

Title	Analysis of lead distribution in avian organs by LA-ICP-MS: Study of experimentally lead-exposed ducks and kites
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Citation	Environmental pollution, 283, 117086 https://doi.org/10.1016/j.envpol.2021.117086
Issue Date	2021-08-15
Doc URL	http://hdl.handle.net/2115/90295
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Rights(URL)	http://creativecommons.org/licenses/by-nc-nd/4.0/
Туре	article (author version)
File Information	Environmental pollution283-117086.pdf



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## 24 ABSTRACT

25 Lead poisoning of wild birds by ingestion of lead ammunition occurs worldwide. 26 Histopathological changes in organs of lead-intoxicated birds are widely known, and lead 27 concentration of each organ is measurable using mass spectrometry. However, detailed lead 28 localization at the suborgan level has remained elusive in lead-exposed birds. Here we investigated 29 the detailed lead localization in organs of experimentally lead-exposed ducks and kites by laser 30 ablation inductively coupled plasma mass spectrometry (LA-ICP-MS). In both the ducks and kites, 31 lead accumulated diffusely in the liver, renal cortex, and brain. Lead accumulation was restricted 32 to the red pulp in the spleen. With regard to species differences in lead distribution patterns, it is 33 noteworthy that intensive lead accumulation was observed in the arterial walls only in the kites. In 34 addition, the distribution of copper in the brain was altered in the lead-exposed ducks. Thus, the 35 present study shows suborgan lead distribution in lead-exposed birds and its differences between avian species for the first time. These findings will provide fundamental information to understand 36 37 the cellular processes of lead poisoning and the mechanisms of species differences in susceptibility 38 to lead exposure.

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40 **KEYWORDS:** Lead; LA-ICP-MS; imaging; waterfowl; raptor

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MAIN FINDING: Suborgan lead distribution in experimentally lead-exposed birds and its species
 differences were revealed.

### 44 INTRODUCTION

45 The use of lead shots and bullets has been regulated in many countries. However, lead poisoning 46 of wild birds is widespread all over the world (Pain et al., 2019). In the field, the ingestion of lead 47 ammunition and fishing sinkers causes many cases of lead poisoning in waterfowls and raptors (Scheuhammer and Norris, 1996; Fisher et al., 2006; Saito, 2009). According to recent statistics, 48 49 lead poisoning is estimated to kill annually a million waterfowls in Europe (Berny *et al.*, 2015; 50 Pain, 2019) and three million birds in the US (De Francisco et al., 2003). Lead poisoning in avian 51 species has been recognized in Japan since 1985 (Honda et al., 1990), and endangered raptors in 52 Japan have also been affected since 1996 (Saito, 2009). In birds, lead exposure exerts toxicity in various organs such as the liver, kidney, cardiovascular system, brain, and bone. The common 53 54 gross lesions in birds are atrophy and brownish discoloration of the liver, distended gallbladder 55 with bile, multifocal pallor areas in the myocardium, multifocal petechial hemorrhage in the 56 cerebellum, and hypoplasia of the bone marrow (Ochiai et al., 1993; Manning et al., 2019). The 57 histological lesions include hepatic hemosiderosis, degeneration and necrosis of the proximal renal 58 tubules, degeneration and necrosis of myocardium, and cerebellar perivascular hemorrhage 59 (Ochiai et al., 1993; Manning et al., 2019). Lead also interferes heme biosynthesis through the 60 inhibition of the activities of  $\delta$ -aminolevulinic acid dehydratase (ALAD) and ferrochelatase 61 (Rogan et al., 1986; Fisher, 2006; Liao et al., 2008). Therefore, the diagnosis of avian lead 62 poisoning is usually made by the characteristic histological changes, the level of ALAD activity 63 in blood, and the lead concentration in blood and organs.

Lead concentration in organs is measurable by inductively coupled plasma-mass spectrometry (ICP-MS) using tissue homogenates (Ishii *et al.*, 2017; Togao *et al.*, 2020). In addition, special staining methods for lead in tissue sections have been developed to detect gunshot residues (Neri *et al.*, 2007; Turillazzi *et al.*, 2013). However, the detailed lead distribution in lead-exposed
animals at the suborgan level cannot be analyzed by these methods. Recently, we showed the tissue
distribution of lead in lead-exposed mice using laser ablation (LA)-ICP-MS (Togao *et al*, 2020).
LA-ICP-MS can identify metals in tissue sections and is useful to reveal the detailed tissue
distribution of metals (Ek *et al.*, 2004; Ishii *et al.*, 2018).

In the present study, we established a model of low-dose lead exposure in waterfowls and raptors by administration of one to three lead pellets, reproducing the low-dose lead exposure found in the field. Then, we investigated the histological distribution of lead in organs of the leadexposed birds using LA-ICP-MS. In addition, we compared the tissue distribution of lead between waterfowls and raptors because the sensitivity to lead poisoning is reported to be different among avian species (De Francisco *et al.*, 2003).

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#### 79 MATERIALS AND METHODS

#### 80 Animal experiments

Animal experiments were performed in strict accordance with the Regulations for Animal Experiments and Related Activities at Hokkaido University. The protocols for animal experiments were approved by the Association for the Assessment and Accreditation of Laboratory Animal Care International and the Institutional Animal Care and Use Committee of Hokkaido University (approval No. 18-0092 and No. 19-0033).

Seven, bred, eight-week-old Muscovy ducks (*Cairina moschata*; body weight, 3.3 - 3.8 kg) were purchased from Sankyo Labo Service. The ducks were randomly divided into two groups; untreated control (n = 3) and lead-treated (n = 4). The ducks were housed individually in cages in controlled light (12 h light/dark cycle) and constant temperature ( $23 \pm 2^{\circ}$ C) with free access to 90food and water. All ducks were acclimated for a week before treatment and kept on a fresh diet.91Lead-treated ducks were given three lead pellets  $(240 \pm 1.7 \text{ mg})$  in a style of oral forced ingestion.92Control and lead-treated groups were euthanized with overdose of pentobarbital sodium on 29 or9330 days after the lead treatment.

94 Five black kites (*Milvus migrans*; body weight, 1.0 - 1.1 kg) were kept in the Institute for Raptor 95 Biomedicine Japan. All kites were unsuitable for release because most had persistent wing 96 damages. These kites were otherwise in good condition. The kites were placed individually in 97 outdoor cages. Each pen was furnished with a log for perching and a pan of water. All kites were 98 acclimated to the pens for a week or more before the treatment and were kept on a diet. The kites 99 were randomly divided into two groups; untreated control (n = 2) and lead-treated (n = 3). Lead-100 treated kites were given one lead pellet (77.9 - 88.4 mg) mixed with venison. To confirm the 101 existence of lead pellet in the gastrointestinal tract, the kites were radiographed every day for 14 102 days after the lead administration. In two kites (Kite-Lead-2 and Kite-Lead-3), the lead pellet 103 disappeared on radiographs probably due to regurgitation or excretion at 7 and 10 days after the 104 lead administration, respectively. Therefore, these kites were dosed one lead pellet again within 105 24 hours. Control and lead-treated groups were euthanized with overdose of pentobarbital sodium 106 on 29 or 30 days after the first lead treatment.

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#### 108 **Blood collection**

Blood samples (5 ml and 4 ml from the ducks and kites, respectively) were obtained from the brachial veins at 28 days after the lead treatment. Blood was quickly heparinized to avoid coagulation and kept on ice until further processing within 2 hours.

### 113 **Tissue sample collection**

Liver, spleen, kidney, heart, lung, cerebrum, midbrain, cerebellum, and bone marrow were collected from the euthanized birds and divided into three pieces. One was used for quantitative analysis of lead by ICP-MS, another was used for histopathological analysis, and the other was used for imaging analysis of lead by LA-ICP-MS.

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# 119 Quantitative analysis of lead by ICP-MS

120 Analyses of lead concentrations in bird organs (liver, spleen, kidney, heart, cerebrum, midbrain, 121 cerebellum, and blood) were performed as reported previously (Yabe et al., 2015). The amounts 122 of samples analyzed are summarized in Appendix A: Table A.1. Samples were digested with nitric 123 acid (HNO<sub>3</sub>) and hydrogen peroxide ( $H_2O_2$ ) in microwave. The concentration of lead was 124 measured with ICP-MS 7700 series (Agilent Technology). Analytical quality control was 125 performed using DOLT-4 (dogfish liver) and DORM-3 (fish protein) (National Research Council 126 of Canada) certified reference materials. Replicate analyses of these reference materials showed 127 good recoveries (95 - 105%); the linearity range of standard solution was 0 - 500  $\mu$ g/L (0, 0.25,  $0.5, 1, 5, 10, 25, 50, 100, 250, 500 \mu g/L, R^2$  of standard curve was more than 0.9999); the limit of 128 129 detection was 0.001  $\mu$ g/kg; and the limit of quantification was 0.003  $\mu$ g/kg. The limit of 130 quantification was determined as  $10 \times$  standard deviation of the intercept / the average of the slope 131 obtained from seven measurements of the standard solutions. For the analysis of lead concentration, Thallium (<sup>205</sup>TI) was used as an internal standard for the lead concentration analysis. 132

133

# 134 Histopathological analysis

For histopathological analysis, the collected organs (liver, spleen, kidney, heart, lung, cerebrum,
midbrain, cerebellum, and bone marrow) were fixed in 10% buffered formalin for 48 hours at room
temperature and embedded in paraffin. The embedded tissues were sectioned at a thickness of 4
µm and were stained with hematoxylin and eosin (H&E).

139

### 140 Lead staining

Lead staining was performed as previously reported (Turillazzi *et al.*, 2013; Neri *et al.*, 2007). Deparaffinized and rehydrated tissue sections were incubated with a staining solution containing 2.5 mg/ml sodium rhodizonate (FUJIFILM Wako Chemicals) and 1.67 mg/ml tartaric acid (Sigma-Aldrich) for 1 minute. The sections were counterstained with hematoxylin for 30 seconds, washed with distilled water, dehydrated and mounted. Kidney sections were also stained with acid-fast stain to detect intranuclear lead inclusion body.

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## 148 Imaging analysis of lead by LA-ICP-MS

149 For LA-ICP-MS, the collected organs (liver, spleen, kidney, heart, cerebrum, midbrain, and 150 cerebellum) were washed with sterilized phosphate-buffered saline (PBS) to remove blood and 151 then embedded in Tissue-Tek O.C.T. Compound (Sakura Finetek). The embedded tissues were 152 frozen in isopentane, which had been cold with dry ice, allowed to dry and then stored at -80°C. 153 The frozen tissues were sectioned at a thickness of 15 µm using a cryostat Leica CM 3500. Some 154 neighboring sections were cut to a thickness of 8 µm for H&E staining. The sections were analyzed 155 using an LA system (NWR213; esi Japan, Tokyo, Japan, working at a wavelength of 213 nm, pulse duration of 4 ns, and fluence of 0.5-0.6 J cm<sup>-2</sup>) associated with an ICP-MS 8800 series (Agilent 156 157 Technology) and scanned by a focused laser beam. Laser spot size, scan speed line and offset

between line were set at 100 µm, 100 µm s<sup>-1</sup> and 100 µm, respectively. ICP-MS conditions were 158 159 the following: RF plasma source, 1600 W; He carrier gas, 0.8 L min<sup>-1</sup>. Measured isotope (dwell time, second) were as follows: <sup>13</sup>C (0.005), <sup>25</sup>Mg (0.005), <sup>31</sup>P (0.005), <sup>43</sup>Ca (0.005), <sup>55</sup>Mn 160 (0.005), <sup>57</sup>Fe (0.005), <sup>65</sup>Cu (0.005), <sup>66</sup>Zn (0.005), <sup>206</sup>Pb (0.01), <sup>207</sup>Pb (0.01), <sup>208</sup>Pb (0.01). In this 161 162 analysis, no quantification of Pb was conducted due to lack of suitable reference materials for calibration, however intensity of Pb (and other elements) was normalized to <sup>13</sup>C (carbon) intensity 163 164 as Wu et al. (2009), Johnston et al. (2019) and others have utilized to normalize the ablation efficiency. From the continuous list of raw pixel values data, elemental images were reconstructed 165 166 using LA-ICP-MS Image generator house-made software iQquant2 (Kawakami et al., 2016).

167

#### 168 **RESULTS**

## 169 Clinical signs

170 Clinical signs of the ducks are summarized in Table 1. Duck-Lead-3 showed mild anorexia and
171 lethargy at 21 days after the lead treatment. The other ducks appeared healthy. The control group
172 did not show any clinical signs.

173 Clinical signs of the kites are summarized in Table 2. Kite-Lead-2 showed moderate anorexia, 174 lethargy and exercise intolerance at 7 days after the lead administration, and Kite-Lead 1 showed 175 mild anorexia and lethargy at one day after the lead treatment. The other kites appeared healthy. 176 The control group did not show any clinical signs.

177

# 178 Necropsy findings

Necropsy findings of the ducks are summarized in Table 1. An eroded lead pellets remained in the
stomach of all the lead-treated ducks except for Duck-Lead-3. Duck-Lead-1, Duck-Lead-2 and

181 Duck-Control-2 showed focal discolored foci on the surface of the liver. In the Duck-Lead-4, mild 182 hepatomegaly and multifocal yellowish foci on the surface of the liver were observed. In addition, 183 the spleen was mildly swollen, and the right metanephros was defective. Duck-Lead-3, Duck-184 Control-1 and Duck-Control-3 did not show any gross pathological changes.

Necropsy findings of the kites are summarized in Table 2. No lead pellet was found in the gastrointestinal tract of all the lead-treated kites. Kite-Lead-1 and Kite-Lead-3 showed focal yellowish-white foci on the surface of the liver. The heart of Kite-Lead-2 was mildly fragile. No gross pathological change was noted in the other organs of the lead-treated kites and the control kites.

190

# 191 Histopathological findings

Histopathologic findings of the ducks are summarized in Table 1. Duck-Lead-1, Duck-Lead-2 and
Dcuk-Control-3 showed hydropic degeneration of hepatocytes, while Duck-Lead-3, Duck-Lead-4
and Dcuk-Control-1 showed vacuolar degeneration of hepatocytes. All ducks except DuckControl-2 showed vacuolar degeneration of the renal tubular epithelia. Deposition of amyloid in
the hepatic portal area, sinusoid of the liver, and white pulp of the spleen was noted in Duck-Lead4.

Histopathological findings of the kites are summarized in Table 2. In Kite-Lead-1, an enlargement of the collecting ducts of the kidney and mild hypoplasia of the bone marrow were observed. Mild myocarditis and mild pulmonary congestion were noted in Kite-Lead-2. All kites showed the deposition of lipofuscin in the renal tubular epithelia. Deposition of amyloid in sinusoid of the liver was noted in Kite-Control-2. Intranuclear lead inclusion body was not found in the acid-fast stained kidney sections of the ducks and kites (data not shown). Lead staining using the sodium rhodizonate reaction was negative in all the ducks and kites (data not shown).

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# 207 ICP-MS analysis

In the quantitative analysis of lead by ICP-MS, Duck-Lead-3 and Kite-Lead-1 showed the highest lead concentrations in each group. The liver and kidneys showed higher lead concentrations compared with the other organs examined (Table 3). The lead concentrations in untreated control groups were less than 0.01 mg/L or mg/kg (data not shown).

212

#### 213 LA-ICP-MS analysis

Tissue distributions of lead were examined in Duck-Lead-3 and Kite-Lead-1 by LA-ICP-MS, as these birds showed the highest lead concentrations. In addition, the cerebrum, midbrain, and cerebellum of Duck-Lead-1, or the cerebrum, midbrain, cerebellum, and heart of Kite-Lead-3 were also examined to confirm the characteristic patterns of lead distribution.

218 In the lead-exposed duck, diffuse lead accumulation except for veins was noted in the liver 219 (Figure 1A). Connective tissues surrounding veins showed slightly higher intensities. The lumen 220 of the gallbladder also showed lead accumulation. In the spleen, lead accumulation was restricted 221 to the red pulp (Figure 1B). In the kidney, diffuse lead accumulation was observed in the cortex 222 (Figure 1C). The cortical areas surrounding the interlobular veins showed higher intensities 223 compared with the areas around the central veins. Notably, the medullary cones did not show lead 224 accumulation. In the brain, lead accumulated diffusely in the cerebrum (Figure 1D). In the 225 cerebellar cortex, the gray matter showed diffuse lead accumulation, with higher intensities in the

Purkinje cell layer (Figure 1E). In the midbrain, the optic tectum (stratum griseum et fibrosum
superficiale, stratum griseum centrale, stratum griseum periventriculare), central gray substance,
and oculomotor nerves showed intensive lead accumulation (Figure 1F). Lead accumulation was
not observed in the untreated control duck (Figures 1G and 1H).

230 In the lead-exposed kite, intensive lead accumulation in the hepatic arterial walls was observed 231 in addition to diffuse accumulation in hepatic parenchyma (Figure 2A). In the spleen, the wall of 232 splenic artery showed much higher amount of lead accumulation than those of the red pulp (Figure 233 2B). In the kidney, the patterns of lead distribution were the same as those of the duck kidney 234 (Figure 2C). Diffuse lead accumulation was observed in the renal cortex, with higher intensities 235 around the interlobular veins. Materials contained in dilated collecting ducts also showed lead 236 signals. In the cerebrum, intensive lead accumulation was noted in the peripheral area of the 237 hyperpallium, hippocampus, and hypothalamus, in addition to diffuse accumulation in 238 parenchyma (Figure 2D). In the cerebellum, lead accumulated diffusely in the gray matter, with 239 higher intensities in the Purkinje cell layer (Figure 2E). In the midbrain, the optic tectum (stratum 240 griseum et fibrosum superficiale, stratum griseum centrale, stratum griseum periventriculare), 241 central gray substance, nucleus mesencephalicus lateralis pars dorsalis, brachium cunjuctivum, 242 and oculomotor nerves showed intensive lead accumulation (Figure 2E). Lead accumulation was 243 not observed in the untreated control kite (Figures 2F and 2G). Thus, the pattens of lead 244 accumulation in the lead-exposed kites were basically similar to those of the lead-exposed ducks, 245 with more prominent lead accumulation in some brain regions, *e.g.*, hyperpallium, hippocampus, 246 hypothalamus, and optic tectum. Meanwhile, the intensive lead accumulation in the arterial wall 247 was characteristic to the lead-exposed kite. The intensive lead accumulation in the arterial wall

248 was also confirmed in the heart of the lead-exposed kite (Figure 2H). The walls of coronary artery 249 and aorta showed high amount of lead accumulation in addition to those of the cardiac cartilage.

Taking advantage of LA-ICP-MS that enables to visualize the distribution of essential elements in tissue sections, we also examined localization of magnesium, phosphorus, calcium, manganese, iron, copper, zinc at the suborgan level in the lead-exposed ducks and kites. Notably, the distribution of copper was altered in the cerebrum of the lead-exposed ducks (Figure 3). Copper accumulated in the entopallium only in the lead-exposed ducks and not in the control duck or the lead-exposed kites. Localization of the other elements in each organ was not altered by the lead administration (data not shown).

257

#### 258 **DISCUSSION**

In the present study, lead distribution in organs of experimentally lead-exposed ducks and kites were investigated at the suborgan level by LA-ICP-MS. Although almost all of the ducks and kites lacked lead-associated pathological changes due to the low-dosage of lead administration, tissue distribution of lead could be clearly identified. In addition, species differences in lead distribution patterns were also revealed.

In the liver, lead accumulated diffusely in parenchyma in both the duck and kite. This distribution pattern is the same as those in lead-exposed mice (Togao *et al.*, 2020). Lead-intoxicated animals show degeneration of hepatocytes and hemosiderosis (Ochiai *et al.*, 1993; Jarrar and Taib, 2012; Hegazy and Fouad, 2014). Thus, the diffuse distribution of lead in hepatocytes is compatible with the histopathological changes in lead-intoxicated animals.

In the spleen, lead accumulated only in the red pulp in both the duck and kite. This finding is in line with the report that 95% of lead in blood accumulates in erythrocytes (Hernández-Avila *et al.*, 271 1998). In addition, hemosiderin-laden (erythrophagocytic) macrophages increase in the red pulp 272 in lead-intoxicated waterfowls (Ochiai *et al.*, 1993). It has been reported that lead specifically 273 affects macrophages in the red pulp in lead-exposed mice (Corsetti *et al.*, 2017). Therefore, the 274 lead accumulation in the red pulp may reflect the lead accumulation in erythrocytes and 275 erythrophagocytic macrophages.

276 In the kidneys of the duck and kite, lead accumulated diffusely in the cortical area, particularly 277 around the interlobular veins, without accumulation in the medullary cones. These distribution 278 patterns are different from those observed in lead-exposed mice, in which corticomedullary 279 boundaries show higher amount of lead accumulation than the cortex (Togao et al., 2020). Birds 280 and mammals have different kidney structures in terms of nephron, portal system, and stratified 281 cortex and medulla (Morild et al., 1985; Harr, 2002). Thus, these structural differences may 282 account for the different lead distribution. Meanwhile, the lack of lead accumulation in the renal 283 medulla is common in both birds and mammals. The appearance of inclusion bodies composed of 284 lead,  $\alpha$ -synuclein and metallothionein in the proximal tubules is a histological hallmark of lead-285 intoxicated animals (Moore and Goyer, 1974; Qu et al., 2002; Zuo et al., 2009). In addition, lead-286 intoxicated animals show degeneration and necrosis of the proximal tubules. Therefore, the diffuse 287 lead distribution in the cortical area is in line with the histopathological changes of the lead-288 intoxicated animals.

In the brain, lead accumulated diffusely in the cerebrum, cerebellar cortex, and midbrain in both the ducks and kites, with higher intensities in the Purkinje cell layer, optic tectum, central gray substance, and oculomotor nerves. In the kites, hippocampus, hyperpallium, and hypothalamus also showed higher amount of lead accumulation. These patterns of lead distribution partially overlap with those of rodents, in which lead preferentially accumulates in hippocampus and the

294 cerebral cortex (Lefauconnier et al., 1983; Al-Shimali et al., 2016; Togao et al., 2020). In lead-295 exposed rodents, neuronal damages are mainly observed in hippocampus, the parietal cortex, and 296 Purkinje cells (Sharifi et al., 2002; Dribben et al., 2011; Gargouri et al., 2012; Owoeye and 297 Onwuka, 2016), and lipid peroxidation is noted in thalamus, hippocampus, the parietal cortex and 298 striatum in rats (Villeda-Hernández et al., 2001). In birds, lead exposure causes neurological 299 dysfunction like blindness, head tilt, and seizures (Fallon et al., 2017). The intensive lead 300 accumulation in the Purkinje cell layer and optic tectum may account for these clinical signs in 301 birds. In addition, the lead accumulation in hippocampus of the kites may associate with the finding 302 that lead exposure has a negative impact on learning and behavior in avian species (Burger and 303 Gochfeld, 2005; Ecke *et al.*, 2017). Further studies will be needed to investigate the relationship 304 between the lead distribution in the brain and neurological signs in lead-exposed birds. In addition, 305 the identification of brain cell types which have higher amount of lead will aid to unveil the 306 mechanisms of lead-induced neurotoxicity. For example, astrocytes generate and store glutathione 307 sulfhydryl enzymes that can bind lead, and the interaction between astrocytes and neurons is 308 inhibited by lead through the prevention of glutamate and glycogen metabolisms (Strużyńska et 309 al., 2005; Liu et al., 2015).

Meanwhile, lead exposure caused copper accumulation in the entopallium of the duck brain. It has been reported that lead administration increases copper concentrations in the brain or in cultured astrocytes (Tiffany-Castiglioni *et al.*, 1987; Sierra *et al.*, 1989). Lead exposure induces copper uptake by up-regulation of the expression of Cu transporter 1 (CTR1) and reduces copper efflux by down-regulation of the expression of ATPase copper transporting alpha (ATP7A) (Zheng *et al.*, 2014). The entopallium is one of the visual centers in the bird brain and a target of the tectofugal visual pathway, *i.e.*, a visual route travels from the eyes to optic tectum to thalamus and 317 then to the entopallium (Karten and Hodos, 1970). The clinical relevance and molecular 318 mechanisms of the copper accumulation in the entopallium of the lead-exposed ducks needs to be 319 investigated in the future.

320 The most striking difference in lead distributions between the ducks and kites was the intensive 321 accumulation in the arterial walls of the kites. In lead-intoxicated eagles, hemorrhage and ischemia 322 caused by fibrinoid necrosis of small and medium caliber arteries are frequently found in the heart, 323 brain, and eyes (Manning et al., 2019). Thus, arterial walls may be one of the target organs of lead-324 poisoning in raptors. Lead exposure causes cardiovascular degeneration also in humans and 325 rodents (Navas-Acien et al., 2007; Fiorim et al., 2011; Ozturk et al., 2014; Nascimento et al., 326 2015). In rats, lead exposure increases the activity of plasma matrix metalloproteinase 9 (MMP9) 327 (92-kDa type IV collagenase) (Nascimento et al., 2015), which can digest type IV collagen in the 328 basement membrane of blood vessels and elastin of the tunica media of blood vessels (Wilhelm et 329 al., 1989; Collier et al., 1988; Yasmin et al., 2005). Further, lead exposure increases the expression 330 of MMP2 (72-kDa type IV collagenase) and MMP9 in hippocampus and the cerebral cortex of 331 mice, resulting in cerebral vascular lesions (Ning et al., 2016). In addition, lead-induced expression 332 of MMP2 and MMP9 affects the blood-brain-barrier permeability through degradation of tight 333 junction proteins (Liu *et al.*, 2017). Although the distribution of MMPs in avian species has not 334 been investigated, lead may bind and activate MMP2 and MMP9 in the arterial walls in raptors. 335 The molecular mechanisms of the predisposition to the lead accumulation in the arterial walls in 336 raptors should be investigated in the future.

Little is known about the toxicity caused by low-level lead exposure in birds. To date, lead toxicity in birds has only been investigated by high-dose lead exposure (Franson *et al.*,1983; Hoffman *et al.*, 1985; Mautino and Bell, 1986; Pain, 1990; Redig *et al.*, 1991; Rocke and Samuel,

340 1991; Ochiai et al., 1993; Hiraga et al., 2008). In mammals, low-dose lead exposure exerts toxic 341 effects (Dribben et al., 2011; Flora et al., 2012; Lanphear et al., 2018; Rahman et al., 2018), and 342 Centers for Disease Control and Prevention (CDC) suggested that the safe blood lead level in 343 humans should be reduced from 10 µg/dL to 5 µg/dL (CDC, 2012). Thus, it is currently considered 344 that previous effect-level 'thresholds' should be abandoned in the field of avian lead poisoning 345 (Pain et al., 2019). To reproduce the low-dose lead exposure found in the field, we established a 346 model of low-dose lead exposure in waterfowls and raptors in the present study. Although these 347 birds lacked apparent lead-associated pathological changes, the imaging analysis using LA-ICP-348 MS clearly identified lead distribution in organs. In addition, the alteration of copper distribution 349 in the brain was also detected by LA-ICP-MS. Thus, the present study will provide useful 350 information to understand the mechanisms of lead poisoning in birds caused by low-level exposure 351 in the field.

352

#### 353 CONCLUSIONS

Here we demonstrate detailed lead distribution in organs of experimentally lead-exposed birds and its differences between avian species for the first time. The present study will pave the way for better understanding the cellular processes of lead poisoning and the mechanisms of species differences in susceptibility to lead exposure.

358

### 359 ACKNOWLEDGEMENT

We would like to express our appreciation to all the members of Laboratory of Comparative
Pathology and Laboratory of Toxicology, Faculty of Veterinary Medicine, Hokkaido University
and the Institute for Raptor Biomedicine Japan for helpful discussions, encouragement and support.

### 364 FUNDING

365 This work was supported by the Grants-in-Aid for Scientific Research from the Ministry of 366 Education, Culture, Sports, Science and Technology of Japan awarded to T. Matsukawa (No. 367 19H01081), M. Ishizuka (No. 16H0177906, 18K1984708, 18KK028708 and JPMXS0420100620), 368 Y. Ikenaka (No. 18H0413208), and S.M.M. Nakayama (No. 17KK0009, 20K20633). This work 369 was also supported by the foundation of JSPS Bilateral Open Partnership Joint Research Projects 370 (JPJSBP120209902) and the Environment Research and Technology Development Fund (SII-1/3-371 2, 4RF-1802/18949907) of the Environmental Restoration and Conservation Agency of Japan. We 372 also acknowledge financial support from the Soroptimist Japan Foundation, the Nakajima 373 Foundation, the Sumitomo Foundation, the Nihon Seimei Foundation, act beyond trust, the Japan 374 Prize Foundation, and Triodos Foundation. This research was also supported by JST/JICA, 375 SATREPS (Science and Technology Research Partnership for Sustainable Development; No. 376 JPMJSA1501). The funders had no role in study design, data collection and analysis, decision to 377 publish, or preparation of the manuscript.

378

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Duck	Clinical signs	Necropsy findings	Histopathological findings	
Lead-1	None	Liver: focal dark red foci	Liver: hydropic degeneration of hepatocytes, diffuse, moderate; subcapsular hemorrhage and edema, focal, mild Kidney: vacuolar degeneration of the renal tubules, diffuse, mild	
Lead-2	None	Liver: focal yellowish- white foci	Liver: hydropic degeneration of hepatocytes, diffuse, moderate Kidney: vacuolar degeneration of the renal tubules, diffuse, mild	
Lead-3	Mild anorexia and lethargy	None	Liver: vacuolar degeneration of hepatocytes, diffuse, moderate Kidney: vacuolar degeneration of the renal tubules, multifocal, mild	
Lead-4	None	Liver: mild hepatomegaly and multifocal yellowish foci Spleen: mild splenomegaly Kidney: defect of the right metanephros	Liver: deposition of amyloid within the hepatic portal area, sinusoid and white pulp, diffuse, moderate; vacuolar degeneration of hepatocytes, diffuse, moderate Kidney: vacuolar degeneration of the renal tubules, diffuse, mild	
Control-1	None	None	Liver: vacuolar degeneration of hepatocytes, diffuse, moderate Kidney: vacuolar degeneration of the renal tubules, multifocal, mild	
Control-2	None	Liver: focal white foci	None	
Control-3	None	None	Liver: vacuolar degeneration of hepatocytes, diffuse, moderate Kidney: vacuolar degeneration of the renal tubules, multifocal, mild	

**Table 1.** Summary of clinical signs, necropsy findings and histopathological findings of the ducks

Kite	Age and	Clinical signs	Necropsy	Histopathological findings
	sex		findings	
Lead-1	6 у,	Mild anorexia	Liver: focal	Kidney: deposition of
	female	and lethargy	yellowish-white	lipofuscin in the renal
			foci	tubules, diffuse, mild;
				enlargement of the collecting
				ducts, moderate
				Bone marrow: hypoplasia,
				mild
Lead-2	2 у,	Moderate	Heart: mild	Kidney: deposition of
	female	anorexia,	fragileness	lipofuscin in the renal
		lethargy and		tubules, diffuse, mild
		exercise		Heart: myocarditis,
		intolerance		lymphocytic, focal, mild
				Lung: pulmonary congestion,
				diffuse, mild
Lead-3	2 у,	None	Liver: focal	Kidney: deposition of
	male		yellowish-white	lipofuscin in the renal
			foci	tubules, diffuse, mild
Control-	6 у,	None	None	Kidney: deposition of
1	female			lipofuscin in the renal
				tubules, diffuse, mild
Control-	3 у,	None	None	Kidney: deposition of
2	female			lipofuscin in the renal
				tubules, diffuse, mild
				Spleen: deposition of amyloid
				within the splenic sinusoid,
				focal

**Table 2.** Summary of clinical signs, necropsy findings and histopathological findings of the kites

Organ	Duck-Lead-3 <sup>a</sup>	Duck-Lead-1	Kite-Lead-1	Kite-Lead-3
Blood <sup>b</sup>	2.95	—	0.99	—
Liver	6.18	—	1.70	—
Spleen	2.80	—	0.35	—
Pronephros	4.89	—	4.90	—
Mesonephros	5.41	—	3.04	—
Metanephros	5.80	—	4.15	—
Cerebrum	0.68	0.18	0.53	0.24
Midbrain	0.81	0.32	0.76	1.52
Cerebellum	0.93	0.73	0.63	0.69
Heart	—	—	—	0.03

**Table 3.** Lead concentrations in the duck and kite organs

<sup>a</sup> Data are expressed as mg/L in blood or mg/kg in wet weight in the other organs.

<sup>b</sup> Lead concentration in blood at 28 d after the lead administration.



Figure 1. Lead distribution in the duck organs. (A) Lead accumulated diffusely in the liver. Lead
signals were also observed in the gallbladder (arrowhead). Duck-Lead-3. (B) Lead accumulated in
the red pulp of the spleen. The white pulp (arrowheads) lacked lead accumulation. Duck-Lead-3.
(C) Lead accumulated diffusely in the cortical area of the kidney, with intensive accumulation
around the interlobular veins (arrowheads). Duck-Lead-3. (D-F) Lead accumulated diffusely in the

582 cerebrum (D, Duck-Lead-3), cerebellar cortex (E, Duck-Lead-1), and midbrain (F, Duck-Lead-3). 583 The Purkinje cell layer, optic tectum, central gray matter, and oculomotor nerves showed higher 584 amount of lead accumulation. (G, H) Lead accumulation was not observed in the organs of the 585 untreated control duck (G, spleen; H, cerebrum; Duck-Control-3). The areas enclosed by the 586 dashed lines are shown in the H&E images. Scale bars: 5 mm in LA-ICP-MS images and 500 μm 587 in H&E images.



588 Figure 2. Lead distribution in the kite organs. (A) Lead accumulated diffusely in the liver, with 589 intensive accumulation in the arterial walls (arrowheads). Kite-Lead-1. (B) Lead accumulated in 590 the red pulp of the spleen, with intensive accumulation in the arterial walls (arrowheads). Kite-591 Lead-1. (C) Lead accumulated diffusely in the cortical area of the kidney. Lead accumulated also 592 in the dilated collecting ducts (arrow). The medullary cones (arrowheads) lacked lead 593 accumulation. Kite-Lead-1. (D) Lead accumulated diffusely in the cerebrum, with intensive 594 accumulation in the periphery of the hyperpallium, hippocampus (arrowhead), and hypothalamus 595 (arrow). Kite-Lead-1. (E) Lead accumulated diffusely in the cerebellar cortex and midbrain, with

intensive accumulation in the Purkinje cell layer, optic tectum, central gray matter, nucleus
mesencephalicus lateralis pars dorsalis, brachium cunjuctivum, and oculomotor nerves. Kite-Lead3. (F, G) Lead accumulation was not observed in the organs of the untreated control kite (F,
cerebrum; G, cerebellum and midbrain; Kite-Control-2). (H) Lead accumulated in the arterial walls
of the heart in addition to the cardiac cartilage (arrowheads). Kite-Lead-3. The areas enclosed by
the dashed lines are shown in the H&E images. Scale bars: 5 mm in LA-ICP-MS images and 500
µm in H&E images.



Figure 3. Copper distribution in the duck brains. (A, B) Copper accumulated in the entopallium
(encircled by the dashed lines) of the lead-exposed ducks (A, Duck-Lead-3; B, Duck-Lead1). (C)

- 605 Copper accumulation in the entopallium was not observed in the untreated control duck (Duck-
- 606 Control-3). Scale bars: 5 mm.