

# Alma Mater Studiorum Università di Bologna Archivio istituzionale della ricerca

Canine indolent and aggressive lymphoma: Clinical spectrum with histologic correlation

This is the final peer-reviewed author's accepted manuscript (postprint) of the following publication:

Published Version:

Aresu L., Martini V., Rossi F., Vignoli M., Sampaolo M., Arico A., et al. (2015). Canine indolent and aggressive lymphoma: Clinical spectrum with histologic correlation. VETERINARY AND COMPARATIVE ONCOLOGY, 13(4), 348-362 [10.1111/vco.12048].

Availability:

This version is available at: https://hdl.handle.net/11585/702699 since: 2019-10-18

Published:

DOI: http://doi.org/10.1111/vco.12048

Terms of use:

Some rights reserved. The terms and conditions for the reuse of this version of the manuscript are specified in the publishing policy. For all terms of use and more information see the publisher's website.

This item was downloaded from IRIS Università di Bologna (https://cris.unibo.it/). When citing, please refer to the published version.

(Article begins on next page)

1	
2	
3	This is the final peer-reviewed accepted manuscript of:
4 5 6 7	Finotello R, Henriques J, Sabattini S, Stefanello D, Felisberto R, Pizzoni S, Ferrari R, Marconato L. A retrospective analysis of chemotherapy switch suggests improved outcome in surgically removed, biologically aggressive canine haemangiosarcoma. Vet Comp Oncol. 2017 Jun;15(2):493-503. doi: 10.1111/vco.12193. Epub 2016 Jan 21. PMID: 26792231.
8	
9	Rights / License:
10 11	The terms and conditions for the reuse of this version of the manuscript are specified in the publishing policy. For all terms of use and more information see the publisher's website.
12	

14	<u>A retrospective analysis of c</u> Chemotherapy switch <u>suggests</u> improve <u>ds</u> outcome in
15	surgically removed, biologically aggressive canine haemangiosarcoma
16	
17	Riccardo Finotello, <sup>1</sup> Joaquim Henriques, <sup>2</sup> Silvia Sabattini, <sup>3</sup> Damiano Stefanello, <sup>4</sup>
18	Ricardo Felisberto, <sup>2</sup> Selene Pizzoni, <sup>5</sup> Roberta Ferrari, <sup>4</sup> Laura Marconato <sup>5</sup>
19	
20	<sup>1</sup> Small Animal Teaching Hospital, School of Veterinary Sciences, University of
21	Liverpool, Neston, UK
22	<sup>2</sup> Centro Veterinario Berna, Lisbon, Portugal
23	<sup>3</sup> Department of Veterinary Medical Sciences, University of Bologna, Italy
24	<sup>4</sup> Department of Veterinary Science and Public Health, University of Milan, Milan, Italy
25	<sup>5</sup> Centro Oncologico Veterinario, Sasso Marconi, Italy
26	
27	Short title: chemotherapy switch in aggressive haemangiosarcoma
28	
29	Keywords: haemangiosarcoma, metronomic, thalidomide, chemotherapy switch, dog
30	
31	
32	Findings of this study were presented in part at the European Society of Veterinary Oncology
33	Annual Meeting, Krakow, Poland, 2015.
34	
35	Corresponding author: Laura Marconato, DVM, Diplomate ECVIM-CA (Oncology), Centro
36	Oncologico Veterinario, via San Lorenzo 1-4, I-40037 Sasso Marconi (Bologna), Italy; email:
37	marconato@centroncologicovet.it

# 39 Abstract

40	Haemangiosarcoma (HSA) has an aggressive biological behaviour and carries a poor
41	prognosis, with less than 10% of treated dogs surviving longer than one year.
42	In this retrospective study a varied metronomic chemotherapy (MC) regimen preceded
43	by standard adjuvant doxorubicin-based maximum-tolerated dose chemotherapy (MTDC) was
44	compared to MTDC, in terms of efficacy (time to metastasis, TTM, and survival time,
45	ST) and safety in dogs with biologically aggressive HSA. Dogs were eligible if they had
46	no metastasis after MTDC and received either no further chemotherapy or MC
47	maintenance.
48	Twenty-twodogswere enrolled: 12 Twelvedogs received MTDC, and 10 received MC thereafter. Median TTM and ST
49	were significantly longer for dogs receiving MTDC-MC (not reached versus 150 days,
50	P=0.028; and not reached versus 168 days, P=0.030, respectively). Treatment was well
51	tolerated.
52	MTDC followed by MC is safe and suggests improveds TTM and ST in dogs with
53	surgically removed, biologically aggressive HSA that are treated in the microscopic
54	setting.
55	

#### Introduction 57

Haemangiosarcoma (HSA) is a common mesenchymal tumour in dogs, arising in three 58 different forms: dermal, subcutaneous/muscular and visceral, the latter mainly involving 59 spleen, right atrium or auricle, and liver.<sup>1-3</sup> With the exception of the dermal form, 60 which may behave in a less aggressive fashion, subcutaneous/intramuscular and visceral 61 62 HSA is a highly malignant cancer, spreading rapidly to lungs, liver, peritoneum and central nervous system.<sup>4,5</sup> Unfortunately, visceral HSA has a silent evolution for a quite 63 64 long time, and is accompanied by non specific clinical signs. As a consequence, when detected, it is usually in an advanced or metastatic stage, therefore precluding cure.<sup>1,2</sup> 65 The mainstay of treatment consists of surgery followed by adjuvant intravenous 66 chemotherapy.6,7 Doxorubicin-based chemotherapy protocols have been administered to 67 dogs with HSA, including doxorubicin as single agent,6 or combined with ifosfamide,8 68 vincristine and cyclophosphamide,7,9-11 and epirubicin as single agent.12 Although a 69 70 three weekly regimen is the commonest schedule administration of doxorubicin, one study attempting to increase dose intensity by more frequent administrations showed 71 72 such strategy to be well tolerated; however, survival time was not improved.13

73 Although the combination of doxorubicin and dacarbazine has provided promising results in a recent clinical trial, it is still common knowledge that < 10% of the dogs 74 75 diagnosed with HSA will survive one year after diagnosis, being attributable to the development of metastatic disease during or after completion of maximum-tolerated 76 dose chemotherapy (MTDC).14 Thus, it appears obvious that MDTC is unlikely to 77 78 provide a durable response in such biologically aggressive solid tumours.

79 Metronomic chemotherapy (MC) refers to the frequent administration of cytotoxic 80 drugs at doses significantly lower than the maximum tolerated dose, with no prolonged drug-free breaks, leading to an anti-angiogenic effect and immune-modulation.<sup>15-16</sup> In 81 82 veterinary oncology, MC has been mainly used in a palliative setting with good 4

response rates and safety profile.<sup>17-18</sup> A continuous low-dose chemotherapy strategy 83 consisting of cyclophosphamide, etoposide, and piroxicam has been proposed as an 84 alternative treatment for dogs with HSA, yielding comparable results to conventional 85 86 MTDC, therefore suggesting a beneficial effect of this regimen in delaying disease progression in canine HSA.19 A more recent study suggested that the combination of 87 both MTDC and MC was more efficacious in dogs with splenic HSA than either type of 88 chemotherapy alone in the early follow-up period; however, no significant prolongation 89 90 of survival time was observed during the late follow-up period when compared with dogs undergoing splenectomy only.20 91

A "chemo-switch schedule" refers to the introduction of a new and potentially noncross-resistant agent after completion of first-line chemotherapy, such as the administration of MC after MTDC.<sup>21</sup> In the current study, we retrospectively compared MC preceded by doxorubicin-based MTDC to MTDC treatment only, in terms of efficacy (time to metastasis, TTM, and survival time, ST) and safety in dogs with biologically aggressive HSA. It was hypothesised that chemo-switch would improve long-term tumour control.

99 100

### 101 Material and methods

102

### 103 Inclusion criteria

104

105 The databases of the Centro Oncologico Veterinario (Bologna, Italy), Centro 106 Veterinario Berna (Lisbon, Portugal) and University of Milan Teaching Hospital 107 (Milan, Italy) were reviewed to identify client-owned dogs with histologically 108 confirmed and biologically aggressive HSA (2011-2014).

Haemangiosarcoma was considered as "biologically aggressive" if arising from anyvisceral, bone and muscular location or, in case of subcutaneous tumours, if the largest

111 diameter was > 6 cm.<sup>1-4</sup>

Eligible dogs for inclusion in the analysis set were those that had no evidence of macroscopic disease after completion of MTDC based on imaging and that received either no further chemotherapy or MC maintenance.

115 Pre-surgical, pre-dosing, and post-dosing investigations included physical examination,

haematology, serum biochemistry, abdominal ultrasound and at least two lateral views
thoracic radiographs or computed tomography (<u>CT</u>) if performed.

118 Dogs were monitored at least every three months after MTDC or during MC119 maintenance, as listed above.

Dogs were staged according to the World Health Organization (WHO) staging system
 for domestic animals.<sup>22</sup>

122

# 123 Treatment protocol

124 Based on owners' and clinicians' preference, dogs received MTDC followed by MC 125 (Group 1) or MTDC only (Group 2). MTDC consisted of a discontinued doxorubicinbased chemotherapy protocol. MC was administered orally and consisted of low-dose 126 127 cyclophosphamide (Endoxan®, Baxter s.r.l., Lurago d'Erba, Como, Italy) administered q24h or q48h at 7-15 mg/m<sup>2</sup>, and the cyclooxygenase-2 (COX-2) inhibitor firocoxib 128 (Previcox®, Merial, Lyon, France), meloxicam (Metacam®, Boehringer Ingelheim, 129 Milan, Italy), or a non-selective COX inhibitor (Piroxicam®, Pfizer Italia s.r.l., Latina, 130 131 Italy) administered daily at the standard recommended dose. The non-steroidal antiinflammatory drug (NSAID) varied depending on clinician's preference. In case of 132 haemorrhagic cystitis, cyclophosphamide was discontinued and dogs received oral 133

chlorambucil (Leukeran®, GlaxoSmithKline S.p.A., Verona, Italy) at the dosage of 4
mg/m<sup>2</sup> q24h or q48h.<sup>23</sup>

Depending on availability, oral thalidomide at 2-3 mg/kg (Thalidomid, Bichsel AG, Interlaken, Switzerland) was also administered q24h or q48h depending on clinician's preference. The dose of thalidomide was arbitrarily chosen based on some of the authors' experience.<sup>24</sup> Owners intending to have thalidomide administered were informed on its known teratogenic effect.<sup>25</sup>

141

### 142 Assessment of toxicity

Toxicity resulting from MTDC was assessed in both groups based on the dog's history, physical examination and complete blood counts (CBCs) 7-10 days after chemotherapy and before the beginning of each next cycle, as stated by the Veterinary Co-operative Oncology Group.<sup>26</sup> In Group 2, urinalysis was also carried out in the case of suspected urothelial toxicity (i.e. haematuria, stranguria, pollachiuria).

148

### 149 Statistical analysis

Follow-up and survival times were calculated from the date of diagnosis to the date of last visit or death. For both groups, ST and TTM (beyond regional lymph nodes) were explored with the Kaplan-Meier product limit method followed by log-rank test. In either group, timing was considered from surgical excision. In the survival analysis, dogs were censored if they were alive at the time of data accrual closure or died of no tumour-related causes, whereas for TTM dogs were censored if, by the last examination, distant metastases had not developed.

157 Causes of death were established reviewing the individual dog clinical histories and 158 through telephone calls to owners and referring veterinarians. Dogs were considered to 159 have died of HSA if the clinical staging work-up was consistent with the presence of

metastatic disease and if symptoms could be linked to HSA progression (i.e recurrence 160 of haemoabdomen); dogs were considered not to have died because of HSA if their last 161 staging work-up (performed no longer than one month before death) revealed no 162 163 evidence of metastatic disease and if death was determined to occur due to an unrelated 164 cause.

165 When appropriate, data sets were tested for normality by use of the D'Agostino and Pearson omnibus normality test. Values were expressed as mean ± standard deviation in 166 167 case of normal distribution, or as median with a range in case of non-normal 168 distribution.

To verify whether features of the two groups differed at admission or during MTDC, the 169 170 T-test (parametric variables) or Mann Whitney U test (non-parametric variables) was used to compare age, body weight, and the time occurred from the diagnosis to the 171 beginning of MTDC. Fisher's exact test was used to compare breed (pure- vs cross-172 breed), sex (male vs female), primary location of the tumour (spleen vs other sites), 173 174 clinical stage, number of doxorubicin cycles (<4 vs 4-6), type of chemotherapy protocol 175 (single agent doxorubicin vs poly-chemotherapy) and MTDC-related toxicity (present 176 vs absent). Data were analysed by use of commercial software programs (SPSS Statistics v. 19, IBM, Somers, NY, and Prism v. 5.0, GraphPad, San Diego, CA). P 177 178 values  $\leq 0.05$  were considered significant.

179

180

#### Results 181

182 Twenty-two dogs met the inclusion criteria and were enrolledincluded in the analysis; 10 (45.5%) of them received MTDC followed by MC (Group 1), whereas the remaining 12 183 (54.5%) were treated with MTDC (Group 2). Dogs' characteristics are listed in Table 1. Dogs 184 185 were not stratified based on prognostic risk, but there was good balance between arms 8

regarding dogs' features and possible outcome variables; however, concerning sex
distribution, there was a statistically significant difference between groups, as males were
more common in Group 1 and females were more common in Group 2 -(P=0.043; Table 2).
For all dogs, pre-surgical, pre-dosing, and post-dosing imaging investigations were performed
through thoracic radiographs and abdominal ultrasound. Two dogs (case 6 and case 22; Table
1) had CT scans repeated throughout the follow-up period.

192

# 193 Group 1 (MTDC-MC)

There were 3 mixed breed dogs, 2 German shepherds, 1 Golden retriever, 1 Labrador retriever, 1 Boxer, 1 Great Dane, and 1 Italian cane Corso. Mean age was  $8.9 (\pm 2.6)$ years and mean weight was  $36.4 (\pm 12.0)$  kg. There were 8 males (n=4 neutered) and 2 spayed female dogs. HSA occurred in the spleen as primary site in 8 dogs; all dogs presented with hemoperitoneum because of splenic rupture. The remaining 2 dogs had subcutaneous (n=1) and osseous (n=1) HSA.

All dogs underwent surgery, consisting of splenectomy, removal of the subcutaneous tumour, or amputation according to cancer location. Histopathological evaluation revealed clean surgical margins in the subcutaneous and osseous HSA; surgical margins were deemed not assessable for dogs presenting with visceral rupture.

According to the WHO classification, 9 dogs had stage II disease, and 1 dog with osseous HSA had stage III disease.

The mean time from surgery to initial MTDC administration was 20.8 ( $\pm$  15.4) days. Eight dogs received doxorubicin as single agent, and 2 dogs received a combination of doxorubicin and dacarbazine. For all dogs, the median number of doxorubicin cycles was 5 (range, 4 to 6 cycles), and the initial dose was 30 mg/m<sup>2</sup> for all dogs. Chemotherapy dose reduction was undertaken in 3 dogs receiving single agent doxorubicin; this was performed at the clinician's discretion after haematological and/or
gastrointestinal toxicity developed: 2 dogs had 10% and 1 had 20% dose reduction. The
median total dose of doxorubicin was 132 mg/m<sup>2</sup> (range, 120 to 180 mg/m<sup>2</sup>).

214 The median time from completion of MTDC to start of MC was 17.5 days (range, 13 to 24 days). Cyclophosphamide was administered q24h in 2 dogs and q48h in the 215 216 remaining 8 dogs. The median single cyclophosphamide dose was 8.5 mg/m<sup>2</sup> (range, 7 to 15 mg/m<sup>2</sup>), and the median weekly cumulative dose was 44 mg/m<sup>2</sup> (range, 28 to 105 217 218 mg/m<sup>2</sup>). Concerning NSAIDs, 5 dogs received piroxicam, 4 had meloxicam and 2 dogs received firocoxib. Thalidomide was given in combination with standard MC in 7 219 (70%) of 10 dogs: 5 dogs received 2 mg/kg q24h, whereas the remaining 2 were treated 220 at 3 mg/kg q24h. 221

222

### 223 Group 2 (MTDC)

There were 5 mixed breed dogs, 2 Labrador retriever and 1 each of Boxer, German shepherd, Pitt Bull, Rottweiler and Yorkshire terrier. Mean age was  $9.8 (\pm 2.2)$  years and mean weight was  $27.2 (\pm 10.4)$  kg. There were 8 female (n=4 spayed) and 4 males (n=1 neutered) dogs. HSA occurred in the spleen as primary site in 11 dogs; 10 of them presented with hemoperitoneum because of splenic rupture. One dog had a subcutaneous HSA.

All dogs underwent surgery, consisting of splenectomy and removal of the subcutaneous tumour according to cancer location. Histopathological evaluation revealed clean surgical margins in the subcutaneous HSA; surgical margins were deemed not assessable for dogs presenting with visceral rupture.

According to WHO, 11 dogs had stage II disease, and 1 had stage I disease. The dogwith stage I disease had a splenic HSA.

The mean time from surgery to initial MTDC administration was 25.0 (± 12.1) days. 236 Nine dogs received doxorubicin as single agent and 3 dogs received a combination of 237 doxorubicin and dacarbazine. The median number of doxorubicin cycles was 4 (range, 2 238 239 to 5 cycles) and all dogs received a starting dose of doxorubicin of 30 mg/m<sup>2</sup>. Chemotherapy dose reduction was performed in 2 dogs receiving single agent 240 241 doxorubicin; this was performed at the clinician's discretion due to haematological and/or gastrointestinal toxicity: one dog had 10% and one had 20% dose reduction. The 242 243 median total dose of doxorubicin was 120 mg/m<sup>2</sup> (range, 60 to 180). In the three dogs receiving doxorubicin and dacarbazine, the protocol was designed as previously 244 reported.14 Cases' data are summarized in Table 1. 245

246

### 247 Clinical outcome

Three (30%) out of the 10 dogs included in Group 1 (MTDC-MC) developed metastatic disease after 119, 151 and 460 days, respectively. Metastases were found in the peritoneum (n=2) and liver and lung (n=1). The two dogs with metastases to the peritoneum developed haemoabdomen.

252

Nine (75%) of the 12 dogs included in Group 2 (MTDC) developed metastatic disease after a median of 134 days (range, 89 to 174 days). Metastases were found in lung (n=3), peritoneum (n=2), liver (n=2), lung and brain (n=1) and lung, stomach and liver (n=1). The two dogs with metastases to the peritoneum developed haemoabdomen.

Overall, median TTM was significantly longer for dogs receiving MTDC-MC compared
to those receiving MTDC only (not reached versus 150 days, respectively; P=0.028;
Figure 1).

262	Six (60%) out of the 10 dogs included in Group 1 (MTDC-MC) were dead at the end of
263	the study. Three (27.2%) dogs with splenic HSA died as a result of disease progression
264	after 152, 191 and 487 days. Three dogs with splenic HSA died of tumour-unrelated
265	causes after 165, 292 and 730 days, respectively, with no evidence of tumour recurrence
266	or metastasis. One dog (splenic HSA) was lost to follow-up after 680 days from the
267	diagnosis; at the last visit this dog had no evidence of macroscopic disease.
268	Three dogs (osseous, $n=1$ , and splenic, $n=2$ ) were still alive with no evidence of disease
269	after 311, 640 and 1280 days, respectively.

271 Ten (83.3%) out of the 12 dogs in Group 2 (MTCD) were dead at data analysis closure:

272 9 (75%) died as a result of HSA progression with a median survival time of 156 days

273 (range, 97 to 341 days). Of these 9 dogs, 7 had splenic stage II HSA, 1 had splenic stage

274 I HSA, and one had subcutaneous stage II HSA. The remaining dog (splenic stage II

275 HSA) died 803 days after the diagnosis because of tumour-unrelated causes.

Two dogs with splenic HSA were still alive with no evidence of disease at 437 and 608

277 days, respectively.

278

Overall, dogs receiving MTDC followed by MC had a significantly longer median ST
than those receiving MTDC only (not reached versus 168 days, respectively; P=0.030;
Figure 2).

282

## 283 Toxicity

During MTDC, neutropenia occurred in 4 (40%) dogs in Group 1. One dog had one episode of grade 1 neutropenia, 2 dogs had one episode of grade 2 neutropenia, whereas 1 dog had 2 episodes of grade 2 neutropenia. In all dogs haematological toxicities resolved without sequel.

In Group 2, 1 (8.3%) dog developed 2 episodes of grade 4 non-febrile neutropenia, and 288 1 (8.3%) dog developed one episode of grade 2 anaemia. 289

Gastrointestinal toxicity was the second most common adverse event in both groups, 290 291 and consisted of vomiting, diarrhoea and decreased appetite of mild to moderate severity. Gastrointestinal toxicity of grade 2 occurred in 2 (20%) dogs in Group 1. 292 293 These dogs had no concurrent episodes of haematological toxicity. In Group 2, 3 (25%) dogs developed gastrointestinal toxicity: 1 dog had one episode of grade 3 anorexia and 294 295 2 dogs had 1 episode of grade 2 vomiting (1 concurrently had grade 2 anaemia).

The overall frequency of MTDC related side effects did not differ between groups 296 (Table 2). 297

298

During MC, 4 (40%) dogs developed gastrointestinal, haematological and/or urothelial 299 adverse events. Two dogs developed grade 2 sterile haemorrhagic cystitis after 180 and 300 301 470 days, respectively; in both cases cyclophosphamide was discontinued and chlorambucil was started; cystitis resolved within 4 weeks in both cases. One dog 302 303 developed grade 1 diarrhoea and in one case grade 1 vomiting and diarrhoea occurred 304 simultaneously. Gastrointestinal signs resolved with symptomatic treatment and did not 305 recur.

306

#### Discussion 307

The treatment of HSA continues to be extremely challenging in veterinary oncology. 308 Unfortunately, little progress has been made over the years, and prognosis for dogs with 309 310 HSA is poor as a result of the aggressive nature of the disease, leading to invasion of 311 nearby organs and vessels, early metastasis and limited treatment options providing durable disease control. Surgery is designed to remove all macroscopic tumours and 312 313 prevent further risk of acute haemorrhage, but is considered purely palliative. The 13

addition of chemotherapy in an effort to treat microscopic disease has been documented 314 to provide a modest improvement in outcome, with reported median survival times in 315 the range of 6-8 months and less than 10% of dogs being alive at 12 months.<sup>1,2</sup> 316 317 The "cell kill" paradigm associated with MTDC has been successful in the treatment of human and canine haematological neoplasia, but unfortunately this has not provided 318 319 long-lasting responses in the majority of advanced solid tumours.<sup>21</sup> Failure of MTDC

may be multifactorial, being attributable to the heterogeneity of cancer cells, genetic 320 321 make-up, and the influence of tumour microenvironment, thereby giving rise to treatment resistance.<sup>21</sup> Based on the above, the Gatenby's hypothesis of controlling 322 tumour growth instead of trying to eradicate it may become a more rational strategy.<sup>27</sup>

Maintenance therapy refers to a treatment that is given to avoid disease progression 324 after the cancer has been successfully controlled with the initial therapy.<sup>21</sup> 325

323

An effective maintenance therapy should accomplish good patient tolerability, lack of 326 327 cumulative toxicities, and cost-effectiveness. Maintenance therapy may consist of "continuation" therapy where one drug of the initial therapy is continued after the 328 induction phase of the protocol, or of "switch" maintenance in which a new agent is 329 330 introduced.21,28

Switch maintenance has been recently investigated in canine stage I-II splenic HSA by 331 332 administering the tyrosine kinase inhibitor toceranib phosphate. Toceranib mainly targets the stem cell factor receptor KIT, platelet derived growth factor receptor and 333 vascular endothelial growth factor receptor (VEGFR), which are typically expressed by 334 canine HSA.29 335

336 As in our study, the switch maintenance was administered in the microscopic disease 337 setting after completion of doxorubicin MTDC. Unfortunately, disease-free interval nor ST were improved when comparing dogs receiving or not receiving maintenance 338 toceranib.29 339

It has become progressively clear that the endothelial cell compartment is an attractive target for anticancer therapy as a result of the evident importance of the tumour vasculature for sustaining tumour growth and metastasis. Also, the endothelial cells are sensitive to the action of conventional cytotoxic drugs, including cyclophosphamide, if the dosing regimen is altered to the so-called anti-angiogenic scheduling.<sup>15</sup>

345 In a previous study, dogs with stage II HSAAS receiving an oral adjuvant therapy 346 consisting of alternating low-dose daily cyclophosphamide and etoposide in combination with piroxicam had comparable survival times to historical controls treated 347 with conventional doxorubicin chemotherapy.<sup>19</sup> Starting from the promising results of 348 349 the mentioned study, Based on the promising results obtained by MC in the treatment of surgically-removed canine HSA,<sup>19</sup>-we hypothesised that outcome might be improved, if 350 a MC schedule is to be administered after MTDC as a consolidation strategy. To this 351 352 end, we retrospectively compared HSA dogs receiving MTDC versus MTDC followed by MC in the microscopic setting. Beside cyclophosphamide and NSAID, thalidomide 353 was added to this combination in the majority of dogs. 354

The results obtained in the current study <u>document suggest</u> an advantage of <u>the addition</u> of maintenance MC over MTDC alone in terms of metastatic control and survival. Indeed, dogs undergoing chemo-switch after dose-intense chemotherapy had a significantly longer TTM and ST compared to dogs receiving MTDC, suggesting that chemo-switch improves long-term tumour control in biologically aggressive canine HSA. These results <u>can-may</u> be explained by the following considerations.

The use of continuous, low-dose cyclophosphamide exerts potent anti-angiogenic properties through the inhibition of proliferation and/or induction of apoptosis of activated endothelial cells, selective inhibition of migration of endothelial cell, increase in the expression of thrombospondin-1, and sustained decrease in levels and viability of bone marrow-derived endothelial progenitor cells.<sup>15</sup> Moreover, it has been shown that metronomic cyclophosphamide can also target the immune system by activating or restoring its antitumor properties, particularly through the inhibition of T regulatory lymphocytes and enhance the cytotoxic T lymphocytes response.<sup>30,31</sup>

Non-selective NSAIDs and COX-2 selective inhibitors such as piroxicam and
 meloxicam are effective in counteracting tumour angiogenesis, by boostering the effect
 of cyclophosphamide.<sup>32-34</sup>

Alongside its teratogenic effect, thalidomide is a potent inhibitor of angiogenesis
through inhibition of VEGF, basic fibroblastic growth factor, and tumour necrosis factor
alpha, and may play a role in anti-angiogenic strategies.<sup>35</sup>

A recent study has suggested that the combination of MTDC and MC may be superior 375 to MTDC alone in the treatment of canine splenic HSA in the early follow-up period,<sup>20</sup> 376 however survival times were modest compared to the group receiving MTDC alone and, 377 importantly, these did not differ substantially from the published literature.<sup>1,2</sup> In the 378 aforementioned study, 13 dogs with splenic HSA received doxorubicin and MC either 379 sequentially (chemo-switch; n=6) or concurrently (n=7). Median survival time for these 380 381 dogs was 4.3 months, and median duration of treatment was 56 days; it was 382 hypothesised that metastatic disease rapidly progressed after chemotherapy was interrupted for whichever reason.<sup>20</sup> In the current study group, the use of MC 383 384 significantly improved outcome, and it may be hypothesised that the difference between our study and Wendelburg's study may be due to the use of the potent antiangiogenic 385 drug thalidomide or to the continuous use of MC. 386

While MTDC can serve to de-bulk HSA by directly targeting the cancer cells, maintenance MC may disrupt crucial angiogenic pathways, impeding the inevitable

389 rebound and regrowth, ultimately translating into significant therapeutic benefits.

390 In agreement with previous studies, MC was well tolerated, and side effects were

mainly gastro-intestinal and of mild severity.<sup>17,18</sup> Haemorrhagic cystitis occurred in 2

dogs, most likely as a consequence of prolonged treatment with cyclophosphamide; however gastrointestinal and haematological adverse event could have also been due to transient and undiagnosed comorbidities and not related to MC.

395 Limitations of this study include its retrospective nature, the low number of cases, the different tumour site origin, the variability of chemotherapy protocols used in the 396 397 MTDC phase and the lack of necropsies. Five dogs received a combination of 398 doxorubicin and dacarbazine, which has recently demonstrated encouraging results 399 providing an increase in the chances of survival for biologically aggressive canine HSA.14,36 Nevertheless, in the present series dogs receiving doxorubicin and 400 dacarbazine were equally distributed among groups, thereby rendering unlikely the 401 chance of having improved outcome in one group only. Dogs' features and possible 402 outcome variables were homogeneously distributed between groups with the exception 403 404 of sex: male dogs were more common in Group 1 whereas females were more common 405 in Group 2. Although this finding is likely to be a bias due to the small sample size, we 406 cannot exclude that the small number of dogs included in this study may have 407 contributed to reach significance for the other variables analysed.

408 This finding, although statistically significant, is likely to be a bias due to the small

409 sample size, and any other explanation might only be speculative.  $\frac{1}{2}$ 

410 Finally, it must be acknowledged that 3 dogs treated with MTDC were censored

- 411 <u>belatedly (after 437, 608 and 803 days), compared to 6 dogs treated with MTDC and</u>
- 412 MC, and among them 3 were censored early (after 165, 292 and 311 days). While this
- 413 may reflect a better outcome, as fewer dogs died due to HSA in Group 1 compared to
- 414 Group 2, it also could have biased the results, as early deaths due to tumour-unrelated
- 415 <u>causes may strongly influence statistics.</u>

416

**Formattato:** Giustificato, Destro 1 cm, SpazioDopo: 0 pt, Interlinea: doppia

Formattato: Tipo di carattere: (Predefinito) Times New Roman, 12 pt, Inglese (Regno Unito)
Formattato: Tipo di carattere: Times New Roman, 12 pt, Inglese (Regno Unito)
Formattato: Tipo di carattere: (Predefinito) Times New Roman, 12 pt
Formattato: Tipo di carattere: (Predefinito) Times New Roman, 12 pt
Formattato: Tipo di carattere: (Predefinito) Times New Roman, 12 pt

417	To conclude, maintenance MC is well tolerated and may prolong TTM and survival
418	time in dogs with biologically aggressive HSA with negative staging after completion of
419	MTDC. Although the role of thalidomide in the treatment of HSA needs further studies,
420	it is possible that this drug used in combination with standard MC plays an important
421	role in controlling the metastatic process of biologically aggressive canine HSA.
422	Prospective studies with larger number of patients are required to confirm these
423	findings.
424	

- 425
- 426 References
- 427

428	1. Thamm DH. Hemangiosarcoma. In: Withrow & MacEwen's Small Animal
429	Clinical Oncology, 5th ed., SJ Withrow, DM Vail and RL Page eds., St Louis,
430	Saunders Elsevier, 2013: 679-688

- 431 2. Smith AN. Hemangiosarcoma in dogs and cats. *Veterinary Clinics of North* 432 *America: Small Animal Practice* 2003; 33: 533-552
- 3. Schultheiss PC. A retrospective study of visceral and nonvisceral
  hemangiosarcoma and hemangiomas in domestic animals. *Journal of Veterinary Diagnostic Investigation* 2004; 16: 522-526
- 4. Shiu KB, Flory AB, Anderson CL, Wypij J, Saba C, Wilson H, Kurzman I and
  Chun R. Predictors of outcome in dogs with subcutaneous or intramuscular
  hemangiosarcoma. *Journal of the American Veterinary Medical Association*2011; 238: 472-479
- 5. Ward H, Fox LE, Calderwood-Mays MB, Hammer AS and Couto CG.
  Cutaneous hemangiosarcoma in 25 dogs: a retrospective study. *Journal of Veterinary Internal Medicine* 1994; 8: 345-348

443	6.	Ogilvie GK, Powers BE, Mallinckrodt CH and Withrow SJ. Surgery and
444		doxorubicin in dogs with hemangiosarcoma. Journal of Veterinary Internal
445		Medicine 1996; 10: 379-384
446	7.	Wiley JL, Rook KA, Clifford CA, Gregor TP and Sorenmo KU. Efficacy of
447		doxorubicin-based chemotherapy for non-resectable canine subcutaneous
448		haemangiosarcoma. Veterinary and Comparative Oncology 2010; 8: 221-233
449	8.	Payne SE, Rassnick KM, Northrup NC, Kristal O, Chretin JD, Cotter SM,
450		Kintzer P, Frimberger AE, Morrison-Collister KE, Wood CA and Moore AS.
451		Treatment of vascular and soft-tissue sarcomas in dogs using an alternating
452		protocol of ifosfamide and doxorubicin. Veterinary and Comparative Oncology
453		2003; 1:171-179
454	9.	Hammer AS, Couto CG, Filppi J, Getzy D and Shank K. Efficacy and toxicity of
455		VAC chemotherapy (vincristine, doxorubicin, and cyclophosphamide) in dogs
456		with hemangiosarcoma. Journal of Veterinary Internal Medicine 1991; 5: 160-
457		166
458	10	. Sorenmo KU, Jeglum KA and Helfand SC. Chemotherapy of canine
459		hemangiosarcoma with doxorubicin and cyclophosphamide. Journal of

Veterinary Internal Medicine 1993; 7: 370-376 460

- 11. Bulakowski EJ, Philibert JC, Siegel S, Clifford CA, Risbon R, Zivin K and 461 Cronin KL: Evaluation of outcome associated with subcutaneous and 462 intramuscular hemangiosarcoma treated with adjuvant doxorubicin in dogs: 21 463 cases (2001-2006). Journal of the American Veterinary Medical Association 464 2008; 233: 122-128 465
- 12. Kim SE, Liptak JM, Gall TT, Monteith GJ and Woods JP. Epirubicin in the 466 adjuvant treatment of splenic hemangiosarcoma in dogs: 59 cases (1997-2004). 467 Journal of the American Veterinary Medical Association 2007; 231: 1550-1557 468

**.**..

469	15. Sorenmo KU, Baez JL, Chilord CA, Mauldin E, Overley B, Skorupski K,	
470	Bachman R, Samluk M and Shofer F. Efficacy and toxicity of a dose-intensified	
471	doxorubicin protocol in canine hemangiosarcoma. Journal of Veterinary	
472	Internal Medicine 2004; 18: 209-213	
473	14. Finotello R, Stefanello D, Zini E and Marconato L. Comparison of doxorubicin-	
474	cyclophosphamide with doxorubicin-dacarbazine for the adjuvant treatment of	
475	canine hemangiosarcoma. Veterinary and Comparative Oncology 2015: 1-11,	
476	doi: 10.1111/vco.12139	
477	15. Maiti R. Metronomic chemotherapy. Journal of Pharmacology and	
478	Pharmacotherapeutics 2014; 5: 186-192	
479	16. Kareva I, Waxman DJ and Klement GL. Metronomic chemotherapy: An	
480	attractive alternative to maximum tolerated dose therapy that can activate anti-	
481	tumor immunity and minimize therapeutic resistance. Cancer Letters 2015; 358:	
482	100-106	
483	17. Biller B. Metronomic chemotherapy in veterinary patients with cancer:	
484	rethinking the targets and strategies of chemotherapy. The Veterinary Clinics of	
485	North America. Small Animal Practice 2014; 44: 817-829	
486	18. Marchetti V, Giorgi M, Fioravanti A, Finotello R, Citi S, Canu B et al. First-line	Formattato: Italiano (Italia)
487	metronomic chemotherapy in a metastatic model of spontaneous canine	
488	tumours: a pilot study. Investigational New Drugs 2012; 30: 1725-1730	
489	19. Lana S, U'ren L, Plaza S, Elmslie R, Gustafson D, Morley P et al. Continuous	
490	low-dose oral chemotherapy for adjuvant therapy of splenic hemangiosarcoma	
491	in dogs. Journal of Veterinary Internal Medicine 2007; 21: 764-769	
492	20. Wendelburg KM, Price LL, Burgess KE, Lyons JA, Lew FH and Berg J.	
493	Survival time of dogs with splenic hemangiosarcoma treated by splenectomy	

**a** a

т,

494	with or without adjuvant chemotherapy: 208 cases (2001-2012). Journal of the
495	American Veterinary Medical Association 2015; 247: 393-403.
496	21. Malik PS, Raina V and André N. Metronomics as maintenance treatment in
497	oncology: time for chemo-switch. Frontiers in Oncology 2014; 4: 76
498	22. Owen LN, ed. TNM classification of tumours in domestic animals. Geneva
499	(Switzerland): World Health Organization; 1980
500	23. Leach TN, Childress MO, Greene SN, Mohamed AS, Moore GE, Schrempp DR,
501	et al. Prospective trial of metronomic chlorambucil chemotherapy in dogs with
502	naturally occurring cancer. Veterinary and Comparative Oncology 2012; 10:
503	102-112
504	24. Marconato L, Buchholz J, Keller M, Bettini G, Valenti P and Kaser-Hotz B.
505	Multimodal therapeutic approach and interdisciplinary challenge for the
506	treatment of unresectable head and neck squamous cell carcinoma in six cats: a
507	pilot study. Veterinary and Comparative Oncology 2013; 11: 101-112
508	25. Vargesson N. Thalidomide-Induced Teratogenesis: History and Mechanisms.
509	Birth Defects Research. Part C, Embryo Today: Reviews 2015; 105: 140-156
510	26. Veterinary Co-operative Oncology Group. Veterinary Co-operative oncology
511	group- common terminology criteria for adverse events (VCOG-CTCAE)
512	following chemotherapy or biological antineoplastic therapy in dogs and cats
513	v1.0. Veterinary and Comparative Oncology 2004; 2: 194-213
514	27. Gatenby RA, Silva AS, Gillies RJ and Frieden BR. Adaptive therapy. Cancer
515	Research 2009; 69: 4894-4903
516	28. Gerber DE and Schiller JH. Maintenance chemotherapy for advanced non-small-
517	cell lung cancer: new life for an old idea. Journal of Clinical Oncology 2013;
518	<b>31</b> : 1009-1020

521	chemotherapy for canine splenic hemangiosarcoma. BMC Veterinary Research
522	2015; 11: 131
523	30. Pasquier E, Kavallaris M and André N. Metronomic chemotherapy: new
524	rationale for new directions. Nature Reviews. Clinical Oncology 2010; 7: 455-
525	465
526	31. Burton JH, Mitchell L, Thamm DH, Dow SW and Biller BJ. Low-dose
527	cyclophosphamide selectively decreases regulatory T cells and inhibits
528	angiogenesis in dogs with soft tissue sarcoma. Journal of Veterinary Internal
529	Medicine 2011; <b>25</b> : 920-926
530	32. Fischer SM, Hawk ET and Lubet RA. Coxibs and other nonsteroidal anti-
531	inflammatory drugs in animal models of cancer chemoprevention. Cancer
532	Prevention Research (Philadelphia, Pa.) 2011; 4: 1728-1735
533	33. Iwase N, Higuchi T, Gonda T, Kobayashi H, Uetake H, Enomoto M et al. The
534	effect of meloxicam, a selective COX-2 inhibitor, on the microvasculature of
535	small metastatic liver tumors in rats. Japanese Journal of Clinical Oncology
536	2007; <b>37</b> : 673-678
537	34. Mohammed SI, Craig BA, Mutsaers AJ, Glickman NW, Snyder PW, de Gortari
538	AE et al. Effects of the cyclooxygenase inhibitor, piroxicam, in combination
539	with chemotherapy on tumor response, apoptosis, and angiogenesis in a canine
540	model of human invasive urinary bladder cancer. Molecular Cancer
541	Therapeutics 2003; 2: 183-188
542	35. Rebuck JA and Fish DN. Thalidomide revisited. The AIDS Reader 1998; 8: 7–9

29. Gardner HL, London CA, Portela RA, Nguyen S, Rosenberg MP, Klein MK, et

al. Maintenance therapy with toceranib following doxorubicin-based

519

543	36. Dervisis NG, Dominguez PA, Newman RG, Cadile CD and Kitchell BE.
544	Treatment with DAV for advanced-stage hemangiosarcoma in dogs. Journal of
545	the American Animal Hospital Association 2011; 47: 170-178
546	
547	
548	

# 550 Captions to figures:

E	E	1
5	5	т

- 552 Figure 1: Time to metastases for dogs treated with MTDC-MC (dots) and MTDC (line).
- 553 In the MTDC-MC group, dogs had a longer time to metastases (not reached versus 150
- 554 days, respectively; P=0.028).
- 555
- 556 Figure 2: Survival time for dogs treated with MTDC-MC (dots) and MTDC (line). In
- 557 the MTDC-MC group, dogs had a longer survival time (not reached versus 168 days,
- 558 respectively; P=0.030).
- 559