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1 **Original Investigation**

2 **CT morphology of anomalous systemic arterial supply to normal lung in dogs**

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17

18

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21

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23 The authors declare no conflicts of interest associated with this manuscript.

24

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26 None of the findings of this study have been presented or published previously elsewhere.

27

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29 An EQUATOR network checklist was not used.

30

31 **Abbreviations:** ASANL, anomalous systemic arterial supply to the normal lung

32

33 **ABSTRACT**

34 Anomalous systemic arterial supply to the normal lung (ASANL) is a rare congenital
35 anomaly in humans, in which the systemic arteries supply the basal segments of the lower lobe.
36 It has a normal bronchial connection, but lacks a normal pulmonary artery. This anomaly has
37 never been reported in the veterinary literature. The objectives of this retrospective descriptive
38 study were to characterize the CT findings and clinical features of ASANL, and to determine
39 the breed predisposition in a population of referral canine cases. Thoracic CT images, in which
40 the caudal lung lobes were fully inflated and the pulmonary artery could be traced to the
41 periphery, were reviewed. A total of 1,950 dogs were enrolled, and the aberrant vasculature
42 equivalent to ASANL in humans was detected in 48 dogs. Shetland Sheepdogs (7/48, OR=8.0,
43 $P<0.00001$), Miniature Dachshunds (19/48, OR=3.9, $P<0.00001$), and Labrador Retrievers
44 (6/48, OR=4.5, $P=0.0009$) were over-represented. The affected lung lobes were the right caudal
45 lobe (24/48, 50%), the left caudal lobe (21/48, 43.8%), and bilateral caudal lobes (3/48, 6.3%).
46 The aberrant vessels originated from the left gastric artery (14/48), descending thoracic aorta
47 (8/48), celiac artery (6/48), and splenic artery (1/48). In the remaining 19 cases, the origin of
48 the aberrant vessels could not be determined. Although the clinical significance of ASANL in
49 dogs remains unclear, surgeons should be aware of this finding prior to lobectomy of the caudal
50 lung lobes to avoid intraoperative systemic arterial bleeding.

51 **INTRODUCTION**

52 The normal lung is supplied by the pulmonary and bronchial arteries. Pulmonary
53 arteries supply the distal portion of the respiratory bronchiole, alveolar duct, alveoli, and the
54 thin pleura, and the bronchial arteries supply the tracheobronchial lymph nodes, peribronchial
55 connective tissue, and the bronchial mucous membrane.¹ The bronchial arteries mostly arise
56 from the bronchoesophageal artery, which arises from the right fifth intercostal artery close to
57 its origin from the aorta, crosses the left face of the esophagus, and branches at the tracheal
58 bifurcation of the left and right bronchial arteries. The bronchial arteries communicate with the
59 pulmonary arteries through the capillary bed at the level of the respiratory bronchioles.^{1,2}
60 Systemic arterial supply to the lungs refers to all abnormal vessels supplying the lung, arising
61 directly or indirectly from the aorta in humans.³ Among several types of systemic arterial
62 supply to lungs, anomalous systemic arterial supply to normal lung (ASANL), also known as
63 anomalous systemic arterial supply to the basal segments of the lung, is the rarest form; in this
64 type, a portion of the lung lacking a normal pulmonary artery receives systemic arterial supply
65 and drains through the normal pulmonary veins.⁴⁻⁶ Based on our review of the literature,
66 ASANL has never been reported in the veterinary literature.

67 ASANL is often asymptomatic and may be diagnosed following detailed examination
68 of an incidental cardiac murmur, for example.⁷ When symptomatic, clinical signs include
69 hemoptysis, heart murmur, exertional dyspnea, tachypnea, and cough.⁸ Although most adult

70 patients are asymptomatic, patients with this disease are recommended to undergo surgery or
71 endovascular embolization, because of the potential risk of hemoptysis due to pulmonary
72 hypertension and heart failure due to left-to-left shunt.⁷ It is very important to differentiate
73 ASANL from other pulmonary vascular anomalies, while selecting the appropriate surgical
74 procedure. Therefore, the computed tomographic morphology of ASANL has been investigated.
75 Reports suggest that Asians are predisposed to have ASANL.⁹ If a pulmonary vascular anomaly
76 equivalent to ASANL exists in dogs, it is important to investigate its clinical findings.

77 The aim of the present retrospective descriptive study was to investigate whether
78 ASANL exists in dogs, and if it does, to evaluate its CT morphology, clinical significance, and
79 breed predisposition in dogs by reviewing CT images in a population of canine referral cases.

80

81 **METHODS**

82 **Selection and Description of Subjects**

83 This retrospective descriptive study was performed at the Hokkaido University
84 Veterinary Teaching Hospital. Approval by the Animal Care and Use Committee of our
85 institution was not required due to the retrospective nature of the study. Consent was obtained
86 from the owners of all dogs recruited for this study. Inclusion criteria were applied to the client-
87 owned dogs that underwent thoracic CT examination for diagnostic and monitoring purposes
88 at a single referral hospital (Hokkaido University Veterinary Teaching Hospital) between April

89 2015 and June 2020. The CT images where the pulmonary artery of the caudal lung lobes could
90 not be traced to the periphery because of inadequate inflation, motion artifacts, or any type of
91 intra-thoracic airway abnormality (e.g., bronchiectasis or bronchial plugging) or parenchymal
92 abnormality (e.g., ground-glass opacity or consolidation) were excluded. The following data
93 were collected from the medical records: signalment, clinical signs, abnormalities detected at
94 the physical examination, purpose of CT studies, and when available, the results of
95 echocardiography. All decisions for exclusion and data collection was made by a veterinary
96 radiology specialist with 10 years of experience in veterinary radiology (G.S.).

97

98 **Data Recording and Analysis**

99 Data on computed tomographic examinations of patients were retrieved and assessed using a
100 PACS workstation DICOM viewer (OsiriX, Pixmeo, Geneva, Switzerland). All thoracic CT
101 images had been reconstructed using 0.5 to 2.0 mm reconstruction slice thicknesses, and the
102 same interslice gap as the slice thickness, as recommended by the attending veterinary
103 radiology specialist; however, the raw data were not available because of the retrospective
104 nature. All CT images were reviewed by the aforementioned veterinary radiology specialist
105 (G.S.). The diagnostic criteria for ASANL were as follows: 1) portions of the caudal lobe(s)
106 had no normal pulmonary arterial supply, 2) an anomalous systemic artery fed that region, 3)
107 bronchial connection of the region was normal, and 4) the venous return to the left atrium

108 occurred via the pulmonary vein.^{10,11} A bronchial connection was considered normal when it
109 connected the normal pulmonary parenchyma that it ventilated to the tracheobronchial tree.

110 The origin of the aberrant vessels were determined where possible.

111

112 **Statistics**

113 All statistical analyses were selected and performed by a veterinarian with a PhD
114 degree in veterinary science (G.S.), using commercial software (JMP Pro version 14, SAS
115 Institute, Inc., Cary, NC, USA). To evaluate the predisposition of ASANL in each breed, the
116 odds ratios and 95% confidence intervals (CIs) were calculated if the number of the control
117 population was at least 20 in each breed category. The control population for breed-specific
118 odds ratio calculations consisted of all dogs included in this study for each breed, with the
119 number of dogs from that breed with a diagnosis of ASANL subtracted from the total. The
120 formula used for the calculation of odds ratios was as follows: odds ratio = [(number of cases
121 for a breed)/(Number of control population for the breed)]/[(number of cases for all other
122 breeds)/(Number of control population for all other breeds)]. Logistic regression analysis was
123 used to determine the 95 % confidence interval, and to calculate P-values for assessing the
124 statistical significance of the predisposition of each breed. A significance level of $P < 0.05$ was
125 modified using a Bonferroni correction, such that $P < 0.0038$ was considered significant. The
126 differences in morbidity among males and females were also examined using the odds ratio.

127

128 **RESULTS**

129 Data on a total of 2,287 dogs who had undergone thoracic scans were found in the
130 hospital database during the study period; 337 dogs were excluded owing to the following
131 imaging findings in the caudal lobes: **evidence of atelectasis (i.e., ground glass opacity [GGO]**
132 **or consolidation associated with a decrease in lung volume often found, but not limited, to the**
133 **dependent portions of the lungs) (170), motion artifacts (84), and lung pathology (83).** Thus, a
134 total of 1,950 dogs (1,075 males and 875 females) were included in this study. The purpose of
135 the thoracic CT studies was to perform staging in neoplastic conditions (1636), to evaluate
136 thoracic diseases (60), and to screen for lung diseases in cases of abdominal diseases (158),
137 head and neck diseases (64), musculoskeletal diseases (19), and intervertebral disc disease (12).

138 All CT examinations were performed using an 80-row multidetector CT scanner
139 (Aquilion Prime, Canon Medical Systems, Tochigi, Japan). Patients were examined in ventral
140 recumbency (1,926) or dorsal recumbency (24) with the dogs anesthetized (1,907), sedated (4),
141 or conscious (39). In cases where CT studies was performed under general anesthesia using a
142 ventilator, apnea was induced during the acquisition by stopping the ventilator (acquisition at
143 the end of expiration). The scan settings included a pitch of 0.813, tube rotation time of 0.5 s,
144 scan slice thickness of 0.5 mm, tube potential of 120 kV, and tube current of 60–500 mA; these
145 were automatically calculated by a commercial software package (Sure Exposure 3D; Canon

146 Medical Systems, Tochigi, Japan). The performance of contrast studies was not standardized.
147 For contrast studies, iohexol (Omnipaque 300, GE Healthcare, Oslo, Norway) was used as a
148 contrast medium; it was administered at a dose of 600 mgI/kg via the cephalic vein, through a
149 power injector (Dual Shot GX7, Nemoto Kyorindo Co., Ltd., Tokyo, Japan). For small and
150 medium sized dogs (approximate weight <30 kg), the injection duration was set to 20 s; it was
151 set to 30 s for larger dogs. Contrast studies consisted of three phases; in cases where the
152 duration of injection was 20 s, the arterial, portal, and delayed phases were acquired at 26, 50,
153 and 180 s after the injection of the contrast medium, respectively. In cases where the duration
154 of injection was 30 s, the arterial, portal, and delayed phases were acquired 36, 60, and 180 s
155 after the injection of the contrast medium, respectively. Total body scans were performed in all
156 patients. When a detailed examination of the lung was included for the purpose of the CT
157 examination, additional thoracic scans were acquired with an airway pressure of 10 cmH₂O,
158 which has been reported to improve lung inflation.¹²

159 Among the 1,950 dogs included in this study, 48 (2.4%) dogs (33 males and 15
160 females) met the diagnostic criteria of ASANL. Among the cases with ASANL, the number of
161 males was twice that of females, but no statistically significant difference (OR=1.8, P=0.058)
162 was found. The median age at the time of the CT study was 11 years (range: 1-16 years). The
163 affected breeds and odds ratios are summarized in Table 1. Shetland Sheepdogs (7 cases,
164 OR=8.0, P<0.0001), Miniature Dachshunds (19 cases, OR=3.9, P<0.0001), and Labrador

165 Retrievers (6 cases, OR=4.5, P=0.0009) were over-represented relative to the control
166 population.

167 The purposes of the CT studies included assessment of the staging in patients with
168 neoplastic disease (n = 35), evaluation of non-neoplastic abdominal disease (10), whole body
169 screening in extremely debilitated patients (2), and evaluation of intervertebral disc disease (1).
170 Clinical signs related to cardiovascular or respiratory diseases were detected in 5 patients; 3
171 had epistaxis with nasal tumors, 1 had nasal discharge with rhinitis, and 1 had upper respiratory
172 obstruction with dynamic nasopharyngeal collapse. On physical examination, 3 cases had a
173 mild heart murmur (murmur intensity of 2/6). Echocardiography was performed in 13 cases; 2
174 had stage B1 myxomatous mitral valve disease (MMVD), 1 had stage B2 MMVD, and the
175 others had no abnormalities. Therefore, only 1 dog (an 11-year-old chihuahua) was found to
176 have left atrial enlargement.

177 In all ASANL cases, the connection of the bronchial tree was normal; however, the
178 pulmonary arteries to the caudomedial portions of the right and/or left lung lobes were lacking.
179 The aberrant vessels were running from within the mediastinum toward the bronchi in the
180 region lacking pulmonary arteries; from there, they ran caudally along the bronchi similar to
181 pulmonary arteries (Figures 1-4). The median diameter of the aberrant vessels at the insertion
182 of the mediastinum was 1.0 mm (range, 0.7-2.6 mm). Aberrant vessels were found in the right
183 caudal lobe in 24 cases, in the left caudal lobe in 21 cases, and in bilateral caudal lobes in 3

184 cases. The origin of the aberrant vessels was determined in 29/48 cases (14 cases, left gastric
185 artery; 8 cases, descending thoracic aorta; 6 cases, celiac artery; and 1 case, splenic artery)
186 (Figures 2, 4). In the remaining 19 cases, the origin of the aberrant vessels could not be
187 determined due to the lack of contrast studies or arterial phase images, the lack of images
188 reconstructed using soft-tissue reconstruction kernels, or the extremely narrow diameter of the
189 aberrant artery. There were 4 cases with 2 aberrant vessels. One case has 2 aberrant vessels
190 originating from thoracic aorta at the level of 11th and 12th thoracic vertebra, both of which
191 supplied the right caudal lobe. In another case, one aberrant vessel originating from the thoracic
192 aorta supplied the right caudal lobe; the other originating from the left gastric artery supplied
193 left caudal lobe. In the other 2 cases, the origin of the aberrant vessels could not be determined.
194 The results of the CT findings are summarized in Table 2.

195

196 **DISCUSSION**

197 This study demonstrated that pulmonary vascular malformations equivalent to
198 ASANL in humans also exist in dogs. Shetland Sheepdogs, Miniature Dachshunds, and
199 Labrador Retrievers were significantly overrepresented; however, it is necessary to note that
200 these results are based on patients that presented to a single referral veterinary institution. At
201 the time of the current study, we did not find any published reports describing the incidence of
202 ASANL in humans; however, Asian populations appear to have a higher incidence of the

203 disease (> 90%), suggesting a probable genetic etiology.^{9,10}

204 In cases where systemic arterial supply to the lung is identified in humans, the
205 differential diagnoses include hypertrophied systemic arteries associated with chronic
206 pulmonary parenchymal inflammation, the hypogenetic lung syndrome, and intra- or
207 extralobar pulmonary sequestration.^{7,11} Diagnoses are obtained based on the description of the
208 venous return and pulmonary arterial supply of the involved lung, and the absence of a normal
209 bronchial tree. CT angiography is the best suited modality, because it allows the evaluation of
210 all components.¹¹ In our cases, there was no evidence of chronic pulmonary parenchymal
211 inflammation, abnormal pulmonary venous return, or abnormal bronchial connections.
212 Furthermore, the aberrant systemic artery solely supplied the normal caudomedial portion of
213 the right or left lobes without normal pulmonary arterial supply to the involved area; these
214 findings are consistent with the diagnosis of ASANL.^{8,13}

215 The most common clinical symptom of ASANL in humans is hemoptysis; however,
216 the majority of patients are asymptomatic, and ASANL is often discovered incidentally.^{10,14}
217 Computed tomography may reveal ground glass opacity (GGO) in the affected area, which
218 indicates an area of relative hypervascularity and/or intra-alveolar hemorrhage.^{7,15} Other
219 manifestations include exertional dyspnea, heart murmur, and congestive heart failure
220 secondary to the cardiac overload produced by left-to-left shunts.^{8,16,17} In this study, none of
221 the cases had hemoptysis or GGO. A heart murmur was present in 3 of the 11 cases in which

222 the results of auscultation were derived from the medical records. However, it was assumed
223 that the cases without any records of auscultation did not have a heart murmur. In addition,
224 among the 13 cases with echocardiography, only an 11-year-old Chihuahua was found to have
225 left atrial enlargement. Since this is a breed with a high incidence of MMVD, it is unlikely that
226 ASANL solely caused the volume overload to the left atrium in this case. Since the CT studies
227 were not performed for cardiovascular or respiratory symptoms, ASANL was presumed to be
228 an incidental finding in all cases; however, further prospective studies are needed to clarify the
229 clinical significance of ASANL in dogs.

230 A possible explanation for these differences in the clinical and imaging findings
231 between humans and dogs is the differences in the diameter of the aberrant arteries. The clinical
232 symptoms and GGO in humans are caused by the volume overload to the heart due to aberrant
233 systemic arteries, and the higher pressure of the systemic circulation compared to the
234 pulmonary circulation.^{15,17} The diameter of the aberrant systemic arteries in humans has been
235 reported to be 6-30 mm.^{8,10,15} Conversely, the median diameter of the aberrant arteries in this
236 study was only 1.0 mm. One hypothesis is that the volume of aberrant arterial flow was
237 considerably low for affecting the cardiovascular system in dogs.

238 In the present study, ASANL had equal incidence in the right and left caudal lobes,
239 and the most common origin of the aberrant arteries was the left gastric artery, followed by the
240 descending thoracic aorta. These findings differ from those reported in humans, where the left

241 lower lobe is most commonly affected, and the right lower and upper and middle lobes are
242 rarely involved.⁷ The aberrant arteries most frequently originate from the descending thoracic
243 aorta, followed by the celiac axis.^{14,17} Although the cause of ASANL is unknown, it is believed
244 that it may result from the abnormal persistence of embryonic aortic postbranchial arches, that
245 supply the lung buds before development of the main pulmonary artery.^{8,18} Differences in the
246 distribution of aberrant arteries between dogs and humans may be attributed to anatomical
247 differences in the lung lobes.

248 One potential clinical significance of ASANL in dogs would be the risk of bleeding
249 during lung lobectomy of the caudal lobes. If the surgeon attempts to resect the caudal lobe
250 without being aware of the presence of ASANL, unexpected systemic arterial bleeding may
251 occur. Therefore, it is recommended to conduct CT in patients scheduled for caudal lung
252 lobectomy to check for the presence of the ASANL. In this instance, it is crucial to perform the
253 CT study while the patient is in a ventral recumbent position to prevent gravity-dependent
254 atelectasis of the caudodorsal area of the caudal lobes, and to obtain arterial phase images,
255 which cover as far as the level of the celiac artery. If the imaging is performed in dorsal
256 recumbency, atelectasis often occurs mostly in part of the periphery of the caudal lobes, even
257 if the airway pressure is positive.¹² Since the aberrant systemic artery is considerably small and
258 supplies the caudal peripheral area of the caudal lobes, even small volumes of atelectasis would
259 mask the aberrant artery, making detection difficult.

260 This study has several limitations. It included a population that presented to a single
261 referral institution; therefore, the results on breed predisposition may reflect the geographical
262 breed distribution or genetic bias. In addition, the CT and echocardiographic procedures were
263 not standardized due to the retrospective nature of the study. The origin of the aberrant systemic
264 arteries could also not be determined in 19 of 48 cases; this was partly due to the absence of
265 images of the arterial phase, or the availability of arterial phase images of only 2 mm thickness.
266 Additionally, echocardiography was not performed in all CT cases, and only 13 of 48 ASANL
267 cases underwent echocardiography. In addition, functional abnormalities could not be
268 completely excluded in dogs with ASANL in this study, even if no parenchymal abnormalities
269 were found on CT images. Therefore, the true incidence and clinical significance of ASANL
270 in dogs could not be determined. Furthermore, since no cases of ASANL underwent lobectomy
271 of the caudal lung lobes in this study, it is not clear whether ASANL actually affects surgical
272 procedures.

273 In conclusion, the incidence of ASANL in dogs is 2.4%, with an equal incidence in
274 the right and left caudal lobes. In this population from a single referral institution, analysis
275 demonstrated significantly higher incidence among Shetland Sheepdogs, Miniature
276 Dachshunds, and Labrador Retrievers. Although the clinical significance of ASANL is not clear,
277 surgeons should be aware of this finding prior to lobectomy of the caudal lung lobes, as it may
278 affect surgical planning. Therefore, when performing CT studies in patients scheduled for

279 caudal lung lobectomy, it is strongly recommended that the arterial phase is obtained in ventral
280 recumbency, including sections from the thorax to the celiac artery.

281

282 **List of Author Contributions**

283 Category 1

284 (a) Conception and design: Shimbo G

285 (b) Acquisition of data: Shimbo G

286 (c) Analysis and Interpretation of Data: Shimbo G, Takiguchi M

287 Category 2

288 (a) Drafting the Article: Shimbo G

289 (b) Revising Article for Intellectual Content: Shimbo G, Takiguchi M

290 Category 3

291 (a) Final Approval of the Completed Article: Shimbo G, Takiguchi M

292

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342

343 **Tables**

Table 1. Odds ratios of dog breeds diagnosed with anomalous systemic arterial supply to normal lung

	Dogs with ASANL (n=48)	Dogs without ASANL (n=924)	Odds Ratio	Lower 95% CI ^a	Upperr 95% CI	P value
Shetland Sheepdog*	7	39	8.0	3.4	19	<.0001
Miniature Dachshund*	19	270	3.9	2.2	7.1	<.0001
Labrador Retriever*	6	57	4.5	1.9	11	.0009
American Cocker Spaniel	3	43	2.9	0.86	9.6	.0860
Chihuahua	3	168	0.66	0.20	2.2	.493
Border Collie	1	25	1.6	0.21	12	.649
Boston Terrier	1	27	1.5	0.20	12	.677
Toy Poodle	2	126	0.6	0.14	2.5	.479

Golden Retriever	2	54	1.5	0.35	6.3	.589
Welsh corgi	1	100	0.36	0.05	2.7	.320
West Highland White Terrier	1	10	ND ^b	ND	ND	ND
Scottish Terrier	1	5	ND	ND	ND	ND
Bolognese	1	0	ND	ND	ND	ND

a: confidence interval

b: not determined

P<.0038 is considered significant

*Breeds significantly predisposed to ASANL

344

345

Table 2. Summary of the distribution, origin, and number of aberrant arteries.

		Number of cases
Affected lung lobe	Right caudal lobe	24
	Left caudal lobe	21
	Bilateral caudal lobes	3
Origin of aberrant artery	Left gastric artery	14
	Descending thoracic aorta	8
	Celiac artery	6
	Splenic artery	1
	Uncertain	19

Number of aberrant arteries	1	44
	2	4

346

347

348 **Figure legends**

349 **Figure 1**

350 The lung window (WL = -150 HU, WW = 1800 HU) (A-E) and the corresponding soft tissue
351 window (WL = 40 HU, WW = 350 HU) (F-J) for post contrast transverse images of the left
352 caudal lung lobe of left-sided ASANL in a 12-year-old spayed female Border Collie (soft-tissue
353 reconstruction algorithm, 0.5 mm slice thickness, ventral recumbency).

354 The bronchus in the caudomedial portion of the left caudal lobe lacks the pulmonary artery (B,
355 C, D: arrowheads). From that level, an aberrant vessel (C, D, E: arrows) arising from the
356 mediastinum supplies that region, as if it were a pulmonary artery.

357

358 **Figure 2**

359 Maximum intensity projections of 3D multiplanar reconstruction images (A – D: transverse, E:
360 oblique sagittal, F: oblique dorsal) of the same case as shown in Figure 1 (WL = -150 HU, WW
361 = 1800 HU, soft tissue reconstruction algorithm, 0.5 mm slice thickness). The aberrant vessel
362 (arrowheads) originates from the left gastric artery (arrow) and supplies the caudal periphery
363 of the left caudal lung lobe. The aorta (open arrow) and the celiac artery (open arrowhead) are

364 indicated.

365

366 **Figure 3**

367 The lung window (WL = -150 HU, WW = 1800 HU) (A-E) and the corresponding soft tissue
368 window (WL = 40 HU, WW = 350 HU) (F-J) for post contrast transverse images of the right
369 caudal lung lobe of right sided ASANL in a 7-year-old male Golden Retriever (soft-tissue
370 reconstruction algorithm, 0.5 mm slice thickness, ventral recumbency). As in the case of figures
371 1 and 2, the bronchus in the caudomedial portion of the right caudal lobe lacks the pulmonary
372 artery (arrowhead), and an aberrant vessel (arrows) supplies the normal lung instead of the
373 normal pulmonary artery.

374

375 **Figure 4**

376 Maximum intensity projection images of the caudal thorax of right sided ASANL in a ten-year-
377 old neutered male Shetland Sheepdog (WL = -150 HU, WW = 1800 HU, soft tissue
378 reconstruction algorithm, 0.5 mm slice thickness, ventral recumbency). The aberrant vessel
379 (arrow) originates from the descending thoracic aorta and supplies the caudal periphery of the
380 right caudal lung lobe (arrowheads).