Polychlorinated biphenyl (118) activates osteoclasts and induces bone resorption in goldfish

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44	Abstract
45	Purpose: To analyze the effect of Polychlorinated biphenyl (PCB118) on fish bone
46	metabolism, we examined osteoclastic and osteoblastic activities, as well as
47	plasma calcium levels, in the scales of PCB (118)-injected goldfish. In addition,
48	effect of PCB (118) on osteoclasts and osteoblasts was investigated in vitro.
49	Methods: Immature goldfish, in which the endogenous effects of sex steroids are
50	negligible, were used. PCB (118) was solubilized in dimethyl sulfoxide at a
51	concentration of 10 ppm. At 1 and 2 days after PCB (118) injection (100 ng/g
52	body weight), both osteoclastic and osteoblastic activities, and plasma calcium
53	levels were measured. In an in vitro study, then, both osteoclastic and osteoblastic
54	activities as well as each marker mRNA expression were examined.
55	Results: At 2 days, scale osteoclastic activity in PCB (118)-injected goldfish
56	increased significantly, while osteoblastic activity did not change significantly.
57	Corresponding to osteoclastic activity, plasma calcium levels increased
58	significantly at 2 days after PCB (118) administration. Osteoclastic activation was
59	also occurred in the marker enzyme activities and mRNA expressions in vitro.
60	Thus, we conclude that PCB (118) disrupts bone metabolism in goldfish both in
61	vivo and in vitro experiments.
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63	Keywords: PCB (118), bone metabolism, fish scales, osteoclasts, osteoblasts,
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## 1. Introduction

It has been reported that polychlorinated biphenyl (PCB) congeners act as endocrine-disrupting compounds (Lind et al. 2004; Bovee et al. 2011; Nakayama et al. 2011; Ju et al. 2012). As bone formation and resorption are controlled by several hormones and vitamins (see a review, Peacock 2010), PCBs might disturb bone metabolism. In some animals, actually, the bone disruption caused by PCB has been reported (rat: Lind et al. 2004a; bear: Sonne et al. 2004; sheep: Gutleb et al. 2010; alligator: Lind et al. 2004b; turtle: Holliday DK and Holliday CM 2012; salmon: Olufsen and Arukwe 2011; zebrafish: Ju et al. 2012). In humans, changes in bone metabolism associated with exposure to PCBs have also been investigated (Hodgson et al. 2008). However, the direct effects of PCBs on osteoclasts and osteoblasts have not yet been elucidated in any animals. The teleost scale is a calcified tissue that contains osteoblasts, osteoclasts, and the bone matrix of two layers (bony layer: a thin, well-calcified external layer; a fibrillary layer: a thick, partially calcified layer) (Bereiter-Hahn and Zylberberg 1993; Suzuki et al. 2000; Yoshikubo et al. 2005; Suzuki et al. 2007; Ohira et al. 2007). The bone matrix, which includes type I collagen (Zylberberg et al. 1992), osteocalcin (Nishimoto et al., 1992), and hydroxyapatite (Onozato and Watabe 1979) is present in the scale as well as in mammalian bone. Recently, we detected both cathepsin K and tartrate-resistant acid phosphatase (TRAP) mRNA expression in scale osteoclasts (Azuma et al. 2007). In osteoblasts, we detected osteoblast-specific markers, such as alkaline phosphatase (ALP), runt-related transcription factor 2, osterix, osteocalcin, type I collagen, and the receptor activator of the NF-kB ligand (Thamamongood et al. 2012). Therefore, the features of osteoclasts and osteoblasts in scales are similar to those in mammals.

In fish as well as mammals, plasma calcium level was regulated by hormones such as parathyroid hormone (Suzuki et al. 2011a) and calcitonin (Suzuki et al. 2000; Suzuki et al. 2004a). In an *in vivo* experiment, fugu parathyroid hormone I induced hypercalcemia resulted from the increase of both osteoblastic and osteoclastic activities in the scale and caused to decrease scale calcium contents (Suzuki et al. 2011a). Scale osteoclastic activation was also observed in the prostaglandin E<sub>2</sub> injected-goldfish (Omori et al. 2012). It is reported that the scales are a better potential internal calcium reservoir than the body skeletons, jaws and otolithes, examined by the <sup>45</sup>Ca-labelling study for the calcified tissues of goldfish and killifish (Mugiya and Watabe 1977). Thus, we conclude that teleost scale is an active and functional calcium reservoir.

In fish, PCB (118) is the highest congeners compared with PCB-105, -156, -167, -123, -157, -114, -189, -77, -126, -81, or -169 (Bhavsar et al. 2007).

In fish, PCB (118) is the highest congeners compared with PCB-105, -156, -167, -123, -157, -114, -189, -77, -126, -81, or -169 (Bhavsar et al. 2007). Furthermore, it has been reported that trabecular bone mineral content was almost 30% lower in the PCB (118) (49 μg/kg body wt/day) at the metaphysis in sheep (Gutleb et al. 2010), although the detail mechanism has not yet been elucidated. We therefore analyzed the effect of PCB (118) on scale osteoclastic and osteoblastic activities, as well as plasma calcium levels, in the goldfish scales. In addition, effect of PCB (118) on osteoclasts and osteoblasts was investigated *in vitro*. This is the first to demonstrate that PCB (118) activates osteoclasts and induced bone resorption in fish.

## 2. Materials and methods

126 Animals

To examine the effect of PCB (118) on the bone metabolism, immature goldfish (4-6 g), in which the endogenous effects of sex steroids are negligible,

were used for the *in vivo* study. A previous study (Suzuki et al. 2000) indicated that the sensitivity for calcemic hormones was higher in mature female than in mature male teleosts. Therefore, female goldfish (Carassius auratus) (30 - 40 g) were purchased from commercial source (Higashikawa Fish Farm, Yamatokoriyama, Japan) and used for the *in vitro* experiments.

All experimental procedures were conducted in accordance with the Guide for the Care and Use of Laboratory Animals prepared by Kanazawa University.

Effects of PCB (118) on scale osteoclastic and osteoblastic activities and the plasma calcium in goldfish at day-1 and -2 after PCB (118) injection (in vivo experiment)

PCB (118) was solubilized in dimethyl sulfoxide (DMSO) at a concentration of 10 ppm. Goldfish (body weight: 4 - 6 g) were anesthetized with ethyl 3-aminobenzoate and methanesulfonic acid salt (Sigma-Aldrich, Inc., MO, USA) and taken the blood (about 100 μl) from caudal vessels of each individual into heparinized syringes just before PCB (118) injection. After centrifugation at 15,000 rpm for 3 min, the plasma was immediately frozen and kept at -80 °C until use. In the experimental group (n = 10), thereafter, PCB (118) was intraperitoneally injected (100 ng/g body weight). The goldfish in the control group (n = 10) were injected with DMSO in the same manner. These goldfish were kept in the aquarium for 1 and 2 days. During the experimental periods, these goldfish were not given any food to exclude intestinal calcium uptake from diets. Each day after injection, the scales were collected from each goldfish. At day-2 after injection, blood samples (about 100 μl) were collected from the gill using a heparinized capillary from individual, anesthetized goldfish. After centrifugation at 15,000 rpm for 3 min, the plasma was also immediately frozen

and kept at -80 °C until use. The plasma total calcium level (mg/100 ml) was determined using an assay kit (Calcium C, Wako Pure Chemical Industries, Ltd., Osaka, Japan). Then, we measured the activities of ALP and TRAP activities as respective indicators of each activity in osteoclasts and osteoblasts (Suzuki et al. 2000; Suzuki et al. 2002; Suzuki et al. 2009). The measurement methods (Suzuki et al. 2009) of ALP and TRAP activities were as follows. The incubated scale was transferred to its own well in a 96-well microplate after washing with saline. An aliquot of 100 µl of an alkaline buffer (100 mM Tris-HCl, pH 9.5; 1 mM MgCl<sub>2</sub>; 0.1 mM ZnCl<sub>2</sub>) for ALP activity or an acid buffer (0.1 M sodium acetate including 20 mM tartrate, pH 5.3) for TRAP activity was added to each well. This microplate was frozen at -80°C immediately and then kept at -20°C until analysis. After thawing, an aliquot of 100 µl of 20 mM para-nitrophenyl-phosphate in an alkaline buffer or an acid buffer was added to each well. This plate was then incubated at 20°C for 30 min with shaking. After incubation, the reaction was stopped by adding 50 µl of a 3 N NaOH-20 mM EDTA solution. Aliquots of 150 μl of a colored solution were transferred to a new plate, and the absorbance was measured at 405 nm. The absorbance was converted into the amount of produced para-nitrophenol (pNP) using a standard curve for pNP. After measurement of the absorbance, the ALP and TRAP activities were normalized by the surface area  $(mm^2)$  of each goldfish scale. The results are shown as the means  $\pm$  SE of eight scales.

*PCB* (118) contents in the scales of goldfish (in vivo experiment)

At day-1 and -2 after PCB (118) injection, the scales were collected from goldfish and then immediately frozen and kept at -80 °C until use. The PCB (118) contents were analyzed by the methods of Hirai et al. (2005). Because a single

sample volume was very small, we conducted three measurements to obtain a pulled sample. Thus, the mean of three measurements was described in the results.

184 Effects of PCB (118) on osteoclastic and osteoblastic activities in the cultured

185 scales of goldfish (in vitro experiment)

Scales collected from goldfish (n = 10) after anesthesia with ethyl 3aminobenzoate and methanesulfonic acid salt (Sigma-Aldrich) and incubated for 6 and 18 h in Leibovitz's L-15 medium (Invitrogen, Grand Island, NY, USA) containing a 1% penicillin-streptomycin mixture (ICN Biomedicals, Inc., OH, USA) supplemented with PCB (118) (0.025, 0.25, and 2.5 ppm). In an in vivo experiment, around 0.1 to 0.05 ppm PCB was detected in the PCB-injected scales. Based on these PCB contents in the scales, we decided the administration doses of PCB in an in vitro experiment. The PCB concentration in one goldfish was performed using 48 scales from each left or right side. The 48 scales used in the present study were considered to use as follows: 1) 8 scales for TRAP analysis by 0.025 ppm, 2) 8 scales for TRAP analysis by 0.25ppm, 3) 8 scales for TRAP analysis by 2.5 ppm, 4) 8 scales for ALP analysis by 0.025 ppm, 5) 8 scales for ALP analysis by 0.25ppm, 6) 8 scales for ALP analysis by 2.5 ppm. The respective mean for TRAP (obtained from 8 individual scales of one goldfish) and ALP (obtained from 8 individual scales of one goldfish) activities from left side (experimental group) was compared with those of right side (control group). Using 10 individual goldfish, same experiment was done repeatedly. The experiments for 0.25 and 2.5 ppm PCB (118) were carried out in the same manner. After incubation, TRAP and ALP activities were measured using the same methods described above (Suzuki et al. 2009). The results are shown as means  $\pm$  SEM (n = 10).

208	Changes in TRAP, cathepsin K and RANKL mRNA expressions in PCB (118)-
209	treated goldfish scales (in vitro experiment)
210	Scales were collected from goldfish under anesthesia with ethyl 3-
211	aminobenzoate and methanesulfonic acid salt (Sigma-Aldrich). To examine
212	changes in TRAP, cathepsin K, and RANKL mRNAs that responded to PCB
213	(118), these scales were incubated for 18 h in Leibovitz's L-15 medium
214	(Invitrogen) containing a 1% penicillin-streptomycin mixture (ICN Biomedicals).
215	In the prostaglandin E <sub>2</sub> -treated scales of goldfish, we previously reported that
216	TRAP, cathepsin K, RANKL mRNA expression increased at 18 h of incubation
217	(Omori et al. 2012). Therefore, this incubation period was adopted. After
218	incubation, the scales were frozen at -80 °C for mRNA analysis.
219	Total RNAs were prepared from goldfish scales using a total RNA isolation
220	kit for fibrous tissue (Qiagen GmbH, Hilden, Germany). Complementary DNA
221	synthesis was performed using a kit (Qiagen GmbH). Gene-specific primers for
222	TRAP (sense: 5'-AACTTCCGCATTCCTCGAACAG-3'; antisense: 5'-
223	GGCCAGCCACCAGGAGATAA-3') (Azuma et al. 2007), cathepsin K (sense:
224	5'-GCTATGGAGCCACACCAAAAGG-3'; antisense: 5'-
225	CTGCGCTTCCAGCTCTCACAT-3') (Azuma et al. 2007), and RANKL (sense:
226	5'-GCGCTTACCTGCGGAATCATATC-3'; antisense: 5'-
227	AAGTGCAACAGAATCGCCACAC-3') (Suzuki et al. 2011a) were used. The
228	amplification of $\beta$ -actin cDNA using a primer set (5':
229	CGAGCGTGGCTACAGCTTCA; 3': GCCCGTCAGGGAGCTCATAG) (Azuma
230	et al. 2007) was performed. The PCR amplification was analyzed by real-time
231	PCR apparatus (Mx3000p, Agilent Technologies, CA, USA) (Suzuki et al.
232	2011a). The annealing temperature of TRAP, cathepsin K, RANKL, and β-actin

233	was 60 °C. The TRAP, cathepsin K and RANKL mRNA levels were normalized
234	to the β-actin mRNA level.
235	
236	Statistical analysis
237	All results are expressed as the means $\pm$ SE (n = 10). The statistical
238	significance between control and experimental group was assessed by Student's t-
239	test (in vivo experiment) or paired t-test (in vitro experiment). In all cases, the
240	selected significance level was $P < 0.05$ .
241	
242	3. Results
243	Effects of PCB (118) on scale osteoclastic and osteoblastic activities and the
244	plasma calcium in goldfish at 1 and 2 days after PCB (118) injection in vivo
245	We measured the activities of ALP and TRAP activities as respective
246	indicators of each activity in osteoclasts and osteoblasts. At day-2, scale TRAP
247	activity in PCB-injected goldfish increased significantly (Fig. 1a), while ALP
248	activity did not change significantly at day-1 and -2 (Fig. 1b).
249	Corresponding to the elevation of osteoclastic activity, plasma calcium levels
250	increased significantly at day-2 after PCB administration (Fig. 2)
251	
252	PCB (118) contents in the scales of goldfish in vivo
253	At day-1 and -2 after PCB (118) injection, PCB (118) was detected in the
254	scales. At day-1, PCB contents in the control and PCB-injected scales were
255	determined as 0.39 and 79 (ng/g-wet), respectively. At day-2, PCB (ng/g-wet) of
256	0.38 and 55 was detected in the control and PCB-injected scales, respectively.
257	

PCB (118) pollution.

258	Effect of PCB (118) on osteoclastic and osteoblastic activities in the cultured
259	scales of goldfish in vitro
260	PCB (118) significantly increased the TRAP activities of the scales by 6 h of
261	incubation ( $P < 0.05$ for 0.25 ppm) (Fig. 3a). At 18 h of incubation, the TRAP
262	activities in the PCB (118)-treated scales also significantly increased ( $P < 0.05$ for
263	0.025 and 2.5 ppm; <i>P</i> <0.001 for 0.25 ppm) (Fig. 4a).
264	In case of the ALP activities, it significantly increased ( $P$ <0.05) only by the
265	concentration of 2.5 ppm at the 6 and 18 h incubation (Figs. 3b and 4b).
266	
267	Changes in TRAP, cathepsin K and RANKL mRNA expressions in PCB (118)-
268	treated goldfish scales in vitro
269	The mRNA expression of osteoclastic markers (TRAP and cathepsin K)
270	increased significantly by PCB (118) (0.25 ppm) treatment (Figs. 5a and 5b).
271	Similar results were obtained in RANKL. The mRNA expression of RANKL,
272	an activating factor of osteoclasts, increased significantly in the osteoblasts in the
273	PCB (118)-treated scales (Fig. 5c).
274	
275	4. Discussion
276	In the present study, we are the first to demonstrate that PCB (118) induced
277	hypercalcemia resulting from increasing osteoclastic activity in vivo. In an in vitro
278	experiment, the data were reproduced and osteoclastic marker mRNA expression
279	as well as enzyme activity increased. In fish, PCB (118) is the highest congeners
280	compared with PCB-105, -156, -167, -123, -157, -114, -189, -77, -126, -81, or -
281	169 (Bhavsar et al. 2007). In aquatic environment, PCB (118) was detected (Hope

2008; Aksoy et al. 2011). Therefore, we paid attention to bone metabolism by

At day-1 and -2 after PCB (118) injection intraperitoneally, we detect PCB (118) in the scale. As described in the Introduction, the scales are potential internal calcium reservoir than the body skeletons, jaws and otolithes. Lake et al. (2006) reported that the correlation between the total mercury concentration of the scales and that of the muscles was high (r = 0.89). In sheep, PCB was accumulated and detected in bone at 2 months after administration (Jan et al. 2006). We therefore suggest that scale PCB content can be used as an environmental PCB monitor to estimate the environmental pollution of PCB. In the present study, we measured hydroxy-PCB which is a kind of metabolites from PCB because hydroxy-PCB possessed specific and competitive interactions with the plasma thyroid hormone transport protein, transthyretin (Lans et al. 1993). In PCB-treated scales, however, hydroxyl-PCB was not detected. Therefore, this phenomenon of osteogenesis seems to be direct action of PCB (118). In an in vivo experiments, osteoblastic activity increased by the high concentration of PCB (118)(2.5ppm). This indicates that PCB (118) is affected on osteoblasts. Osteogenesis is regulated by osteoblasts (Suda et al. 1999; Teitelbaum 2000; Lacey et al. 2012). RANKL produced by cells in the osteoblast lineage binds to the receptor activator of NF-κB (RANK) in mononuclear hemopoietic precursors and promotes the formation and activity of multinucleated osteoclasts (Suda et al. 1999; Teitelbaum 2000; Lacey et al. 2012). Our present study indicated that RANKL mRNA expression was promoted by PCB (118) treatment. In addition, osteoclastic marker (TRAP and cathepsin K) mRNA expression also increased significantly. Therefore, we strongly suggest that PCB (118) promotes osteoclastogenesis by the RANK-RANKL pathway.

In the present study, we succeeded to analysis the PCB (118) on osteoclasts and osteoblasts. Our results suggest that scale is a good model for analysis of bone metabolism. We previously demonstrated that the osteogenesis of regenerating scale is very similar to that of mammalian membrane bone and a good model of osteogenesis (Yoshikubo et al. 2005). Using this system, furthermore, we first demonstrated that calcitonin, a hypocalcemic hormone, suppressed osteoclastic activity in teleosts as well as in mammals (Suzuki et al. 2000) and that melatonin, a major hormone secreted from the pineal gland, suppressed the functions in both osteoclasts and osteoblasts (Suzuki and Hattori 2002). Osteoblasts in the scale responded to estrogen as they do in mammalian bone (Yoshikubo et al. 2005). In addition, the effects of endocrine disrupters, such as bisphenol-A (Suzuki and Hattori 2003) and tributyltin (Suzuki et al. 2006), and heavy metals, i.e., cadmium and mercury (Suzuki et al. 2004b; Suzuki et al. 2011b), on osteoblasts and osteoclasts have been examined. Moreover, we indicated that cadmium (even at  $10^{-13}$  M) responded to TRAP activity in the scale (Suzuki et al. 2004b).

In conclusion, PCB (118) disrupts bone metabolism in goldfish both *in vivo* and *in vitro* experiments. Our results suggest that PCB (118) promotes osteoclastogenesis by the RANK-RANKL pathway. Furthermore, our previous and present results indicate that scale assay system will be useful for analysis of environmental contaminant on bone metabolism and findings of PCB (118) on bone in fish may be tied in to an overall health issue for mammals in general.

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512	FIGURE LEGENDS
513	
514	Fig. 1. Effects of PCB (118) injection on scale TRAP (a) and ALP (b) activities
515	in goldfish. Each column and the vertical line represent the mean $\pm$ _SEM (n = 10
516	samples; one sample from one fish). ** indicates statistically significant
517	difference at $P < 0.01$ from the values in the control.
518	
519	Fig. 2. Effects of PCB (118) injection on plasma calcium level (mg/100 ml) in
520	goldfish. Each column and the vertical line represent the mean $\pm$ _SEM (n = 10
521	samples; one sample from one fish). ** indicates statistically significant
522	difference at $P < 0.01$ from the values in the control.
523	
524	Fig. 3. Effects of PCB (118) administration on TRAP (a) and ALP (b) activities
525	in the scales of goldfish at the 6 h of incubation. Each column and the vertical line
526	represent the mean $\pm$ _SEM (n = 10 samples; one sample from one fish).
527	* indicates statistically significant difference at $P<0.05$ from the values in the
528	control.
529	
530	Fig. 4. Effects of PCB (118) administration on TRAP (a) and ALP (b) activities
531	in the scales of goldfish at the 18 h of incubation. Each column and the vertical
532	line represent the mean $\pm$ _SEM (n = 10 samples; one sample from one fish). * and
533	*** indicate statistically significant differences at $P<0.05$ and $P<0.001$ ,
534	respectively, from the values in the control.
535	
536	
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Fig. 5. Effect of PCB (118) (0.25 ppm) in the expression of osteoclastic markers: cathepsin K (a), TRAP (b), and RANKL (c) mRNAs in the scale. The cathepsin K, TRAP and RANKL mRNA levels were normalized by the β-actin mRNA level. The values of ordinate indicate relative ratio of cathepsin K/β-actin (a), TRAP/β-actin (b), and RANKL/β-actin (c) respectively. Each column and the vertical line represent the mean  $\pm$  SEM (n = 10 samples; one sample from one fish). \* and \*\* indicate statistically significant differences at P<0.05 and *P*<0.01, respectively, from the values in the control.

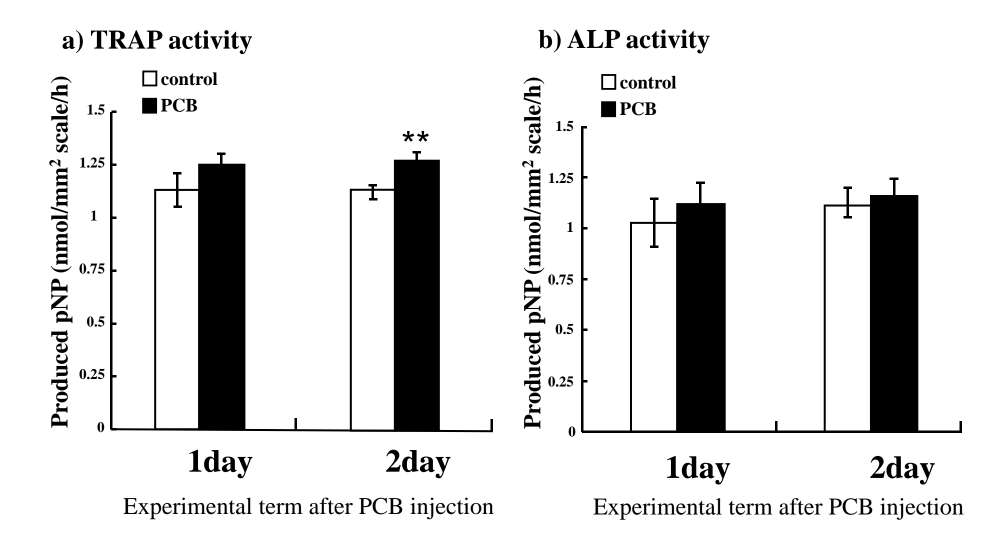


Figure 1 Yachiguchi et al.

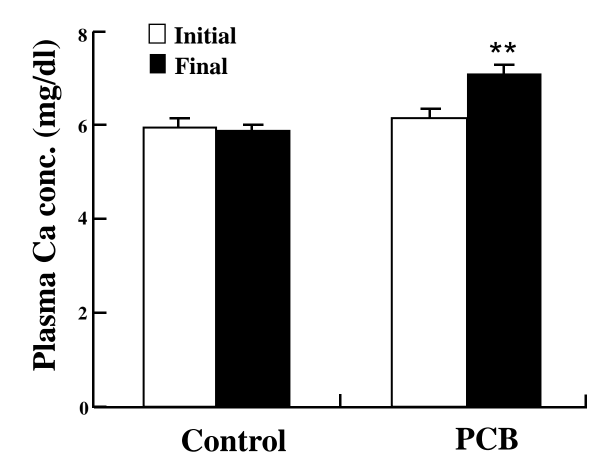


Figure 2 Yachiguchi et al.

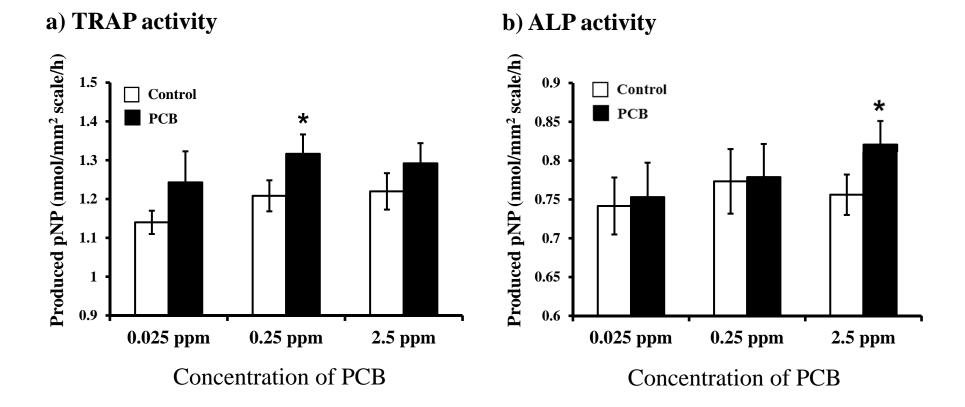


Figure 3 Yachiguchi et al.

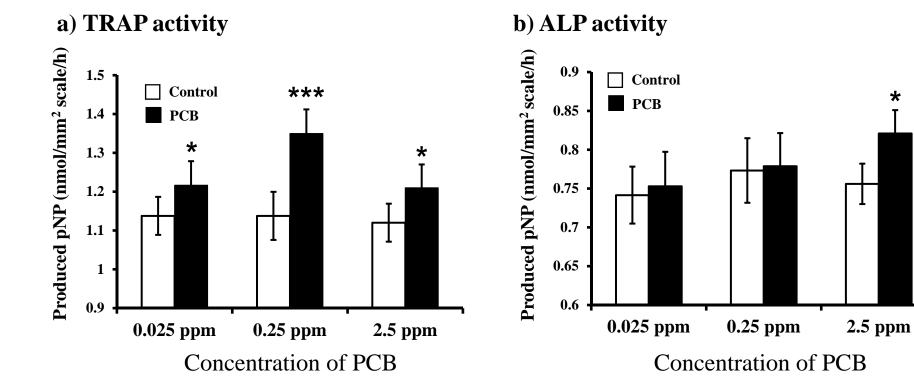


Figure 4 Yachiguchi et al.

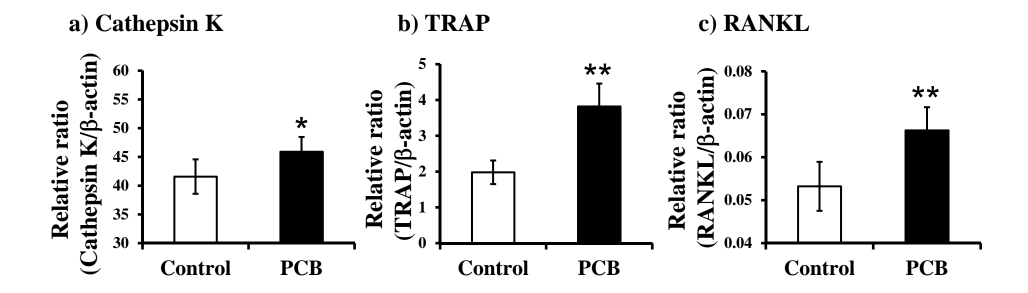


Figure 5 Yachiguchi et al.