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# Longitudinal Analysis of Quality of Life, Clinical, Radiographic, Echocardiographic, and Laboratory Variables in Dogs with Preclinical Myxomatous Mitral Valve Disease Receiving Pimobendan or Placebo: The EPIC Study

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**Background:** Changes in clinical variables associated with the administration of pimobendan to dogs with preclinical myxomatous mitral valve disease (MMVD) and cardiomegaly have not been described.

**Objectives:** To investigate the effect of pimobendan on clinical variables and the relationship between a change in heart size and the time to congestive heart failure (CHF) or cardiac-related death (CRD) in dogs with MMVD and cardiomegaly. To determine whether pimobendan-treated dogs differ from dogs receiving placebo at onset of CHF.

Animals: Three hundred and fifty-four dogs with MMVD and cardiomegaly.

Materials and Methods: Prospective, blinded study with dogs randomized (ratio 1:1) to pimobendan (0.4–0.6 mg/kg/d) or placebo. Clinical, laboratory, and heart-size variables in both groups were measured and compared at different time points (day 35 and onset of CHF) and over the study duration. Relationships between short-term changes in echocardiographic variables and time to CHF or CRD were explored.

**Results:** At day 35, heart size had reduced in the pimobendan group: median change in ( $\Delta$ ) LVIDDN -0.06 (IQR: -0.15 to +0.02), P < 0.0001, and LA:Ao -0.08 (IQR: -0.23 to +0.03), P < 0.0001. Reduction in heart size was associated with increased time to CHF or CRD. Hazard ratio for a 0.1 increase in  $\Delta$ LVIDDN was 1.26, P = 0.0003. Hazard ratio for a 0.1 increase in  $\Delta$ LA:Ao was 1.14, P = 0.0002. At onset of CHF, groups were similar.

Conclusions and Clinical Importance: Pimobendan treatment reduces heart size. Reduced heart size is associated with improved outcome. At the onset of CHF, dogs treated with pimobendan were indistinguishable from those receiving placebo. Key words: Cardiology; Cardiovascular; Echocardiography; Endocardiosis; Heart Failure; Mitral regurgitation.

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The "Evaluation of Pimobendan In dogs with Cardiomegaly caused by preclinical mitral valve disease" (EPIC) study¹ was a multicenter, blinded, randomized, placebo-controlled clinical trial evaluating the effect of pimobendan in delaying the onset of clinical signs in dogs with cardiac enlargement secondary to myxomatous mitral valve disease (MMVD). The main findings have been published and demonstrated a significant benefit of treatment in prolonging the time to the primary endpoint, which was a composite of the onset of congestive heart failure (CHF) or cardiac-related death.¹ Although the beneficial effects of the treatment were clear, none of the longitudinal effects of the drug on clinical, radiographic, and echocardiographic variables have yet been reported in dogs with preclinical MMVD.

Dogs with MMVD go through various stages of the disease.<sup>2</sup> In the first stage of the disease (stage B1), dogs have insufficiency of their mitral valve, but there is no evidence of cardiac enlargement. In those dogs in which the disease progresses, heart size enlarges as a consequence of the left-sided volume overload. Dogs with cardiac enlargement that have not yet developed signs of CHF are considered to be in stage B2. Signs of CHF develop in some dogs, typically after a period during which the heart undergoes progressively more rapid enlargement.<sup>3–5</sup>

Dogs with increased heart size and with the most rapid rate of change in heart size are at the greatest risk of development of CHF. 4.5 Increased heart size in dogs at all stages is predictive of an increased risk of cardiac-related death. 6-8

Previous studies of dogs with cardiomegaly secondary to heart disease have demonstrated a reduction in heart size associated with the administration of pimobendan. Polar In Doberman pinschers with preclinical dilated cardiomyopathy (DCM), the degree to which the heart size reduced in response to the administration of pimobendan was predictive of a decreased likelihood of experiencing the primary endpoint of the study, which was a composite of CHF or cardiac-related death.

Longitudinal effects of pimobendan treatment have been described in dogs with stage C MMVD<sup>9</sup> but have not been reported in dogs with stage B disease. Some dogs show subtle signs associated with their cardiac disease before overt signs of CHF develop. Such signs might include cough attributable to cardiomegaly, <sup>13</sup> intolerance of exercise, increasing resting respiratory rate, and weight loss. No previous prospective study has described the change in clinical signs and other variables in a large cohort of dogs as they progress from stage B2 to stage C MMVD. Of the dogs enrolled in the EPIC study, 135 were confirmed to have developed CHF, <sup>1</sup> and this population allows us to characterize the changes that occur as dogs progress from a preclinical stage of the disease, stage B2, into CHF.

The aims of this study were to investigate, in a population of dogs with stage B2 MMVD, the influence of treatment with pimobendan on clinical, radiographic, and echocardiographic variables in both the short and long term, and whether short-term changes in

#### Abbreviations:

Abbreviatio	ns:
2D	2-dimensional
ALT	alanine aminotransferase
AUC	area under the curve
BCS	body condition score
BPM	beats per minute
CHF	congestive heart failure
CI	confidence intervals
CKCS	Cavalier King Charles Spaniels
CRD	cardiac-related death
ECG	electrocardiogram
EPIC	Evaluation of Pimobendan in dogs with Cardiomegaly
	Caused by Preclinical Mitral Valve Disease
F	female
FS%	fractional shortening
FS	female spayed
GPT	glutamic-pyruvate transaminase
HR	hazard ratio
IQR	interquartile range
K +	potassium
LA/Ao	left atrial-to-aortic ratio
LVIDd	left ventricular internal diameter in diastole
LVIDDN	normalized left ventricular internal diameter in diastole
LVIDs	left ventricular internal diameter in systole
LVIDSN	normalized left ventricular internal diameter in systole
MC	male castrated
M	male
MMVD	myxomatous mitral valve disease
MR	mitral regurgitation
Na+	sodium
NA	not able to calculate
PCV	packed cell volume
PP	per protocol
SAP	systolic arterial blood pressure
TPC	total protein concentration
UK	United Kingdom
USA	United States of America

echocardiographic variables might predict the time to the onset of CHF or cardiac-related death. Additionally, we aimed to describe a large group of dogs as they progress from stage B2 to stage C MMVD and to determine whether dogs that develop CHF while receiving pimobendan differ from those that develop CHF while receiving placebo.

vertebral heart sum

#### **Materials and Methods**

# Trial Design

The EPIC trial was a prospective multicenter, blinded, randomized, placebo-controlled study. Complete and detailed description of the study has been published. 1

#### Dogs

#### Enrollment Criteria

VHS

Dogs were eligible for participation in the study provided that the owner had given informed consent.

To be eligible for inclusion, a dog had to be 6 years of age or older, have a body weight  $\geq$ 4.1 and  $\leq$ 15 kg, have a characteristic systolic heart murmur of moderate-to-high intensity ( $\geq$ grade 3/6) with maximal intensity over the mitral area, have echocardiographic evidence of advanced MMVD defined as characteristic valvular lesions of the mitral valve apparatus, mitral regurgitation (MR) on the color Doppler echocardiogram, and have echocardiographic and radiographic evidence of cardiomegaly defined as a left atrial-to-aortic root ratio (LA/Ao)  $\geq$ 1.6 measured in a short-axis view, <sup>14</sup> body weight normalized left ventricular internal diameter in diastole (LVIDDN)<sup>15</sup>  $\geq$ 1.7, and a vertebral heart sum (VHS) >10.5. <sup>16</sup>

#### **Exclusion Criteria**

Dogs were excluded from the study if they had any of the following: known clinically important systemic or other organ-related disease that was expected to limit the dog's life expectancy or required chronic administration of cardiovascular medication precluded as part of the trial. Dogs with hypothyroidism could be included provided the investigator deemed them clinically stable on treatment. Dogs with current or previous evidence of cardiogenic pulmonary edema, pulmonary venous congestion or both, cardiac disease other than MMVD, clinically significant supraventricular, ventricular tachyarrhythmias or both (ie, requiring antiarrhythmic treatment), or evidence of pulmonary hypertension considered to be clinically relevant (RV:RA pressure gradient >65 mmHg) were excluded. Dogs with a history of chronic or recent administration (>14 days of duration or within 30 days of intended enrollment) of any precluded medication were excluded. Dogs that were pregnant or lactating were not eligible for enrollment. In the event that before study enrollment, if a dog had received short-term treatment (<14 days) with a precluded agent, but was no longer receiving treatment and had not received it within 30 days of intended enrollment, then the dog was eligible

Details of study sites, randomization and blinding, concomitant treatment, and data management have previously been described.<sup>1</sup>

# Trial Medication

Pimobendan verum<sup>a</sup> was administered PO at a target dose of 0.4–0.6 mg/kg/d as per registered label instructions in most countries where the product was licensed. Placebo was administered in a similar manner. Placebo and verum tablets and packaging containers were visually identical.

## Schedule of Events

Before inclusion, a case history was taken for each dog. The dog then underwent a physical examination, echocardiography, thoracic radiography, and routine hematology and blood biochemistry (performed at laboratories local to each investigator) with a minimum database consisting of packed cell volume (PCV), alanine aminotransferase (ALT), glutamic pyruvate transaminase (GPT), total protein concentration (TPC), creatinine, potassium (K+), and sodium (Na+) concentrations. Re-examinations were scheduled at day 35 and approximately 4 months after inclusion. Thereafter, the dogs were scheduled for re-examination every 4 months. Details of examinations that were undertaken on each visit are provided in Table 1.

## Clinical Evaluation

At inclusion, dog characteristics such as breed, age, sex, and neutering status were noted. The body weight, body condition score (BCS), rectal temperature, and respiratory rate were measured at each visit.

# ${\it Echocardiography}$

Echocardiography was performed on unsedated dogs. The following measurements were each taken over at least 3 cardiac cycles, and the mean was recorded as follows: the LA/Ao obtained from the right parasternal short-axis 2D view as previously described, <sup>14</sup> the left ventricular internal diameter at end-diastole (LVIDd), and left ventricular internal diameter at end-systole

(LVIDs) measured on the M-mode echocardiogram, obtained from the right parasternal short-axis view. \(^{17}\) M-mode values were used to derive the fractional shortening (FS%). Normalized dimensions \(^{15}\) were calculated according to the following formulae: normalized LVIDd (LVIDDN) = LVIDd(cm)/(BW (kg))^{0.294}; normalized LVIDs (LVIDSN) = LVIDs(cm)/(BW(kg))^{0.315}.

#### Thoracic Radiography

Thoracic radiography was performed at inclusion, 8 months after inclusion and every 8 months thereafter for as long as the dog remained in the per-protocol (PP) population. It was also performed at the time a dog was considered to have developed signs of CHF. Right lateral and dorsoventral projections were used to evaluate the thorax. Cardiac size was assessed by the VHS method, and pulmonary edema and congestion were recorded, when considered to be present, by the attending cardiologist.

#### Primary Endpoint

The primary endpoint was a composite of the development of left-sided CHF verified by an endpoint committee, euthanasia for a cardiac reason, or death presumed to be cardiac in origin. The primary outcome variable of the study was the time period from inclusion until the primary endpoint was reached.

Visit data for the day 35 visit were included in the analyses provided the dog was re-examined between 28 and 42 days (day  $35\pm7$  days) after initiation of treatment. Visit data for subsequent visits were included in between-group comparisons for each visit providing the visit took place within 30 days of the scheduled date of re-examination.

Quality of life variables were assessed at baseline by an ordinal scoring system (Table S1). After baseline assessment of quality of life variables, at each subsequent visit, owners were asked to score the following characteristics on a 5-point scale (Table 2): appetite, demeanor, exercise tolerance, fainting, respiratory effort, cough, and nocturnal dyspnea/coughing.

Owners were also asked to measure resting respiratory rate before every re-examination, and the average resting respiratory rate during that period was noted. Instruction was provided to owners who were advised to count the respiratory rate over one minute within the week before each re-examination. Ideally, this was performed on several days, and the average of the measurements obtained was recorded as a single value at the corresponding visit.

#### Statistical Methods

Any dog in the study population that was confirmed to have met all inclusion criteria (and none of the exclusion criteria) was included in the PP population until one of the following occurred: The dog reached the primary endpoint, the dog was censored from the primary endpoint analysis due to the occurrence of an event that precluded continuation in that population, or the end of the study was reached. All analyses described herein were conducted on dogs that remained in the PP population.

For continuous variables that were measured at both baseline and day 35, a "change from baseline" variable was created by subtracting the value obtained at baseline from the day 35 value.

For every dog for which any of the clinical or radiographic continuous variables (ie, body weight, rectal temperature, respiratory rate, owner-measured resting respiratory rate, heart rate, and VHS) were measured on at least one additional visit after baseline assessment (2 occasions after baseline for owner-measured respiratory rate), a curve was constructed by plotting each data point on a graph with the unit of measurement on the vertical axis and time on the horizontal axis. The area under the curve (AUC) averaged for the number of days between the first and last observation was calculated, thus giving an average value for that variable that incorporated all of the observations made and was independent of the duration of time the dog remained in the

**Table 1.** Schedule of procedures undergone by animals remaining in the per-protocol population of the study at different examinations.

	Day 0	Day 35 (±7 days)	4 months <sup>a</sup> (±30 days)	8 months <sup>b</sup>	Event
Questionnaire	X	X	X	X	X
Physical examination	X	X	X	X	X
Thoracic radiograph	X			X	X
Echocardiography	X	X			X
Hematology and clinical chemistry	X	X			X

<sup>&</sup>lt;sup>a</sup>4 months from day 0 and every 8 months thereafter.

**Table 2.** Ordinal scoring system used for quality of life observations; appetite, demeanor, exercise tolerance, fainting, respiratory effort, cough, and nocturnal dyspnea/coughing.

Owner's Perception of Quality of Life	Ordinal Score Ascribed	Summary Category for Between-Group Comparisons
Greatly improved	1	Improved
Improved	2	Score = +1
Unchanged	3	Unchanged
		Score = 0
Deteriorated	4	Deteriorated
Severely deteriorated	5	Score = $-1$

study. 18 These summary values were then compared between treatment groups.

For comparison of quality of life characteristics between groups at specific visits, dogs in each group were classified on an ordinal scale to be improved (+1), unchanged (0), or deteriorated (-1) (Table 2).

For ordinal variables measured on the same scale at baseline and all subsequent visits (ie, BCS, heart murmur intensity, and heart disease class), dogs were classified as having increased (+1), remained unchanged (0), or decreased (-1).

Between-group comparisons of the change in ordinal variables were undertaken at each scheduled visit up to the visit at which fewer than 50% of dogs remained in each group.

For all continuous and ordinal variables, between-group comparisons were made by Mann-Whitney tests. For within-group paired comparisons of all ordinal and continuous variables, the Wilcoxon signed ranks test was used.

The results of multivariable Cox proportional hazards analysis analyzing the association between treatment and clinical variables measured at baseline and the time to the primary endpoint have been described in this population. In this longitudinal analysis, the effects of a change in echocardiographic variables on time to the primary endpoint were explored by multivariable Cox proportional hazards analysis. Separate multivariable models were constructed for each of 4 echocardiographic variables; LVIDDN, LVIDSN, LA/Ao, and FS%. In each model, the baseline value of the variable, the change from baseline at day 35, and the treatment effect were entered as explanatory variables. Time to the primary endpoint was the dependent variable. For each model, all dogs in the PP population for which paired observations were available to derive a value for the change from baseline at day 35 were entered into each analysis. Dogs leaving the PP population for reasons other than reaching the primary endpoint were right-censored at the time of leaving the population. Variables entered into each

multivariable model were assessed for collinearity with other variables in the same model and to ensure that the proportional hazards and linearity assumptions were met.

The threshold of significance was defined as P < 0.05. All analyses were 2-tailed. Statistical analyses were performed with a commercially available software program.<sup>b</sup>

#### Results

A flow diagram outlining the number of dogs in the PP population in each group at different time points and the number of dogs reaching the primary endpoint is illustrated in Figure 1.

Relevant baseline characteristics of the 2 treatment groups are summarized in Table 3.

# Comparisons at Day 35

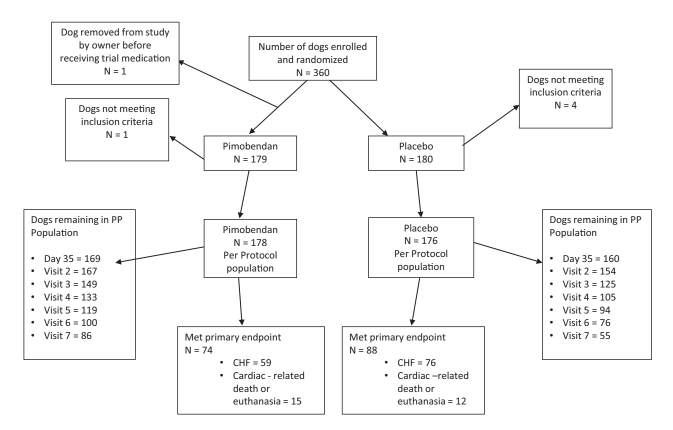
Between-group comparisons of quality of life variables as perceived by owners at day 35 showed that the pimobendan and placebo groups were not significantly different in the extent to which any of the quality of life variables had changed (Table 4). Within-group comparisons in the pimobendan group showed the following quality of life scores to have significantly improved: appetite, demeanor, exercise tolerance, and cough (Table 4). The scores for owner perceptions of demeanor and exercise tolerance were also significantly improved in the placebo group (Table 4).

For the physical examination, diagnostic imaging, and laboratory variables (Table 5), in the pimobendan group, there were significant decreases in LVIDDN, LA/Ao, LVIDSN, plasma Na<sup>+</sup> concentration, and creatinine. There was a significant increase in FS% in the pimobendan group. In the placebo group, the only significant change was a decrease in LA/Ao. Betweengroup comparisons of the changes showed a significantly greater change in FS% in the pimobendan group, and greater reductions in LVIDDN, LA/Ao, and LVIDSN in the pimobendan group (Fig 2a–d).

# Prognostic Value of Change in LVIDDN, LA/Ao, LVIDSN, and FS% Between Baseline and Day 35 (Multivariable Cox Proportional Hazard Analysis)

Changes in echocardiographic variables between baseline and day 35 were significantly predictive of outcome

<sup>&</sup>lt;sup>b</sup>8 months from day 0 and every 8 months thereafter.



**Fig. 1.** A flowchart illustrating the randomization of the 360 dogs in the study, the numbers of dogs reaching the different components of the primary endpoint in each treatment group, and the number of dogs in each treatment group remaining in the per-protocol population at different scheduled visits. **PP**, per protocol; CHF, congestive heart failure.

in all multivariable models, except for change in FS% (Table 6). The hazard ratio (HR) for the treatment effect remained comparably constant and significant in all the models, except for the multivariable analysis including LVIDSN and change in LVIDSN, where the observed treatment effect was apparently reduced (and no longer significant). The 4 multivariable Cox proportional hazards models are summarized in Table 6. (A Forest plot for each model is illustrated in Fig S1a–d.) Proportional hazards and linearity assumptions were not violated, and no pair of variables entered into the same model showed unacceptable collinearity.

# Effect of Treatment Over Time

The AUCs adjusted for number of days in the study for: body weight, rectal temperature, respiratory rate, owner-measured resting respiratory rate, heart rate, and VHS for each study group are summarized and compared between groups in Table 7. The median AUCs adjusted for number of days in the study for respiratory rate, heart rate, and VHS were significantly lower in the group receiving pimobendan.

There were no consistent differences observed between treatment groups in quality of life scores, change in heart murmur intensity, change in heart disease stage, and change in BCS when they were compared at each visit up to the scheduled visit at 24 months, although heart failure score was worse at

20 months, and respiratory effort score was worse at 24 months in the placebo group (Table S2).

# Comparison of Groups at the Onset of CHF

At the onset of CHF, there were no significant differences between treatment groups for any of the physical examination, diagnostic imaging, or laboratory variables recorded (Table 8). For all dogs that went on to develop CHF, there was, with the exception of serum Na+, K+, and PCV, a significant change from baseline to the onset of CHF in both the pimobendan and placebo group in all of the continuous clinical, diagnostic imaging, and laboratory variables recorded. The magnitude of the change from baseline of these variables for dogs from both groups combined is summarized in Table 9. Likewise, there was a significant worsening of quality of life variables, BCS, and heart murmur intensity in their score as all dogs progressed from baseline to CHF (Table 10). Only the difference in change in exercise tolerance was significantly different between groups, with a greater number of dogs in the pimobendan group experiencing deterioration in exercise tolerance at the onset of CHF (Table 10).

# Discussion

Our study has demonstrated a number of changes in dogs with preclinical MMVD administered pimobendan,

**Table 3.** Baseline characteristics of the 2 treatment groups in the per-protocol population.

		Treatmer	nt Groups
	Variable	Pimobendan n = 178	Placebo n = 176
Dog characteristics	Age (years) Sex (M/F/MC/FS) (%) Breed (CKCS/Dachshund/Miniature Schnauzer/ Poodle/Yorkshire terrier/mixed	9.0 (8.0–11.0) 36/6/75/61 (20/3/42/34) 77/12/5/4/0/26/54 (43/7/3/2/0/15/30)	9.0 (7.0–11.0) 35/12/66/63 (20/7/38/36) 84/9/8/4/8/19/44 (47/5/5/2/5/11/25)
Dose of test medication Quality of life and respiratory variables	breed/other) (%) Dose pimobendan (mg/d) Appetite (decreased/normal/ increased) (%)	0.49 (0.44–0.53) 4/165/9 (2/93/5)	NA 3/166/7 (2/94/4)
(see Table S1 for details)	Demeanor (Alert/mildly lethargic/moderately lethargic) (%)	175/3/0 (98/3/0)	168/7/1 (95/4/1)
	Exercise tolerance (very good/good/decreased) (%)	118/53/7 (66/30/4)	99/70/7 (56/40/4)
	Fainting (none/rarely/occasional) (%) Respiratory effort (normal/mildly increased/moderately increased) (%)	175/3/0 (98/2/0) 172/5/1 (97/3/1)	170/4/2(97/2/1) 164/10/2 (93/6/1)
	Cough (none/occasional/ frequent/persistent) (%)	108/48/20/2 (61/27/11/1)	123/35/17/1 (70/20/10/1)
	Nocturnal coughing (none/slight/moderate) (%)	157/20/0 (89/11/0)	163/10/2 (93/6/1)
Physical examination variables	Body weight (kg) Body condition score (underweight (1–3)/normal (4–6)/overweight (7–9)) (%)	8.6 (6.9–10.6) 0/166/12 (0/93/7)	9.0 (7.1–10.5) 5/148/23 (3/84/13)
	Rectal temperature (°C)	38.7 (38.4–39.1)	38.7 (38.4–39.0)
	Heart rate (BPM)	124 (110–140)	122 (112–140)
	Respiratory rate (breaths/min)	28 (24–36)	28 (24–36)
	Heart murmur intensity (moderate (grade 3–4)/severe (grade 5–6) (%)	133/45 (75/25)	133/43 (76/24)
Diagnostic imaging variables	VHS	11.3 (10.9–12.0)	11.5 (11.0–11.9)
	LVIDSN	1.03 (0.93–1.14)	1.02 (0.93–1.10)
	LVIDDN	1.9 (1.8–2.1)	1.9 (1.8–2.0)
	FS% (%)	43 (39–48)	44 (41–49)
	LA/Ao	1.89 (1.73–2.10)	1.86 (1.72–2.06)
Laboratory variables	Na+ (mmol/L)	148 (145–150)	148 (146–149)
	K+ (mmol/L)	4.4 (4.1–4.8)	4.4 (4.1–4.7)
	PCV (%)	44.0 (41.0–48.0)	44.0 (40.0–48.1)
	Creatinine (µmol/L)	70.7 (60.0–88.4)	70.7 (61.0–82.2)
	TPC (g/L)	65 (61–69)	66 (62–70)
	GPT (ALT) (U/L)	43 (29–68)	42 (30–66)

ALT, alanine aminotransferase; BCS, body condition score; BPM, beats per minute; CKCS, Cavalier King Charles Spaniels; F, female; FS, female neutered; FS%, fractional shortening; GPT, glutamic-pyruvate transaminase; K+, potassium concentration; LA/Ao, left atrial-to-aortic root ratio; LVIDd, left ventricular internal diameter in diastole; LVIDDN, normalized left ventricular internal diameter in diastole; LVIDs, left ventricular internal diameter in systole; LVIDSN, normalized left ventricular internal diameter in systole; M, male; MN, male neutered; Na+, sodium concentration; PCV, packed cell volume; TPC, total protein concentration; VHS, vertebral heart sum. Continuous variables are reported as median (interquartile range). Categorical variables are reported as number (%).

which are different to those that occur in dogs that receive a placebo. Within the first month of treatment, dogs in the pimobendan group demonstrated significant reductions in 3 measures of heart size. The degree to which heart size reduced was predictive of outcome with a greater reduction in heart size associated with a prolongation of the time to the primary endpoint of the study. Heart size was on average lower for the duration of the study for dogs in the pimobendan group compared to those in the placebo group. Quality of life was perceived by owners to be improved after 1 month of

treatment for both the placebo and pimobendan groups. At the onset of CHF, dogs receiving pimobendan were indistinguishable from those receiving placebo, and both groups had larger hearts and worse quality of life compared to baseline.

There is a reduction in heart size in dogs with MMVD<sup>9,10,12</sup> and DCM<sup>11</sup> administered pimobendan. Earlier studies have only evaluated the effect of initiation of pimobendan on dogs' heart size after the onset of heart failure in dogs with MMVD. This is therefore the first study to show an effect of pimobendan in

**Table 4.** Within-group paired comparisons and between-group unpaired comparisons of the change from baseline to day 35 of various quality of life and physical examination variables.

Categorical Variables	Treatment Group	N	Deteriorated	Unchanged	Improved	Within-Group Comparison (Wilcoxon Signed Ranks)	Between-Group Comparison (Mann-Whitney)
Appetite	Placebo	160	6	140	14	0.12	0.33
**	Pimobendan	169	5	144	20	0.0012	
Demeanor	Placebo	160	3	135	22	< 0.0001	0.34
	Pimobendan	169	2	138	29	< 0.0001	
Exercise tolerance	Placebo	160	4	132	24	< 0.0001	0.75
	Pimobendan	169	3	144	22	< 0.0001	
Fainting	Placebo	160	2	157	1	1.0000	0.65
-	Pimobendan	169	1	167	1	1.0000	
Respiratory effort	Placebo	160	3	148	9	0.15	0.75
	Pimobendan	169	1	162	6	0.13	
Cough	Placebo	160	9	134	17	0.12	0.46
	Pimobendan	169	6	143	20	0.0038	
Nocturnal dyspnea or	Placebo	160	3	152	5	0.73	0.65
cough	Pimobendan	169	2	161	6	0.29	
Body condition score	Placebo	160	24	123	13	0.070	0.55
•	Pimobendan	169	24	127	18	0.36	
Heart failure stage	Placebo	160	_	157	3	0.25	0.76
-	Pimobendan	169	-	165	4	0.13	
			Decreased	Unchanged	Increased		
Auscultation	Placebo	160	12	130	18	0.28	0.22
(change in murmur grade)	Pimobendan	169	21	131	17	0.52	

All *P*-values that appear in bold are < 0.05.

reducing the heart size of dogs with MMVD and cardiac enlargement before the onset of clinical signs. The reduction in heart size was evident by 3 different indices: LVIDDN, LVIDSN, and LA/Ao. Similar changes in left ventricular diameter have been shown in Dobermans with preclinical DCM.<sup>11</sup> There was also an increase in the FS% of dogs receiving pimobendan. A small but significant reduction in left atrial size (LA/ Ao) was also seen in the placebo group. One possible explanation for this would be "regression to the mean." Regression to the mean can occur when a clinical measurement must exceed a threshold in order for a patient to be recruited to a study. Patients with "true" values for the measurement just below the threshold but an erroneously high measurement at the time of screening will enter the study, whereas those with true values just above the threshold with an erroneously low measurement will not. On a second measurement, despite the "true" value not having changed, if the measurement made is more reflective of the true measurement, it will appear to have fallen or "regressed to the mean." In this case, as LA/Ao was one of the more stringent entry criteria, regression to the mean in some recruited patients makes it appear as though the value has fallen in the placebo group. Importantly the reduction in left atrial size was significantly greater in the pimobendan group compared to the placebo group (median -0.08compared to -0.027), indicating that there was a true treatment effect as well as any change attributable to variation in measurement.

The change in echocardiographic variables at day 35  $(\pm 7 \text{ days})$  was associated with outcome. Those dogs that had greater reductions in LA/Ao, LVIDDN, and LVIDSN tended to have longer periods before reaching the primary endpoint of the study. This effect was independent of the effect of pimobendan administration as shown by the multivariable analyses, in which the change in the echocardiographic variable was significantly associated with outcome even when the effect of pimobendan was included in the model. For instance, a 0.1-unit increase in LA/Ao increased the risk of reaching the primary endpoint by 14% (HR = 1.14). A reduction in the LA/Ao by 0.1 units decreased the risk to 100/1.14%, that is, 87.7% representing a 12.3% reduction in risk. When the change in systolic diameter was included in the multivariable model, it appeared to diminish the effect of pimobendan administration to the greatest extent, making it appear nonsignificant (HR 0.71 [95% CI 0.51–1.00] P = 0.051). The same phenomenon was observed in the PROTECT study<sup>11</sup> where in a multivariable analysis, the inclusion of the change in systolic diameter led to a widening of the 95% CI of the estimate of the treatment effect resulting in a nonsignificant P-value for the treatment effect. This might indicate that the change in systolic diameter is an "intervening variable"20 and therefore on the causal pathway by which pimobendan mediates its favorable effect on outcome. The administration of pimobendan brings about a reduction in systolic diameter. If the reduction in systolic diameter is one of the major ways in which

**Table 5.** Within-group paired comparisons and between-group unpaired comparisons of the change from baseline to day 35 of physical examination, echocardiographic, and laboratory variables.

	Continuous Variables	Treatment Group	N	Absolute Change from Baseline Median (Interquartile Range)	Within-Group Comparison (Wilcoxon Signed Ranks)	Between-Group Comparison (Mann-Whitney)
Physical examination	Body weight (kg)	Placebo	159	0.00 (-0.20 to +0.20)	0.90	0.86
variables		Pimobendan	168	0.00 (-0.20 to +0.20)	0.99	
	Rectal temperature (°C)	Placebo	160	0.00 (-0.30  to  +0.30)	0.47	0.98
	•	Pimobendan	164	0.00 (-0.30  to  +0.30)	0.80	
	Respiratory rate	Placebo	152	0.0 (-4  to  +6)	0.24	0.51
	(breaths/min)	Pimobendan	154	0.0 (-4  to  +4)	0.60	
	Heart rate (BPM)	Placebo	160	$0.0 \ (-10.0 \ \text{to} \ +12.0)$	0.37	0.26
		Pimobendan	169	$0.0 \ (-12.0 \ \text{to} \ +10.0)$	0.32	
Echocardiographic	FS% (%)	Placebo	160	0.00 (-3.75  to  +3.20)	0.55	< 0.0001
variables		Pimobendan	169	+2.00 (-1.00 to +5.00)	< 0.0001	
	LVIDDN	Placebo	159	+0.01 (-0.07 to +0.07)	0.71	< 0.0001
		Pimobendan	168	-0.06 (-0.15  to  +0.02)	< 0.0001	
	LA/Ao	Placebo	160	-0.027 (-0.182  to  +0.084)	0.036	0.0076
		Pimobendan	169	-0.080 (-0.229  to  +0.028)	< 0.0001	
	LVIDSN	Placebo	159	+0.0144 (-0.0722 to +0.0715)	0.33	< 0.0001
		Pimobendan	168	-0.0614 ( $-0.1555$ to $-0.0033$ )	< 0.0001	
Laboratory variables	Na+ (mmol/L)	Placebo	159	0.000 (-2.000  to  +2.000)	0.82	0.21
		Pimobendan	167	0.000 (-2.000  to  +1.000)	0.030	
	K+ (mmol/L)	Placebo	159	0.000 (-0.200  to  +0.300)	0.24	0.24
		Pimobendan	168	0.000 (-0.300 to +0.200)	0.67	
	PCV (%)	Placebo	158	0.000 (-1.900 to +2.100)	0.17	0.59
		Pimobendan	168	+0.500 (-2.000  to  +2.750)	0.060	
	Creatinine (µmol/L)	Placebo	159	0.000 (-8.840 to +8.840)	0.86	0.094
		Pimobendan	168	0.000 (-8.840 to +4.210)	0.014	
	TPC (g/L)	Placebo	158	0.000 (-2.000  to  +2.500)	0.72	0.19
		Pimobendan	168	+0.415 (-2.000 to +4.000)	0.054	
	GPT (ALT) (U/L)	Placebo	159	0.000 (-6.000 to +8.000)	0.52	0.38
		Pimobendan	168	0.000 (-9.000 to +6.000)	0.48	

ALT, alanine aminotransferase; BPM, beats per minute; FS%, fractional shortening; GPT, glutamic-pyruvate transaminase; K+, potassium concentration; LA/Ao, left atrial-to-aortic root ratio; LVIDDN, normalized left ventricular internal diameter in diastole; LVIDSN, normalized left ventricular internal diameter in systole; Na+, sodium concentration; PCV, packed cell volume; TPC, total protein concentration; VHS, vertebral heart sum.

All P-values that appear in bold are < 0.05.

pimobendan mediates its favorable effect (ie, it intervenes on the "causal pathway") once the reduction in systolic diameter is taken into account in a multivariable analysis, the association between treatment and the outcome is no longer apparent. These findings add further weight to the existing literature which suggests that, at least in part, the benefit of pimobendan treatment in canine heart disease is mediated through the reduction in heart size that it brings about. This speculation about a potential mechanism of action in dogs with MR is supported by experimental findings in dogs with induced MR which suggest that reduced heart size and enhanced contractile function are associated with a reduction in mitral regurgitant orifice area and therefore reduced MR.<sup>21,22</sup>

The observed reduction in heart size appeared to be maintained over the duration of the study with the average heart size, as indicated by the AUC for the VHS, of dogs in the pimobendan group being smaller compared to dogs in the placebo group.

Change in quality of life variables did not differ significantly between groups at day 35. Although there were 2 quality of life variables, appetite and cough, that appeared to improve significantly in the pimobendan group and not the placebo group, between-group comparisons did not reveal significant differences between groups. Both groups appeared to show a highly significant (P < 0.0001) improvement in demeanor and exercise tolerance at day 35, yet the difference between groups was not significant. These observed changes suggest a possible "placebo effect" with owners perceiving improvement in their dogs because they believed they were receiving medication. This observation emphasizes the importance of including a placebo group in clinical trials in which subjective judgments are made about responses to treatment. Almost all the dogs recruited to the study were perceived by their owners to be normal at the time of entry to the study; for instance, 96% of owners rated their dog's exercise tolerance as good or very good on entry to the study. It would therefore be less likely that a difference in improvement between groups could be shown given that the dogs were not considered to be showing clinical signs as a consequence of their disease. The absence of differences between

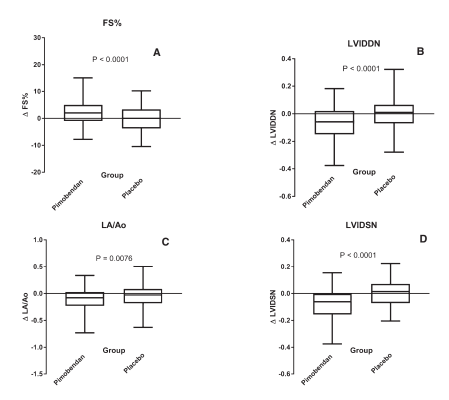


Fig. 2. Four box and whiskers plots illustrating absolute values of the change in 4 different echocardiographic variables between baseline and day 35 in the 2 different treatment groups for (A) fractional shortening, (B) normalized left ventricular internal diameter in diastole, (C) left atrial-to-aortic ratio and (D) normalized left ventricular internal diameter in systole. The bold horizontal line indicates the median change, the boxes extend from the 25th to the 75th percentile. The whiskers extend from the 2.5th to the 97.5th percentile. P-values are for between-group comparisons by a Mann-Whitney test. Δ, change in; FS%, fractional shortening; LA/Ao, left atrial-to-aortic root ratio; LVIDDN, normalized left ventricular internal diameter in diastole; LVIDSN, normalized left ventricular internal diameter in systole.

groups may also be a reflection of the relatively crude means of rating quality of life scores that were used, by comparison to validated tools such as the FETCH score.<sup>23</sup>

Two clinical variables that were seen to be lower in the pimobendan group when compared to the placebo group over the duration of the study by the AUC method were the heart rate on physical examination and the owner-measured resting respiratory rate. The median AUC of resting respiratory rate was 2 breaths per minute lower in the pimobendan group, and the median AUC of heart rate was 6 beats per minute lower. Given that higher heart rate and higher resting respiratory rate tend to be associated with more advanced MMVD<sup>6</sup> and the onset of CHF,<sup>24</sup> the lower rates in dogs treated with pimobendan might be an indication of improved stability of these dogs even though they were considered to be preclinical at the time of recruitment to the trial.

The magnitude of the differences observed in heart rate, resting respiratory rate, and VHS between groups as a consequence of treatment was considerably smaller than the observed variability within the population. For instance, the IQR for heart rate in the population as a whole at baseline was 30 beats per minute, whereas the difference observed between the medians for the AUCs observed in the 2 treatment groups was only 6 beats per minute. Thus, the changes were not sufficiently profound

to allow the treatment groups to be discriminated, or to act as reliable indicators of a response to treatment in individual dogs. Similarly, although the difference observed between groups in the change in echocardiographic variables at day 35 was statistically highly significant, the magnitude of the absolute changes was relatively small (the median change was <7% of median baseline values). The within-group variability of the measured changes was relatively large, and the magnitude of the changes in the 2 groups overlapped considerably (Fig 2). This means that individual dogs on pimobendan could not be reliably discriminated from those on placebo on the basis of the changes observed.

Not all dogs recruited to the EPIC study went on to develop CHF; however, the relatively large number of dogs (135 in total) that did develop CHF provides us with an opportunity to describe the changes that occur in a number of clinical, radiographic, and echocardiographic variables as dogs develop heart failure. In particular, we have paired observations in dogs obtained at baseline and at the time of onset of CHF. It is notable that the 2 treatment groups did not differ significantly in any of the measured characteristics at the onset of CHF; among other characteristics, heart size, heart rate, and respiratory rate were very similar in both groups at that time. The similar heart size seen in both groups suggests that by the onset of CHF, the heart size in dogs receiving pimobendan had increased up to a level

**Table 6.** Multivariable Cox proportional hazards models exploring the effect of treatment, change in an echocardiographic variable, and the value of the same variable recorded at baseline on time to the primary endpoint constructed for 4 echocardiographic variables; LVIDDN, LA/Ao, LVIDSN, and FS%.

	Hazard Ratio	95% Confidence Intervals of Hazard Ratio	<i>P</i> -Value
Change in LVIDDN model	(n = 327)		
Treatment (pimobendan)	0.64	0.46-0.90	0.010
LVIDDN (per 0.1 unit)	1.30	1.20-1.41	< 0.0001
Δ LVIDDN	1.26	1.11-1.43	0.0003
(per 0.1 unit change)			
Change in LA/Ao model (n	= 329)		
Treatment (pimobendan)	0.63	0.46 - 0.88	0.0060
LA/Ao (per 0.1 unit)	1.21	1.15-1.28	< 0.0001
Δ LA/Ao	1.14	1.06-1.22	0.0002
(per 0.1 unit change)			
Change in LVIDSN model (	(n = 327)		
Treatment (pimobendan)	0.71	0.51 - 1.00	0.051
LVIDSN (per 0.1 unit)	1.07	0.97 - 1.18	0.16
$\Delta$ LVIDSN	1.22	1.05-1.41	0.0084
(per 0.1 unit change)			
Change in FS% model (n =	329)		
Treatment (pimobendan)	0.64	0.46-0.90	0.011
FS % (per 10 units)	1.47	1.16-1.85	0.0014
ΔFS%	1.05	0.77 - 1.44	0.75
(per 10 units change)			

Δ, Change in; FS%, fractional shortening; LA/Ao, left atrial-to-aortic root ratio; LVIDDN, normalized left ventricular internal diameter in diastole; LVIDSN, normalized left ventricular internal diameter in systole.

All *P*-values that appear in bold are < 0.05.

comparable to that observed in the placebo group, despite being, on average, lower for the duration of the study as determined by the AUC comparison of VHS. If there is a critical heart size for a dog, which when reached is likely to be associated with the development of signs of CHF, it is possible that reducing heart size at the onset of treatment is one of the ways in which pimobendan delays the onset of CHF. We know from the main results of the EPIC study that dogs receiving pimobendan take longer to develop CHF, which may argue again in favor of the reduction in heart size being one of the primary ways in which this effect is mediated.

Our analysis also showed that a number of clinical and quality of life variables change in dogs as they develop CHF. In the population of dogs that developed CHF, body weight and rectal temperature decreased. Some of the biochemical variables also changed with creatinine and ALT increasing (although the median change for creatinine was zero) and total protein decreasing. No change in electrolyte concentrations or PCV was observed.

As might be expected, the quality of life variables measured universally showed evidence of deterioration at the onset of CHF with only the change in exercise tolerance differing between groups (with more

pimobendan dogs showing evidence of deterioration). There was also a reduction in BCS and an increase in murmur intensity in both groups.

#### Limitations

The analyses described in this study are exploratory analyses of longitudinal data acquired from dogs in a randomized clinical trial. They do not relate directly to the primary hypothesis of the original study and should therefore be regarded as hypothesis generating. Statistical analyses are not corrected for multiple comparisons, and therefore, P-values close to the threshold of significance should be interpreted with caution. For many of the more important conclusions of the study, for example, the reduction in heart size after 35 days, the *P*-values associated with the comparisons are <0.0001 and therefore very unlikely to represent a type I statistical error.

Comparing groups in longitudinal studies is challenging, particularly when the rate at which individuals leave those groups is unequal, not at random and potentially related to what is being measured.<sup>25</sup> For example, dogs with bigger hearts are more likely to be missing from the PP population because they have met the primary endpoint of the study. This unequal time spent in the study and unequal numbers of observations in the 2 groups can be partially compensated for by methods such as the AUC method we employ.<sup>18</sup> This method is only likely to be appropriate for use with continuous variables where the values can be meaningfully averaged. When variables such as the quality of life variables we recorded are compared, such a method is probably not appropriate. Comparing groups at individual visits is problematic because lower numbers of animals will be left in the group that meets the primary endpoint of the study more quickly (in this study the placebo group). This means that after a given point in the study, those animals that remain are likely to be in some way exceptional rather than being representative of the population as a whole. For this reason, we did not compare quality of life variables between groups after 24 months when fewer than 50% of the dogs remained in each group.

Comparing quality of life between groups is also challenging because those dogs with a poor quality of life are likely to have left the study through either having met the primary endpoint or having been censored from the PP population for another reason. These types of events tended to occur sooner in the group receiving placebo, as was reflected by the results of the time to first event analysis we have already described and is indicated by the lower number of dogs remaining in the PP population in the placebo group. Thus, the populations being compared are those remaining in the study that continue to have a good quality of life, and differences between groups are not observed as is seen in Table S2. As acknowledged above, the method we used for assessing quality of life was a relatively insensitive, subjective, and unvalidated tool and may also have contributed to the absence of observed differences between

**Table 7.** Comparison between groups of the areas under the curve averaged over the number of days in the study for each dog for those continuous variables that were recorded regularly throughout the monitoring period and on at least 2 occasions.

Continuous Variables	Treatment Group	N	Median AUC Adjusted for Days in Study (IQR)	Between-Group Comparison
Body weight (kg)	Placebo	173	8.5 (6.9–10.4)	0.54
	Pimobendan	176	8.8 (6.9–10.5)	
Rectal temperature (°C)	Placebo	173	38.6 (38.4–38.8)	0.53
	Pimobendan	176	38.6 (38.4–38.8)	
Respiratory rate (breaths/min)	Placebo	170	32 (26–40)	0.49
	Pimobendan	173	31 (26–38)	
Owner-measured resting	Placebo	156	24 (20–28)	0.0051
respiratory rate (breaths/min)	Pimobendan	163	22 (19–26)	
Heart rate (BPM)	Placebo	173	130 (119–140)	0.038
, ,	Pimobendan	176	124 (115–137)	
Vertebral heart score (VHS)	Placebo	157	11.9 (11.4–12.4)	0.0009
,	Pimobendan	163	11.5 (11.0–12.2)	

BPM, beats per minute; VHS, vertebral heart sum. All *P*-values that appear in bold are < 0.05.

**Table 8.** Values of physical examination, diagnostic imaging, and laboratory variables for those dogs that developed congestive heart failure, measured at the time of onset of heart failure summarized for the 2 treatment groups.

	Continuous Variables	Treatment Group	N	Median Value at CHF and IQR	Between-Group Comparison
Physical examination	Body weight (kg)	Placebo	73	8.7 (6.9–10.6)	0.96
variables		Pimobendan	58	9.2 (7.1–10.0)	
	Rectal temperature (°C)	Placebo	66	38.6 (38.3–38.8)	0.11
		Pimobendan	56	38.4 (38.0–38.8)	
	Respiratory rate (breaths/min)	Placebo	69	44 (40–58)	0.25
		Pimobendan	52	50 (36–66)	
	Owner-measured resting	Placebo	58	36 (30–50)	0.50
	respiratory rate	Pimobendan	50	39 (34–50)	
	(breaths/min)				
	Heart rate (BPM)	Placebo	71	150 (140–162)	0.60
		Pimobendan	58	150 (140–164)	
Diagnostic imaging	FS% (%)	Placebo	67	46.6 (42.5–50.1)	0.63
variables		Pimobendan	55	46 (42.0–53.8)	
	LVIDDN	Placebo	67	2.26 (2.14–2.51)	0.94
		Pimobendan	55	2.27 (2.14–2.47)	
	LA/Ao	Placebo	67	2.41 (2.15–2.80)	1.00
		Pimobendan	55	2.47 (2.17–2.71)	
	LVIDSN	Placebo	67	1.16 (1.03–1.33)	0.62
		Pimobendan	55	1.13 (0.98–1.36)	
	VHS	Placebo	72	13.0 (12.2–13.5)	0.45
		Pimobendan	59	12.8 (12.1–13.5)	
Laboratory variables	Na+ (mmol/L)	Placebo	68	147.0 (145.4–150.0)	0.66
		Pimobendan	55	147.0 (146.0–149.0)	
	K+ (mmol/L)	Placebo	68	4.50 (4.25-4.80)	0.40
		Pimobendan	52	4.50 (4.00-4.80)	
	PCV (%)	Placebo	66	41.5 (37.4–48.0)	0.12
		Pimobendan	50	44.0 (40.0–47.0)	
	Creatinine (µmol/L)	Placebo	68	78.8 (61.9–90.0)	0.90
		Pimobendan	52	79.6 (64.9–87.2)	
	TPC (g/L)	Placebo	66	64.0 (58.0–70.0)	0.77
		Pimobendan	52	66.0 (60.5–68.0)	
	GPT (ALT) (U/L)	Placebo	62	52.5 (35.0–77.0)	0.68
	. , , , ,	Pimobendan	51	54.0 (34.0–95.0)	

ALT, alanine aminotransferase; BPM, beats per minute; FS%, fractional shortening; GPT, glutamic-pyruvate transaminase; K+, potassium concentration; LA/Ao, left atrial-to-aortic root ratio; LVIDDN, normalized left ventricular internal diameter in diastole; LVIDSN, normalized left ventricular internal diameter in systole; Na+, sodium concentration; PCV, packed cell volume; TPC, total protein concentration; VHS, vertebral heart sum.

**Table 9.** Change from baseline to the onset of heart failure for values of physical examination, diagnostic imaging, and laboratory variables for those dogs that developed congestive heart failure summarized for the population as a whole.

	Continuous Variables	N	Absolute Change from Baseline Median (Interquartile Range)	Comparison to Baseline
Physical examination variables	Body weight (kg)	131	-0.30 (-0.7 to 0.1)	< 0.001
	Rectal temperature (°C)	122	-0.3 (-0.7  to  0.1)	< 0.001
	Respiratory rate (breaths/min)	118	16 (4–29)	< 0.001
	Heart rate (BPM)	129	22 (8–38)	< 0.001
Diagnostic imaging variables	FS% (%)	122	2.6 (-2.6  to  7.7)	0.003
	LVIDDN	122	0.35 (0.14–0.49)	< 0.001
	LA/Ao	122	0.47 (0.24–0.75)	< 0.001
	LVIDSN	122	0.13 (-0.04 to 0.27)	< 0.001
	VHS	131	1.3 (0.80–1.9)	< 0.001
Laboratory variables	Na+ (mmol/L)	120	0.00 (-3.0  to  2.0)	0.24
•	K+ (mmol/L)	119	$0.10 \ (-0.20 \ \text{to} \ 0.40)$	0.067
	PCV (%)	115	-1.0 (-3.9  to  3.0)	0.095
	Creatinine (µmol/L)	119	0.0 (-8.8  to  17.7)	0.018
	TPC (g/L)	116	-2.0 (-5.5  to  3.0)	0.027
	GPT (ALT) (U/L)	112	10.0 (-6.0 to 43.0)	< 0.001

ALT, alanine aminotransferase; BPM, beats per minute; FS%, fractional shortening; GPT, glutamic-pyruvate transaminase; K+, potassium concentration; LA/Ao, left atrial-to-aortic root ratio; LVIDDN, normalized left ventricular internal diameter in diastole; LVIDSN, normalized left ventricular internal diameter in systole; Na+, sodium concentration; PCV, packed cell volume; TPC, total protein concentration; VHS, vertebral heart sum.

All *P*-values that appear in bold are < 0.05.

**Table 10.** A summary of the number of dogs in each group that experienced a change in various clinical variables at the time of the onset of congestive heart failure.

Categorical Variables	Treatment Group	N	Deteriorated	Unchanged	Improved	Within-Group Paired Longitudinal Comparisons (Wilcoxon signed ranks)	Between-Group Comparison (Mann-Whitney)
Appetite	Placebo	69	25	44	0	< 0.0001	0.26
**	Pimobendan	57	27	29	1		
Demeanor	Placebo	69	41	28	0	< 0.0001	0.45
	Pimobendan	57	38	18	1		
Exercise tolerance	Placebo	69	41	28	0	< 0.0001	0.020
	Pimobendan	57	45	12	0		
Fainting	Placebo	69	14	55	0	< 0.0001	0.52
•	Pimobendan	57	9	48	0		
Respiratory effort	Placebo	69	55	14	0	< 0.0001	0.074
	Pimobendan	57	52	5	0		
Cough	Placebo	69	60	6	3	< 0.0001	0.82
· ·	Pimobendan	57	50	6	1		
Nocturnal dyspnea	Placebo	67	46	21	0	< 0.0001	0.76
or cough	Pimobendan	56	37	19	0		
			Decreased	Unchanged	Increased		
Body condition score	Placebo	72	26	37	9	0.0002	0.84
-	Pimobendan	58	21	28	9		
Heart murmur intensity	Placebo	72	8	30	34	< 0.0001	0.17
•	Pimobendan	58	2	23	33		

P-values for paired comparisons with the same characteristics recorded from the same dogs at baseline are given in the penultimate column. All P-values that appear in bold are < 0.05.

groups. Having acknowledged these weaknesses in the method of measurement, it was still possible by this method for us to show a clear deterioration in quality of life associated with the onset of CHF.

#### **Conclusions**

In dogs with preclinical MMVD, pimobendan treatment results in a smaller heart size both in the short

and long term. The degree to which heart size reduces in the short term is predictive of outcome with a greater reduction in heart size associated with a prolongation of the time to CHF or cardiac-related death. In particular, the therapeutic benefit of pimobendan appears to be associated with the reduction in LV end-systolic dimension. There were no consistent differences observed between treatment groups in quality of life variables, either in the short or long term as might be expected in a study of dogs with preclinical disease where deterioration in quality of life often signified reaching the primary endpoint or resulted in censoring. At the onset of CHF, dogs treated with pimobendan were indistinguishable from those receiving placebo showing clear evidence of a deterioration in quality of life and significant changes in a number of clinical, diagnostic imaging, and laboratory variables.

### **Footnotes**

<sup>a</sup> Vetmedin 2.5 mg chewable tablets

<sup>b</sup> SAS Version 8.2; SAS Institute Inc., Cary, NC, USA

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### **Supporting Information**

Additional Supporting Information may be found online in the supporting information tab for this article:

**Figure S1.** Four Forest plots illustrating the hazard ratios and their 95% confidence intervals obtained from the four multivariable Cox proportional hazards analysis models.

**Table S1.** Ordinal scoring system for clinical variables recorded at baseline.

**Table S2.** *P*-values for the between-group comparison of ordinal quality of life and clinical variables at each of the visits up to and including the visit at 24 months.