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17 **Running Title**: Red cell distribution width in dogs with pulmonary hypertension

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- 25

26 Abstract

Objectives: To compare red cell distribution width (RDW) between dogs with different 27 causes of pulmonary hypertension (PH) and a control dog population to determine whether 28 29 RDW was correlated with severity of PH as measured by echocardiography. A further aim was to determine the prognostic significance of increased RDW for dogs with PH. 30 Animals: Forty-four client-owned dogs with PH and 79 control dogs presented to a single 31 tertiary referral institution. 32 Methods: Signalment, clinical pathological and echocardiographic data were obtained 33 34 retrospectively from the medical records of dogs with PH, and RDW measured on a Cell-Dyn 3500 was compared between dogs with pre- and post-capillary PH and a control population. 35 Referring veterinary surgeons were contacted for follow-up information and Kaplan-Meier 36 37 analysis was conducted to investigate differences in survival time between affected dogs with 38 different RDW values. **Results:** The RDW was significantly greater in dogs with pre-capillary PH compared to 39 40 control dogs. There was no difference in median survival times between dogs with PH divided according to RDW values. The RDW was positively correlated with mean 41 42 corpuscular volume and haematocrit in dogs with PH, but did not correlate with echocardiographic variables. 43 **Conclusions:** An association was found between dogs with PH and increased RDW; however 44 45 there was considerable overlap in values between control dogs and dogs with PH. The RDW was not associated with survival in this study. 46 47 48 Keywords: Angiostrongylus vasorum, canine, erythrocyte, tricuspid regurgitation, sildenafil 49 50

51 Abbreviations

cTnI	cardiac troponin I
IQR	inter-quartile range
LA:Ao	ratio of left atrial to aortic root diameter
NT-proBNP	N-terminal pro-brain natriuretic peptide
РН	pulmonary hypertension
PR	pulmonic regurgitation
PRPG	peak diastolic pulmonic regurgitant pressure gradient
RDW	red cell distribution width
ROC	receiver operator characteristic
TR	tricuspid regurgitation
TRPG	peak systolic tricuspid regurgitant pressure gradient

55 Introduction

Red cell distribution width (RDW) is a measure of degree of anisocytosis in the erythrocyte population and is expressed as the coefficient of variation of the erythrocyte size distribution data.^a The RDW is regularly reported by modern haematology analysers.¹⁻³ A number of recent publications have identified RDW as an independent predictor of outcome in a wide range of different human diseases.⁴⁻⁹ The ability of RDW to predict outcome independently in human patients with pre- and post-capillary pulmonary hypertension (PH) has been widely investigated and found to be clinically useful.¹⁰⁻¹⁵

63 Pulmonary hypertension is recognised in dogs with increased frequency due to growing access to Doppler echocardiography enabling non-invasive measurement of 64 tricuspid and pulmonic insufficiency jet velocities, which are surrogate measures of systolic 65 and diastolic pulmonary artery pressure gradients, respectively.¹⁶ Pulmonary hypertension 66 can be classified as pre-capillary or post-capillary, or a five point classification system can be 67 used based on the pathological process underlying the PH. These five categories of PH 68 69 include pulmonary arterial hypertension associated with parasite infestation or congenital systemic-to-pulmonary shunts, PH due to left-sided cardiac disease, PH related to diseases of 70 71 the pulmonary parenchyma, PH resulting from thromboembolic events involving the pulmonary vasculature, and miscellaneous causes.^{17,18} 72

A number of studies have explored the diagnostic utility of circulating biomarkers, including natriuretic peptides and cardiac troponin I (cTnI), in dogs with pre- and postcapillary PH.¹⁹⁻²² To date, the relationship between PH and RDW in dogs has been described in a single study, which suggested that RDW was increased in dogs with pre-capillary PH when compared to controls.^b There remains a paucity of information regarding the diagnostic and prognostic utility of RDW in dogs with PH due to a variety of causes.

The aims of the study were to determine if RDW differed between dogs with pre- and post-capillary PH and clinically normal dogs, to ascertain if RDW is associated with severity of PH determined by Doppler echocardiography, and to evaluate the prognostic value of a single measurement of RDW at time of presentation in dogs with PH.

84

85 Animals, materials and methods

86 Selection of cases and controls

The computerised medical record system of a tertiary referral hospital was searched to 87 88 identify dogs that had a diagnosis of PH between February 2008 and February 2012. Cases were selected if a complete medical history and physical examination were available, routine 89 90 diagnostic samples were submitted for complete haematology and serum biochemistry 91 analysis at the same visit, and a full echocardiographic examination was also performed to 92 confirm the presence of PH. Other diagnostic procedures were instituted based on the presenting signs and initial assessment. Cases were excluded if the haematocrit value was 93 94 below the lower reference limit, if the complete medical record was not available for review, or if the dog had concurrent systemic disease including cancer, diabetes mellitus, 95 portosystemic shunt or had undergone surgical procedures or blood transfusion within the 96 previous 90 days. 97

A control group of dogs was also identified, that was comprised of blood donor dogs and clinical cases treated at the same institution over the same time period. All dogs underwent complete physical examination and had blood samples submitted for complete haematology and serum biochemistry analysis either as part of clinical investigations or prior to blood donation. Blood donors were included if they had not donated blood for at least 90 days prior to submission of blood samples, if there were no abnormalities on physical examination and no reported health problems, and if blood sample results were unremarkable. 105 Blood sample results were considered to be unremarkable if the values for all parameters were within the laboratory reference intervals, or, if outside these intervals, were judged to be 106 clinically unimportant by two of the authors (DJC, AH). Clinical cases were included if their 107 108 clinical signs were not associated with systemic inflammatory or infectious diseases and if their blood sample findings were unremarkable, as described above. These cases largely 109 consisted of dogs presented for investigation of orthopaedic or ophthalmological problems. 110 111 Clinical cases were excluded if they had undergone surgical procedures or blood transfusion prior to sampling or had clinical or historical evidence of haemorrhage within the previous 90 112 113 days.

114

115 **Data collection**

116 Data relating to signalment, clinical presentation, and blood sample results were extracted from the medical records of each case with PH. Referring veterinary surgeons were contacted 117 by telephone to obtain follow-up data in January 2014 using a protocol that conformed to 118 good research practice policy at the institution. The endpoint of the study was death, 119 categorised as cardiorespiratory or non-cardiac. Cardiorespiratory causes included death or 120 euthanasia due to clinical signs of congestive heart failure, worsening breathlessness, signs 121 attributable to polycythaemia or sudden death unrelated to known systemic disease or trauma. 122 123 Non-cardiac causes were death or euthanasia following clinical signs not attributable to 124 cardiorespiratory disease (e.g. immune mediated disease, trauma, and neoplasia other than pulmonary neoplasia). The end of the study period was February 2014. 125

126

127 Red cell distribution width

128 The RDW was measured using a single haematological analyser^a previously validated for

129 canine haematology as part of a routine haematology analysis^c as previously reported.³

Erythrocyte histograms, blood smears, or blood smear reports were reviewed where available,
and individuals were excluded if histogram separation appeared poor or if platelet clumps
were identified. Blood samples were processed within 24 hours of collection at the same site
as the hospital, and serum biochemical analyses were conducted with the same analyser in all
cases.^d

135

136 Echocardiography

All patients with PH underwent complete echocardiographic examination^e and all 137 echocardiographic measurements were made off-line^f by one observer (DJC). Standard two-138 dimensional views were obtained²³ and Doppler studies were also completed using a 2.5-5.5 139 140 or 5.5-7.5 MHz transducer. Dogs were positioned in left and right lateral recumbency with 141 echocardiography performed on the dependent side. For Doppler evaluation, the right- or left-142 sided view that allowed for optimal alignment of the continuous wave Doppler interrogation beam through the regurgitant flow across the tricuspid valve and/or pulmonic valve was used 143 144 to measure instantaneous peak systolic tricuspid regurgitation (TR) or peak diastolic pulmonic regurgitation (PR) velocities. Pulmonic stenosis was excluded by confirming 145 normal valvular anatomy and mobility on two-dimensional echocardiography and identifying 146 laminar pulmonic flow profile via pulsed-wave Doppler echocardiography with peak 147 pulmonary artery flow velocities less than 1.5 m/s. Doppler flow interrogations of TR and PR 148 149 jets provided estimates of systolic and diastolic pulmonary artery pressure respectively, allowing diagnosis of PH and estimation of severity. A peak systolic TR flow velocity > 2.8150 m/s or a peak diastolic PR flow velocity ≥ 2.2 m/s was considered compatible with PH.¹⁶ 151 The modified Bernoulli equation was applied to the peak systolic TR and peak diastolic PR 152 flow velocity to estimate systolic and diastolic pulmonary artery pressure. 153

155 Statistical analysis

All statistical analyses were performed using a commercially available software package^g and 156 an alpha value of 0.05 was used throughout. Variables were assessed for normality by visual 157 inspection of histograms and using Shapiro-Wilks tests. Parametrically and non-158 parametrically distributed variables were compared using Student's t test or Mann-Whitney U 159 test, respectively. Categorical variables were compared using Fisher's exact test or Chi square 160 161 test. Data from parametric variables were presented as mean (\pm standard error), and those from non-parametric data were presented as median (\pm inter-quartile range (IQR)). Pearson 162 163 correlation coefficients were calculated to determine whether RDW was correlated with age, haematocrit, mean corpuscular volume (MCV), total white blood cell count, platelet count, 164 serum creatinine concentration, TR pressure gradient (TRPG), PR pressure gradient (PRPG) 165 166 or ratio of left atrial to aortic root diameter (LA:Ao) in dogs with PH. For the purpose of further analysis, cases with PH were divided into those with post-167 capillary hypertension (due to left-sided cardiac disease) or pre-capillary hypertension (due to 168 primary respiratory disease, idiopathic PH, or pulmonary vascular disease). Cases with PH 169 were also divided according to the severity of the PH. Dogs with a peak systolic TRPG 170 regurgitation pressure gradient of 31-50 mmHg were considered to have mild PH, dogs with 171 a gradient of 51-75 mmHg were considered to have moderate PH and dogs with a gradient 172 greater than 76 mmHg were considered to have severe PH.²⁰ 173 174 To determine whether RDW was a useful predictor of moderate or severe PH (TRPG > 50 mmHg), a receiver operator characteristic (ROC) curve was constructed using RDW 175 values. The area under the curve was determined, and a suitable cut off value was chosen to 176

177 estimate sensitivity and specificity values.

Dogs with PH were divided into balanced terciles according to RDW values (lowest
to 15.7, 15.8 to 17.3 and 17.4 to highest). Average survival times were compared between

these groups using Kaplan-Meier product limit estimates and log-rank test. This analysis was also conducted separately for dogs with known cardiorespiratory death. Dogs that were still alive at the study end point were censored. Similar survival analyses were also conducted to compare dogs with pre- and post-capillary PH and according to the severity of PH.

184

185 **Results**

186 **Study populations**

187 Forty-four dogs were diagnosed with PH during the study period. The median age of affected

dogs was 9.0 years (IQR, 4.3-11.4). The group was comprised of 18 intact males, 9 neutered

males, 4 intact females and 13 neutered females. Twenty-five different breeds were

190 represented, and the 5 most common were Jack Russell terrier (n=5), cross-breed (n=4),

191 Border Collie (n=3), West Highland White terrier (n=3), and Yorkshire terrier (n=3).

The control group was composed of 79 dogs, of which 42 were blood donors and 37were dogs presented for problem investigation. The median age of this group was 2.9 years

194 (IQR, 2.0–6.0), which was significantly lower than the median age of the dogs with PH (P <

195 0.001). This group was composed of 18 intact males, 30 neutered males, 7 intact females and

196 24 neutered females. Twenty-six different breeds were represented, and the most common

197 were cross-breeds (n=17), Labrador retrievers (n=13), Golden retrievers (n=7), German

198 Shepherd dogs (n=6), and English springer spaniels (n=4).

199

200 Pulmonary hypertension

All of the dogs diagnosed with PH had systolic TR flow velocities > 2.8 m/s, and PR flow

velocities were > 2.2 m/s in 24/44 dogs in which it was present. The median TRPG in dogs

with PH was 71.0 mmHg (IQR, 51.1-98.7; range, 37.5-238.8). Eleven dogs had mild PH, 14

204	had moderate PH, and 19 had severe PH. Among the 24 dogs for which measurements were
205	available, the median PRPG was 38.1 mmHg (IQR, 33.8-49.5; range, 14.4-63.4).
206	Twenty-eight dogs had pre-capillary PH caused by Angiostrongylus vasorum
207	infestation (n=7), primary PH with no detected underlying cause (n=7), idiopathic pulmonary
208	fibrosis (n=5), suspected or confirmed pulmonary thromboembolism (n=3), other
209	parenchymal pulmonary disease (n=2), systemic-to-pulmonary shunt (n=4) including 2 dogs
210	with right-to-left shunting patent ductus arteriosus. Sixteen dogs had post-capillary PH,
211	caused by degenerative mitral valve disease (n=12), dilated cardiomyopathy (n=3), and aortic
212	stenosis with systolic failure (n=1). The LA/Ao was significantly greater in dogs with post-
213	capillary PH compared to dogs with pre-capillary PH, and TRPG, PRPG and total white
214	blood cell count were all significantly greater in dogs with pre-capillary PH (Table 1).
215	Once the diagnosis was established, dogs with PH were treated with one or more of
216	the following drugs at the discretion of the attending clinician: angiotensin converting
217	enzyme inhibitor (n=44), pimobendan (n=24), furosemide (n=16), fenbendazole (n=10),
218	sildenafil (n=7), spironolactone (n=7), and atenolol (n=1).
219	
220	Red cell distribution width
221	The median RDW in control dogs was 15.7% (IQR, 15.0-16.4; range, 13.5-18.3), which was
222	significantly lower than in dogs with pre-capillary PH (16.3%; IQR, 15.6-17.9; range, 12.7-
223	19.6; <i>P</i> = 0.008) but not post-capillary PH (16.9; IQR, 15.0-17.6; range, 14.5-21.1; <i>P</i> =
224	0.063). There was no difference in RDW between dogs with pre- and post-capillary PH ($P =$
225	0.714). Distributions of RDW in each group are shown in Figure 1.
226	There was no difference in RDW between dogs with mild, moderate or severe PH
227	(Fig. 2, $P = 0.384$). The area under the ROC curve constructed using RDW values for

prediction of a TRPG value greater than 50 mmHg was 0.612 (95% confidence interval,

0.429–0.794) (Fig. 3). Using a cut-off value of 15.9%, RDW had a sensitivity of 60.6% and
specificity of 45.5% for differentiation of mild from moderate or severe PH. The RDW was

significantly correlated with mean corpuscular volume (P < 0.001) and haematocrit (P =

232 0.045) in dogs with PH but was not associated with LA:Ao, TRPG, PRPG, total white blood

- cell count, platelet count or serum creatinine concentration (Table 2).
- 234

235 Survival times

There was no significant difference in median survival times between dogs with PH divided into terciles according to RDW values (P = 0.071), though there was a trend for greater mortality with higher RDW values (Fig. 4A). Survival curves for dogs with PH stratified according to cause of PH and severity of peak systolic TR are shown in Figures 4B and 4C, respectively. There was no difference in survival between these groups (P = 0.948 and P = 0.622, respectively).

242

243 Discussion

This study demonstrates that RDW differs significantly between dogs with pre-capillary PH and control dogs, but not between control dogs and those with post-capillary PH. There was no difference in survival between dogs grouped according to RDW values, and RDW was not a useful predictor of the severity of PH. The RDW was correlated with MCV and haematocrit in dogs with PH.

The findings presented here are in agreement with retrospective and prospective studies in human patients, which indicate that RDW can provide additional diagnostic information in patients with PH.²⁴ In a recent study in dogs, RDW was significantly increased with pre-capillary PH compared to controls, but not in dogs with post-capillary PH,^b similar to the findings of this study. There was no difference in RDW between healthy dogs and those with either compensated or decompensated heart failure due to degenerative mitral
valve disease.²⁵

The reasons for alteration in RDW in patients with PH remain unclear. Because RDW 256 is an indicator of anisocytosis, it varies with iron deficiency anaemia, and iron status appears 257 to be essential in the regulation of pulmonary vascular tone.^{26,27} In human patients, iron 258 deficiency has been identified in idiopathic PH.^{28,29} In heart failure patients with post-259 capillary PH, RDW is thought to be affected by chronic inflammation, renal dysfunction, 260 altered erythropoiesis, and oxidative and nutritional stress.^{9,14,30} It is therefore likely that 261 262 RDW is influenced by numerous factors involved in the pathogenesis of PH. Angiostrongylus vasorum infestation is known to cause variable effects on erythroid 263 parameters,^{31,32} though, to the authors' knowledge, RDW values have not been assessed in 264

265 infected dogs.

Significant correlations between RDW and serum urea and creatinine concentration 266 were previously reported in dogs with PH.^b No such correlation was detected in the present 267 study, but the relationship between RDW and serum urea and creatinine concentration in 268 human patients with PH is contradictory, with one study showing a significant correlation 269 with serum urea and a strong trend towards significance with serum creatinine¹⁵ whereas in 270 other studies no such correlation was identified.^{13,14} Interestingly, in contrast to studies in 271 human patients^{10,12} but in line with the canine study by Poser,^b we found no significant 272 correlation between RDW and peak TR or PR jet velocity, which may reflect differences in 273 sample size between canine studies and the larger investigations in human patients. To date, 274 the only circulating biomarker shown to be associated with peak TR gradient in dogs with PH 275 is N-terminal pro-brain natriuretic peptide (NT-proBNP).²⁰ The RDW was found to be 276 significantly correlated with haematocrit and mean corpuscular volume in dogs with PH, as 277 was also found in dogs with degenerative valve disease.²⁵ 278

279 The RDW has been shown to predict outcome in human patients with pre- and postcapillary PH in numerous studies.¹⁰⁻¹⁵ In the present study however, RDW did not provide 280 clinically useful prognostic information in dogs with PH. This result may reflect the small 281 282 sample size of affected dogs, particularly as there was a trend towards increased mortality with higher RDW values as observed in people,¹⁵ or the influence of treatments such as 283 furosemide, sildenafil and pimobendan instituted following diagnosis of PH. Studies in 284 humans indicate incremental increases in serial RDW measurements are associated with 285 decreased survival in heart failure^{33,34} therefore serial RDW measurements in dogs may be an 286 287 area of future study.

A number of publications have explored the diagnostic utility of circulating 288 biomarkers including natriuretic peptides and cardiac troponin I (cTnI) in dogs with pre- and 289 post-capillary PH.¹⁹⁻²² One report indicated that measurement of circulating NT-proBNP but 290 not N-terminal pro-atrial natriuretic peptide (NT-proANP) or cTnI was able to stratify dogs 291 with pre-capillary PH into mild, moderate and severe.²⁰ In an experimental canine model of 292 pre-capillary PH, circulating NT-proANP and NT-proBNP could identify dogs with moderate 293 and severe but not mild PH.¹⁹ In the present study, a trend was identified with increasing 294 RDW values in dogs with moderate and severe PH, but this did not reach significance which 295 may reflect the relatively small number of cases in each group. 296

In a study of 162 human patients of which 62% had pulmonary arterial hypertension, RDW was found to be independently associated with death in patients with severe PH and performed better as a prognostic indicator than NT-proBNP.¹⁵ The RDW was also shown to add significant prognostic value to measurements of NT-proBNP in human patients with idiopathic pulmonary hypertension.¹⁴ Unfortunately, in the retrospective report presented here, NT-proBNP data was not available and it was not possible to compare the ability of the two biomarkers to identify dogs with PH and predict survival. 304 The present study is the first to indicate that RDW, a simple parameter routinely measured as part of a haematological profile, may be useful in the diagnosis of dogs with PH 305 with a number of different aetiologies. The sensitivity and specificity of RDW in the 306 307 diagnosis of PH are based on the performance of the parameter in the whole study sample. It is clear for our results that some overlap in RDW values occurs between the control 308 population and dogs with pre-capillary PH, which suggests that RDW may not be a useful 309 parameter for determining the disease status of single individuals. The performance of a 310 screening test depends not only on its intrinsic ability to identify affected individuals but also 311 312 on the prevalence of disease within the population. In this retrospective study, populations were preselected on the basis of the presence of confirmed PH or control dogs representing 313 314 groups with 100% or 0% disease prevalence, respectively. Under these circumstances, a 315 screening test is likely to exhibit superior performance than would the same test used in a 316 more heterogeneous population encountered in the clinic. Furthermore, RDW may be increased due to other pathological processes, so this indicator may also lack specificity in 317 clinical settings. Further studies will be required to determine whether RDW values may 318 contribute useful prognostic information, as they do in humans. 319

320 This study has a number of limitations as a result of its retrospective nature. Due to the strict inclusion criteria, the study population of affected animals was small which reduces 321 statistical power. The study was not designed to assess the influence of different treatments 322 323 for dog with pre- and post-capillary PH on survival or RDW. Affected dogs with PH were significantly older than the control population. An age related increase in red blood cell 324 membrane stability and RDW has been reported in women³⁵ but no information is available 325 for dogs. The RDW was assessed as a single prognostic factor in the present study as the 326 small sample size and low event (mortality) rate within the study period precluded more 327 sophisticated analysis, such as Cox proportional hazards regression. Future studies of a larger 328

329 size will be required to establish whether the prognostic value of this parameter is altered after other variables have been accounted for. The retrospective nature of the study meant 330 that patient and control RDW values were obtained from a Cell-Dyn 3500, although the 331 majority of the human literature refers to more recent haematology analyser models. 332 Prospective studies using newer generation analysers may provide better discrimination for 333 RDW value prognostication. 334 The results of this study show that RDW differed significantly between dogs with pre-335 capillary PH and control dogs, but RDW was not a significant prognostic marker in dogs with 336 PH. Further studies will be required to determine whether RDW has prognostic significance 337 in larger groups of dogs with PH, or in dogs with other forms of cardiorespiratory disease. 338 339

341 Footnotes

- ^a Cell-Dyn 3500 System Operators Manual, Abbott Laboratories, Abbott Park, Illinois, USA..
- ^b Poser H, Mazzotta E, Menciotti G, Contiero B, Baron Toaldo M, Guglielmini C. Red blood
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- ^c RVC Diagnostic Laboratories, Royal Veterinary College, London, UK.
- ^d ILab600 Instrumentation Laboratory, Werfen Life Group, Barcelona, Spain.
- ^eGE Vivid 7, GE Healthcare, Hatfield, UK
- ^fGE EchoPAC PC, GE Healthcare, Hatfield, UK.
- ^g IBM SPSS Statistics for Windows, Version 20.0, Released 2011, IBM Corp. Armonk, NY,

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- **353 Conflict of Interest:** The authors declare no conflict of interest.
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- 355

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Figure Legends 477

- Figure 1: Red cell distribution width values in control dogs and dogs with pre-capillary or 478
- post-capillary pulmonary hypertension. Lines represent the median for each group. 479

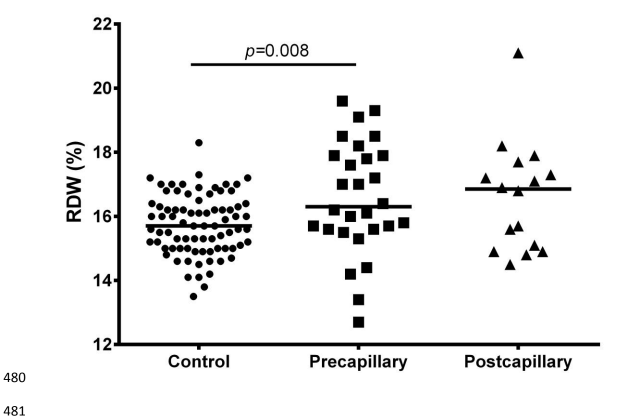
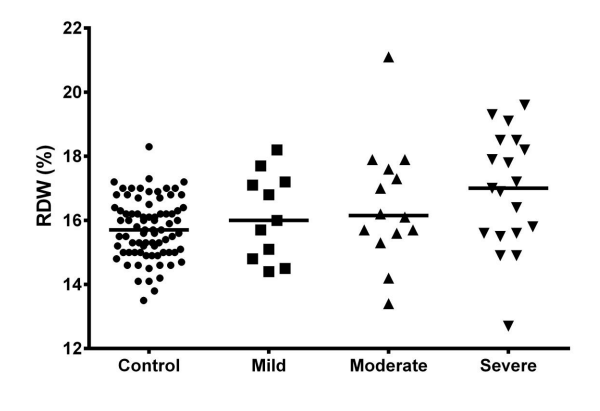
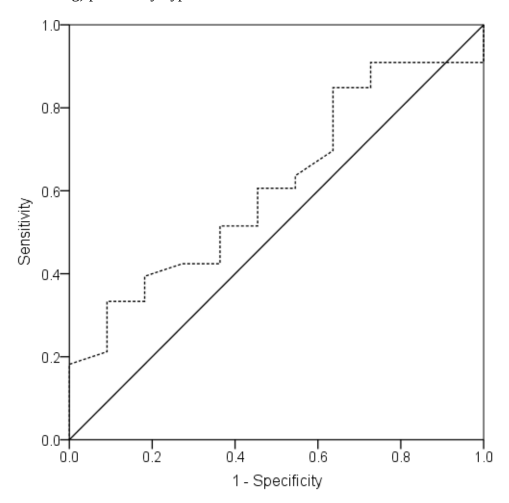


Figure 2: Red cell distribution width values in control dogs and dogs with mild, moderate
and severe pulmonary hypertension defined by peak systolic tricuspid regurgitant pressure
gradient. Lines represent the median for each group.



487 **Figure 3:** Receiver operator characteristic curve constructed to determine whether red cell 488 distribution width values can predict mild (peak systolic tricuspid regurgitant pressure 489 gradient < 50 mmHg) or moderate to severe (peak systolic tricuspid regurgitant pressure 490 gradient \geq 50 mmHg) pulmonary hypertension.



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Figure 4: Kaplan-Meier survival curve in dogs with (A) pulmonary hypertension divided into
terciles according to red cell distribution width values: lowest value to 15.7 (solid line), 15.8
to17.3 (dotted line), 17.4 to highest (dashed line), (B) pre-capillary pulmonary hypertension
(dotted line) or post-capillary pulmonary hypertension (solid line), and (C) mild (solid line),
moderate (dotted line) and severe pulmonary hypertension (dashed line). Tick marks indicate
censored cases.

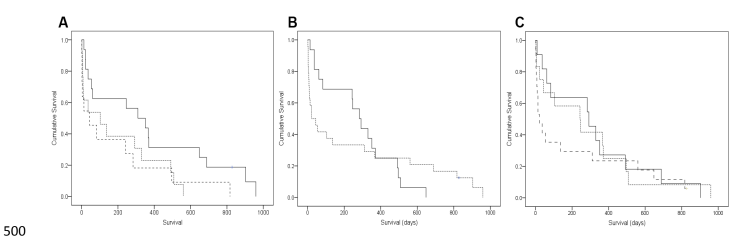


Table 1: Echocardiographic, haematological and biochemical variables in dogs with pre-502

capillary and post-capillary pulmonary hypertension and control dogs. P values represent 503

comparison between dogs with pre-capillary and post-capillary pulmonary hypertension. 504

Parameter	Pre-capillary PH ^a	Post-capillary	<i>P</i> value	Control
	(median, IQR ^b)	PH ^a (median,		
		IQR ^b)		
LA:Ao ^c ratio	1.33 (1.13-1.55)	2.21 (1.16-1.54)	< 0.001	
TRPG ^d (mmHg)	80.4 (56.3-108.8)	52.5 (43.2-70.8)	0.001	
PRPG ^e (mmHg)	40.1 (34.9-52.1)	33.9 (22.3-38.5)	0.040	
RDW ^f (%)	16.3 (15.6-17.9)	16.9 (15.0-17.6)	0.714	15.6 (15.0-16.2)
Haematocrit (%)	43.7 (39.0–49.5)	47.6 (43.7–50.9)	0.205	46.8 (44.5-50.0)
MCV ^g (µl)	69.1 (66.8–71.1)	70.7 (68.7–71.7)	0.317	71.5 (70.1-73.8)
WBCC ^h (x10 ⁹ /l)	16.2 (11.2-19.8)	10.6 (8.3-12.8)	0.010	9.3 (8.2-10.4)
Platelet count	302 (187-377)	290 (227-519)	0.317	229 (193-260)
(x10 ⁹ /l)				
Serum creatinine	93.0 (81.0-120.5)	108.5 (90.8-	0.432	104.0 (95.0-
concentration		120.0)		119.0)
(µmol/l)				

PH, pulmonary hypertension; IQR, inter-quartile range; LA/Ao, ratio of left atrial to aortic 505

506 root diameter; TRPG, peak systolic tricuspid regurgitation pressure gradient; PRPG, peak

diastolic pulmonic regurgitation pressure gradient; RDW, red cell distribution width; MCV, 507

mean corpuscular volume; WBCC, total white blood cell count. 508

510 Table 2: Results of correlations between red cell distribution width and clinical pathological

511 variables in dogs with pulmonary hypertension.

Variable	Correlation coefficient	P value
Age (years)	-0.249	0.103
LA:Ao ^a	0.054	0.777
TRPG ^b	0.113	0.465
PRPG ^c	-0.020	0.927
Haematocrit	0.304	0.045
MCV ^d	-0.664	<0.001
WBCC ^e	-0.066	0.668
Platelet count	-0.268	0.079
Serum creatinine concentration	0.147	0.385

512 LA/Ao, ratio of left atrial to aortic root diameter; TRPG, peak systolic tricuspid regurgitation

513 pressure gradient; PRPG, peak diastolic pulmonic regurgitation pressure gradient; RDW, red

cell distribution width; MCV, mean corpuscular volume; WBCC, total white blood cell

515 count.