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- Connexin Hemichannels with Prostaglandin Release in Anabolic Function of Bone to
 Mechanical Loading
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20 Abstract

21	Mechanical stimulation, such as physical exercise, is essential for bone formation and health. Here,
22	we demonstrate the critical role of osteocytic Cx43 hemichannels in anabolic function of bone in
23	response to mechanical loading. Two transgenic mouse models, R76W and Δ 130-136, expressing
24	dominant-negative Cx43 mutants in osteocytes were adopted. Mechanical loading of tibial bone
25	increased cortical bone mass and mechanical properties in wild-type and gap junction-impaired
26	R76W mice through increased PGE ₂ , endosteal osteoblast activity, and decreased sclerostin. These
27	anabolic responses were impeded in gap junction/hemichannel-impaired $\Delta 130-136$ mice and
28	accompanied by increased endosteal osteoclast activity. Specific inhibition of Cx43 hemichannels by
29	Cx43(M1) antibody suppressed PGE ₂ secretion and impeded loading-induced endosteal osteoblast
30	activity, bone formation and anabolic gene expression. PGE2 administration rescued the osteogenic
31	response to mechanical loading impeded by impaired hemichannels. Together, osteocytic Cx43
32	hemichannels could be a potential new therapeutic target for treating bone loss and osteoporosis.
33	

34 Introduction

Bone as a mechanosensitive tissue is adaptive to mechanical stimuli, which are essential for 35 bone homeostasis, formation, and remodeling (Bonewald, 2011). Reduced mechanical stimulation 36 37 leads to bone loss and elevated risk of fracture (Lang et al., 2004), while enhanced mechanical stimulation, such as physical exercise, has positive, anabolic impacts on bone tissue, even following 38 39 a prolonged cessation of stimulation (Erlandson et al., 2012; Warden et al., 2007). The osteocytes 40 embedded in the bone mineral matrix comprise over 90–95% of all bone cells and are thought to be a major mechanoreceptor in the adult skeleton (Bonewald, 2011). Osteocytes detect the mechanical 41 loading-induced alterations of the bone matrix microenvironment and translate them into biological 42 43 responses to regulate osteoblast and osteoclast activity on the bone surface (Bonewald, 2011; Bonewald and Johnson, 2008). 44

Connexin (Cx)-forming gap junctions and hemichannels permit small molecules (≤ 1 kDa) to 45 pass through the cellular membrane, such as prostaglandin E_2 (PGE₂) and ATP (Loiselle et al., 2012). 46 47 Among Cx family members, Cx43 is the predominant Cx subtype expressed in osteocytes (Civitelli, 2008). Cx43 gap junctions allow cell-cell communication between osteocytes or between osteocytes 48 49 and other bone cell types (Ishihara et al., 2008), and mechanical stimuli increase communication 50 between two adjacent cells through gap junctions (Alford et al., 2003; Cheng et al., 2001). However, osteocytic Cx43 gap junctions are only active at the tips of osteocyte dendritic processes and remain 51 open even without mechanical stimulation (Cusato et al., 2006). In contrast, Cx43 hemichannels, 52 53 which mediate the communication between the intracellular and the extracellular microenvironment, 54 are highly responsive to mechanical stimulation in osteocytes (Cherian et al., 2005; Jiang and

55	Cherian, 2003). Our previous studies have shown that in vitro mechanical stimulation, through fluid
56	flow shear stress (FFSS), increases cell surface expression of Cx43 hemichannels (Cherian et al.,
57	2005; Jiang and Cherian, 2003; Siller-Jackson et al., 2008), and opens Cx43 hemichannels, leading
58	to the release of anabolic factor, PGE_2 in osteocytes (Cherian et al., 2005; Siller-Jackson et al., 2008).
59	Activation of integrins and PI3K-Akt signaling by FFSS plays an essential role in activating Cx43
60	hemichannels (Batra et al., 2012; Batra et al., 2014). PGE ₂ released by Cx43 hemichannels acts in an
61	autocrine/paracrine manner to promote gap junction communication through transcriptional
62	regulation of Cx43 (Xia et al., 2010) and blocks glucocorticoid-induced osteocyte apoptosis (Kitase
63	et al., 2010). The opening of Cx43 hemichannels by FFSS also triggers the release of ATP by a
64	protein kinase C-mediated pathway in osteocytes (Genetos et al., 2007). Extracellular PGE2
65	accumulation caused by continuous FFSS exerts a negative feedback, leading to hemichannel closure
66	(Riquelme et al., 2015). However, the biological role of osteocytic Cx43 hemichannels in the
67	anabolic function of mechanical loading has remained largely elusive.
68	Several bone cell type-specific Cx43 conditional knockout (cKO) mouse models have been

Several bone cell type-specific Cx43 conditional knockout (cKO) mouse models have been 68 reported. Deletion of Cx43 from osteoblasts and osteocytes driven by the Col-2.3-kb a1(I) collagen 69 promoter (Col-2.3kb-Cre; Cx43^{-/flx}) attenuated tibial endosteal response to non-physiological 70 71 mechanical loading, induced by four-point (Grimston et al., 2006) or three-point tibial bending 72 (Grimston et al., 2008). However, deletion of Cx43 in osteochondroprogenitors driven by the (Dermo1-Cre; $Cx43^{-/flx}$)(Grimston et 73 collagen promoter Dermo1 al., 2012) or in osteoblasts/osteocytes driven by the osteocalcin promoter (OCN-Cre; Cx43^{flx/flx})(Zhang et al., 2011) 74 75 showed an enhanced tibial periosteal response to tibial axial compression (Grimston et al., 2012) or

76 tibial cantilever bending (Zhang et al., 2011). Similarly, deletion of Cx43 in osteocytes driven by an 8-kb dentin matrix protein 1 (DMP1) promoter (DMP1-8kb-Cre; Cx43^{flx/flx}) showed enhanced 77 β-catenin levels and correspondingly increased periosteal response to ulna compression (Bivi et al., 78 2013). Interestingly, endosteal bone formation decreased more in Dermo1-Cre; Cx43^{-/flx} mice 79 (Grimston et al., 2012), but did not change in DMP1-8kb-Cre; Cx43^{flx/flx} mice (Bivi et al., 2013) 80 81 during mechanical loading. Together, these findings suggest that Cx43 plays a distinct role in the 82 adaptive response to bone loading. However, since Cx43 forms both gap junctions and hemichannels, 83 it has remained largely elusive whether the responses in knockout models could be attributed to either or both types of Cx43-forming channels. Here we dissect the distinctive roles of Cx43 gap 84 85 junctions and hemichannels using two transgenic mouse models that overexpress dominant-negative Cx43 mutants primarily in osteocytes with the 10 kb DMP1 promoter. The R76W transgenic mouse 86 inhibits gap junctions with enhanced hemichannel function, whereas, $\Delta 130-136$ inhibits both gap 87 junctions and hemichannels (Xu et al., 2015). To further delineate the role of osteocytic Cx43 88 hemichannels under mechanical loading in vivo, a monoclonal Cx43(M1) antibody that specifically 89 90 blocks Cx43 hemichannels was developed. In this study, we unveil a novel role of Cx43 91 hemichannels in osteocytes and their release of PGE₂ in mediating anabolic function of the bone in 92 response to mechanical loading.

93

94 **Results**

Impairment of Cx43 hemichannels attenuate anabolic responses of tibial bone to mechanical
 loading

In this study, we used two transgenic mouse models to distinguish the roles of osteocytic Cx43-gap junction channels and hemichannels in osteocytes in bone response to mechanical loading. We injected EB dye into mouse tail veins to determine the activity of Cx43 hemichannels in WT and transgenic mice in response to axial tibial loading. Bone tissue sections around the tibial midshaft region showed that tibial loading increased EB dye uptake in the osteocytes of WT and R76W mice,

102 but not in the osteocytes of Δ 130-136 mice (**Figure 1-figure supplement 2A, B**).

103 We subjected WT and transgenic mice with similar body weights (Figure 1-figure supplement 104 **3A**) to a 2-week cyclic tibial loading regime. μ CT analyses of tibial metaphyseal trabecular bone 105 showed that loading increased bone volume fractions (BV/TV) in WT and R76W mice (Figure 1A). 106 In contrast, compared to contralateral, unloaded controls, tibial loading of Δ 130-136 mice exhibited a 107 significant reduction of trabecular number (Tb.N) and bone mineral density (BMD), as well as 108 increased trabecular separation (Tb.Sp) during mechanical loading (Figure 1B, C, E). However, 109 compared to contralateral, unloaded tibias, trabecular thickness (Tb.Th) was increased in loaded 110 tibias of WT and two transgenic mice (Figure 1D). There was no change of structural model index 111 (SMI), indicating that loading did not affect the shape of trabecular bone (Figure 1F). Representative 112 μ CT images of trabecular bone are shown in **Figure 1G**.

Similar attenuation of anabolic responses to tibial loading was also observed in cortical bone. μ CT analysis was conducted at the midshaft cortical bone (50% site). Loading increased bone area (B.Ar), bone area fraction (B.Ar/T.Ar), and cortical thickness (Ct.Th) in WT and R76W mice (**Figure 2B, C, E**). Although T.Ar was increased by mechanical loading in Δ 130-136 (**Figure 2A**), enlarged bone marrow area (M.Ar) (**Figure 2D**) attenuated the ratio of B.Ar/T.Ar (**Figure 2C**). The

118	increased Ct.Th. due to tibial loading was not observed in $\Delta 130-136$ mice (Figure 2E). Interestingly,
119	the loading caused a decrease of BMD in R76W mice (Figure 2F). Torsional strength, predicted by
120	polar moment of inertia (pMOI), was increased as a result of mechanical loading in WT and R76W,
121	but not in Δ 130-136 mice (Figure 2G). Representative images of cortical bone are shown in Figure
122	2H. Together these data suggested that osteocytic Cx43 hemichannels, not gap junctions, play an
123	important role in anabolic responses of both trabecular and cortical bones to mechanical loading.

124

125 Cx43 hemichannels mediate endosteal osteogenic responses to mechanical loading.

126 Dynamic histomorphometric analyses were performed to evaluate periosteal and endosteal bone 127 formation in response to tibial loading. Loading caused a significant increase of endosteal MAR, 128 MS/BS, and BRF/BS compared to contralateral tibias in WT and R76W, but such an increase was 129 not observed in $\triangle 130-136$ (Figure 3A-D). The decreased endosteal bone formation may partially 130 account for the enlarged bone marrow. Contrary to the endosteal surface, Δ 130-136 mice showed a 131 statistically significant osteogenic response on the periosteal surface compared to WT and R76W mice, manifesting a threefold increase in MAR, MS/BS, and BRF/BS during loading (Figure 3E-H). 132 133 Together, these data suggested that impaired osteocytic Cx43 hemichannels in Δ 130-136 mice 134 attenuated endosteal bone formation, and enhanced periosteal bone formation upon mechanical 135 loading.

136

137 Impaired Cx43 hemichannels inhibit the loading-induced PGE₂ secretion and osteoblast
 138 activity, and promote osteoclast activity.

139	Cx43 hemichannels mediate PGE ₂ release from osteocytes induced by FFSS in vitro (Cherian et al.,
140	2005), and extracellular PGE_2 is reported to play a key role in the anabolic response to mechanical
141	loading of bone tissue (Jee et al., 1985; Thorsen et al., 1996). We measured PGE ₂ levels in the tibial
142	bone diaphysis and found PGE ₂ levels in mechanically loaded tibias were significantly increased in
143	WT and R76W mice compared to those in contralateral, non-loaded tibias (Figure 4A). However,
144	loading had minimal effect on PGE ₂ levels in Δ 130-136 mice. Immunohistochemical staining
145	showed that the expression of cyclooxygenase-2 (COX-2), a key enzyme that catalyzes the
146	conversion of arachidonic acid to prostaglandins, was significantly increased in the osteocytes of
147	loaded tibias of WT and R76W mice compared to contralateral, unloaded tibias (Figure 4B, C and
148	Figure 4-figure supplement 1A). However, the increase of COX-2 in loaded tibias was not
149	observed in Δ 130-136 mice. Sost-positive osteocytes decreased significantly in WT and R76W in
150	response to tibial loading, while such decrease in loaded tibias was absent in Δ 130-136 mice (Figure
151	4D, E and Figure 4-figure supplement 1B). A similar reduction at the mRNA level of the bone was
152	also found in WT and R76W, but absent in Δ 130-136 mice (Figure 4F). Since Sost, a Wnt receptor
153	antagonist, is a potent inhibitor of osteoblastic activity, we examined osteoblasts on the endosteal
154	surface. WT and R76W mice exhibited an increase of osteoblast numbers on the endosteal surface; in
155	contrast, this increase was absent in Δ 130-136 mice (Figure 4G, H). Moreover, the levels of mRNA
156	of osteoblastic markers Runx2 and Bglap were higher in the bone of loaded WT and R76W than
157	loaded Δ 130-136 mice (Figure 4I, J). The mRNA expression of the osteocytic marker Dmp1 in the
158	bone of Δ 130-136 mice showed a similar trend of reduction compared to that of control and R76W
159	mice (Figure 4K). Contrary to osteoblasts, osteoclast number on the endosteal bone surface was

significantly increased in loaded tibias in $\Delta 130-136$ mice (**Figure 4L-N**). The data suggested that osteocytic Cx43 hemichannels influence PGE₂ secretion, key bone marker expression, and osteoblastic and osteoclastic activities on endosteal surfaces in response to mechanical loading.

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164 Cx43 hemichannel-blocking antibody impairs the anabolic effects of mechanical loading on 165 trabecular and cortical bones.

166 We have developed a polyclonal antibody, Cx43(E2), that targets an extracellular loop domain of 167 Cx43 and specifically blocks osteocytic Cx43 hemichannels (Siller-Jackson et al., 2008). We 168 recently developed a specific mouse monoclonal blocking antibody Cx43(M1) to investigate the 169 roles of hemichannels in vivo. Gap junction channels and hemichannels were assayed using dye 170 coupling (Figure 5-figure supplement 1A) and dye uptake assays (Figure 5-figure supplement 171 **1B**), respectively. Both Cx43(E2) and Cx43(M1) had minimal effects on gap junction channels as indicated by comparable levels of dye transfer with red-orange AM dye in MLO-Y4 cells (Figure 172 173 5-figure supplement 1A). Conversely, FFSS-induced hemichannel opening, as determined by EtBr 174 uptake, was inhibited by Cx43(E2) and Cx43(M1) antibodies in MLO-Y4 cells (Figure 5-figure 175 supplement 1B). The extent of inhibition is comparable between Cx43(E1) and Cx43(M1) 176 antibodies. To ensure antibody delivery to osteocytes, we labeled tibial bone sections with a 177 rhodamine-conjugated anti-mouse secondary antibody. Strong antibody signals were primarily 178 detected in osteocytes in cortical bone for Cx43(M1)-injected mice, but not in IgG-injected ones 179 (Figure 5-figure supplement 1C). Interestingly, low levels of Cx43(M1) were detected in trabecular 180 bone (Figure 5-figure supplement 1D). The hemichannel opening detected by EB fluorescence was

found only in loaded tibial bone, and this uptake was almost completely blocked by the Cx43(M1) antibody (**Figure 5-figure supplement 1E, F**). Hence, these studies established the feasibility of using Cx43(M1) antibody to assess the role of Cx43 hemichannels *in vivo* using the tibial loading model.

WT mice with similar body weight (Figure 5-figure supplement 2A) were randomly allocated 185 to Cx43(M1) or vehicle groups. A slight decline in body weight was found during the first week of 186 187 the study, but was stabilized by the second week (Figure 5-figure supplement 2A). The antibody 188 had a negligible effect on body weight (Figure 5-figure supplement 2A). µCT analysis of tibial 189 metaphyseal trabecular bone showed that Cx43(M1) treatment significantly abated the 190 loading-induced increase of Tb.N and decrease of Tb.Sp (Figure 5B, C) as compared to the 191 vehicle-treated group, while Cx43(M1) did not manifest significant differences in the Tb.Th, BV/TV, 192 and BMD, in response to loading (Figure 5D-F). There was no change of SMI in both Cx43(M1) or 193 vehicle groups (Figure 5G). Representative images of trabecular bone are shown in Figure 5A. 194 Cx43(M1) treatment attenuated the anabolic response to mechanical loading in midshaft cortical

bone. The increase of B.Ar, B.Ar/T.Ar and Ct.Th by tibial loading was attenuated in the Cx43(M1)-treated group (**Figure 5J, K, M**), and consequently, the increase of pMOI was also attenuated in this group (**Figure 5O**). Similar to Δ130-136 mice, larger M.Ar and lower B.Ar/T.Ar ratios were found in loaded tibias of the Cx43(M1)-treated group when compared to the vehicle group (**Figure 5K, L**). Although Cx43(M1) further increased T.Ar by mechanical loading (**Figure 5I**), enlarged M.Ar (**Figure 5L**) significantly reduced the ratio of B.Ar/T.Ar (**Figure 5K**). Thus, Ct.Th did not increase and was even lower in loaded tibia compared to vehicle loaded tibias (**Figure**

202	5M). However, BMD was not changed by mechanical loading (Figure 5N). Three-point bending
203	analyses revealed a significant increase of elastic modulus and stiffness only in the vehicle group
204	(Figure 5P, Q). For loaded tibias, the elastic modulus and ultimate stress in the vehicle group were
205	greater than the Cx43(M1) group (Figure 5P, R). Mechanical loading did not change ultimate force
206	in either vehicle or Cx43(M1)-treated group (Figure 5S). Representative images of cortical bone are
207	shown in Figure 5H. These results are consistent with those obtained from $\Delta 130-136$ mice,
208	suggesting the critical roles of Cx43 hemichannels in the anabolic effects of mechanical loading of
209	cortical bone.
210	Blocking Cx43 hemichannels by Cx43(M1) inhibits the load-induced increase in midshaft
211	endosteal osteogenesis.
212	Bone formation in response to tibial loading was evaluated in vehicle and Cx43(M1)-treated
213	mice. The vehicle group exhibited increased endosteal MAR, MS/BS, and BRF/BS compared to
214	contralateral, unloaded controls, whereas this response was absent in the Cx43(M1) group (Figure
215	6A-D). In contrast, loading increased bone formation in vehicle and Cx43(M1)-treated groups on the
216	periosteal surface (Figure 6E-H). Cx43(M1) treatment induced a greater increase in periosteal
217	MS/BS (Figure 6G). The results showed that impaired Cx43 hemichannels attenuated endosteal
218	bone formation, but enhanced periosteal bone formation, induced by mechanical loading.
219	
220	Blocking Cx43 hemichannels by Cx43(M1) impedes the loading-induced increased PGE ₂
221	secretion, bone marker expression, and endosteal osteoblastic activity, and decreased

222 osteoclastic activity.

223	Tibial loading significantly increased PGE ₂ expression in tibial bone, and this increase was not
224	observed with Cx43(M1) antibody treatment (Figure 7A). Immunohistochemical staining of tibial
225	cortical bone showed a significant increase of COX-2 positive osteocytes by mechanical loading, and
226	such increase was not detected in the Cx43(M1)-treated group (Figure 7B, C and Figure 7-figure
227	supplement 1A). COX-2 bone mRNA levels detected by RT-qPCR exhibited close to a 5-fold
228	increase due to tibial loading, and this increase was not detected in Cx43(M1) treated bone samples
229	(Figure 7D). Loading caused a significant decrease of Sost-positive osteocytes in tibial bone in the
230	vehicle group, and Cx43(M1) antibody abated the load-induced decrease of Sost-positive osteocytes
231	(Figure 7E, F and Figure 7-figure supplement 1B). Sost bone mRNA also showed a significant
232	decrease due to loading in the vehicle group compared to the Cx43(M1)-treated group (Figure 7G).
233	We next determined another mechanical response protein, β -catenin expression, in osteoblasts. Tibial
234	loading resulted in a robust increase in endosteal β -catenin-positive osteoblasts and tibial β -catenin
235	gene expression in the vehicle group; in contrast, such increase was absent in the Cx43(M1) group
236	(Figure 7H-J and Figure 7-figure supplement 1C). Consistent with β -catenin expression,
237	endosteal osteoblast number only increased in the vehicle group (Figure 7K, L). Moreover, the
238	increase of gene expression of the osteoblastic marker, Bglap, was greater in the bone of the vehicle
239	group compared to the Cx43(M1) group (Figure 7M). Interestingly, increased osteoclast activity was
240	also found in the Cx43(M1) group (Figure 7K, N, O). The results showed that under mechanical
241	loading, inhibition of Cx43 hemichannels by Cx43(M1) antibody impedes the PGE ₂ release and Sost
242	decrease in osteocytes. This was associated with inhibited β -catenin and Bglap expression and
243	osteoblast activity on the endosteal surface.

244

PGE₂ rescues impeded osteogenic responses to mechanical loading by impaired Cx43 hemichannels.

247 Intermittent PGE₂ treatment has been reported to increase both trabecular and cortical bone mass (Jee et al., 1985; Tian et al., 2007). To explore whether the attenuated anabolic function of bone to 248 249 mechanical loading with Cx43(M1) is caused by inhibited PGE₂ released by Cx43 hemichannels, 250 PGE₂ was IP injected into the vehicle control and Cx43(M1) treated mice. The mice in the control 251 and treated groups had comparable body weights to minimize variations in tibial bone sizes before 252 loading (Figure 8-figure supplement 1A). µCT analysis showed that there was no difference in 253 trabecular morphometric parameters among the four groups (Figure 8-figure supplement 2). 254 Contrary to trabecular bone, PGE₂ treatment impeded the significant reduction of B.Ar/T.Ar ratio, 255 decreased M.Ar and increased Ct.Th in Cx43(M1)-treated loaded tibias, although there were no significant differences in T.Ar, B.Ar, and BMD in all four groups (Figure 8B-G). Representative 256 257 images of cortical bone are shown in Figure 8A. Interestingly, PGE₂ did not further enhance 258 anabolic bone responses in control, loaded mice. Together, these results demonstrate that 259 administration of PGE₂ significantly rescues impeded anabolic responses of cortical bone to 260 mechanical loading as a result of Cx43 hemichannel inhibition.

261

262 Discussion

In this study, we unveiled the distinctive roles of two types of Cx43 channels in responses to mechanical loading using both transgenic mouse models and Cx43 hemichannel-blocking antibodies,

and demonstrated the critical role of Cx43 hemichannels in mediating the anabolic, or bone forming, function of bone upon mechanical loading. Moreover, PGE_2 , a factor released by Cx43 hemichannels in response to mechanical stimulation, rescued the impeded anabolic effects on bone by tibial loading as a result of impaired hemichannels.

We determined bone structural and biomechanical properties, new bone formation, and 269 270 metabolism using an established bone mechanical loading model, axial tibial compression, in two 271 transgenic mouse models expressing Cx43 dominant negative mutants R76W and Δ 130-136 in osteocytes. We observed that $\Delta 130-136$ mice, with inhibited osteocytic Cx43 hemichannels. 272 273 attenuated anabolic bone responses to mechanical loading in both trabecular and cortical bones. The 274 attenuation of bone formation and osteoblastic activities primarily occurred on the tibial endosteal surface, while increased bone formation was observed on the periosteal surface. The activities of 275 276 osteoblasts and osteoclasts on periosteal and endosteal surfaces, respectively, lead to an enlarged bone marrow area and decreased bone to tissue area ratio. Due to impaired osteocytic gap junction 277 channels and hemichannels in Δ 130-136 mice, but only impaired gap junction channels in R76W 278 279 mice, we postulated that osteocytic Cx43 hemichannels, not gap junctions in osteocytes, are likely to 280 play a predominant role in anabolic bone response to mechanical loading.

The role of Cx43 hemichannels was further validated by the Cx43(M1) antibody, a potent monoclonal antibody that effectively inhibits osteocytic Cx43 hemichannels both *in vitro* and *in vivo*. Remarkably, treatment with Cx43(M1) only twice in a span of two weeks significantly attenuated anabolic effects to mechanical loading, with greater effects in cortical bones. Interestingly, Cx43(M1) treatment not only attenuated, but even reversed anabolic effects in cortical bone, similar to

 Δ 130-136 mice. A similar impediment of the rate of bone formation and mineral apposition to tibial 286 287 loading was observed on the endosteal surface with Cx43(M1) treatment. Previous studies have shown that cortical bone modeling/remodeling is more pronounced at the endosteal surface (Birkhold 288 289 et al., 2017), and mature bones respond to mechanical loading through changes on endosteal surfaces (Bass et al., 2002). Similar findings were also noted in Cx43 cKO mouse models; mice lacking Cx43 290 291 in osteoblasts and osteocytes showed an attenuated increase in endosteal bone formation during 292 four-point or three-point tibial bending (Grimston et al., 2008; Grimston et al., 2006). Mice lacking 293 Cx43 in osteochondroprogenitors showed a greater extent of decrease in endosteal formation during 294 tibial compression loading (Grimston et al., 2012). In our study, notably, endosteal MAR and 295 BFR/BS were not responsive to tibial loading in Δ 130-136 mice and the Cx43(M1) group, suggesting not only increased osteoblastic activity, but also that increased osteoblast number was 296 impeded by the impairment of Cx43 hemichannels. Moreover, histomorphometric analysis further 297 confirmed a lack of response in osteoblast number and marker genes in $\Delta 130-136$ mice and 298 299 Cx43(M1)-treated mice. The difference between Cx43 cKO and our transgenic models and Cx43(M1) 300 treatment could be caused by aberrant, compensatory effect of other pathways as a result of Cx43 301 deletion. Thus, our results suggest that axial compression loading promotes osteoblast recruitment 302 and differentiation on the endosteal surface, an anabolic effect likely mediated by mechanosensitive 303 Cx43 hemichannels.

Interestingly, contrary to our hypothesis, axial load increased periosteal bone formation and total tissue area on the tibial midshaft in Δ 130-136 mice. This observation was also reported in cKO mouse models with Cx43 deletion in osteoblasts and osteocytes under tibial axial compression

307	(Grimston et al., 2012) and tibial cantilever bending (Zhang et al., 2011). Similarly, deletion of Cx43
308	in osteocytes also showed an enhanced periosteal response to ulnar compression (Bivi et al., 2013).
309	The enhanced periosteal bone formation was further observed when hemichannels were inhibited by
310	Cx43(M1). These results posit the role of Cx43 hemichannels in periosteal bone formation during
311	mechanical loading. Our previous study showed that $\Delta 130-136$ mice have more periosteal bone
312	apposition than WT and R76W mice, suggesting that periosteal osteoblasts in Δ 130-136 mice are
313	more active and sensitive than WT and R76W mice (Xu et al., 2015). Thus, osteocytic Cx43
314	hemichannels exert differential roles in controlling osteogenic osteoblastic activities on periosteal
315	and bone resorping osteoclastic activity on endosteal surfaces, respectively. The consequence of the
316	disruption of coordinated activities by hemichannel inhibition results in an enlarged bone marrow
317	cavity. This is likely an adaptive response due to compromised cortical bones resulting from
318	impaired hemichannels and consequently lower extracellular PGE ₂ . From a mechanical point of view,
319	increased bone marrow area and cortical bone size allow the bone to respond to high stress levels
320	(Sharir et al., 2008). Thus, the role of osteocytic Cx43 hemichannels in regulating the osteogenic
321	response on endosteal surfaces is distinct from its role on periosteal surfaces.

Besides cortical bone, the anabolic response of trabecular bone to tibial loading in $\Delta 130-136$ mice was also attenuated or even reversed, as observed in trabecular number, separation and bone density. However, the compromised response of trabecular bone to mechanical loading was less evident in Cx43(M1) treated mice, except for trabecular number and trabecular separation. One possible explanation is that the accessibility and binding of the antibody to osteocytes in trabecular bone may not be as efficient as in cortical bone. Indeed, we could clearly detect Cx43(M1) on the

surface of osteocytes in cortical bone. However, the binding in trabecular bone is much weaker than that in cortical bone. Since Haversian canals containing blood vessels provide supply to the osteocytes in cortical, but not in trabecular bone (Dahl and Thompson, 2011), it is plausible that the delivery of Cx43(M1) to the bone is mediated primarily by the Haversian canal system.

Previous studies have reported that in vitro Cx43 hemichannel opening induced by FFSS 332 333 mediates the release of PGE₂ (Cherian et al., 2005), a critical factor for anabolic function of bone in 334 response to mechanical loading (Jee et al., 1985; Thorsen et al., 1996). On the contrary, PGE₂ release 335 by FFSS is inhibited by a potent hemichannel-blocking rabbit polyclonal antibody Cx43(E2) (Siller-Jackson et al., 2008). Here, we found that PGE₂ levels and osteocytic COX-2 expression were 336 337 increased by tibial loading in WT and R76W mice, but such an increase was not detected in Δ 130-136 mice. The use of hemichannel-blocking monoclonal Cx43(M1) antibody further confirmed 338 339 the role of the hemichannels in the release of PGE₂ in bone in situ. In accordance with our observation, the reduced release of PGE₂ was reported in calvarial cells isolated from Cx43 cKO 340 341 mice driven by the Col-2.3-kb α 1(I) collagen promoter after mechanical stretching (Grimston et al., 342 2006). The increase in Cox-2 gene expression in Cx43 cKO mice driven by the 8-kb DMP1 promoter 343 is attenuated after axial tibial compression (Grimston et al., 2012).

We showed that inhibited release of PGE_2 in $\Delta 130-136$ mice and in the Cx43 (M1) group was accompanied by an attenuated endosteal bone response to mechanical loading. Moreover, PGE_2 injection rescued the anabolic responses of cortical bone to mechanical loading impeded by Cx43(M1), including the ratio of bone area to tissue area, cortical thickness, and bone marrow area. PGE₂ is a skeletal anabolic factor, and its synthesis and release are highly responsive to mechanical

349	stimulation in osteocytes (Cherian et al., 2005; Jiang and Cherian, 2003). Using the microdialysis
350	technique, a rapid and significant increase of PGE ₂ levels in the proximal tibial metaphysis was
351	observed in response to dynamic mechanical loading in healthy women (Thorsen et al., 1996).
352	Furthermore, intermittent PGE_2 treatment increases endosteal bone formation (Jee et al., 1985) and
353	bone mass (Tian et al., 2007). Conversely, inhibition of PGE ₂ by a COX-2 inhibitor blocks endosteal
354	tibial bone formation induced by mechanical loading in rats (Forwood, 1996). Here, we demonstrate
355	that PGE_2 is indeed involved in the anabolic action of hemichannels in response to mechanical
356	loading. Interestingly, PGE ₂ administration did not provide an additional increase in the cortical bone
357	of mice in loaded vehicle control group. It is likely that extracellular PGE_2 released by osteocytes by
358	normal exercise (mechanical loading) is sufficient to promote bone formation and additional
359	extracellular PGE ₂ would not further increase cortical bone mass.

360 Increased PGE₂ by mechanical stimuli is reported to bind to EP4 receptor and reduce Sost 361 expression (Galea et al., 2011). Sost, a Wnt signaling antagonist (Semenov et al., 2005), is a key regulator of mechanotransduction in bone. Sost, secreted primarily by osteocytes, acts upon 362 363 osteoblasts in a paracrine manner to inhibit bone formation (Poole et al., 2005) through its binding to 364 the Wnt co-receptor Lrp5/6 (Li et al., 2005) and suppressing β -catenin (Sawakami et al., 2006). Sost 365 gene and protein expression is suppressed by mechanical loading, and is accompanied by increased 366 bone formation(Moustafa et al., 2012; Robling et al., 2008). We observed suppressed Sost expression 367 in WT and R76W mice by tibial loading; however, the suppressive effect of Sost disappeared in 368 Δ 130-136 mice and the Cx43(M1)-treated mice. Correspondingly, the increased β -catenin expression 369 and osteoblast activity observed in WT and R76W mice was abated in Δ 130-136 and the Cx43(M1)

370 mice. These results indicate that PGE₂ released by Cx43 hemichannels in osteocytes is a likely factor 371 that participates in the bone anabolic response to mechanical stimuli. We previously showed that PGE₂ released from osteocytes via Cx43 hemichannels exerts autocrine effects via the EP2/4 372 373 receptor during mechanical stimulation (Xia et al., 2010). This study indicates that the increased β -catenin in osteoblasts by tibial loading is attenuated in Δ 130-136 and the Cx43(M1)-treated mice. 374 375 These results establish a close functional relationship between Cx43 hemichannel-released PGE₂ and 376 decreased Sost, and thereby increased β -catenin expression in osteoblasts, ultimately leading to 377 enhanced osteoblast activity and endosteal bone formation (Figure 8H).

There are possible limitations in this study. First, analysis of cortical bone changes at additional proximal or distal sites may provide a more comprehensive understanding of the role of Cx43 hemichannels in anabolic responses to mechanical loading, although a previous study has reported that cortical bone located at 25%, 37% and 50% of the tibia's length had similar responses to tibial loading (Yang et al., 2017). Second, the monoclonal Cx43(M1) antibody blocks hemichannels not only in osteocytes, but also, possibly other cells, such as osteoblasts. However, in our study, Cx43(M1) was primarily detected in osteocytes, not in osteoblasts or other bone cells.

In summary, this study, for the first time, unveils the crucial role of osteocytic Cx43 hemichannels in mediating the anabolic function of mechanical loading on endosteal bone surfaces and trabecular bone. Cx43 hemichannels activated by mechanical stimulation release PGE_2 from osteocytes, which suppresses Sost expression in osteocytes, and enhances osteoblast activity and bone formation on endosteal surfaces. These results suggest that osteocytic Cx43 hemichannels could be established as a *de novo* new therapeutic target, and activation of these channels may

- 391 potentially aid in treating bone loss, in particular, in the elder population with the lost sensitivity to
 392 anabolic responses to mechanical stimulation (Lanyon and Skerry, 2001).
- 393

394 Materials and Methods

395 Mouse models

396 Two transgenic models expressing dominant-negative mutants of Cx43 in osteocytes, R76W and 397 Δ 130-136, were generated as previously described (Xu et al., 2015). The two transgenes were driven 398 by a 10-kb DMP1 promoter and expressed predominantly in osteocytes. The WT and transgenic 399 mice in C57BL/6J background were housed in a temperature-controlled room with a light/dark cycle 400 of 12 hrs at the University of Texas Health Science Center at San Antonio (UTHSCSA) Institutional Lab Animal Research facility under specific pathogen-free conditions. Food and water were freely 401 402 available. 15-week-old male WT and homozygous transgenic were sedated under isoflurane and euthanized by cervical dislocation. All animal protocols were performed following the National 403 404 Institutes of Health guidelines for care and use of laboratory animals and approved by the 405 UTHSCSA Institutional Animal Care and Use Committee (IACUC).

406

407 **Tibial mid-diaphyseal strain measurements and cyclic tibial loading**

The relationship between applied compressive loading and bone tissue deformation of the left tibia was established for 15-week-old mice *in vivo* following a previously reported protocol (De Souza et al., 2005; Lynch et al., 2010). Briefly, a strain gauge (EA-06-015DJ-120, Vishay Measurements Group) was attached on the tibial diaphyseal medial mid-shaft of a euthanized mouse and load

417	comparable periosteal strain in Δ 130-136 mice (Figure 1-figure supplement 1C).
416	tibia ($R^2 > 0.99$). Compared to WT mice, a higher compressive force was required to generate
415	compliance relationship between applied load and the resulting strain was determined for each left
414	and a conditioner/amplifier system. Mechanical load-induced strain was measured, and the
413	gauge was connected to a bridge completion module (MR1-350-127, Vishay Measurements Group)
412	applied from 0 to 9.5 N at the ends of the left tibia using a loading machine (LM1, Bose). The strain

418 The cyclic axial compressive load was applied to the left tibia of each mouse using a custom 419 loading device based on previous studies (De Souza et al., 2005; Lynch et al., 2010). Briefly, the left 420 tibia of anesthetized mice was positioned into a custom-made apparatus (Figure 1-figure 421 supplement 1A). The upper padded cup containing the knee was connected to the loading device 422 (7528-10, Masterflex L/S, Vernon Hills, IL, USA), and the lower cup held the heel. The left tibia 423 was held in place by a 0.5 N continuous static preload, loaded for 600 cycles (5 min) at 2-Hz frequency, with a sinusoidal waveform (Figure 1-figure supplement 1B). Compressive load was 424 425 performed 5 days/week for 2 weeks to determine bone structural and anabolic response, or 5 426 consecutive days to assess PGE₂ level. Based on the load-strain relationship, peak force was selected 427 to generate peak periosteal strains of 1200 $\mu\epsilon$ at the cortical midshaft for WT, R76W, and Δ 130-136 428 mice, respectively. This strain level has been previously shown to elicit an anabolic response at this 429 region(Melville et al., 2015). The right tibia was used as a contralateral, non-loaded control.

430

431 **M1** antibody generation and treatment

432	A monoclonal Cx43(M1) antibody targeting the second extracellular loop domains of Cx43 was
433	originally generated by Abmart (Tulsa, OK, USA) and described previously (Zhang et al., 2021).
434	Briefly, mice were immunized with a Cx43 extracellular domain peptide, and after functional
435	characterization of the hybridoma clones, genes that encode the antibody heavy and light chain
436	variable region were cloned from the mouse hybridoma cell line HC1 by reverse transcription
437	quantitative PCR (RT-qPCR), using a combination of a group of cloning PCR primers. The heavy
438	and light chain constructs were co-transfected into human embryonic kidney freestyle 293
439	(HEK293F) cells, supernatants were harvested, and antibodies were purified by affinity
440	chromatography using protein A resin.
441	The day before tibial loading, randomly allocated WT mice based on the body weight were
442	intraperitoneally (IP) injected with 25 mg/kg Cx43(M1) or vehicle (phosphate-buffered saline (PBS),
443	pH 7.4). A second dose was administered the day before the start of loading in the 2nd week. The
444	dosage of antibody was based on our data with Cx43(M1) antibody (Figure S5) and a previous study
445	using an anti-sclerostin (Sost) antibody (Spatz et al., 2013).
446	
447	In vitro dye uptake and gap junction coupling assays
448	The osteocyte-like MLO-Y4 cells were cultured in a-modified essential medium (a-MEM) with 2.5%
449	fetal bovine serum (FBS) and 2.5% calf serum (CS) in a 5% CO ₂ incubator at 37 $^{\circ}$ C. MLO-Y4 cells
450	were grown at a low initial cell density on glass slides coated with type I collagen (rat tail collagen

451

452 in contact. The cells were preincubated with Cx43(E2) or Cx43(M1) (2 μ g/ml) for 30 min and then

type I, Corning, Bedford, MA, USA, 0.15 mg/ml) to ensure that most of the cells were not physically

453	subjected to FFSS at 4 dynes/cm ² for 15 min in the presence of 25 mM ethidium bromide (EtBr) in
454	the recording media (HCO $_3^-$ -free a-MEM medium buffered with 10 mM HEPES). These cells were
455	then fixed with 1% paraformaldehyde (PFA) for 10 min. The intensity of EtBr fluorescence in cells
456	was measured and quantified by NIH Image J software (NIH, USA). Primary osteocytes were
457	microinjected using an Eppendorf micromanipulator InjectManNI 2 and Femtojet (Eppendorf) at
458	37°C with 10 mM Oregon green 488 BAPTA-AM (Mr: 1751 Da) as a cell tracker probe, and calcein
459	red-orange AM (Mr: 789 Da) as a probe for detecting gap junction coupling. Images were captured
460	using an inverted microscope equipped with a Lambda DG4 device (Sutter Instrument Co, Novato,
461	CA, USA), a mercury arc lamp illumination, and a Nikon Eclipse microscope (Nikon, Tokyo, Japan)
462	using a rhodamine filter. Loaded cells (dye donor) were "parachuted" over acceptor cells. The cells
463	(acceptors and donors) were pre-incubated for 20 min with Cx43 antibodies before the parachuting
464	assay. Donor cells were then incubated with acceptor cells for 90 min, the time duration sufficient to
465	detect dye transfer.

466

467 *In vivo* dye uptake assay

468 We developed an approach to assess hemichannel activity in osteocytes in the bone in vivo 469 (Riquelme et al., 2021). Briefly, 20 mg/ml Evans blue (EB) dye dissolved in sterile saline solution 470 (previously used to study hemichannel activity in muscle cells in vivo (Cea et al., 2013)) was injected 471 into the mouse tail vein. For the vehicle or Cx43(M1) treated group, mice were IP injected with 472 mouse IgG or Cx43 (M1) (25 mg/kg) 4 hrs before dye injection. After the dye injection, mice were 473 kept in cages for 20 min, and the left tibias were then loaded for 10 min. Mice were sacrificed and

474	perfused with PBS and 4% PFA 40 min after tibial loading. Tibias were isolated and fixed in 4%
475	PFA for 2 days, decalcified in 10% ethylenediaminetetraacetic acid (EDTA) for 3 weeks, and then
476	12-µm-thick frozen sections were prepared. The cell nuclei were stained with
477	4',6-diami-dino-2-phenylindole (DAPI). Images were captured using an optical microscope
478	(BZ-X710, KEYENCE, Itasca, IL, USA) and EB fluorescence intensity in osteocytes was quantified
479	by NIH Image J software (NIH, USA).

480

481 **PGE**₂ measurement and treatment

482 The level of PGE_2 in the tibia bone was determined according to the manufacturer's protocol (PGE_2) 483 ELISA kit, #514010, Cayman Chemical, Ann Arbor, MI, USA). Briefly, 4 hrs after the final round of 484 five-day tibial loading, bone marrow-flushed tibias were isolated free of soft tissues, and bone shafts 485 were prepared by removing proximal and distal ends of the bone. Bone tissue was homogenized in 486 liquid nitrogen with a frozen mortar and pestle. The concentration of PGE₂ was normalized by total 487 protein concentration using a BCA assay (#23225, Thermo Scientific, Rockford, IL, USA). 488 PGE₂ powder (#2296, Tocris Bioscience, Bristol, UK) was dissolved in 10% ethanol and stock 489 prepared at the concentration of 0.15 mg/ml. Wild-type mice randomly allocated based on the body 490 weight were injected with 1 mg/kg/day of PGE₂ solution or 6.7µl/kg/day 10% ethanol (vehicle) for 491 two weeks.

492

493 Micro-computed tomography

494 Tibias were dissected and frozen in saline-soaked gauze at -20°C until scanning. Samples in PBS 495 were imaged using a high-resolution micro-computed tomography (µCT) scanner (1172, SkyScan, 496 Brüker microCT, Kontich, Belgium) with the following settings: 59 Kvp, 167 µA beam intensity, 0.5 mm aluminum filter, 800 ms exposure, 1024 x 1024 pixel matrix, and a 10 µm isotropic voxel 497 dimension. The background noise was removed from the images by eliminating disconnected objects 498 499 smaller than 4 pixels in size. Two bone volumes of interest (VOI) were selected in the metaphyseal 500 and midshaft regions. The analyses were conducted excluding the fibula. In the proximal tibial 501 metaphysis, the trabecular bone VOI was positioned 0.44 mm distal to the proximal growth plate and 502 extended 0.65 mm in the distal direction, excluding the primary spongiosa. Gravscale values of 503 80-256 were set as the threshold for trabecular bone. For the cortical region, a 0.3 mm distance was 504 centered at 50% tibial length (proximal to distal). Automated contouring was used to select the 505 cortex. A threshold of 106-256 was applied to all of the cortical slices for analysis. The structural morphometric properties of cortical and trabecular regions were analyzed using the CT Analyser 506 507 software (CTAn 1.18.8.0, Bruker Skyscan).

508

509 Mechanical testing

510 Three-point bending tests were performed after μ CT scanning. The tibia was thawed to room 511 temperature before testing. Any remaining muscles and the fibula were carefully removed. The tibia 512 was subjected to a three-point bending test along the medial-lateral direction in a micromechanical 513 testing system (Mach-1 V500CST, Biomomentum, Laval, Canada). The span distance for the 514 three-point bending test was 8 mm, and the loading pin was placed at the midpoint of the span. The

515	test was performed in a displacement control mode at a constant rate of 0.01 mm/sec, and the data
516	was collected at a 200-Hz sampling rate for all measurements. The accurate cross-sectional areas
517	were determined from μ CT and used to calculate mechanical properties (Jepsen et al., 2015).

518

519 Histomorphometry, immunohistochemistry, and dynamic bone histomorphometry

520 Tibias were collected and fixed in 4% PFA for 2 days and decalcified using 10% EDTA (pH 7.5) for 521 21 days. These tibial samples were embedded in paraffin, and 5-mm-thick sections were collected 522 and mounted onto glass slides. For static bone histomorphometry, tartrate resistant acid phosphatase 523 (TRAP) and toluidine blue staining was used to determine the osteoclast activity (Xu et al., 2015) 524 and osteoblast numbers, respectively. For immunohistochemistry, paraffin sections were rehydrated 525 and antigen site was retrived with 10 mM citrate buffer (pH 6.0) at 60°C for 2 hrs (for sclerostin or 526 β-catenin) or trypsin buffer (pH 7.8) at 37°C for 30 min (for COX-2), and then probed with an 527 anti-sclerostin (AF1589, 1:400, R&D systems, Minneapolis, MN, USA), anti-COX-2 (12375-1-AP, 528 1:200, Proteintech, Rosemont, IL, USA) or an anti-β-catenin antibody (ab16051, 1:200, Abcam, 529 Waltham, MA, USA) overnight at 4°C. The sections were probed with a biotin-labeled secondary 530 antibody and ABC Reagent (VECTASTAIN, Burlingame, CA, USA). Staining was visualized with 531 DAB Chromogen (SK-4100, Vector Laboratories, Burlingame, CA, USA). Hematoxylin was used as 532 a counterstain. Images were captured using an optical microscope (BZ-X710, KEYENCE). 533 Osteoclast surface (Oc.S), osteoclast number (N.Oc), osteoblast number (N.Ob), and bone surface 534 (BS) on the tibial midshaft cortical bone along the endosteal surface were counted and positive cells 535 quantified using ImageJ software (NIH, USA).

536 Mice were IP injected with calcein (C0875, Sigma-Aldrich, St. Louis, MO, USA) at 20 mg/kg of 537 body weight 1 day before tibial loading, and followed by alizarin red injection (A5533, Sigma-Aldrich, St. Louis, MO, USA) at 30 mg/kg of body weight 3 days before euthanization. Tibias 538 539 were dissected, fixed in 70% ethanol, and then embedded in methylmethacrylate for 10-µm thick longitudinal plastic sections. Two-color fluorescent images were obtained using a fluorescence 540 541 microscope (BZ-X710, KEYENCE). Single label was defined as any bone surface with green, red, or 542 yellow (no separation between green and red). The distance between green and red fluorescence 543 signals was measured along the bone surface (Grimston et al., 2012). The following parameters were 544 quantified at tibial midshaft using NIH ImageJ software (NIH, USA): total perimeter (BS); single 545 label perimeter (sLS); double label perimeter (dLS), and double-label area (dL.Ar). The following values were then calculated: mineralizing surface [MS/BS = (sLS/2 + dLS)/BS], mineral apposition 546 547 rate [MAR = dL.Ar/dLS/12], and bone formation rate (BFR/BS = MAR × MS/BS).

548

549 **RNA extraction and RT-qPCR**

The long tibial bone was isolated free of soft tissues, and bone marrow was removed by flushing with RNase-free PBS after two weeks tibial loading. The bone shaft was prepared by removing the proximal and distal ends of the bone and pulverizing it using a frozen mortar and pestling in liquid nitrogen. Total RNA was isolated by using TRIzol (Molecular Research Center, Cincinnati, OH, USA) according to the manufacturer's protocol. cDNA was synthesized by a high-capacity cDNA reverse transcription kit (#4388950, Applied Biosystems, Carlsbad, CA, USA). mRNA level was analyzed by real-time RT-qPCR using an ABI 7900 PCR device (Applied Biosystems, Bedford, MA,

557	USA) and SYBR Green (#1725124, Bio-Rad Laboratories, Hercules, CA, USA) with a two-step
558	protocol (94°C for 10 sec, and 65°C for 30 sec for 40 cycles). The relative gene expression in each
559	loaded tibia was represented by normalizing to GAPDH and then normalized to the control tibia
560	(2 ^{-ΔΔCt}) (Kenneth J. Livak and Schmittgen, 2001). The primers for Sost, COX-2, Runx2, Bgalp,
561	β -catenin, and Dmp1 are provided in Table S1.

562

563 Statistical analysis

564 Data collection and analysis were conducted in a blind manner. Statistical analysis was performed 565 using IBM SPSS Statistics 24 (SPSS Inc., Chicago, IL, USA) and graphed with GraphPad Prism 7 566 (GraphPad Software; La Jolla, CA, USA). For in vitro studies, each experiment had three technical 567 replicates and was repeated at least three times. Normal distribution of the data was evaluated by the 568 Shapiro-Wilk test, and homogeneity of variance was assessed by the Levene test. The paired t-test 569 was used for comparisons of the loaded and contralateral tibias within the same group. One-way 570 ANOVA with Tukey test was used for multiple group comparisons. Student unpaired t-test was used 571 to compare between vehicle and Cx43 (M1)-treated groups. All data are presented as means \pm SD. 572 P<0.05 was considered significant.

573

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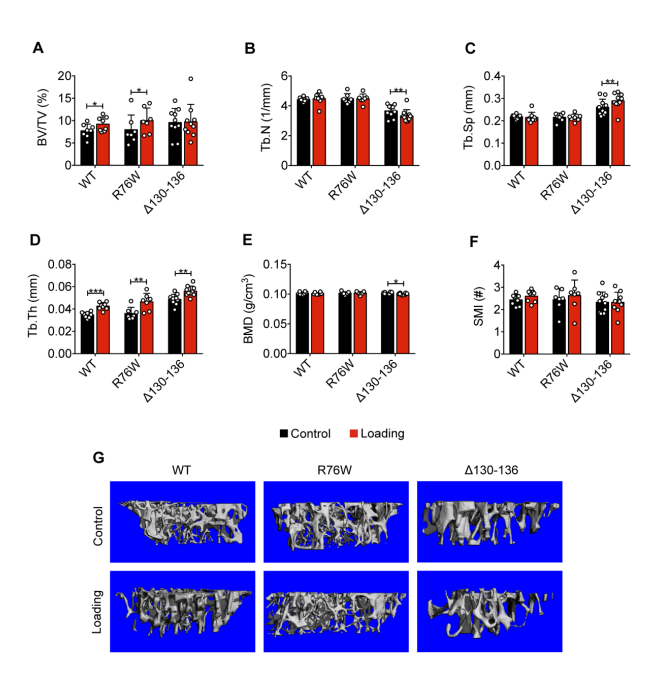
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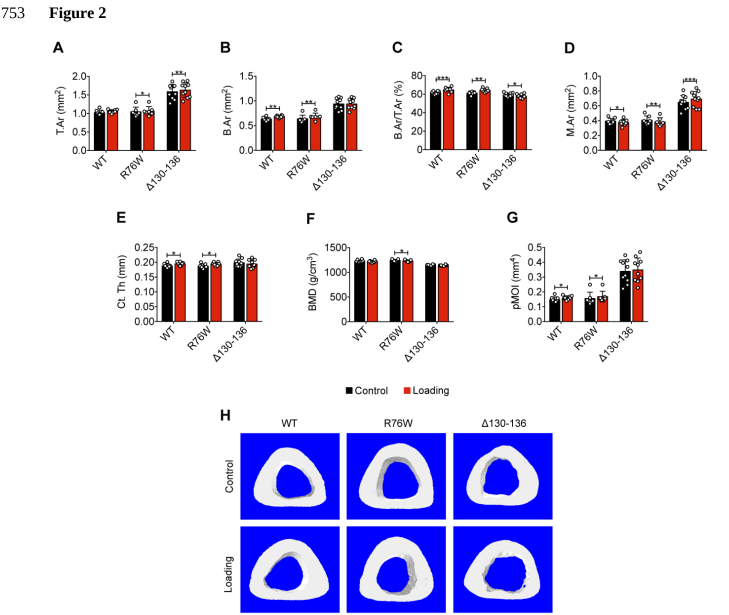
742 **Figure 1**



743

Figure 1. Attenuation or reversal of anabolic responses to mechanical loading in tibial metaphyseal trabecular bone of Δ 130-136 mice.

 μ CT was used to assess metaphyseal trabecular bone of WT, R76W, and Δ 130-136 mice; (A) bone volume fraction, (B) trabecular number, (C) trabecular separation, (D) trabecular thickness, (E) bone mineral density, and (F) structure model index. n=7-10/group. (G) Representative 3D models of the metaphyseal trabecular bone for all groups. Data are expressed as mean ± SD. *, P<0.05; **, P<0.01; ***, P<0.001. Statistical analysis was performed using paired t-test for loaded and contralateral, unloaded tibias.



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Figure 2. Attenuation or reversal of anabolic responses to mechanical loading in midshaft cortical bone of Δ 130-136 mice.

40 μCT was used to assess tibial midshaft cortical bone (50% site) of WT, R76W, and Δ130-136 mice; 41 total area, (**B**) bone area, (**C**) bone area fraction, (**D**) bone marrow area, (**E**) cortical thickness, 45 (**F**) bone mineral density, and (**G**) polar moment of inertia. n=7-10/group. (**H**) Representative 3D 46 models of the tibial midshaft cortical bone for all groups. Data are expressed as mean \pm SD. *, 47 P<0.05; **, P<0.01; **, P<0.001. Statistical analysis was performed using paired t-test for loaded 47 and contralateral, unloaded tibias.

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Figure 3

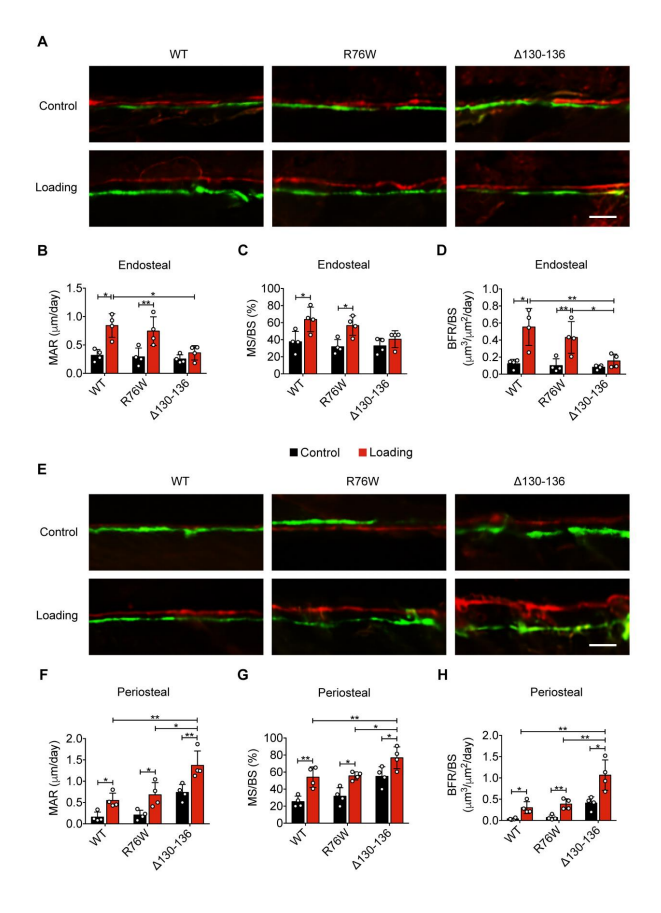
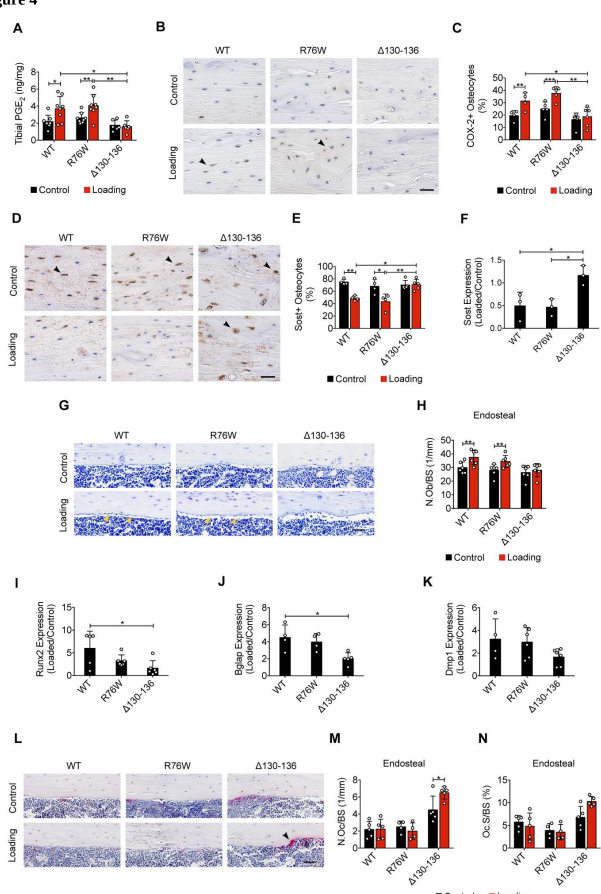


Figure 3. Reduced midshaft endosteal osteogenic responses to mechanical loading in Δ130-136 mice.

- 770 Dynamic histomorphometric analyses were performed on the tibial midshaft cortical endosteal and
- periosteal surfaces after 2 weeks of tibial loading of WT, R76W, and Δ 130-136 mice. Representative
- images of calcein (green) alizarin (red) double labeling on (A) endosteal and (E) periosteal surface.
- 773 Scale bar: 50 μm. Mineral apposition rate (MAR) (**B and F**), mineralizing surface/bone surface
- (MS/BS) (C and G), and bone formation rate (BFR/BS) (D and H) were assessed for endosteal
- (B-D) and periosteal (F-H) surfaces n=4/group. Data are expressed as mean \pm SD. *, P<0.05; **,
- P<0.01. Statistical analysis was performed using paired t-test for loaded and contralateral, unloaded
- tibias or one-way ANOVA with Tukey test for loaded tibias among different genotypes.





Loading

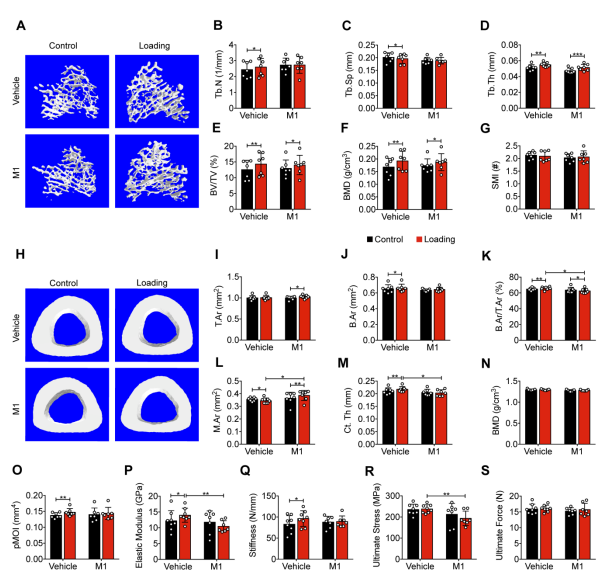
Control

Figure 4. Inhibition of the loading-induced PGE₂ secretion and osteoblast activity, and promotion of osteoclast activity in Δ 130-136 mice.

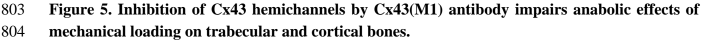
782 (A) ELISA analysis of PGE₂ in bone marrow-flushed tibial diaphysis after 5 days of tibial loading, in WT, R76W, and Δ 130-136 mice. n=6-8/group. (**B** and **C**) Representative images and quantitative 783 784 analysis of COX-2-postive osteocytes (black arrows) in the tibial midshaft cortical bone after 2 weeks of loading in WT, R76W, and Δ 130-136 mice. Scale bar: 30 µm. n=4-6/group. (**D** and **E**) 785 786 Representative images and quantitative analysis of the Sost-positive osteocytes (black arrows) in 787 tibial midshaft cortical bone after 2 weeks of tibial loading in WT, R76W, and Δ 130-136 mice. Scale bar: 30 µm. n=4-5/group. (F) Gene expression of Sost in bone marrow-flushed tibial diaphysis of 788 789 WT, R76W, and Δ 130-136 mice. n=3/group. (G and H) Toluidine blue staining was used to 790 determine the number of endosteal osteoblasts (yellow arrows) on tibial midshaft cortical bone in 791 WT, R76W, and Δ 130-136 mice after 2 weeks of loading. Scale bar: 30 µm; n=6/group. mRNA 792 expression of osteoblast markers, Runx2 (I) and Bglap (J), and osteocyte marker, Dmp1 (K) in bone 793 marrow-flushed tibial diaphysis of WT, R76W, and Δ 130-136 mice. n=4-6/group. (L) Representative images of tibial midshaft endosteal surface stained for TRAP (black arrows). Scale bar: 30 µm. (M 794 795 and N) Histomorphometric quantitation of osteoclasts per bone perimeter (M) and osteoclast surface 796 per bone perimeter (N) (n=4-5/group). All quantitative data are expressed as mean \pm SD.^{*}, P<0.05; **, P<0.01; ***, P<0.001. Statistical analysis was performed using paired t-test for loaded and 797 contralateral tibias or one-way ANOVA with Tukey test for loaded tibias among different genotypes. 798

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801 **Figure 5**

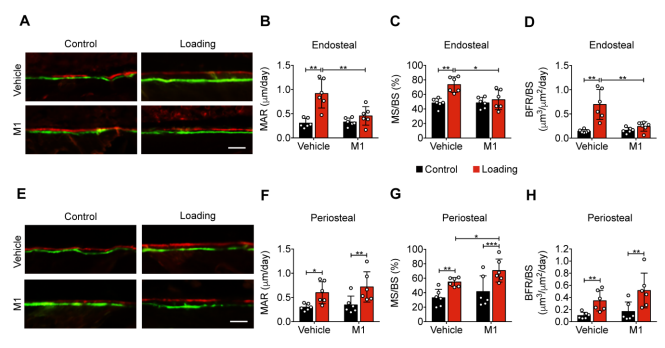


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(A) Representative 3D models of the metaphyseal trabecular bone of vehicle and Cx43(M1)-treated 805 806 mice. (B-G) μ CT was used to assess structural parameters of trabecular bone; (B) trabecular number, (C) trabecular separation, (D) trabecular thickness, (E) bone volume fraction, (F) bone mineral 807 density, and (G) structure model index in vehicle and Cx43 (M1)-treated mice. n=7/group. (H) 808 809 Representative 3D models of the tibial midshaft cortical bone (50% site) in vehicle and 810 Cx43(M1)-treated mice. (I-N) µCT was used to assess structural parameters of cortical bone; (I) total area, (J) bone area, (K) bone area fraction, (L) bone marrow area, (M) cortical thickness, (N) bone 811 mineral density and (O) polar moment of inertia in vehicle and Cx43(M1)-treated mice. n=7/group. 812 813 (P-S) The three-point bending assay was performed for tibial bone of vehicle and Cx43 (M1)-treated mice; (P) elastic modulus, (Q) stiffness, (R) ultimate stress, and (S) ultimate force. n=7-8/group. 814 Data are expressed as mean ± SD. *, P<0.05; **, P<0.01; ***, P<0.001. Statistical analysis was 815 816 performed using paired t-test for loaded and contralateral or unpaired t-test for loaded tibias between 817 vehicle- and Cx43(M1)-treated groups.





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821 Figure 6. Cx43(M1) inhibits the load-induced increase in midshaft endosteal osteogenesis. Dynamic histomorphometric analyses were performed on the tibial midshaft cortical endosteal (A-D) 822 and periosteal (E-H) surfaces after 2 weeks of loading in vehicle and Cx43(M1)-treated mice. (A 823 824 and E) Representative images of calcein (green) alizarin (red) double labeling on (a) endosteal and 825 (E) periosteal surface Scale bar: 50 µm. Mineral apposition rate (MAR), mineralizing surface/bone surface (MS/BS), and bone formation rate (BFR/BS) were assessed for (B-D) endosteal and (F-H) 826 periosteal surfaces (n=6/group). Data are expressed as mean ± SD. *, P<0.05; **, P<0.01; ***, 827 P<0.001. Statistical analysis was performed using paired t-test for loaded and contralateral, unloaded 828 829 tibias or unpaired t-test for loaded tibias between vehicle and Cx43(M1)-treated groups.

832 Figure 7

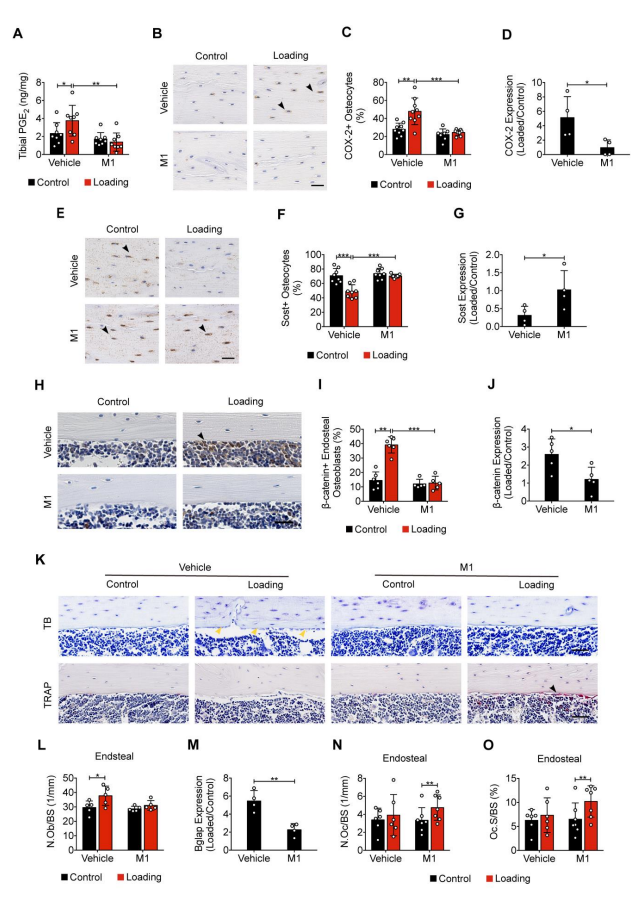
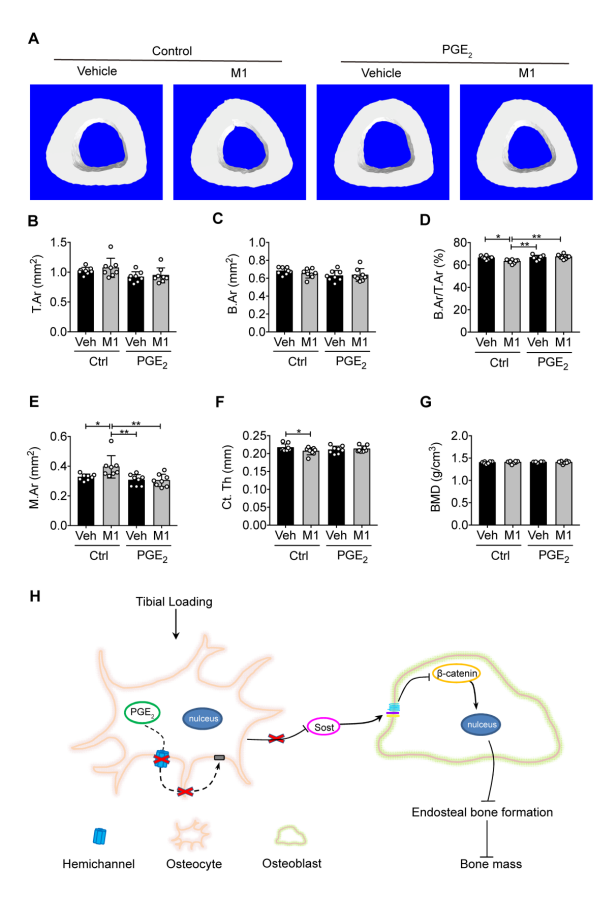


Figure 7. Cx43(M1) impedes the loading-induced increased PGE₂ secretion and osteoblast activity, and decreased osteoclast activity.

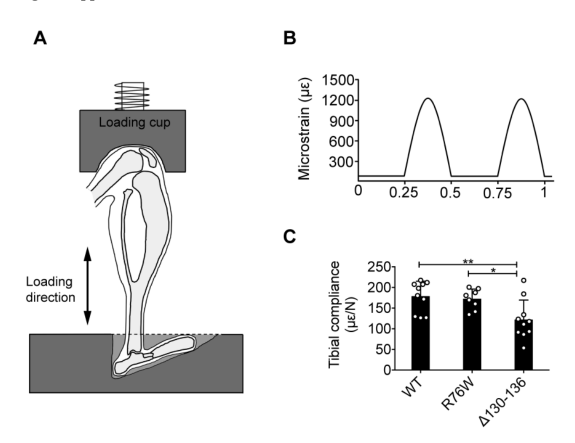
836 (A) ELISA analysis of PGE₂ in bone marrow-flushed tibial diaphysis after 5 days of mechanical loading in vehicle and Cx43(M1)-treated mice (n=8/group). (B and C) Representative images and 837 838 quantitative analysis of COX-2-postive osteocytes (vellow arrows) in tibial midshaft cortical bone after 2 weeks of loading in vehicle and Cx43(M1)-treated mice. Scale bar: 30 µm. n=8-9/group. (D) 839 840 COX-2 mRNA determined by RT-qPCR in bone marrow-flushed tibial diaphysis of vehicle and 841 Cx43(M1)-treated mice. n=4/group. (E-F) Representative images and quantitative analysis of the Sost-positive osteocytes (yellow arrows) in tibial midshaft cortical bone after 2 weeks of mechanical 842 843 loading in vehicle and Cx43(M1)-treated mice. Scale bar: 30 µm (n=8/group). (G) Sost mRNA 844 determined by RT-qPCR from bone marrow-flushed tibial diaphysis of vehicle and 845 Cx43(M1)-treated mice. n=4/group. (H and I) Representative images and quantitative analysis of the β-catenin positive periosteal cells (black arrows) on tibial midshaft endosteal surface after 2 weeks of 846 847 loading in vehicle and Cx43(M1)-treated mice. Scale bar: 20 μm; n=5-6/group. (J) β-catenin mRNA determined by RT-qPCR in bone marrow-flushed tibial diaphysis of vehicle and Cx43(M1)-treated 848 849 mice. n=4/group. (K) Representative images of tibial midshaft endosteal surface stained for toluidine 850 blue (top panel) or TRAP (low panel). The vellow arrows indicate osteoblasts and the black arrows 851 indicate the TRAP-positive osteoclasts. Scale bar: 30 µm. (L) Histomorphometric quantitation of 852 osteoblast per bone perimeter (n=5-7/group). (M) Bglap mRNA determined by RT-qPCR in bone marrow-flushed tibial diaphysis of vehicle and Cx43(M1)-treated mice. n=4/group. (N and O) 853 854 Histomorphometric quantitation of osteoclast per bone perimeter (N) and osteoclast surface per bone perimeter (O) (n=5-7/group). Data are expressed as mean \pm SD. *, P<0.05; **, P<0.01; ***, 855 P<0.001. Statistical analysis was performed using paired t-test for loaded and contralateral tibias, 856 857 unpaired t-test for loaded tibias between vehicle and Cx43(M1)-treated groups.

Figure 8



861 Figure 8. PGE_2 rescues the osteogenic response to mechanical loading with the impairment of 862 Cx43 hemichannels in cortical bone. (A) Representative 3D models of the tibial midshaft cortical bone (50% site) in vehicle and Cx43(M1)-treated mice treated with 1 mg/kg/day PGE₂ or vehicle 863 control. (B-G) µCT was used to assess tibial midshaft cortical bone; (B) total area, (C) bone area, 864 865 (D) bone area fraction, (E) bone marrow area, (F) cortical thickness, and (G) bone mineral density. n=8/group. Data are expressed as mean ± SD. *, P<0.05; **, P<0.01; ***, P<0.001. Statistical analysis 866 was performed using one-way ANOVA with Tukey test. (H) Schematic diagram illustrating the 867 868 mechanistic roles of osteocytic Cx43 hemichannels in mediating anabolic responses to tibial loading. Briefly, Cx43 hemichannels mediate the release of PGE₂ by mechanical loading, leading to 869 870 suppression of Sost expression with enhanced β-catenin expression and osteogenesis on the 871 endosteal surface. The inhibition of Cx43 hemichannels impedes the loading-induced PGE₂ secretion 872 and anabolic function of mechanical loading on bone tissue.

873 Figure 1-figure supplement 1



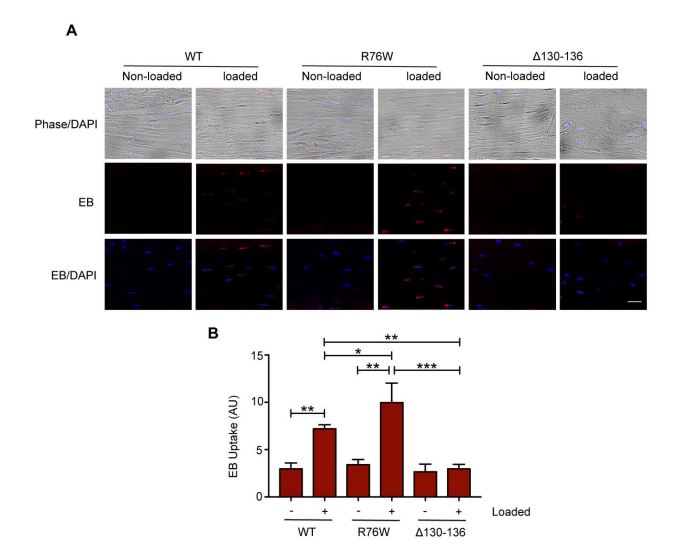
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Figure 1-figure supplement 1. Experimental setup for *in vivo* axial loading.

(A) Diagram of the left tibia positioned at the loading device and the direction of loading. (B) Schematic graph of 1 s of the daily 5 min loading signal. Approximately 1200 microstrain was detected on the medial mid-shaft surface of the tibia. (C) The average compliance of the relationship between applied load and resulting strain on the medial mid-shaft of WT, R76W and Δ 130-136 mice. n=8-10/group. Data are expressed as mean \pm SD. *, P<0.05; **, P<0.01. Statistical analysis was performed using one-way ANOVA with Tukey test among groups with different genotypes.

882

884 Figure 1-figure supplement 2

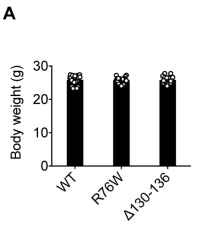


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Figure 1-figure supplement 2. Hemichannel opening is Inhibited in Δ130-136 mice.

(A) Representative images of Evans blue (EB) dye uptake in control and loaded tibial bone in WT, R76W and Δ 130-136 mice. Scale bar, 60 µm. (B) Quantitative analysis of Evans blue (EB) dye uptake. n=3/group. Data are represented as mean ± SD. *, P<0.05; **, P<0.01; ***, P<0.001. Statistical analysis was performed using one-way ANOVA with Tukey test among different groups.

892 Figure 1-figure supplement 3

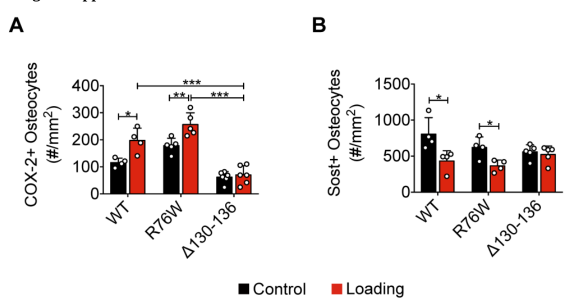


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Figure 1-figure supplement 3. Body weights of transgenic mice. (A) The body weights of WT, R76W and Δ 130-136 mice at the beginning of mechanical loading. n=22/group. Data are expressed

as mean \pm SD. Statistical analysis was performed using one-way ANOVA with Tukey test among different genotypes.

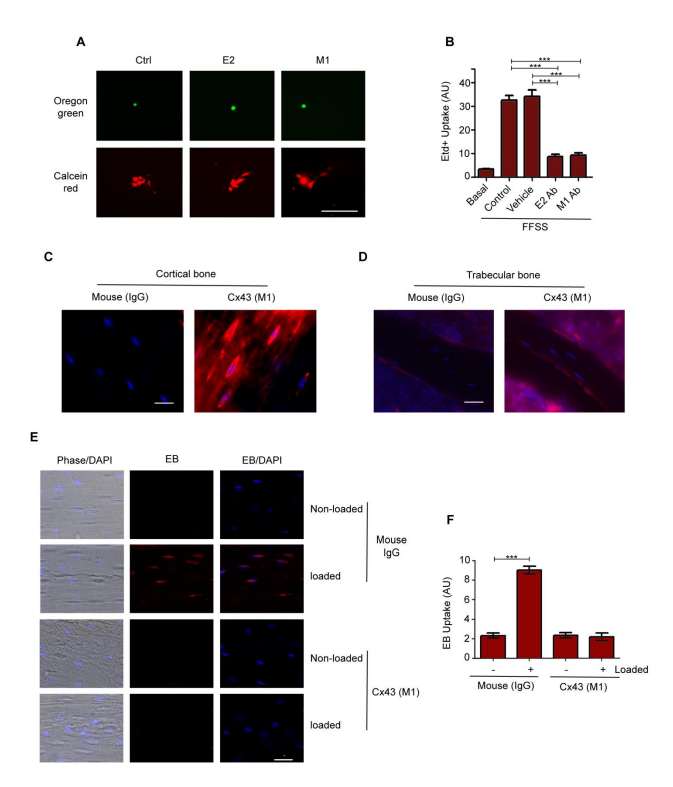
899 Figure 4-figure supplement 1



901 Figure 4-figure supplement 1. Bone marker protein expression in WT, R76W and \triangle 130-136 mice. (A) Quantitative analysis of the COX-2-positive osteocytes per bone area in tibial midshaft 902 cortical bone after 2 weeks of mechanical loading in WT, R76W and Δ 130-136 mice. n=4-6/group. 903 904 (B) Quantitative analysis of the Sost -positive osteocytes per bone area in tibial midshaft cortical bone after 2 weeks of mechanical loading in WT, R76W and ∆130-136 mice. n=4-6/group. Data are 905 expressed as mean ± SD. *, P<0.05; **, P<0.01; ***, P<0.001. Statistical analysis was performed using 906 907 paired t test for loaded and contralateral tibias, or one-way ANOVA with Tukey test for loaded tibias 908 among groups with different genotypes.

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910 Figure 5-figure supplement 1



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Figure 5-figure supplement 1. Monoclonal antibody of Cx43 inhibits hemichannel opening induced by mechanical stress *in vitro and in vivo*.

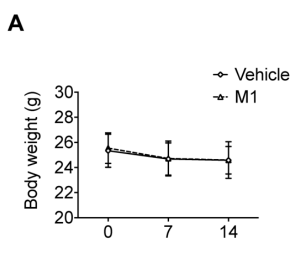
914 (A) Parachuting dye coupling assay was conducted to determine gap junction coupling in MLO-Y4

915 cells loaded with Oregon green 488 BAPTA-AM (Mr: 1751 Da) as a cell tracker probe and calcein

916 red-orange AM (Mr: 789 Da). Scale bar, 100 µm. (B) MLO-Y4 cells were preincubated with

917 Cx43(E2), Cx43(M1), PBS (vehicle) or rhodamine-conjugated anti-mouse IgG (control) and then subjected to fluid flow shear stress (FFSS) (8 dynes/cm²) for 10 min and followed by ethidium 918 bromide (Etd⁺) dye uptake assay. n=4/group. (C and D) Representative images of Cx43(M1) 919 920 detected with rhodamine-conjugated anti-mouse IgG in tibial midshaft cortical bone and trabecular 921 bone. Bar, 50 µm. (E) Representative images of Evans blue (EB) dye uptake in control and tibial 922 loaded bone in the absence or presence of Cx43(M1) antibody. Scale bar, 40 µm. (F) Quantitative analysis of Evans blue (EB) dye uptake. n=3/group. Data are expressed as mean \pm SD. *, P<0.05; **, 923 924 P<0.01; ***, P<0.001. Statistical analysis was performed using one-way ANOVA with Tukey test 925 among different groups. 926

927 Figure 5-figure supplement 2



928

Figure 5-figure supplement 2. Body weights of mice during 2 weeks of tibial loading. (A)
Weekly body weights of vehicle and Cx43(M1)-treated groups. n=18/group. Data are expressed as

 $mean \pm SD$. Statistical analysis was performed using one-way ANOVA with Tukey test among different genotypes.

934 Figure 7-figure supplement 1

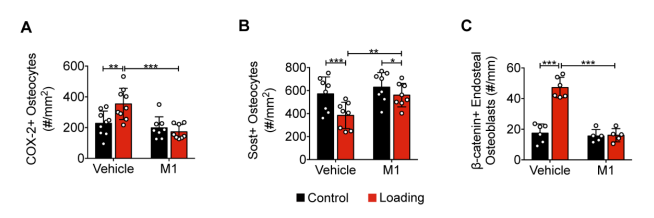




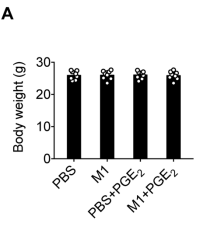
Figure 7-figure supplement 1. Bone marker protein expression in vehicle- and Cx43(M1)-treated mice.

938 (A, B) Quantitative analysis of the COX-2-positive or Sost-positive osteocytes per bone area in tibial

midshaft cortical bone after 2 weeks of loading in vehicle and Cx43(M1)-treated mice. n=8-9/group. (C) Quantitative analysis of the β -catenin-positive osteoblasts per bone perimeter in tibial midshaft cortical bone after 2 weeks of mechanical loading in vehicle and Cx43(M1)-treated mice. n=4-6/group. Data are expressed as mean \pm SD. *, P<0.05; **, P<0.01; ***, P<0.001. Statistical analysis was performed using paired t test for loaded and contralateral tibias, unpaired t test for

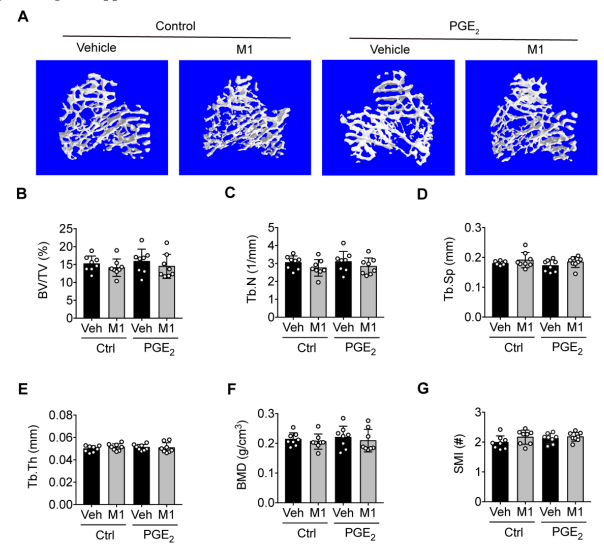
loaded tibias between vehicle- and Cx43(M1)-treated groups.

946 Figure 8-figure supplement 1



- Figure 8-figure supplement 1. Body weights of mice of vehicle- and Cx43(M1)-treated mice
 treated with 1mg/kg/day PGE2 or vehicle control.
- 950 (A) The body weights of vehicle- and Cx43(M1)-treated mice treated with 1mg/kg/day PGE₂ or
- vehicle control at the beginning of mechanical loading. n=8/group. Data are expressed as mean \pm SD.
- 952 Statistical analysis was performed using one-way ANOVA with Tukey test among different 953 genotypes.
- 954

955 Figure 8-figure supplement 2



956

Figure 8-figure supplement 2. PGE₂ does not exert additional trabecular osteogenic response to mechanical loading.

959 (A) Representative 3D models of metaphyseal trabecular bone in vehicle and Cx43(M1)-treated mice 960 treated with 1 mg/kg/day PGE₂ or vehicle control. (**B-G**) μ CT was used to assess tibial midshaft 961 cortical bone; (**B**) bone volume fraction, (**C**) trabecular number, (**D**) trabecular separation, (**E**) 962 trabecular thickness, (**F**) bone mineral density and (**G**) structure model index. n=8/group. Data are 963 expressed as mean \pm SD. *, P<0.05; **, P<0.01; ***, P<0.001. Statistical analysis was performed 964 using one-way ANOVA with Tukey test among different groups.

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- 970 **Figure 1-Source Data 1.** Trabecular micro-CT data of transgenic and wild-type mice.
- 971 Figure 1-figure supplement 1-source data 1. Raw data of compliances for Figure 1-figure972 supplement 1C.
- 973 Figure1-figure supplement 2-source data 1. Raw data of dye uptake for Figure 1-figure974 supplement 2B.
- 975 Figure1-figure supplement 3-source data 1. Raw data of body weight for Figure1-figure976 supplement 3A.
- 977 **Figure 2-Source Data 1**. Cortical micro-CT data of transgenic and wild-type mice.
- Figure 3-Source Data 1. Raw data of periosteal and endosteal bone formation of transgenic andwild-type mice.
- **Figure 4-Source Data 1.** Raw data of PGE₂ level for Figure 4A.
- Figure 4-Source Data 2. Raw data of immumohistochemical, TRAP and toluidine blue staining of
 transgenic and wild-type mice.
- 983 **Figure 4-Source Data 3**. Raw data of RT-qPCR of transgenic and wild-type mice.
- Figure 4-figure supplement 1-source data 1. Raw data of COX-2 and Sost quantification for
 Figure 4-figure supplement 1A, B.
- **Figure 5-Source Data 1.** Micro-CT data of vehicle and Cx43(M1)-treated mice.
- **Figure 5-Source Data 2.** Three-point bending data of vehicle and Cx43(M1)-treated mice.
- Figure 5-figure supplement 1-source data 1 Raw data of dye uptake for Figure 5-figuresupplement 1B and F.
- Figure 5-figure supplement 2-source data 1. Raw data of body weight for Figure 5-figuresupplement 2A.
- Figure 6-Source Data 1. Raw data of periosteal and endosteal bone formation of vehicle andCx43(M1)-treated mice.
- **Figure 7-Source Data 1.** Raw data of PGE₂ level for Figure 7A.
- Figure 7–Source Data 2. Raw data of immumohistochemical, TRAP and toluidine blue staining of
 vehicle and Cx43(M1)-treated mice.
- **Figure 7-Source Data 3.** Raw data of RT-qPCR of vehicle and Cx43(M1)-treated mice.
- Figure 7-figure supplement 1-source data 1. Raw data of COX-2 Sost and β-catenin quantification
 for Figure 7-figure supplement 1A-C.

- 1000 Figure 8-Source Data 1. Cortical micro-CT data of vehicle and Cx43(M1)-treated mice treated with
- 1001 1 mg/kg/day PGE2 or vehicle control.
- 1002 **Figure 8-figure supplement 1-source data 1.** Raw data of body weight for Figure8-figure
- 1003 supplement 1A.
- 1004 Figure 8-figure supplement 2-source data 1. Trabecular micro-CT data of vehicle and
- 1005 Cx43(M1)-treated mice treated with 1 mg/kg/day PGE2 or vehicle control.