferometry (BLI)

All measurements were made using a ForteBio BLItz instrument at 25°C (ForteBio Corp., USA). The peptide (5 μM) was immobilized onto the amine-reactive biosensors using N-hydroxysuccinimide and N-(3-dimethylaminopropyl)-N-ethylcarbodiimide hydrochloride (Sigma Aldrich). Amine coupling was performed in 10 mM MES (pH 5.8) after pH optimization. Unoccupied amine groups were quenched with 1 M of ethanolamine. The binding affinity of peptide was calculated using Rmax values that were estimated with 0.2, 1, 5, 10, and 15 μM purified RANKL. In each cycle of measurements, the baseline was adjusted with 10 mM 2-ethanesulfonic acid (MES; pH 5.8) for 30 s, followed by biosensor activation and peptide immobilization for 150 s. After ethanolamine quenching for 150 s, the baseline was set again for 30 s, and association and dissociation steps of 300 s were each performed sequentially. Kinetic parameters were calculated using ForteBio and SigmaPlot software (Systat Software Inc., USA).

Western blot analysis

Bone marrow-derived macrophage-like cells (BMMs) were washed with ice-cold phosphate-buffered saline (PBS) and lysed in extraction buffer [50 mM Tris-HCl (pH 8.0), 150 mM NaCl, 1 mM ethylenediaminetetraacetic acid, 0.5% Nonidet P-40, and protease inhibitors]. Cell lysates were separated by sodium dodecyl sulfate polyacrylamide gel electrophoresis and transferred to a polyvinylidene difluoride membrane (Millipore, USA). Membranes were blocked with TBS-T [10 mM Tri-HCl (pH 7.6), 150 mM NaCl, and 0.1% Tween 20] containing 5% skim milk and probed with the primary antibodies against phospho-P38, phospho-JNK, P38, AKT, JNK, and actin (Cell Signaling Technology, USA). Blots were subsequently probed with horse-radish peroxidase-conjugated secondary antibodies (Abcam, UK). Signals were detected with enhanced chemiluminescence (Millipore) and analyzed with an LAS3000 luminescent image analyzer (GE Healthcare).

Modeling and docking of the protein-peptide interaction

The coordinates of the structure of a trimetric biological unit of the RANK-RANKL complex (PDB: 3QBQ) were used as templates in modeling and docking analyses (Ta et al., 2010). The binding mode of Loop3 to the binding cavity of the RANKL trimer guided the homology modeling of each peptide complexed with RANKL. The homology-based multichain-modeling tool of MODELLER version 9.13 was used for this purpose (Eswar et al., 2006). The peptide-RANKL complexes were supplied as a docking input to the Rosetta FlexPepDock ab initio docking server (http://flexpepdock.furmanlab.cs.huji.ac.il/index.php) with default parameters (London et al., 2011). No specific constraints were given to the peptide during docking, so that the peptide was in constraint-free and fully flexible conditions. In subsequent steps of molecular dynamics simulations, the highest-scoring docked structure was used as input for each peptide.

Molecular dynamics simulation

Molecular dynamics simulation was performed using the dock-
ing solution obtained from the FlexPepDock docking protocols (London et al., 2011). GROMACS version 5.0.2 was used (Tate, 2014) for molecular dynamics simulations using the DGIST supercomputing cluster facility (Daegu Gyeongbuk Institute of Science and Technology, Korea). An OPLS-AA/L force field was used to generate topology files (George et al., 2001). To define the simulation box, the RANKL-peptide complex was extended by 1 nm using the tool editconf. Water and ions were added to the simulation box and the protein surface using the solvate and genion tools, respectively, in GROMACS. The protein surface was neutralized automatically with the neutral option of genion. The peptide-RANKL complex was energy minimized for several cycles (50,000 steps/cycle) and applied for iso-thermal-isochoric equilibration (NVT-equilibration) for 100 ps to stabilize the temperature of the system to 300 K. The output of the NVT-equilibration was subjected to constant temperature (300 K) and pressure equilibration (NPT-equilibration) to stabilize the pressure of the system. The system was stabilized under a constant average pressure of 1 bar. Next, using the model from the NPT-equilibration, molecular dynamics simulation was performed for 30 ns. For all peptides, a temperature of 300 K and a predetermined constant pressure were applied. Pressure coupling was maintained at the reference pressure of 1 bar and isothermal compressibility of 4.5e-5. Finally, the root-mean-square deviation (RMSD) values of both separate peptides and peptide-protein complexes were calculated.

RESULTS AND DISCUSSION

Design of Loop3-mimic peptides
In our previous work, the crystal structure of the RANKL/RANK complex and biochemical assays of RANK mutants revealed that Loop3 of RANK (118-GYWNSDCECCRR-131) was essential for binding to RANKL (Ta et al., 2010). From these observations, we designed several peptides that mimic Loop3 and investigated their inhibitory effects on complex formation with RANKL and osteoclastogenesis. Among the tested peptides, L3-3 (1-YCWNSDCECC YRR-13; Table 1) was shown to be the most potent inhibitor and had higher affinity for RANKL compared to OP3-4, a previously characterized OPG-mimic inhibitor (Ta et al., 2010). In L3-3, H120 of RANK loop3 was exchanged with a cysteine (C2) for the purpose of cyclization through the C2-C10 disulfide bond, and a tyrosine (Y11) was inserted between C10 and R12 (R129 in RANK) to enable formation of a phi-phi interaction with Y1 (Y119 in RANK). L3-3 was expected to retain an overall loop structure similar to Loop3; however, it was desirable to reduce its size and improve the inhibitory efficiency for clinical applications, given that elucidation of the structural features required for inhibitory function will facilitate the rational design of peptides. Therefore, we modified specific residues in L3-3 (Table 1) and tested the activities of the derived peptides.

First, W3, a tryptophan corresponding to W121 in RANK, was replaced with a serine in L3-3S in order to relieve the steric hindrance of the indole group. In L3-3N, Y11, the C-terminal tyrosine that was introduced in L3-3 for a phi-phi interaction, was deleted to see how phi-phi stacking affects activity. Two shorter peptides, L3-3A and L3-3B, were designed to determine the minimum requirements for interaction with the Loop3-binding pocket in the RANKL trimer. L3-3A and L3-3B both lacked the N-terminal cysteine residue needed for cyclization (corresponding to C2 in L3-3); however, the formation of an internal disulfide bond (corresponding to C7-C9 in L3-3 and C125-C127 in RANK), which is the critical feature of Loop3, would not be affected. Moreover, the cysteine corresponding to C10 of L3-3 (C128 in RANK) was replaced with a serine to preclude formation of undesired linkages with either cysteine residue (C7 or C9 in L3-3). In L3-3A, two arginine residues at the N-terminus (RR) were shown to interact with E225 and D266 of RANKL through ionic and hydrogen bonding, respectively, in the crystal structure of the complex (Ta et al., 2010).

Inhibition of RANKL-induced osteoclastogenesis

MTT- and TRAP-activity assays were performed for the newly designed peptides, as well as L3-3 as a control, in order to estimate their cytotoxicity and inhibitory effects on osteoclastogenesis, respectively. In the MTT assay, none of the peptides displayed noticeable cytotoxicity on mouse osteoclast-precursor cells (Fig. 1A). The results from the TRAP activity assay indicated that L3-3S and L3-3B were more effective relative to the original L3-3 peptide at inhibiting the maturation of osteoclast precursors (Fig. 1B). The elimination of steric hindrance by the tryptophan residue in L3-3S enhanced the inhibitory effect of the peptide. Most noticeably, the activity of L3-3B was improved significantly, causing almost complete cessation of osteoclastogenesis according to the TRAP-staining assay (Fig. 2).
However, inhibitory activity was diminished by the deletion of the RR residues in L3-3A, and no substantial change was observed for L3-3n. These results showed the potential value of L3-3B for therapeutic applications, indicated that overall cyclization was not absolutely required, and that the RR residues contributed to RANKL binding.

**L3-3B binding affinity for RANKL**

We analyzed the binding affinity of L3-3B for RANKL using BLI to corroborate the differentiation assay results. After conjugating a fixed concentration of L3-3B to an amine-reactive biosensor chip, different concentrations of RANKL were applied for association, followed by the dissociation step. The representative graphs at three concentrations are shown in Fig 3A. The \( R_{\text{max}} \) value is defined as the peak point where association ends and dissociation begins at a given concentration of RANKL (Fig. 3A). The steady-state dissociation constant (\( K_d \)) of L3-3B for RANKL was calculated using \( R_{\text{max}} \) values at five different concentration points (0.2-15 \( \mu \)M; Fig. 3B). The obtained value (\( K_d = 6.08 \, \mu \text{M} \)) was significantly lower than that of L3-3, which was reported in our previous work (\( K_d = 17.3 \, \mu \text{M} \); Ta et al., 2010), thereby confirming its improved potency to block formation of the RANK-RANKL complex.

**Specific interference of RANKL signaling by L3-3B**

RANKL and macrophage colony-stimulating factor (M-CSF) are two essential cytokines that activate respective intracellular signaling pathways during osteoclast differentiation. In order to investigate the specificity of RANKL inhibition by L3-3B, we performed western blot analyses to examine the interference of RANKL- and M-CSF-signaling pathways in BMM cells. When BMM cells were treated with RANKL that was pre-incubated with L3-3B for 30 min, phosphorylation of p38 and AKT was significantly diminished (Fig. 4A). However, M-CSF-mediated activation of AKT and JNK in BMM cells was not effectively blocked by treatment with L3-3B (Fig. 4B). These results indicated that L3-3B specifically interfered with the RANKL-signaling pathway in BMM cells.

**Docking of the modeled peptides**

The osteoclast differentiation and BLI analyses highlighted the importance of the sharp turn induced by the C-C disulfide linkage (C125 and C127 in RANK) and the RR residues (R129 and R130 in RANK). In order to investigate their interaction modes in the Loop3-binding pocket, peptides were modeled in the RANKL trimer using homology modeling and docking. The binding affinity of each peptide to RANKL was estimated using...
To verify the binding modes of the peptides to RANKL sug-

Molecular dynamics simulation
To verify the binding modes of the peptides to RANKL sug-

Table 2. The docking scores for five inhibitory peptides.

| Peptide | FlexPepDock score (REU) | Interface score (I_sc) |
|---------|--------------------------|------------------------|
| L3-3    | -182.367                 | -19.05                 |
| L3-3A   | -231.901                 | -7.803                 |
| L3-3B   | -233.223                 | -15.62                 |
| L3-3n   | -170.286                 | -14.76                 |
| L3-3S   | -203.767                 | -16.02                 |

Fig. 5. Docking and molecular dynamics simulation of RANKL inhibitory peptides. (A) Docking simulation of the L3-3B peptide. L3-3B is depicted in green, and monomers of the RANKL trimer are colored light brown, light pink, and sky blue. Residues are labeled using the same color scheme with darker shading. (B) Comparison of RMSD graphs from 30-ns molecular dynamics simulation results for four peptides: L3-3, L3-3B, L3-3n, and L3-3S. (C) Overlap of six coordinates of the L3-3B-RANKL complex captured over the course of a 50-ns molecular dynamics simulation. Different backbone orientations of the L3-3B peptide are shown in different colors, with all RANKL backbones colored grey. (D) Binding mode of the L3-3B peptide on the RANKL surface. The RANKL surface is mapped using Coulombic surface-coloring gradient (Chimera 1.10.1), where deep red and deep blue indicate surface charges of −10 and +10, respectively, and white indicates a neutral surface charge. The L3-3B peptide backbone is shown in green, and positive and negative side-chain atoms are depicted in blue and red, respectively. The C3-C5 disulfide bond is colored yellow.

suggested from the docking analyses, further investigation of bind-

ing stability was performed using molecular dynamics simula-

tions. The docking model of each peptide bound to RANKL with the highest docking score was used to generate a 30-ns molecular dynamics simulation under an OPLS-AA/L force field. L3-3A showed a high degree of displacement from its initial docking position, the Loop3-binding pocket. Specifically, changes occurring in the first 0.6 ns were the most drastic, with RMSD values close to 5 Å. Throughout the remainder of the simulation, a high degree of fluctuation was observed, indicating the instability of the peptide in the Loop3-binding pocket (Fig. 5B). It is believed that this instability was largely due to the lack of the RR residues, which mediate key interactions involved in RANKL binding (Fig. 5A). However, L3-3B appeared to maintain stable interactions in the binding pocket, which was represented by the minimal changes in RMSD of backbone positions throughout most of the simulation period (30 ns; Fig. 5B). Other peptides showed considerable fluctuation throughout the simulation. These simulation results demonstrated the stable nature of the L3-3B backbone bound to RANKL. These results were further corroborated in a 50-ns simulation, showing the superposition of six L3-3B orientations captured during the simulation and demonstrating the rigidity of its backbone over time (Fig. 5C).

In the molecular dynamics simulation, the interaction between L3-3B and the RANKL trimer was stabilized by interactions between positively charged R7 and R8 residues in L3-3B (R129 and R130 in RANK) and negatively charged residues in
RANKL (D73 and E108; Figs. 5A and 5D). Additionally, the sharp loop region, which is maintained by a C3-C5 disulfide bond (C125-C127 in RANKL) provided stable packing onto the pocket, consisting of Y74, E76, T100, and F109 in RANKL. Taken together, we concluded that peptide binding to RANKL largely depended upon the packing of the C-E-C loop in RANKL and electrostatic interactions provided by the RR residues.

**CONCLUSION**

In this study, we designed a new peptide inhibitor of RANKL based on the crystal structure of a RANK loop bound to RANKL. From these efforts, we identified L3-3B as the most promising inhibitor in terms of its length and inhibitory activity. The *in vitro* binding and osteoclast differentiation assays demonstrated that its binding and inhibitory activities, respectively, were significantly improved as compared to the original peptide, L3-3. The specific inhibition of RANKL by L3-3B was confirmed based on western blot analysis showing that L3-3B was capable of interfering with RANKL-induced phosphorylation of p38 and AKT in BMM cells.

Additionally, the proposed L3-3B binding mode to RANK according to molecular docking and molecular dynamics simulations was consistent with the experimental results. Our studies confirmed that the two cysteine residues that constitute a sharp turn via a disulfide bond that fits into the Loop3-binding pocket, as well as RR residues that form salt bridges with the negatively charged residues of RANKL, were crucial for stable binding. These interactions appear to be essential requirements for RANKL binding and inhibition of RANKL-mediated osteoclastogenesis. We expect that these findings and strategies will be useful for the rational development of antiresorptive therapeutics specific to RANKL modulation.

**ACKNOWLEDGMENTS**

We acknowledge the DGIST Supercomputing & Big Data Center for the allocation of supercomputing time. This work was supported by the Postdoctoral Research Program of Sungkyunkwan University to HK (2014).

**REFERENCES**

Burgess, T.L., Qian, Y., Kaufman, S., Ring, B.D., Van, G., Capperelli, C., Kelley, M., Hsu, H., Boyle, W.J., Dunstan, C.R., et al. (1999). The ligand for osteoprotegerin (OPGL) directly activates mature osteoclasts. *J. Cell Biol.*, 145, 527-538.

Cohen, S.B., Dore, R.K., Lane, N.E., Ory, P.A., Peterfy, C.G., Sharp, J.T., van der Heijde, D., Zhou, L., Tsuji, W., Newmark, R., et al. (2002). Osteosub and osteoblasts provide growth factors essential for periodontal ligament fibre formation. *J. Bone Miner. Res.*, 20281-20286.

Tanaka, S. (2013). Regulation of bone destruction in rheumatoid arthritis through RANKL-RANK pathways. *World J. Orthop.*, 4, 1-6.

Tate, A., Aarniala, B., Sambrook, P., Delavault, C., Poon, W., McQuinn, J., and Pennington, J. (2006). Tumor necrosis factor receptor superfamily member 11a (TNFRSF11A) and osteoprotegerin (OPG): novel targets for the treatment of osteoporosis. *Biochim. Biophys. Acta Mol. Cell. Biol. Org.*, 1763, 521-536.

Wada, T., Nakashima, T., Hiroshi, N., and Pennington, J. (2006). RANKL-RANK signaling in osteoclastogenesis and bone disease. *Trends Mol. Med.*, 12, 17-25.

Wittrant, Y., Theoleyre, S., Chipoy, C., Padrines, M., Blanchard, F., Heymann, D., and Redini, F. (2004). RANKL/RANK/OPG: new therapeutic targets in bone tumours and associated osteolysis. *Biochim. Biophys. Acta*, 1704, 49-57.