1 Research article

2 Dimeric ^{R25C}PTH(1-34) Activates the Parathyroid Hormone-1 Receptor *in vitro*

and Stimulates Bone Formation in Osteoporotic Female Mice

- 4 Minsoo Noh^{1, 2, *}, Xiangguo Che^{3, *}, Xian Jin³, Dong-Kyo Lee³, Hyun-Ju Kim³, Doo Ri Park⁴, Soo
- 5 Young Lee⁴, Hunsang Lee², Thomas Gardella⁵, Je-Yong Choi^{3, #}, Sihoon Lee^{1, #}
- ⁶ ¹Department of Internal Medicine and Laboratory of Molecular Endocrinology, Gachon University
- 7 School of Medicine, Incheon 21565, Republic of Korea
- ⁸ ²Department of Life Sciences, Korea University, Seoul 02841, Korea
- ⁹ ³Department of Biochemistry and Cell Biology, Cell and Matrix Research Institute, School of Medicine,
- 10 Kyungpook National University, Daegu 41944, Republic of Korea.
- ⁴Department of Life Sciences, Multitasking Macrophage Research Center, Ewha Womans University,
- 12 Seoul 03760, Korea
- ¹³ ⁵Endocrine Unit, Massachusetts General Hospital and Harvard Medical School, Boston, MA 02114,
- 14 USA
- 15
- 16 *These two authors are equally contributed to this manuscript.
- 17
- 18 [#]Correspondence:
- 19 Je-Yong Choi D.D.S., Ph.D., Department of Biochemistry and Cell Biology, Cell and Matrix Research
- 20 Institute, School of Medicine, Kyungpook National University, Daegu 41944, Republic of Korea. Tel.:
- 21 +82-53-420-4823; Fax: +82-53-422-1466; E-mail: jechoi@knu.ac.kr
- 22 **&**
- 23 Sihoon Lee, M.D., Ph.D., Department of Internal Medicine and Laboratory of Molecular Endocrinology,
- 24 Gachon University School of Medicine, Incheon 21565, Republic of Korea. Tel.: +82-32-458-2646;
- 25 Fax: +82-32-460-2381; E-mail: <u>shleemd@gachon.ac.kr</u>
- 26

27 Abstract

28 Osteoporosis, characterized by reduced bone density and strength, increases fracture risk, pain, 29 and limits mobility. Established therapies of Parathyroid hormone (PTH) analogs effectively promote 30 bone formation and reduce fractures in severe osteoporosis, their use is limited by potential adverse effects. In the pursuit of safer osteoporosis treatments, we investigated ^{R25C}PTH, a PTH variant 31 32 wherein the native arginine at position 25 is substituted by cysteine. These studies were prompted by 33 our finding of high bone mineral density in a hypoparathyroidism patient with the R25C homozygous 34 mutation, we explored its effects on PTH type-1 receptor (PTH1R) signaling in cells and bone metabolism in mice. Our findings indicate that ^{R25C}PTH(1-84) forms dimers both intracellularly and 35 extracellularly, and the synthetic dimeric peptide, ^{R25C}PTH(1-34), exhibiting altered activity in PTH1R-36 mediated cAMP response. Upon a single injection in mice, dimeric ^{R25C}PTH(1-34) induced acute 37 calcemic and phosphaturic responses comparable to PTH(1-34). Furthermore, repeated daily 38 39 injections increased calvarial bone thickness in intact mice and improved trabecular and cortical bone 40 parameters in ovariectomized (OVX) mice, akin to PTH(1-34). The overall results reveal a surprising 41 capacity of a dimeric PTH peptide ligand to activate the PTH1R in vitro and in vivo, suggesting a 42 potential new path of therapeutic PTH analog development.

43

44 Key words: Parathyroid hormone, PTH type 1 receptor, Bone formation, Osteoporosis

46 Introduction

47 Osteoporosis is a prevalent global bone disorder characterized by low bone mineral density (BMD), causing weakened bones prone leading to fragility fractures, particularly in the spine, hip, and 48 49 wrist. The development of osteoporosis is influenced by factors including gender, being more 50 prevalent in women; hormonal changes, like decreased estrogen levels during menopause; and age, 51 with heightened susceptibility post-menopause in women contributing to bone loss. Recent meta-52 analysis of previous studies indicates a global osteoporosis prevalence of 23.1 % among women and 53 11.7 % among men (1, 2). Osteoporosis stands as a noteworthy risk factor that poses challenges to 54 the preservation of autonomous mobility and overall well-being within an aging society. There is thus a 55 pressing need for safe and efficacious therapies for osteoporosis that mitigate fractures, alleviate 56 associated symptoms, and preserve physical functionality.

57 Anti-resorptive agents (e.g., bisphosphonates, denosumab, and romosozumab) encompass 58 one therapeutic approach that aims to specifically counteract the declines in bone mass by tempering 59 the balance between bone resorption and formation (3-5). It is pertinent to acknowledge, however, 60 that the prolonged use of most of such agents is limited due to potential long-term side effects. 61 Furthermore, most anti-resorptive therapies cannot stimulate new bone formation. In contrast, bone 62 anabolic agents, such as parathyroid hormone (PTH) and its analogs, such as teriparatide 63 (recombinant human PTH(1-34)), increase BMD by stimulating bone formation more than bone 64 resorption (6). PTH has an exceptionally short half-life in the blood of approximately 2-4 minutes (7, 8), 65 which helps in avoiding excessive increases in blood calcium levels that can otherwise limit the utility 66 of PTH-related medications, while yet inducing a desired anabolic effect on bone. It is also worth 67 noting that studies in rodents reveal that long-term administration of a PTH anabolic agent can lead to 68 bone overgrowth, osteosarcoma, as well as hypercalcemia (9, 10). Consequently, a goal of ongoing 69 research is to uncover the underlying molecular mechanisms driving the anabolic and catabolic 70 effects of PTH to thereby secure more effective therapeutic alternatives for osteoporosis (11).

PTH is produced and secreted by the parathyroid glands as a straight-chain monomeric polypeptide of 84 amino acids (12-14). It plays a vital role in maintaining calcium and phosphate equilibrium by acting on the PTH1R, a class B G protein-coupled receptor (GPCR) family (15-17) that is expressed primarily in cells of bone and kidney (18). The orchestrated downstream effects of PTH

in target cells act to ensure optimal bone health and the maintenance of the ambient blood calcium
 and phosphate concentrations required for proper nerve conduction, muscle activity, and systemic
 cellular communication, whereas disturbances in this system can lead to multiple disorders.

78 Central to the PTH signaling cascade is the activation of intracellular G proteins (19), most 79 prominently Gsa, which in turn activates adenylyl cyclase leading to the synthesis of the second 80 messenger cyclic AMP (cAMP), and the activation of cAMP-dependent protein kinase A (PKA) (20, 81 21). PTH can also activate other second messenger cascades, including the Gαg/phospholipase C 82 (PLC) / inositol 1,4,5-trisphosphate (IP3), diacylglycerol (DAG), protein kinase C (PKC) signaling 83 pathway (22, 23), highlighting the diverse biology of PTH and the PTH1R (24). Adding to this, the 84 PTH1R also mediates the actions of parathyroid hormone-related protein (PTHrP), a development 85 protein that acts in the formation of bones and other tissues. PTHrP shares homology with PTH in the 86 first 34 amino acids, which encompass the receptor-binding portions of the two respective ligands. 87 The PTH1R thus has an intrinsic capacity for dual ligand recognition, which opens possibilities for 88 exploring new modes of therapeutic development for diseases such as osteoporosis and 89 hypothyroidism (18, 19, 25-28). Pharmacologically, the PTH1R can adopt at least two distinct ligandbinding conformations, RG and R⁰, the selectivity for which can lead to altered modes of signaling *in* 90 91 vitro and in vivo for peptides such as PTH, PTHrP, and various hybrid analogs (29-33).

92 The current study extends our prior investigation in which we identified in a patient with 93 chronic hypocalcemia and hyperphosphatemia a mutation that changes the arginine at position 25 in the mature PTH(1-84) polypeptide to cysteine (R25CPTH) (34, 35). Antibody assays revealed the 94 95 ^{R25C}PTH mutant protein to be present in the patient's blood at markedly elevated levels. We now have found that this patient expressing the ^{R25C}PTH variant has higher-than-normal BMD. We further 96 97 characterize the ^{R25C}PTH protein and find that it can manifest in two distinct molecular forms: as a monomer and as a dimer, and we demonstrate that a dimeric ^{R25C}PTH(1-34) synthetic peptide retains 98 agonistic properties on the PTH1R that are driven by a moderate selectivity for the RG vs. R⁰ receptor 99 conformation. Finally, we demonstrate in mice that ^{R25C}PTH(1-34) can induce skeletal responses that 100 101 are similar to those induced by PTH(1-34), but without triggering an excessive hypercalcemic 102 response. Considering the proven bone-anabolic capacity of several established PTH agonist ligands, 103 and the need for safe, long-term treatments for skeletal disorders, our studies on dimeric ^{R25C}PTH(1-

104 34) suggest alternative strategies to consider in such drug development programs.

106 Results

107 Dimerization of ^{R25C}PTH(1-84)

In our previous studies, we showed that synthetic monomeric peptide, ^{R25C}PTH(1-34), as 108 109 compared to PTH(1-34), exhibits a moderately diminished PTH1R-binding affinity and decreased cAMP signaling potency *in vitro*, and that with long-term infusion in mice, ^{R25C}PTH(1-34) leads to only 110 111 minimal calcemic and phosphaturic effects, which corroborates the hypocalcemia seen in the original patient, despite the significantly elevated levels of ^{R25C}PTH in the plasma (34). On subsequent 112 analysis of this patient, we found a particularly high BMD (Supplementary File 1), which prompted us 113 to further characterize the functional properties of ^{R25C}PTH, as described herein. We produced 114 115 recombinant PTH(1-84) with or without the R25C mutation by expression of the corresponding cDNA 116 in HEK293T cells (Figure 1). We considered the possibility that the introduction of a sole new cysteine within the polypeptide chain of ^{R25C}PTH(1-84) might induce homologous bimolecular dimerization 117 through a disulfide bond involving the thiol functional group in each monomer (36, 37). To specifically 118 investigate this, we designed the cDNA constructs to express either pre-pro-PTH(1-115)-3xFLAG or 119 ^{R56C} pre-pro-PTH(1-115)-3xFLAG, such that after intracellular processing and cleavage of the pre-pro 120 regions, the mature PTH(1-84)-3xFLAG or R25CPTH(1-84)-3xFLAG peptides would be generated upon 121 122 transfection in HEK293T cells (Figure 1C) (38-40). We performed western blot analysis of total cell 123 lysates and conditioned culture media collected from the transfected cells to specifically assess the 124 possible presence of a disulfide-bonded dimeric form. Each sample was thus prepared in either 125 reduced or non-reduced form and proteins were detected using an anti-flag antibody. The results 126 demonstrated the presence of both a low molecular weight monomeric, and in the non-reduced samples, a higher molecular weight dimeric form of the ^{R25C}PTH(1-84)-3XFLAG protein in both the cell 127 128 lysate and extracellular conditioned medium fractions, and the dimer appeared to be at an elevated 129 proportion in the medium relative to the lysate (Figure 1D). To control for potential artifact effects 130 attributed to the 3xFLAG tag, we utilized plasmid constructs pcDNA3.0-(pre-pro-PTH)-IRES and pcDNA3.0-(^{R56C}pre-pro-PTH)-IRES encoding non-tagged PTH variants and an anti-PTH(39-84) 131 132 antibody for Western blot analysis, which again confirmed the presence of the dimer in total HEK293T 133 cell lysates (Supplementary File 2).

134

We observed in the above studies that the expression level of ^{R25C}PTH(1-84) was higher 135 136 than that of wild-type PTH(1-84) in both cell lysate and medium, which we considered might be due to 137 an intrinsic enhancement in protein stability and resistance to protein degradation in the dimeric molecule. To address this, we treated the cells with the proteasome inhibitor MG132, which acts by 138 139 forming a hemiacetal with the hydroxyl groups of active site threonine residues, and compared the expression levels of wild-type PTH(1-84) and ^{R25C}PTH(1-84) in the treated vs. untreated cells. The 140 141 results indicated that while the expression level of wild-type PTH was increased by MG132 treatment, it did not reach the level of R25CPTH, suggesting that the difference in expression is not related to a 142 difference in sensitivity to proteasome-mediated degradation (Supplementary File 3). 143

Overall, we have confirmed that ^{R25C}PTH(1-84) can form a dimeric structure, and the ^{R25C}PTH(1-84) secreted outside the cells predominantly exists in dimeric form. Thus, utilizing dimer R^{25C}PTH(1-84) in the analysis would be more relevant to understanding the actual function of ^{R25C}PTH. Consequently, we aim to conduct further validation using dimeric ^{R25C}PTH in our subsequent investigations.

149

150 Functional characterization of dimeric ^{R25C}PTH(1-34) *in vitro*

To explore the functional properties of dimeric ^{R25C}PTH, we conducted experiments using 151 synthetic peptides of PTH(1-34), ^{R25C}PTH(1-34) (monomeric) and disulfide-bonded dimeric ^{R25C}PTH(1-152 153 34). First, we examined the receptor-binding affinity of these ligands by performing competition 154 experiments using membranes prepared from HEK293-derived GP-2.3 cells that stably express the human PTH1R and assay formats designed to assess binding to either the G protein-uncoupled R⁰ or 155 G protein-coupled RG receptor conformation. Each experiment was replicated four times (n = 4). The 156 results revealed that monomeric ^{R25C}PTH(1-34) bound to both the R⁰ and RG conformations with 157 comparable, albeit slightly weaker affinity as compared to PTH(1-34), while dimeric ^{R25C}PTH(1-34) 158 159 bound to each conformation with weaker affinity than did the monomeric form while showing an apparent selectivity for higher binding to the RG vs R⁰ conformation of PTH1R (Figure 2A). 160

161 To investigate the signaling properties of the ligands, we measured the changes increases in 162 intracellular levels of cAMP induced by each ligand in an osteoblastic SaOS-2 derived cell line (SGS-

163 72 cells) that stably expresses the Glosensor cAMP reporter. Each measurement was replicated four 164 times (n = 4). These assays revealed that each ligand dose-dependently increased the cAMP levels in 165 the cells, detected as an increase in luminescence in an Envision plate reader, and while the 166 potencies were moderately and more substantially reduced for the monomeric and dimeric forms of 167 the ligand, respectively, as compared to PTH(1-34), the maximum response attained by each ligand were comparable (Figure 2B). Dimeric ^{R25C}PTH(1-34) thus retains signaling functionality at the PTH1R 168 169 that is characterized by a potency approximately commensurate with its affinity for binding to the RG 170 PTH1R conformation.

171

172 Effect of single injection of dimeric ^{R25C}PTH(1-34) on calcium and phosphate regulation in mice

To assess whether dimeric ^{R25C}PTH can function *in vivo*, we injected the ligand, and in 173 174 parallel either vehicle or PTH(1-34) (each peptide at a dose of 50 nmol/kg) into CD1 female mice and 175 measured levels of ionized calcium (Ca²⁺) in the blood (n = 6 mice/group), inorganic phosphate (Pi) in plasma (n = 12 mice/group), and the excretion of Pi into urine (n = 6 mice/group). Blood Ca^{2+} levels 176 177 were measured at serial time points of pre-injection, 1, 2, 4, and 6 hours post-injection. Both PTH(1-34) and dimeric ^{R25C}PTH(1-34) induced increases in blood Ca²⁺ levels that were significant, relative to 178 179 the levels in vehicle-injected mice, at 1 and 2 hours post-injection, and the levels then returned to the 180 baseline levels of vehicle control mice by 4 hours (Figure 3A). Plasma Pi levels were measured in 181 samples obtained pre-injection, at 6 minutes, and 1, 2, and 6 hours post-injection. PTH(1-34) induced 182 a significant decrease in plasma Pi at 1-hour post-injection, and the levels subsequently returned to baseline by 2 hours. Injection of dimeric ^{R25C}PTH(1-34) resulted in a slight decrease in plasma Pi at 2 183 184 hours post-injection, but the effect was not significant (Figure 3B). Consistent with this trend, Pi levels in the urine of mice injected with dimeric ^{R25C}PTH(1-34) were increased significantly at 2 hours post-185 injection and then gradually returned to baseline levels (Figure 3C). These results thus indicate that 186 dimeric ^{R25C}PTH can elicit calcemic and phosphaturic responses in vivo that are fully with those 187 188 expected for an injected PTH1R agonist ligand. We did not detect an increase in plasma cAMP in response to dimeric ^{R25C}PTH(1-34) injected at a dose of either 50 nmol/kg (n = 12) or 100 nmol/kg (n 189 190 = 2), whereas a significant increase was observed at 15 minutes post-injection of PTH(1-34) at a dose 191 of 50 nmol/kg (Figure 4A and B). This result likely reflects at least in part a reduced cAMP-stimulating

potency of dimeric ^{R25C}PTH(1-34) ligand, relative to PTH(1-34), as revealed by our cell-based
Glosensor assays, but other possibilities, such as a reduced stability of the dimeric ligand in the blood,
might also account for the weaker cAMP response to the dimeric peptide *in vivo*.

195

196 Effect of dimeric ^{R25C}PTH(1-34) on bone calvariae in mice

197 To initially assess the effects that short-term treatment of dimeric ^{R25C}PTH(1-34) can have on 198 bone, we injected 8-week-old male C57BL/6 mice once a day for six days (days 1-6), with either 199 dimeric R25C PTH(1-34) (160 µg/kg/day), PTH(1-34) (80 µg/kg/day) or vehicle (N = 6 for group), and 200 after 10 days without treatment (day 16) followed by euthanasia, we isolated the calvariae for 201 histological analysis of new bone formation. Specifically, we examined sections stained with 202 hematoxylin and eosin (H&E) to assess the width of newly formed bone areas along the edge of each 203 sample. These regions exhibited a more vivid coloration compared to the surrounding existing bone 204 tissue, demarcated by a dotted line for clarity (Figure 5A). Measurements were taken below and 205 above the dissection area where new bone had formed, and these measurements were then utilized 206 to calculate the mean values for further analysis. These analyses revealed that both PTH(1-34) and dimeric ^{R25C}PTH(1-34) significantly increased the width of the new bone area by approximately four-207 208 fold, as compared to the vehicle group, and the effects of two ligands were not significantly different from each other (Figure 5B). These findings thus support a capacity of dimeric ^{R25C}PTH(1-34) to 209 210 induce new bone formation in vivo, which is of interest given the high BMD observed in the patient 211 with the ^{R25C}PTH mutation, despite the presumably continuous exposure of the bone to the mutant 212 ligand.

213

214 Effect of dimeric ^{R25C}PTH on bone mass in osteoporotic mice

To more directly assess the impact of dimeric R25C PTH(1-34) on bone mass, we administered it to ovariectomized (OVX) mice, which serve as a well-established model for postmenopausal osteoporosis. The OVX mice were injected daily for a duration of 4 weeks with either the dimeric ligand (OVX + dimeric R25C PTH(1-34), PTH(1-34) (OVX + PTH(1-34)) or vehicle (OVX + vehicle, OVXcontrols), and sham-operated (Sham) mice were used as further controls. Mice were euthanized at

220 the end of the injection period and tissue samples were isolated for analysis. Quantitative micro-221 computed tomography (µ-CT) analysis of the femurs obtained from each group revealed that, as 222 compared to OVX + vehicle controls, treatment with PTH(1-34) increased femoral trabecular bone 223 volume fraction (Tb.BV/TV) by 121%, cortical bone volume fraction (Ct.BV/TV) by 128%, cortical 224 thickness (Ct.Th) by 115%, and cortical area fraction (Ct.Ar/Tt.Ar) by 118% (Figure 6A). Treatment 225 with dimeric ^{R25C}PTH(1-34) resulted in similar effects on the femoral cortical bone parameters, as it 226 increased Ct.BMD by 104%, Ct.BV/TV by 125%, Ct.Th by 107%, and Ct.Ar/Tt.Ar by 116% (Figure 6B). The increase in cortical bone BMD was significant with dimeric ^{R25C}PTH(1-34) but not with PTH(1-34), 227 whereas an increase in femoral trabecular bone was only observed with PTH(1-34). 228

The effects of the treatments on bone biomechanical properties were assessed by conducting a three-point bending test on femurs isolated from the mice. The maximum load parameter was significantly decreased in the femurs from OVX-control versus sham mice, and was significantly increased, relative to the OVX-controls, by treatment with the PTH(1-34) but not dimeric ^{R25C}PTH(1-34) (Figure 6C). The slope parameter was significantly decreased in the femurs from OVX-control versus sham mice, and tended to increase versus OVX-controls by treatment with either PTH(1-34) or ^{R25C}PTH(1-34) but the changes were not significant.

236 We further analyzed the levels of bone metabolism markers in the serum obtained from the 237 mice at the study endpoint (Figure 6D). The levels of serum calcium were within the normal range in 238 all treatment groups, while serum phosphate levels were modestly increased in the OVX mice treated with PTH(1-34) or dimeric ^{R25C}PTH(1-34) as compared to with vehicle, but the effect was significant 239 240 only with PTH(1-34). The serum levels of CTX1, a bone resorption marker (41), were elevated in 241 each of the OVX groups versus the sham group, and tended to be lower in the OVX-PTH(1-34) 242 treatment group but the change relative to OVX-vehicle was not significant (Figure 6D). Serum P1NP 243 and alkaline phosphatase levels were each significantly increased in both the PTH(1-34)- and dimeric ^{R25C}PTH(1-34)-treated groups, compared to the OVX-vehicle group, consistent with an enhancing 244 245 effect of the ligands on bone formation (42, 43). Histological staining of proximal tibial sections for 246 tartrate-resistant acid phosphatase (TRAP) activity, a marker of osteoclast-mediated bone resorption, 247 revealed an apparent increase in this activity in bones of the OVX mice, as compared to those in 248 sham control mice, reflecting a heightened rate of bone turnover, as also suggested by the increased levels of serum CTX1 in the OVX mice, and the TRAP staining appeared to be reduced in the tibiae of the OVX mice treated with PTH(1-34) (Figure 6D and E). Further histomorphometric analysis confirmed a significant increase in the osteoclast surface area relative to bone surface area (Oc.S/BS) in the proximal tibiae of the OVX-vehicle mice, relative to that in the Sham-control mice, and this parameter was significantly decreased by treatment with PTH(1-34) but not dimeric ^{R25C}PTH(1-34) (Figure 6F).

We then analyzed the bone microstructure in the lumbar vertebrae through von Kossa staining of histological sections and histomorphometric quantification (Figure 7A). The trabecular bone volume fraction (TB; V, %), and trabecular number (Tb N) were significantly reduced in the OVXvehicle group, as compared to the Sham group, and treatment with either PTH(1-34) or dimeric ^{R25C}PTH(1-34) resulted in a significant increase in each of these parameters, as well as a concomitant reduction in trabecular separation (Tb Sp), as compared to the respective parameters in the OVXvehicle group (Figure 7B).

Dynamic bone histomorphometry was also performed on the vertebrae to evaluate rates of bone formation (Figure 7C). The trabecular mineral apposition rate (MAR) and cortical MAR were each significantly increased in both the OVX-PTH(1-34) and OVX-dimeric ^{R25C}PTH(1-34) groups, as compared to in the OVX-vehicle group, and although there was a tendency for an increase in the bone formation rate (BFR/BS) in both the trabecular and cortical bone with either PTH(1-34) or dimeric ^{R25C}PTH(1-34) treatment, the differences were not statistically significant, as compared to the OVX control (Figure 7D).

In summary, injection of dimeric ^{R25C}PTH into osteoporotic OVX mice resulted in significant increases in cortical bone in the femurs and trabecular bone in the vertebrae, as well as significant increases in the trabecular MAR and serum markers of the bone formation markers, ALP and PINP, without inducing excessive bone resorption or hypercalcemia.

273 Discussion

274 In this study, we show the introduction of a cysteine mutation at the 25th amino acid position 275 of mature parathyroid hormone (PTH) facilitates the formation of homodimers comprised of the resulting ^{R25C}PTH peptide. This dimerization surprisingly was compatible with receptor binding affinity, 276 277 and lead to relatively minor deviations in functional behavior as assessed in our cell-based assays and compared to the standard monomeric control PTH peptide. The homozygous ^{R25C}PTH mutation 278 279 was identified in patients that presented with hypocalcemia and hyperphosphatemia, despite elevated 280 PTH levels (34, 44), and the mutation was found to impact the bioactive region of PTH (34, 45-48). Our initial research on this ^{R25C}PTH mutant focused on the monomeric state of the peptide, and these 281 282 studies revealed relatively moderate decreases in PTH1R binding affinity and cAMP stimulating 283 potency in vitro and moderately impaired calcemic effects upon infusion in mice. Additional patient 284 observations, however, revealed the patients to have a higher BMD than anticipated for age-matched 285 averages. This elevated bone mass, coupled with the elevated serum PTH levels, prompted our 286 further investigations into the properties of the mutant PTH, as described herein.

287 Our investigations brought to light the capacity of the cysteine-25 mutation to induce dimer 288 formation in the otherwise monomeric PTH polypeptide as produced in transfected cells. This result 289 was established by comparing western blots of transfected cell lysates analyzed under reducing 290 versus non-reducing conditions of gel electrophoresis. Subsequently, we employed synthetic peptides to further explore the functional properties of dimeric ^{R25C}PTH(1-34). The results of our current studies 291 292 show some divergence from our previous findings obtained using the monomeric counterpart, ^{R25C}PTH (1-34). Compared to the monomer, dimeric ^{R25C}PTH(1-34) exhibited a more preferential 293 binding affinity for the RG versus R⁰ PTH1R conformation, despite a diminished affinity for either 294 295 conformation. We also observed that the potency of cAMP production in cells was lower for dimeric ^{R25C}PTH as compared to the monomeric ^{R25C}PTH, in accordance with a lower PTH1R-binding affinity. 296

We further pursued *in vivo* applications in mice. Initially, we assessed the calcemic and phosphatemic responses to a single injection of synthetic peptides of either PTH(1-34) or dimeric R^{25C}PTH(1-34) in intact mice. We found the dimer could induce increases in plasma calcium levels that were at least as robust and as sustained as those induced by PTH(1-34), and was similarly in phosphaturic (Figure 3). Intriguingly, we did not detect an increase in plasma cAMP levels upon injection of dimeric ^{R25C}PTH, whereas PTH(1-34) injection resulted in the expected rapid and transient increase in blood cAMP (Figure 4). Determining whether this difference is due to a difference in the capacity of the two ligands to activate the presumably renal pool of PTH receptors that mediate the blood cAMP response to PTH, a difference in the pharmacokinetic properties of the ligands, or some other such effects will require further investigation.

307 Activation of the canonical Gas-cAMP-PKA signaling pathway is generally thought to underly 308 most of the biological responses induced by PTH1R activation, and our studies in SAOS2-derived 309 osteoblastic cells confirm that dimeric PTH can activate this pathway, albeit not as efficiently as a 310 monomeric PTH peptide. Arg25 resides in the 20-34 (C-terminal region) of PTH(1-34), which plays a 311 significant role in the binding of the ligand to the extracellular domain (ECD) of the PTH1R. In concert 312 with the binding of the 20-24 region of PTH to the ECD, the N-terminal portion of PTH engages the 313 transmembrane domain (TMD) of the PTH1R to then induce the conformational changes involved in 314 G protein coupling and cAMP production (33, 49-52). The precise binding mode used by dimeric ^{R25C}PTH to the PTH1R is unknown, but it may be anticipated that it differs to some extent from that 315 316 used by the monomeric peptide, due, for example, to the changes in bulk molecular size and display 317 of accessible functional groups. Consequently, the receptor conformational changes and the modes of 318 coupling to downstream effectors may differ for the monomeric versus dimeric ligands, which could 319 potentially lead to altered signaling and biological responses in vivo. Whether such changes account for the increased bone density observed in the patient with the homozygous ^{R25C}PTH mutation is 320 321 unknown but cannot be presently ruled out.

The results of practical assessments of dimeric ^{R25C}PTH(1-34) for effects on calvarial bone 322 after short-term (6-days) injection into normal mice, and for effects on bone mass after long-term (4-323 324 weeks) daily injection in osteoporotic OVX mice demonstrate the dimer can mediate significant 325 influences on bone metabolism. Moreover, the results of these studies in OVX mice provided 326 evidence to suggest that the effects of dimeric PTH may favor bone anabolism versus bone 327 catabolism more effectively than a monomeric PTH peptide. While further investigation is necessary, the current experimental data strongly imply that the dimeric form of ^{R25C}PTH can serve as a new 328 329 peptide with distinct functionalities compared to the wild-type PTH.

Furthermore, the recent identification of a young patient in Denmark displaying homozygous R^{25C}PTH has opened avenues for observing the direct impacts of ^{R25C}PTH within the human biological system (44). The continual monitoring and observation of patients will contribute to a more profound comprehension of the long-term consequences associated with ^{R25C}PTH exposure. This extensive observation is crucial in delineating the extended effects of this compound on individuals. Consequently, by conducting thorough investigations to confirm the potential bone anabolic effect of ^{R25C}PTH, we hope to develop a novel bone anabolic agent with a targeted focus on the PTH1R.

337 Materials and Methods

338 Plasmid construction

The coding sequences (CDS) of pre-pro-PTH and the mutated form, ^{R56C} pre-pro-339 PTH[^{R25C}PTH], were amplified using primers containing the attB site. These CDS fragments were 340 obtained from pcDNA3.0-(hpre-pro-PTH)-IRES and pcDNA3.0-(h^{R56C}pre-pro-PTH)-IRES, which were 341 used in the previous research conducted by Lee, et al. (34). Both CDS were introduced into donor 342 343 vector pDONR223 with Gateway™ BP Clonase™ II Enzyme mix kit (Invitrogen, USA). Then pre-pro-344 PTH and ^{R56C} pre-pro-PTH were each cloned into pcDNA3.1-ccdB-3xFLAG-V5 with LR Gateway™ LR Clonase™ II Enzyme mix (Invitrogen, USA) to construct pcDNA3.1-(pre-pro-PTH)-3xFLAG-V5, and 345 pcDNA3.1-(R56C pre-pro-PTH) -3xFLAG-V5. All experimental procedures were done with the 346 manufacturer's instruction. pDONR223 was a gift from Kim Lab (Roswell Park Comprehensive 347 348 Cancer Center, USA), and pcDNA3.1-ccdB-3xFLAG-V5 was a gift from Taipale Lab (Donnelly Centre, University of Toronto, Canada). 349

350

351 Cell culture

All cell lines were grown at 37°C in 5 % CO₂. HEK293T cells were cultured in Dulbecco's Modified Eagle Medium (Cytiva[™], HyClone DMEM/High glucose with L-Glutamine and HEPES, Cat No. SH30243.01, USA) with 10 % FBS (Gibco[™], Fetal Bovine Serum, certified, Origin: USA, Cat No. 16000044, Lot No. 2522247RP, USA) and 1 % penicillin-streptomycin (Cytiva[™], HyClone Penicillin-Streptomycin 100X solution, Cat No. SV30010, USA). Cell lines obtained from commercial sources were not further authenticated.

358

359 Transfection

pcDNA3.1-(pre-pro-PTH)-3xFLAG-V5 and pcDNA3.1-(^{R56C}pre-pro-PTH)-3xFLAG-V5 were
 each transfected into HEK293T with Lipofectamine[™] 3000 Transfection Reagent (Invitrogen[™], Cat
 No. L3000001, USA) according to the manufacturer's instruction. After 48 hours of transfection, a
 culture medium was collected and used for western blot as a secreted protein sample. The rest of the

364 cells were lysed by RIPA buffer (Thermo Scientific[™], RIPA Lysis and Extraction Buffer, Cat No. 89900,
 365 USA) following the manufacturer's instruction and used for western blot as total cell lysate sample.

366

367 Western blot

368 Protein samples were prepared in two types, reduced sample and non-reduced sample. 369 Reduced samples were a mixture of protein (secreted protein or cell lysate), sample buffer 370 (Invitrogen™, NuPAGE™ LDS Sample Buffer (4X), Cat No. NP0007, USA), with reducing agent (Invitrogen[™], 10X Bolt[™] Sample Reducing Agent, Cat No. B0009, USA) and heated for 5 min at 95°C. 371 372 Non-reduced samples were a mixture of protein, and sample buffer, without reducing agent, and not 373 heated. The protein samples were loaded onto 4 - 12 % Bis-Tris protein gels (GeneSTAR, StarPAGE 374 Bis-Tris 4-12 %/15well, Cat No. GPG4115), and ran with MOPS/SDS running buffer (GeneSTAR, 20X 375 MOPS / SDS Running Buffer, Cat No. GMB0080). Transfer to the membrane was done by iBlot™ 2 376 Dry Blotting System (Invitrogen[™], iBlot[™] 2 Gel Transfer Device, Cat No. IB21001, USA) with PVDF 377 transfer stack (Invitrogen™, iBlot™ 2 Transfer Stacks-PVDF-mini, Cat No. IB24002, USA) according 378 to manufacturer's instruction. The membranes were blocked for 1 hour at room temperature (RT) in 5 % 379 skim-milk solution in Tris-buffered saline (TBS; 20 mM Tris-base, 500 mM NaCl, pH 7.5) and then 380 washed three times for 10 min each with tris-buffered saline with 0.05 % tween-20 (TBST). The 381 membranes were incubated with primary antibody for 1 hour at RT, then washed three times for 382 10 min each with TBST. If needed, the membranes were incubated with horseradish peroxidase (HRP)-conjugated secondary antibody for 1 hour at RT after primary antibody incubation, then 383 384 washed three times for 10 min each with TBST. After washing, the membranes were rinsed and 385 soaked in TBS. To develop a blot image, the membranes were treated with a chemiluminescent 386 substrate solution (Merck Millipore, Immobilon ECL Ultra Western HRP Substrate, Cat No. WBULS0500, USA) according to the manufacturer's instruction. The blot images were obtained by 387 388 LAS 4000 mini (Cytiva, ImageQuant™ LAS 4000 mini, USA). The dilution condition for the anti-FLAG 389 with hHRP conjugated antibody (Sigma-Aldrich, Monoclonal ANTI-FLAG® M2-Peroxidase (HRP) 390 antibody produced in mouse, Cat No. A8592, USA) was 1:2,000. The dilution condition for the anti-391 HSP90 primary antibody (Santa Cruz Biotechnology Inc., HSP 90 Antibody (AC-16), Cat No. sc-

- 101494, USA) was 1:5,000. The dilution condition for the anti-mouse secondary antibody was 1:5,000.
 Each antibody was diluted in TBST with 1 % BSA solution.
- 394

395 **Proteasome inhibition assay**

396 HEK293T cells were seeded in culture dishes at approximately 60 % confluence, and they 397 were allowed to grow for about 20 to 24 hours prior to transfection. The transfection of pcDNA3.1-(pre-pro-PTH)-3xFLAG-V5 and pcDNA3.1-(^{R56C}pre-pro-PTH)-3xFLAG-V5 was conducted following the 398 399 method mentioned earlier. After 24 hours of transfection, MG132, dissolved in DMSO, was added to 400 the cells to achieve a final concentration of 10 µM. For the mock treatment, DMSO alone was added. 401 The cells were then incubated for an additional 24 hours after MG132 treatment. Both the culture 402 medium and cell lysate were prepared for western blot analysis to assess the restored protein levels. 403 The western blot procedure was carried out as described in the previous section.

404

405 **Peptide synthesis and quantification**

406 Human PTH(1-34), ^{R25C}PTH(1-34), and dimeric ^{R25C}PTH(1-34) were chemically synthesized 407 by Anygen (Gwangju, Republic of Korea). The purity and mass of each peptide were analyzed by 408 high-performance liquid chromatography (HPLC) and matrix-assisted laser desorption ionization-time 409 of flight mass spectrometry (MALDI-TOF MS) in the manufacturer.

410

411 **PTH1R competition binding assay**

The binding of PTH and its analogs to G protein-uncoupled PTH1R (R⁰ conformation) and G protein-coupled PTH1R (RG conformation) was assessed using a competition method with membranes prepared from GP-2.3 cells (HEK-293 cells stably expressing the hPTH1R). For tracer radioligands, we utilized ¹²⁵I-PTH(1-34) and ¹²⁵I-MPTH(1-15). The unlabeled ligands tested were PTH(1–34), ^{R25C}PTH(1-34), and dimeric ^{R25C}PTH(1-34). Binding to the R⁰ conformation was assessed using ¹²⁵I-PTH(1-34) as the tracer radioligand, while binding to the RG conformation was assessed

using ¹²⁵I-MPTH(1-15). The addition of unlabeled ligands PTH(1–34), ^{R25C}PTH(1-34), or dimeric
 ^{R25C}PTH(1-34) caused dissociation of the tracer radioligand from each receptor if it had affinity to the
 receptors. Measurement of the dissociated ratio of the radioligand indicated the binding affinity
 between PTH1R and each unlabeled ligand.

422

423 cAMP assay

To measure intracellular cAMP production, SGS-72 cells, derived from SaOS-2 cells and stably expressing the Glosensor cAMP reporter, were utilized to measure intracellular cAMP production. The detection of cAMP-dependent expression was performed using an Envision plate reader (PerkinElmer, Waltham, MA, USA), based on luciferase-based luminescence, as previously described by Maeda, *et al.* (53).

429

430 Animal model used in the study

431 CD1 female mice were purchased from Charles River Laboratories (Massachusetts, USA), 432 and all animal care and experimental procedures were conducted under the guidelines set by the 433 Institutional Animal Care and Use Committee (IACUC) at Massachusetts General Hospital (MGH). 434 The mice were housed in a specific pathogen-free environment, with 4-5 mice per cage, under a 12-435 hour light cycle at a temperature of 22±2°C.

436 C57BL/6N female mice were purchased from KOATECH (Gyeonggi-do, Republic of Korea), 437 and all animal care and experimental procedures were conducted under the guidelines set by the 438 Institutional Animal Care and Use Committees of Kyungpook National University (KNU-2021-0101). 439 The mice were housed in a specific pathogen-free environment, with 4-5 mice per cage, under a 12-440 hour light cycle at a temperature of 22 \pm 2°C. They were provided with standard rodent chow and 441 water ad libitum.

442 C57BL6 female mice aged 8-10 weeks were used to establish an ovariectomizing (OVX) 443 mouse model of osteoporosis. The mice were divided into the following four groups (n = 6 mice/group) 444 as follows: sham, OVX control group, OVX + PTH (1-34) treated group (40 µg/kg/day), and OVX +

dimeric ^{R25C}PTH treated group (40-80 µg/kg/day). The OVX mice were allowed to recover for 4 weeks.
Subsequently, PTH (1-34) and ^{R25C}PTH were subcutaneously administered 5 days per week for 4
weeks. After the treatment period, the mice were sacrificed, and µ-CT and histological analyses were
conducted.

449

450 Acute injections

The peptides PTH(1-34) and dimeric ^{R25C}PTH(1-34) were diluted in a solution comprising 0.05% Tween 80, 10 mM citric acid, and 150 mM NaCl at a pH of 5.0. Intravenous injections of these peptides were administered at doses ranging from 50-100 µg/kg into 9-week-old CD1 female mice. As a control, mice received only the vehicle. Plasma cAMP levels were assessed before and after peptide or vehicle injection at 15-minute intervals, following the methodology detailed by Maeda, *et al.* (53). Blood-ionized calcium, plasma phosphate, and urine phosphate levels were measured before and after the injection at 1-hour intervals.

458

459 Calvarial injection mouse model

460 C57BL/6 male mice (8-week-old) were divided into the following three groups (n = 6mice/group): control, PTH (1-34) treated group (80 µg/kg/day), and ^{R25C}PTH treated group (160 461 µg/kg/day). Subcutaneous injections of the respective drugs were administered once daily for 6 days. 462 463 On the sixteenth day, the mice were sacrificed, and their bone tissues were harvested and fixed in 10 % 464 formaldehyde at 4°C. The fixed bone tissues were then decalcified in PBS (pH 7.4) containing 0.5 465 moles of ethylenediaminetetraacetic acid (EDTA). Following decalcification, the tissues were 466 embedded in paraffin, and paraffinized tissues were sectioned to a thickness of 5-7 µm. Histological 467 analysis was performed using the sectioned tissue slides stained with hematoxylin and eosin (H&E). 468 The area of new bone formation, which displays a more intense coloration compared to the existing 469 bone tissue, was examined.

470

471 μ-CT analysis

472 Mouse femurs were fixed in a 4 % paraformaldehyde solution for 24 hours at 4°C. In µ-CT, 473 we used the Quantum FX µ-CT (Perkin Elmer, Waltham, MA, USA). The images were acquired at a 474 9.7 µm voxel resolution, with settings of 90 kV and 200 µA, a 10 mm field of view, and a 3-minute 475 exposure time. Serial cross-sectional images were reconstructed using the Analyze 12.0 software 476 (Overland Park, KS, USA). To ensure consistent analysis, identical regions of interest (ROIs) were selected for the trabecular and cortical bones. The ROIs were positioned 0.3 mm from the bottom of 477 478 the growth plate. All bone parameters were evaluated according to the guidelines of the American 479 Society for Bone and Mineral Research (54).

480 Three-point bending test

The left femur of the mouse was immersed in 0.9 % NaCl solution, wrapped in gauze, and stored at -20°C until ready for a three-point bending test. In this test, we placed the mouse femurs on a suitable mold and set the pressure sensor at a distance that allowed for the maximum allowable pressure without interfering with the test (20.0 mm for the femur). A miniature material testing machine (Instron, MA, USA) was used for this test. The crosshead speed decreased at 1 mm/min. During the test, force-displacement data were collected to determine the maximum load and slope of the bones.

487

488 Serum biochemistry analysis

Serum bone resorption and osteogenesis marker levels, specifically the C-terminal telopeptide of type I collagen (CTX) and procollagen type I N-terminal propeptide (P1NP), were assessed in mice from the sham, OVX-control, PTH(1-34), and dimeric ^{R25C}PTH(1-34) groups by using the enzyme-linked immunosorbent assay (ELISA) according to the manufacturer's instructions. Additionally, their concentrations were determined using specific mouse CTX-1 and P1NP ELISA kits (Cloud Clone, Wuhan, China) respectively.

495

496 Bone histological analyses

497 The tibiae were initially fixed in 4% paraformaldehyde at 4°C overnight. The following day, 498 samples were decalcified using 10% ethylenediaminetetraacetic acid (EDTA) solution (pH 7.4) for 4

499 weeks at 4°C. The decalcified tibiae were then embedded in paraffin and sectioned at 3 µm thick. For 500 TRAP staining, dehydrated paraffin sections were fixed in an acetone/ethanol mixture (1:1) for 1 501 minute, followed by complete air-drying at RT for 20 minutes. Thereafter, the sections were immersed 502 in TRAP reagent for 30 minutes at 37°C. Osteoclast surface per bone surface (Oc.S/BS) and 503 Osteoclast number per bone surface (Oc.N/BS) analysis followed the ASBMR guidelines (55).

504

505 **Dynamic bone histomorphometric analysis**

506 To conduct dynamic histomorphometry analysis, we injected the mice with 25 mg/kg body 507 weight of calcein (Sigma-Aldrich) or Alizarin Red S (Sigma-Aldrich) intraperitoneally before sacrifice, 508 with 3- or 10-day intervals between injections. Briefly, femurs or vertebrates were fixed in 4 % 509 paraformaldehyde solution for 24 hours at 4°C. The following day, the samples were washed with 510 phosphate-buffered saline (PBS) solution and then dehydrated using a gradient of ethanol (50 %, 511 70 %, 85 %, 90 %, and 100 %). Subsequently, we embedded the dehydrated femurs or vertebrates in 512 methyl methacrylate (Sigma) to prepare resin blocks. The resin blocks were sectioned at 6 µm thick by using a Leica SP1600 microtome (Leica Microsystems, Germany). The fluorescence signals of 513 514 calcein (green) and Alizarin Red S (red) from ROIs were captured using a fluorescence microscope 515 (Leica, Wetzlar, Germany). For vertebral bone analysis, bone mineralization was evaluated by von 516 Kossa staining. The sections were placed in 2-methoxyethyl acetate (Sigma-Aldrich) for 20 minutes, 517 followed by rehydration with serial ethanol solutions (100 %, 90 %, 80 %, 70 %, and 50 %) and 518 distilled water for 2 minutes each. The sections were subsequently dipped in a 1 % AgNO₃ solution 519 (Sigma-Aldrich) for 5 minutes under ultraviolet (UV) light photons, washed in distilled water for 5 520 minutes, and dipped in 5 % sodium thiosulfate solution for 5 minutes to remove nonspecific binding. 521 Finally, we covered the sections with mounting solution and captured images by using a Leica 522 microscope. The parameters of dynamic bone histomorphometry were analyzed using the Bioquant 523 Osteo 2019ME program (Bioquant Osteo, Nashville, TN, USA).

524

525 Statistical analysis

526 Statistical analysis was performed in GraphPad Prism 10.1.2. The data are presented as the 527 mean \pm standard error of the means (SEM). Statistically significant differences between the two 528 groups were determined using an unpaired *t*-test. A *p*-value less than 0.05 was considered statistically 529 significant.

530 Author contributions

- 531 Conceptualization, Methodology: M.N., X.C., X.J., D-K.L., H-J.K., D.P., S.Y.L., H.L., T.J.G., J.-Y.C.,
- and S.L.; Validation, Formal Analysis: M.N., and X.C.; Investigation, Resource: M.N., X.C., X.J., D.-
- 533 K.L., and D.P.; writing Original Draft Preparation: M.N., and X.C.; Writing Review & Editing: M.N.,
- 534 X.C., T.J.G., H.L., J.-Y.C., and S.L.; Visualization: M.N., and X.C.; Supervision: H.-J.K., H.L., J.-Y.C.,
- and S.L.; Project Administration: J.-Y.C., and S.L.; Funding Acquisition: S.Y.L., H.L., J.-Y.C., and S.L.

536

537 Acknowledgments

538 This work was supported by the National Research Foundation of Korea (NRF) grant funded by the 539 Korea government (MSIT) (2022R1A2C3006002 S.L.; 2022R1F1A1074610 and to 540 2022R1A4A1025913 H.L.; RS-2023-00217798 2021R1A2C3003675 to and S.Y.L.; 541 2022R1A2C1006105 to J.-Y.C.), Gachon University Gil Medical Center (FRD2023-12 to S.L.), and 542 Korea University Grants.

543

545 **References**

- T. Sozen, L. Ozisik, N. C. Basaran, An overview and management of osteoporosis. *Eur J Rheumatol* 4, 46-56 (2017).
- 548 2. N. Salari *et al.*, The global prevalence of osteoporosis in the world: a comprehensive 549 systematic review and meta-analysis. *J Orthop Surg Res* **16**, 609 (2021).
- S. Papapoulos *et al.*, Five years of denosumab exposure in women with postmenopausal
 osteoporosis: results from the first two years of the FREEDOM extension. *J Bone Miner Res*27, 694-701 (2012).
- 553 4. D. M. Reid *et al.*, Zoledronic acid and risedronate in the prevention and treatment of 554 glucocorticoid-induced osteoporosis (HORIZON): a multicentre, double-blind, double-dummy, 555 randomised controlled trial. *Lancet* **373**, 1253-1263 (2009).
- 556 5. M. R. McClung *et al.*, Romosozumab in postmenopausal women with low bone mineral 557 density. *N Engl J Med* **370**, 412-420 (2014).
- 558 6. T. J. Martin, N. A. Sims, E. Seeman, Physiological and pharmacological roles of PTH and 559 PTHrP in bone using their shared receptor, PTH1R. *Endocrine reviews* **42**, 383-406 (2021).
- 560 7. C. Bieglmayer, G. Prager, B. Niederle, Kinetic analyses of parathyroid hormone clearance as
 561 measured by three rapid immunoassays during parathyroidectomy. *Clinical Chemistry* 48,
 562 1731-1738 (2002).
- 563 8. G. W. Maier *et al.*, Parathyroid hormone after adenectomy for primary hyperparathyroidism. A
 564 study of peptide hormone elimination kinetics in humans. *J Clin Endocrinol Metab* 83, 3852565 3856 (1998).
- J. L. Vahle *et al.*, Skeletal changes in rats given daily subcutaneous injections of recombinant
 human parathyroid hormone (1-34) for 2 years and relevance to human safety. *Toxicol Pathol*30, 312-321 (2002).
- 569 10. N. M. Appelman-Dijkstra, S. E. Papapoulos, From disease to treatment: from rare skeletal 570 disorders to treatments for osteoporosis. *Endocrine* **52**, 414-426 (2016).
- 571 11. S. Khosla, L. C. Hofbauer, Osteoporosis treatment: recent developments and ongoing
 572 challenges. *Lancet Diabetes Endocrinol* 5, 898-907 (2017).
- 573 12. E. Moallem, R. Kilav, J. Silver, T. Naveh-Many, RNA-Protein binding and post-transcriptional

574 regulation of parathyroid hormone gene expression by calcium and phosphate. *J Biol Chem*575 **273**, 5253-5259 (1998).

- J. Silver, J. Russell, L. M. Sherwood, Regulation by vitamin D metabolites of messenger
 ribonucleic acid for preproparathyroid hormone in isolated bovine parathyroid cells. *Proc Natl Acad Sci U S A* 82, 4270-4273 (1985).
- 579 14. R. A. Chen, W. G. Goodman, Role of the calcium-sensing receptor in parathyroid gland 580 physiology. *Am J Physiol Renal Physiol* **286**, F1005-1011 (2004).
- 581 15. J. F. Habener, M. Rosenblatt, J. T. Potts, Jr., Parathyroid hormone: biochemical aspects of 582 biosynthesis, secretion, action, and metabolism. *Physiol Rev* **64**, 985-1053 (1984).
- 583 16. V. Veldurthy *et al.*, Vitamin D, calcium homeostasis and aging. *Bone Res* **4**, 16041 (2016).
- 584 17. P. R. Kiela, F. K. Ghishan, Recent advances in the renal-skeletal-gut axis that controls 585 phosphate homeostasis. *Lab Invest* **89**, 7-14 (2009).
- 18. H. Jüppner *et al.*, AG protein-linked receptor for parathyroid hormone and parathyroid
 hormone-related peptide. *Science* 254, 1024-1026 (1991).
- T. J. Gardella, H. Ju[°] ppner, Interaction of PTH and PTHrP with their receptors. *Reviews in Endocrine and Metabolic Disorders* 1, 317-329 (2000).
- 590 20. L. S. Weinstein, S. Yu, D. R. Warner, J. Liu, Endocrine manifestations of stimulatory G protein 591 alpha-subunit mutations and the role of genomic imprinting. *Endocr Rev* **22**, 675-705 (2001).
- T. N. Feinstein *et al.*, Retromer terminates the generation of cAMP by internalized PTH
 receptors. *Nat Chem Biol* **7**, 278-284 (2011).
- A. lida-Klein *et al.*, Mutations in the second cytoplasmic loop of the rat parathyroid hormone
 (PTH)/PTH-related protein receptor result in selective loss of PTH-stimulated phospholipase
 C activity. *J Biol Chem* 272, 6882-6889 (1997).
- R. Dunlay, K. Hruska, PTH receptor coupling to phospholipase C is an alternate pathway of
 signal transduction in bone and kidney. *Am J Physiol* **258**, F223-231 (1990).
- A.-B. Abou-Samra *et al.*, Expression cloning of a common receptor for parathyroid hormone
 and parathyroid hormone-related peptide from rat osteoblast-like cells: a single receptor
 stimulates intracellular accumulation of both cAMP and inositol trisphosphates and increases
 intracellular free calcium. *Proceedings of the National Academy of Sciences* **89**, 2732-2736
 (1992).

H. G. Bone *et al.*, Ten years' experience with alendronate for osteoporosis in postmenopausal
women. *N Engl J Med* **350**, 1189-1199 (2004).

- M. R. Rubin *et al.*, Therapy of Hypoparathyroidism With PTH(1-84): A Prospective Six Year
 Investigation of Efficacy and Safety. *J Clin Endocrinol Metab* **101**, 2742-2750 (2016).
- 608 27. K. K. Winer, J. A. Yanovski, G. B. Cutler, Jr., Synthetic human parathyroid hormone 1-34 vs 609 calcitriol and calcium in the treatment of hypoparathyroidism. *JAMA* **276**, 631-636 (1996).
- K. K. Winer *et al.*, Long-Term Parathyroid Hormone 1-34 Replacement Therapy in Children
 with Hypoparathyroidism. *J Pediatr* 203, 391-399 e391 (2018).
- M. Okazaki *et al.*, Prolonged signaling at the parathyroid hormone receptor by peptide ligands
 targeted to a specific receptor conformation. *Proceedings of the National Academy of Sciences* 105, 16525-16530 (2008).
- A. A. Pioszak, N. R. Parker, T. J. Gardella, H. E. Xu, Structural basis for parathyroid hormonerelated protein binding to the parathyroid hormone receptor and design of conformationselective peptides. *J Biol Chem* 284, 28382-28391 (2009).
- 81. R. W. Cheloha, S. H. Gellman, J. P. Vilardaga, T. J. Gardella, PTH receptor-1 signallingmechanistic insights and therapeutic prospects. *Nat Rev Endocrinol* **11**, 712-724 (2015).
- S. R. Hoare, T. J. Gardella, T. B. Usdin, Evaluating the signal transduction mechanism of the
 parathyroid hormone 1 receptor. Effect of receptor-G-protein interaction on the ligand binding
 mechanism and receptor conformation. *J Biol Chem* 276, 7741-7753 (2001).
- 33. T. Dean, J. P. Vilardaga, J. T. Potts, Jr., T. J. Gardella, Altered selectivity of parathyroid
 hormone (PTH) and PTH-related protein (PTHrP) for distinct conformations of the PTH/PTHrP
 receptor. *Mol Endocrinol* 22, 156-166 (2008).
- S. Lee *et al.*, A Homozygous [Cys25]PTH(1-84) Mutation That Impairs PTH/PTHrP Receptor
 Activation Defines a Novel Form of Hypoparathyroidism. *J Bone Miner Res* 30, 1803-1813
 (2015).
- 629 35. C. H. Bae *et al.*, A novel human PTH analog [Cys25] hPTH (1–34) restores bone mass in
 630 ovariectomized mice. *The Journal of Clinical Endocrinology & Metabolism* **101**, 3700-3708
 631 (2016).
- 83. R. R. Banerjee, M. A. Lazar, Dimerization of resistin and resistin-like molecules is determined
 by a single cysteine. *J Biol Chem* 276, 25970-25973 (2001).

M. V. Trivedi, J. S. Laurence, T. J. Siahaan, The role of thiols and disulfides on protein stability.
 Curr Protein Pept Sci 10, 614-625 (2009).

- 636 38. B. Kemper, J. F. Habener, R. C. Mulligan, J. T. Potts, Jr., A. Rich, Pre-proparathyroid hormone:
- a direct translation product of parathyroid messenger RNA. *Proc Natl Acad Sci U S A* **71**,
 3731-3735 (1974).
- 639 39. T. J. Vasicek *et al.*, Nucleotide sequence of the human parathyroid hormone gene. *Proc Natl*640 *Acad Sci U S A* **80**, 2127-2131 (1983).
- 641 40. K. M. Wiren *et al.*, Mutations in signal sequence cleavage domain of preproparathyroid
 642 hormone alter protein translocation, signal sequence cleavage, and membrane-binding
 643 properties. *Mol Endocrinol* **3**, 240-250 (1989).
- H. N. Rosen *et al.*, Serum CTX: a new marker of bone resorption that shows treatment effect
 more often than other markers because of low coefficient of variability and large changes with
 bisphosphonate therapy. *Calcif Tissue Int* 66, 100-103 (2000).
- F. Pagani, C. M. Francucci, L. Moro, Markers of bone turnover: biochemical and clinical
 perspectives. *J Endocrinol Invest* 28, 8-13 (2005).
- G. Wheater, M. Elshahaly, S. P. Tuck, H. K. Datta, J. M. van Laar, The clinical utility of bone
 marker measurements in osteoporosis. *J Transl Med* **11**, 201 (2013).
- 44. S. L. Andersen, A. L. Frederiksen, A. B. Rasmussen, M. Madsen, A. R. Christensen,
 Homozygous missense variant of PTH (c.166C>T, p.(Arg56Cys)) as the cause of familial
 isolated hypoparathyroidism in a three-year-old child. *J Pediatr Endocrinol Metab* 35, 691-694
 (2022).
- A. Arnold *et al.*, Mutation of the signal peptide-encoding region of the preproparathyroid
 hormone gene in familial isolated hypoparathyroidism. *J Clin Invest* **86**, 1084-1087 (1990).
- b. B. Parkinson, R. V. Thakker, A donor splice site mutation in the parathyroid hormone gene
 is associated with autosomal recessive hypoparathyroidism. *Nat Genet* 1, 149-152 (1992).
- T. Sunthornthepvarakul, S. Churesigaew, S. Ngowngarmratana, A novel mutation of the signal
 peptide of the preproparathyroid hormone gene associated with autosomal recessive familial
 isolated hypoparathyroidism. *J Clin Endocrinol Metab* 84, 3792-3796 (1999).
- 48. D. A. Ertl, S. Stary, B. Streubel, A. Raimann, G. Haeusler, A novel homozygous mutation in
 the parathyroid hormone gene (PTH) in a girl with isolated hypoparathyroidism. *Bone* **51**, 629-

- 664 632 (2012).
- 49. L. J. Clark *et al.*, Allosteric interactions in the parathyroid hormone GPCR-arrestin complex
 formation. *Nat Chem Biol* 16, 1096-1104 (2020).
- 667 50. A. Kumar, M. Baumann, J. Balbach, Small Molecule Inhibited Parathyroid Hormone Mediated
 668 cAMP Response by N-Terminal Peptide Binding. *Sci Rep* 6, 22533 (2016).
- L. H. Zhao *et al.*, Structure and dynamics of the active human parathyroid hormone receptor-1. *Science* **364**, 148-153 (2019).
- 52. X. Zhai *et al.*, Molecular insights into the distinct signaling duration for the peptide-induced
 PTH1R activation. *Nat Commun* 13, 6276 (2022).
- A. Maeda *et al.*, Critical role of parathyroid hormone (PTH) receptor-1 phosphorylation in
 regulating acute responses to PTH. *Proc Natl Acad Sci U S A* **110**, 5864-5869 (2013).
- 675 54. M. L. Bouxsein *et al.*, Guidelines for assessment of bone microstructure in rodents using 676 micro-computed tomography. *J Bone Miner Res* **25**, 1468-1486 (2010).
- 55. D. W. Dempster *et al.*, Standardized nomenclature, symbols, and units for bone
 histomorphometry: a 2012 update of the report of the ASBMR Histomorphometry
 Nomenclature Committee. *J Bone Miner Res* 28, 2-17 (2013).
- 680

682 Figures and Tables

683



Figure 1. Formation of R25C mutant PTH(1-84) dimer.

(A) Schematic representation of PTH gene structure and expression. (B) Schematic representation of
 ^{R56C}pre-pro-PTH(1-115) (In mature form, ^{R25C}PTH(1-84)) gene structure and expression (C) Schematic
 representation of recombinant PTH proteins (D) *In vitro* dimerization of ^{R25C}PTH. Recombinant protein
 constructs were transfected into HEK293T cells, and expression of PTH-3xFLAG and
 ^{R25C}PTH3xFLAG in culture medium or cell lysate was demonstrated by western blot. The result
 confirms the presence of dimeric R25CPTH. (*bp: base pairs; *AA: amino acids; *RA: reducing agent)

692

684



694

695 Figure 2. Effect of PTH, monomeric ^{R25C}PTH, and dimeric ^{R25C}PTH to the PTH1R *in vitro*.

(A) The binding of PTH(1-34), monomeric ^{R25C}PTH(1-34), and dimeric ^{R25C}PTH(1-34) to the PTH1R in
R⁰ conformation of RG conformation was assessed by competition methods using ¹²⁵I-PTH(1-34) and
¹²⁵I-MPTH(1-15) as radioligand. (B) Ligand potency for cAMP signaling was assessed in SGS-72 cells,
which were derived from SaOS2 cells modified to express Glosensor cAMP reporter. The cells were
preloaded with luciferin and treated with varying concentrations of PTH(1-34), monomeric ^{R25C}PTH(1-34), and dimeric ^{R25C}PTH(1-34).



703

704 Figure 3. Calcemic and phosphatemic responses by PTH injection in CD1 female mice.

(A) Plasma Calcemic Response after Injection (n = 6). Both PTH(1-34) and dimeric R25C PTH(1-34) 705 706 significantly elevate ionized calcium levels in plasma at 1 to 2 hours post-injection. After 2 hours post-707 injection, plasma ionized calcium level gradually restored to baseline levels similar to those of the 708 vehicle group. (B) Plasma Phosphatemic Response after Injection (n = 12). Following PTH(1-34) 709 injection, plasma phosphate levels significantly decrease at 1-hour post-injection, subsequently 710 returning to baseline akin to those of the vehicle group. Conversely, dimeric ^{R25C}PTH(1-34) injection 711 shows no significant alteration in phosphatemic response but demonstrates a tendency towards a 712 slight decrease in phosphate levels, gradually restoring to baseline levels akin to those of the vehicle 713 group. (C) Urine Phosphatemic Response after Injection (n = 6). The urine phosphate levels markedly increased at 1 hour post-injection for both PTH(1-34) and dimeric ^{R25C}PTH(1-34), followed by a return 714 715 to baseline levels akin to those of the vehicle group. This analysis was conducted using 9-week-old 716 female CD1 mice. The mice were administered PTH(1-34) and dimeric ^{R25C}PTH(1-34) at a 717 concentration of 50 nmol/kg for each compound. Error bars represent mean ± standard error. p-values 718 were determined using the t-test. * denotes p-value < 0.05 for PTH(1-34) compared to vehicle, ** 719 denotes p-value < 0.01 for PTH(1-34) compared to vehicle, # denotes p-value < 0.05 for dimeric ^{R25C}PTH(1-34) compared to vehicle, ## denotes p-value < 0.01 for dimeric ^{R25C}PTH(1-34) compared to 720 vehicle, ### denotes *p*-value < 0.001 for dimeric R25C PTH(1-34) compared to vehicle. 721



722

Figure 4. Effect of cAMP production by PTH injection in CD1 female mice.

724 (A) Comparison of cAMP levels induced following the injection of PTH(1-34) (50 nmol/kg) and two concentrations (50, 100 nmol/kg) of dimeric ^{R25C}PTH(1-34). cAMP levels were assessed both before 725 726 and 15 minutes after injection, confirming induced cAMP production exclusively by PTH(1-34). (B) To compare the amount of cAMP generated after injecting PTH(1-34) and two concentrations of dimeric 727 R25C PTH(1-34) more precisely, we measured cAMP levels in a group of 6 mice (n = 6) before injection 728 729 and at 6, 15, 30, and 60 minutes after injection. We observed that cAMP production was induced only 730 by PTH(1-34), and at the 15-minute time point, the cAMP levels reached their peak and gradually 731 decreased thereafter. Female CD1 mice at 9 weeks old were used for each analysis. The error bars 732 indicate mean \pm standard error. p-values were obtained using the t-test. * indicates p-value < 0.05 733 against vehicle.

734



736

737 Figure 5. Effect of dimeric ^{R25C}PTH(1-34) in calvarial injection model

738 (A) Dissections of the calvarial bones. Calvarial injections were performed on eight-week-old male 739 C57BL/6 mice (N = 6 per group) that received daily administrations of vehicle, PTH(1-34), or dimeric R25CPTH(1-34) for six days. Following a 10-day treatment period, histological sections of calvariae, 740 741 stained with hematoxylin (pink; representing bone matrix) and eosin (blue-purple; indicating cell 742 nuclei), were obtained. The area of new bone formation, with more intense staining compared to the existing bone tissue, is denoted by the dotted line. The bar indicates 50 µm. (B) Quantification of new 743 744 bone width in calvarial injection model. The result showed a significant increase in new bone width following injections of both PTH(1-34) and dimeric ^{R25C}PTH(1-34) compared to the vehicle group. ** 745 746 indicates p-value <0.01 against vehicle, *** indicates p-value <0.001 against vehicle.



748

Figure 6. Impact of ^{R25C}PTH(1-34) on Bone Turnover.

The effects of Sham, OVX-Control (OVX + vehicle), OVX treated with PTH(1-34) (OVX + PTH(1-34)), 750 and OVX treated with dimeric ^{R25C}PTH(1-34) (OVX + dimeric ^{R25C}PTH(1-34)) on bone turnover in mice. 751 752 (A) Femurs obtained from mice in each group were subjected to µCT analyses for the assessment of 753 bone mass. (B) Several parameters of (A) were quantified using µCT measurements, including 754 trabecular bone mineral density (Tb.BMD), trabecular bone volume to tissue volume (Tb.BV/TV), 755 trabecular bone thickness (Tb.Th), trabecular number (Tb.N), trabecular separation (Tb.Sp), cortical bone mineral density (Ct.BMD), cortical bone volume to tissue volume (Ct.BV/TV), cortical thickness 756 757 (Ct.Th), and cortical area to total tissue area (Ct.Ar/Tt.Ar). (C) A 3D-point bending test was conducted 758 with femurs obtained from mice in each group. The left panel describes a schematic model of a 3D-

759 point bending test. The middle and right panels each indicate the maximum bending load (kgf) and 760 slope (kgf/mm). (D) Serum levels of calcium, phosphorus, CTX, P1NP, and ALP were measured for 761 each group using an ELISA assay. (E) TRAP staining of histological sections of proximal tibias was 762 carried out to visualize osteoclast activity. The scale bars represent 100 µm (F) Quantification of 763 osteoclast number per bone surface (Oc.N/BS), and osteoclast surface per bone surface (Oc.S/BS) 764 was performed. Each group consisted of six samples (n = 6). The error bars indicate mean \pm standard 765 error. *p*-values were obtained using the *t*-test to compare the mean of each column with the mean of a 766 control column. * indicates p-value < 0.05, ** indicates p-value < 0.01, *** indicates p-value < 0.001, 767 **** indicates *p*-value < 0.0001.



770 Figure 7. Impact of ^{R25C}PTH(1-34) on Osteoblast Function in Vertebrae

769

771 The effects of Sham, OVX-Control (OVX + vehicle), OVX treated with PTH(1-34) (OVX + PTH(1-34)), and OVX treated with dimeric ^{R25C}PTH(1-34) (OVX + dimeric ^{R25C}PTH(1-34)) on osteoblast function in 772 773 mice. (A) Mineralization of vertebrae obtained from each group was assessed through Von Kossa staining. (B) Quantification of trabecular bone parameters including trabecular bone volume to tissue 774 775 volume (Tb.BV/TV), trabecular number (Tb.N), and trabecular separation (Tb.Sp) was performed 776 using the Bioquant Osteo 2019 v19.9.60 program. Each bar represents 500 µm. (C) Fluorescent 777 microscopic observations of trabecular and cortical bone sections from each group demonstrate the 778 apposition of xylenol (red) and calcein (green) labels. (D) Quantification of trabecular bone 779 parameters such as trabecular bone mineral apposition rate (Tb.MAR), trabecular bone formation rate 780 to bone surface (Tb.BFR/BS), cortical bone MAR (Ct.MAR), and trabecular MAR (Tb.MAR) was 781 carried out using the Bioquant Osteo 2019 v19.9.60 program. Each group consisted of five samples

- 782 (n = 5). The error bars indicate mean \pm standard error. *p*-values were obtained using the *t*-test
- compare the mean of each column with the mean of a control column. * indicates *p*-value < 0.05, **
- indicates *p*-value < 0.01, *** indicates *p*-value < 0.001, **** indicates *p*-value < 0.0001.





- PTH(1-34)
 Monomeric ^{R25C}PTH(1-34)
- Dimeric ^{R25C}PTH(1-34)



Α



-- Vehicle PTH(1-34) Dimeric ^{R25C}PTH(1-34)

0:45

1:00



Α





В



Vehicle

PTH(1-34)

Dimeric ^{R25C}PTH(1-34)















