Comparison of two melphalan protocols and evaluation of outcome and prognostic factors in multiple myeloma in dogs

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Background: Multiple myeloma (MM) in dogs typically is treated with melphalan. A daily melphalan dosing schedule reportedly is well tolerated and associated with favorable outcome. Although anecdotaly a pulse dose regimen has resulted in successful responses, little long-term outcome and safety data is available regarding this dosing regimen for dogs with MM.

Hypothesis/objectives: (1) To compare outcome and adverse event profiles between pulse dose and daily dose melphalan schedules and (2) to report prognostic factors in dogs with MM treated with melphalan. We hypothesized that both protocols would have similar outcomes and tolerability.

Animals: Thirty-eight client-owned dogs diagnosed with MM receiving pulse dose (n = 17) or daily dose (n = 21) melphalan.

Methods: Retrospective cohort study assessing outcome and adverse events in dogs receiving either protocol. Risk factors were evaluated for their prognostic relevance.

Results: Both regimens were well tolerated and similarly effective, with an overall median survival time of 930 days. Renal disease and neutrophil-to-lymphocyte ratio (NLR) were negative prognostic factors, whereas hypercalcemia and osteolytic lesions were not prognostic factors in this study population.

Conclusions and Clinical Importance: Positive results support the use of either dosing regimen for the treatment of dogs with MM, and renal disease and NLR were negative prognostic factors. Prospective, controlled, and randomized studies are warranted to confirm these findings.

KEYWORDS
daily dose, dogs, pulse dose, neutrophil lymphocyte ratio, renal

INTRODUCTION

Multiple myeloma (MM) is a systemic proliferation of malignant plasma cells or their precursors.1 In dogs, MM accounts for approximately 8% of all hematopoietic malignancies.2 A diagnosis of MM in dogs typically is made by identification of bone marrow plasmacytosis, myeloma proteins in the serum or urine, and osteolytic lesions.3 Although visceral organ involvement can aid in the diagnosis of MM in other species, such as in cats,3-6 intra-abdominal infiltration does not occur.
commonly in dogs, with its frequency and effects on prognosis not previously reported.\(^1\)

Melphalan is a cell cycle phase-nonspecific alkylating agent, often given in combination with prednisone to treat dogs with MM.\(^1\)\(^2\) A therapeutic regimen of daily melphalan and prednisone was associated with a median survival time of 540 days and overall response rate of 92%.\(^2\) An alternative, pulse dose schedule also has been used to treat dogs with MM,\(^1\) especially when delayed thrombocytopenia has limited continuous daily dose therapy.\(^1\) Although successful responses have been obtained using the pulse dose protocol to treat dogs with MM (personal communication, D. Vail, August, 2017), long-term response and safety data currently are lacking.

Hypercalcemia, osteolytic lesions, and Bence-Jones proteinuria were reported to be negative prognostic factors for MM in dogs.\(^2\) Nonazotemic dogs also had a longer survival compared with azotemic dogs, but this difference was not statistically significant.\(^2\) In contrast, studies of MM in humans have not consistently found hypercalcemia, osteolytic lesions, and Bence-Jones proteinuria to be prognostic\(^7\)\(^8\) and instead use risk stratification models that primarily rely on cytogenetics, gene expression profiling, the International Staging System, and serum lactate dehydrogenase activity as prognostic markers.\(^9\)\(^11\) Other studies have evaluated factors such as renal disease,\(^7\)\(^8\)\(^12\)\(^18\) neutrophil-to-lymphocyte ratio (NLR),\(^19\)\(^–\)\(^24\) platelet-to-lymphocyte ratio (PLR),\(^23\)\(^24\) and anemia,\(^25\)\(^–\)\(^27\) which have negatively affected outcome. Several studies in veterinary oncology have evaluated NLR as a prognostic factor in dog,\(^28\)\(^–\)\(^31\) but no studies, to the authors’ knowledge, have evaluated NLR or PLR in dogs with MM. Additionally, factors such as anemia, neutropenia, thrombocytopenia, and abdominal involvement, have not been specifically evaluated for prognostic relevance in dogs with MM.

The primary objective of our study was to compare outcome and adverse event profiles between pulse dose and daily dose melphalan protocols in dogs with MM. Our secondary objective was to report prognostic factors. We hypothesized that both melphalan-based protocols would be associated with similar outcomes and be tolerated well.

2 | MATERIAL AND METHODS

2.1 | Study design

Retrospective cohort study performed at the School of Veterinary Medicine at the University of Wisconsin-Madison.

2.2 | Cohort identification

A search of medical records of dogs diagnosed with MM that had received melphalan at the University of Wisconsin-Madison Veterinary Care Hospital between January 1998 and April 2016 was performed. Additional cases were contributed by other veterinary medical oncologists during the same time period in response to a call posted on the ACVIM listserv. Dogs were included in the study if they were diagnosed with MM and received PO melphalan, either pulse dose or daily dose. A diagnosis of MM was reached based on evidence of ≥2 of the following criteria: bone marrow plasmacytosis, osteolytic lesions, other organ involvement, and presence of myeloma proteins in blood or urine. Dogs presenting solely with polyostotic lytic lesions with cytologically or histologically diagnosed plasma cell neoplasia from ≥1 of these lesions also were included. Dogs were excluded if they were lost to follow-up immediately after initiation of therapy (ie, did not have at least 1 clinical reevaluation after initiating melphalan treatment). Required follow-up information included CBC, physical examination, and history. Information provided by primary care veterinarians was included.

2.3 | Data collection

Presenting information obtained from the medical records included patient demographics (breed, age, sex, and neuter status), clinical signs and physical examination findings. Results of diagnostic tests (CBC, biochemistry panel, urinalysis, bone marrow aspiration [BMA], thoracic and abdominal radiographs, abdominal ultrasound examination, cytology or histopathology, and immunoglobulin type as determined by radial immunodiffusion) were recorded. Other collected information included other chemotherapeutic agents administered for treatment of MM, response to treatment (complete response [CR], partial response, stable disease, or progressive disease [PD]), remission status, adverse events, concomitant medications and concurrent diseases, follow-up visits (dates, tests, and results), duration of remission, date of relapse, method to determine relapse, rescue chemotherapy protocols (if received), cause and date of death, and necropsy results (if performed). Categories for presenting clinical signs and clinicopathologic abnormalities (based on the performed diagnostic tests) are described in Table 1. All dogs with renal disease had resolution of azotemia after initiation of treatment, and their renal disease was retrospectively graded (solely by serum creatinine concentration) according to International Renal Interest Society (IRIS) guidelines for acute kidney injury (AKI).\(^32\)\(^–\)\(^35\)

Outcome was reported as: overall survival time (OST), defined as the interval from diagnosis to death; progression-free survival (PFS), defined as the interval from treatment initiation to onset of PD; disease-free interval (DFI), defined as the interval from a CR to relapse; survival from remission (SFR), defined as the interval from a CR until death; and, time to remission (TR), defined as the interval from treatment initiation to a CR. Dogs that died of a cause other than MM, were lost to follow-up or were still alive at the end of data collection were censored. Dogs that remained alive at the end of data collection were censored as of the last date they were reported to be alive.

The NLR was defined as the absolute neutrophil count divided by the absolute lymphocyte count. The PLR was defined as the platelet count divided by the absolute lymphocyte count.\(^24\) All laboratory variables used to calculate NLR and PLR were obtained within 4 weeks before or after the diagnosis of MM, in accordance with a study of humans with MM.\(^24\) A contemporaneous control group was randomly selected from hospital dogs and used for the receiver operating characteristic (ROC) curve.\(^24\) Exclusion criteria for the control group included the following: acute or chronic infections, acute or chronic liver disease, other concomitant malignancies, thrombocytopenia, inflammation-promoting diseases (eg, osteoarthritis, colitis), or dogs receiving anti-inflammatory medications for >2 weeks.
TABLE 1 Descriptive baseline data of dogs in pulse dose cohort, daily dose cohort, and all dogs combined

| Variable                     | Pulse cohort | Daily cohort | All dogs |
|------------------------------|--------------|-------------|---------|
| Number of dogs               | 17           | 21          | 38      |
| Median age (years)           | 9 (range 5–16) | 9 (range 4–12) | 9 (range 4–16) |
| Sex                          |              |             |         |
| Male intact                  | 1            | 3           | 4       |
| Male neutered                | 8            | 12          | 20      |
| Female intact                | 0            | 0           | 0       |
| Female spayed                | 8            | 6           | 14      |
| Female: Male                 | 0.9:1        | 0.4:1       | 0.6:1   |
| Presenting clinicopathologic abnormalities | | | |
| Lethargy/weakness            | 5            | 15          | 20/38 (53%) |
| Polyuria/polydipsia          | 8            | 6           | 14/38 (37%) |
| Inapetence                   | 3            | 12          | 15/38 (39%) |
| Weight loss                  | 5            | 7           | 12/38 (32%) |
| Ocular abnormalities         | 4            | 3           | 7/38 (18%) |
| Laminence/pain               | 3            | 3           | 6/38 (16%) |
| Nausea/vomiting              | 2            | 4           | 6/38 (16%) |
| Diarrhea                     | 3            | 3           | 3/38 (8%) |
| Paraparesis                  | 1            | 2           | 3/38 (8%) |
| Fever                        | 2            | 1           | 3/38 (8%) |
| Vision loss                  | 2            | 1           | 3/38 (8%) |
| Peripheral lymphadenopathy   | 1            | 1           | 2/38 (5%) |
| Cutaneous lesions            | 1            | 1           | 2/38 (5%) |
| CNS abnormalitiesa            | 0            | 0           | 0/38 (3%) |

TABLE 1 (Continued)

| Variable                     | Pulse cohort | Daily cohort | All dogs |
|------------------------------|--------------|-------------|---------|
| Hypertension                 | 6            | 6           | 12/38 (32%) |
| Increased M                  | 14           | 17          | 31/31 (100%) |
| Monoclonal                   | 12           | 13          | 25/31 (81%) |
| Biclonal                     | 2            | 4           | 6/31 (19%) |
| IgA                          | 2            | 9           | 11/14 (79%) |
| IgG                          | 1            | 2           | 3/14 (21%) |

* Dogs with abnormal mentation, cranial nerve deficits or seizure activity were categorized as having CNS abnormalities.

Overall response rate and biologic response rate were evaluated using an adaptation of the International Uniform Response Criteria for MM36 in the majority of cases; in 2 dogs responses were determined from radiographic changes (ie, osteolytic lesions). Survival rates were calculated using the indirect method as previously described.37 Adverse events and presenting cytopenias were graded retrospectively according to the Veterinary Cooperative Oncology Group Common Terminology Criteria for Adverse Events v1.1,38 based on client history, physical examination, CBC, biochemistry profile, and urinalysis. Thrombocytopenia was further categorized by duration of melphalan therapy into 4 arbitrary groups: <6 months, 6 months-1 year, 1–2 years, and 2–4 years.

2.4 Statistical analysis

All statistical analyses were performed using commercial statistical software (GraphPad Prism 7.03. GraphPad Software Inc, La Jolla, California; R studio v1.0.132. R Studio®, Boston, Massachusetts). Outcome analysis was conducted using Kaplan-Meier curves (GraphPad Prism 7.03. GraphPad Software Inc). The log-rank test (R studio v1.0.132. R Studio®) was used to compare outcome between treatment cohorts as well as to assess the prognostic relevance of hypercalcemia, osteolytic lesions, concurrent hypercalcemia, and osteolytic lesions, renal disease, high NLR, high PLR, anemia, thrombocytopenia, neutropenia, proteinuria, hypoalbuminemia, hyperviscosity syndrome, abdominal involvement, and cyclophosphamide as a concurrent chemotherapeutic given
within 10 days of melphalan initiation. A Fisher’s exact test (R studio v1.0.132, R Studio™) was performed to assess differences in frequency of potential prognostic factors between treatment cohorts. Optimal cut-off points for NLR and PLR as predictors of OST were based on the ROC curve (GraphPad Prism 7.03, GraphPad Software Inc), as previously described. Multivariate and univariate analyses deliberately were not performed because of the small sample size. Values of $P < .05$ were considered statistically significant.

### RESULTS

#### 3.1 | Patient signalment and cohort assignment

Sixty-one dogs were identified in the initial search, and 38 dogs met the inclusion criteria. Golden Retrievers ($n = 8$), Labrador Retrievers ($n = 6$), mixed breed dogs ($n = 5$), and Doberman Pinschers ($n = 4$) were the most common breeds. There also were 2 German Shepherds and 1 each of the following breeds: Pug, Siberian Husky, Airedale Terrier, Rottweiler, Bassett Hound, Bearded Collie, French Bulldog, American Bulldog, Standard Poodle, Boxer, Dalmatian, Samoyed, and Pembroke Welsh Corgi.

#### 3.2 | Clinical and clinicopathologic findings

The presenting clinical and clinicopathologic findings are summarized in Table 1. Concurrent malignancies are summarized in Table 2.

#### 3.3 | Diagnostic imaging

Abdominal ultrasound examination was performed at the time of diagnosis in 28 dogs. Sixteen dogs (57%) had cytologically confirmed abdominal involvement (Table 1). Among the 14 dogs with confirmed splenic involvement, ultrasonographic findings included splenomegaly ($n = 4$), hypoechoic nodules ($n = 3$), normal appearance ($n = 4$), mottled ($n = 1$), mottled with hypoechoic nodules ($n = 1$), and splenic mass effect ($n = 1$). Among the 5 dogs with confirmed liver involvement, ultrasonographic findings included hepatomegaly ($n = 2$), hepatic mass effect ($n = 1$), mottled with hepatomegaly ($n = 1$) and normal appearance ($n = 1$). In the 1 dog with jejunal lymph node involvement, the jejunal lymph node was mildly enlarged and hypoechoic. Eleven dogs did not have cytologic or histologic confirmation of intra-abdominal plasma cell neoplasia. Of these 12 dogs, 7 dogs had a normal appearance to the intra-abdominal organs whereas 5 had ultrasonographic changes suggestive of abdominal involvement including hypoechoic splenic nodules ($n = 2$), splenomegaly ($n = 1$), and hypoechoic liver nodules ($n = 2$).

Thoracic radiographs were performed at the time of diagnosis in 35/38 (92%) dogs. Thoracic radiographic findings included normal thorax ($n = 16$), osteolytic lesions ($n = 13$) and 1 each of the following: pulmonary nodules, hepatomegaly, alveolar pattern consistent with pneumonia or fluid overload, sternal lymphadenopathy, enlarged cardiac silhouette suggestive of compensated cardiomyopathy, and mild unstructured pulmonary pattern consistent with previous insult.

Radiographically diagnosed osteolytic lesions were documented in 16/38 (42%) dogs. Ribs, vertebrae, and dorsal spinous processes (13/16 [81%]) were the most common locations for lytic lesions, whereas long bones were less frequently affected (3/16 [19%]). Notably, only dogs presenting with lameness had long-bone radiography performed. Contrast myelography, performed in 2 dogs that were presented with paraparesis, identified extradural compression at the level of T12/T13 and L5. In both dogs, hemilaminectomies with subsequent histologic examination identified plasma cell tumor and resulted in resolution of paraparesis. Cytologic examination of osteolytic lesions (in the proximal tibia, rib and T12/T13 vertebrae) was performed in 3/16 (19%) dogs and was consistent with MM in all. Histologic evaluation of osteolytic lesions (in the distal femur, and L5 and T12/T13 vertebrae) was performed in 3/16 (19%) dogs and disclosed MM in all, with 1 already confirmed cytologically before biopsy (T12/T13 osteolytic lesion).

#### 3.4 | Treatment with melphalan

Dogs in the daily dose cohort received melphalan at a dosage of 0.1 mg/kg/day for 10 days and 0.05 mg/kg/day thereafter for a median of 267 days (range, 64–1108 days). Dogs in the pulse dose cohort received melphalan at a dosage of 7 mg/m²/day (rounded to the

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**Table 1** Concurrent malignancies

| Malignancy                                | Pulse cohort | Daily cohort | Dx in relation to MM | Comments                                                                 |
|-------------------------------------------|--------------|--------------|----------------------|--------------------------------------------------------------------------|
| Peripheral nerve sheath tumor (grade I, incompletely excised, right front foot) | X            | At MM work-up |                      | Euthanized 13 mo after dx because of pulmonary metastasis; in MM remission (normal globulins) |
| Suspect cardiac hemangiosarcoma with pulmonary metastasis | X            | 24 months after MM tx start |                      | Euthanized at dx (in MM remission based on hyperglobulinemia, not necropsy-confirmed) |
| Anaplastic carcinoma on right side of neck | X            | 6 months after MM tx start |                      | Lost to follow-up ~2 weeks after carcinoma dx. Progressive disease noted at the time of carcinoma dx and received one dose of doxorubicin before being lost to follow-up |
| Metastatic AGASACA                         | X            | 25 months after MM tx start |                      | Euthanized from causes attributed to AGASACA whereas in MM remission (necropsy-confirmed) |
| Oral malignant melanoma                    | X            | 28 months before MM dx |                      |                                                                          |
| Dermal malignant melanoma                  | X            | 18 months before MM dx |                      |                                                                          |

Abbreviations: AGASACA, apocrine gland anal sac adenocarcinoma; MM, multiple myeloma; mo, months; dx, diagnosis; tx, treatment.
nearest whole 2 mg tablet; Alkeran®, Apopharma, Weston, Florida) for 5 consecutive days every 21 days for a median of 342 days (range, 67–1481 days). Prednisone was administered at a dosage of 0.5 mg/kg/day for 10 days and 0.5 mg/kg every other day for 50 days in 16 dogs. Nineteen dogs received prednisone on variable schedules, whereas 3 dogs did not receive prednisone. Six dogs received 1–2 doses of cyclophosphamide within 10 days of initiating melphalan therapy with the intention of achieving a more rapid remission (based on individual clinicians’ judgment). Of these 6 dogs, 4 were in the pulse dose cohort and 2 were in the daily dose cohort.

3.5 Follow-up diagnostic testing

For the majority of dogs in the daily dose cohort, a CBC and biochemistry panel were performed monthly for the first 2 months, every other month for 2 months, and then every 3 months thereafter as long as no dose-limiting toxicities were observed. The median CBC follow-up time for dogs in the daily dose cohort was 276 days (range, 64–1481 days). Dogs in the pulse dose cohort had a CBC performed 7 days after the fifth consecutive dose of melphalan and every 21 days before each treatment cycle initiation. The median CBC follow-up time for dogs in the pulse dose cohort was 277 days (range, 56–1407 days). In the majority of dogs, a biochemistry panel was performed every other cycle at the beginning of therapy and every 2–3 cycles thereafter. The median CBC follow-up time including dogs in both cohorts was 277 days (range, 56-1481 days).

All 9 dogs that presented with renal disease had resolution of their azotemia a median of 52 days (range, 1–276 days) after initiation of melphalan therapy. Seven dogs had resolution of azotemia within 65 days; the other 2 dogs had resolution noted at 264 and 273 days, which is the first time their biochemistry results were reevaluated after diagnosis. In this renal disease group, median serum creatinine concentration was 2 mg/dL (range, 1.6–3.1 mg/dL) in the 8 dogs for which serum creatinine concentration was reported (the remaining dog had only increased BUN concentration reported). Concurrent urine specific gravity (USG) was determined only in 2 dogs (USG, 1.016 and 1.017), with both dogs receiving IV fluids at the time of USG evaluation.

3.6 Adverse events

Both treatment protocols were well tolerated. Thrombocytopenia was the most common adverse event in both cohorts. The highest grade of thrombocytopenia reported while receiving therapy is presented in Table 3. Aside from 1 dog with grade 1 thrombocytopenia receiving 10 months of daily melphalan therapy that was euthanized for progressive MM, all other grade 1 and 2 thrombocytopenic events did not result in dose reductions, dose delays, or discontinuation of melphalan therapy.

Neutropenia and anemia were common and low-grade in both cohorts. The daily dose melphalan cohort had one grade 1 neutropenic event and seven grade 1 and one grade 2 anemic events. The pulse dose melphalan protocol had one grade 1, two grade 2, and two grade 3 neutropenic events and six grade 1 anemic events.

No adverse gastrointestinal effects were reported.

3.7 Outcome

Response and outcome data are summarized in Table 4. Of the 38 dogs evaluated, 10 were lost to follow-up and 6 were still alive at the end of the follow-up period. Of the remaining 22 dogs, 14 dogs were suspected to have died from MM, 2 of which were confirmed on necropsy. Eight dogs died from causes most likely unrelated to MM, 1 of which was necropsy-confirmed to have renal disease and widespread metastatic apocrine gland anal sac adenocarcinoma with no evidence of MM. One dog died from traumatic wounds (ie, hit by car), although necropsy disclosed evidence of plasma cell neoplasia localized to the right popliteal lymph node only. The median follow-up period was 499 days (range, 70–2262 days).

Twenty-four dogs were censored from the OST analysis (Figure 1); 14, 21, 26, and 10 dogs were censored from the PFS, DFI, SFR, and TR analyses, respectively. No significant differences were found between the treatment cohorts for any of the outcome variables.

3.8 Treatment at relapse

Three dogs (8%) initially were treated with the daily dose protocol and later were switched to the pulse dose protocol. One of these dogs achieved SD after starting daily dose melphalan. Eighty-five days later,
PD was noted and this dog was switched to pulse dose melphalan. The pulse dose regimen resulted in a CR, which was sustained for 1481 days. Pulse dose melphalan was discontinued thereafter, and this dog continued to be free of MM or any myelosuppressive effects at the end of the follow-up period. The second dog achieved a CR with daily dose melphalan for 258 days, at which time PD was noted and pulse dose melphalan was initiated. This dog maintained SD with the pulse dose regimen for 190 days, after which time a single dose of lomustine was given and no subsequent treatment was pursued. The third dog maintained SD for 70 days on daily dose treatment, after which pulse dose treatment was initiated. This dog maintained SD for 48 days, at which time progression was noted and lomustine was given as rescue treatment. All 3 dogs were included in the daily dose cohort for analysis, because the initial intent was to treat them with the daily schedule. The dog that received pulse dose treatment for 1481 days was included in the pulse dose group only for the analysis of thrombocytopenia.

Dogs that developed PD in either treatment cohort received a variety of rescue chemotherapy protocols including: single agent cyclophosphamide (n = 3), single agent doxorubicin (n = 4), vincristine, doxorubicin, and dexamethasone (VAD; n = 2), rabacfosadine (Tanovea®-CA1, VetDC, Fort Collins, Colorado; n = 1), chlorambucil (n = 1), lomustine (n = 2), pegylated liposomal doxorubicin (n = 1), cyclophosphamide with doxorubicin (n = 1), vincristine, doxorubicin, cyclophosphamide (VAC; n = 1), and single agent vincristine (n = 1). No difference was observed in the frequency of possible prognostic factors between treatment cohorts. Fourteen dogs, including 9 dogs in the daily dose cohort and 5 dogs in the pulse dose cohort, received at least 1 rescue protocol with a median of 1 protocol per dog (range, 1–3). Three dogs received > 1 rescue protocol, 2 of which were in the daily dose cohort and 1 in the pulse dose cohort. The median length of rescue treatment was 54 days (range, 7–761 days).

### 3.9 NLR and PLR

Twenty-six dogs (68%) were included in the NLR and PLR analyses because only these dogs had complete CBCs available for review within 4 weeks of diagnosis. The remaining 12 dogs (32%) had CBCs performed and reported to be unremarkable. Hence, specific results were not reported, and these dogs therefore were not included in the NLR and PLR analyses. The median NLR was 4.03 (range, 1.58–28.67), and the median PLR was 146.70 (range, 1–1000).
The optimal cut-off points for the NLR and PLR as predictors of OST were 4.28 (sensitivity, 57.59%; specificity, 55.26%) and 216.2 (sensitivity, 88.46%; specificity, 50%), respectively. Based on the cut-off points for the NLR and PLR, dogs were divided into the following groups: high NLR (n = 11; NLR > 4.28), low NLR (n = 15; NLR ≤ 4.28), high PLR (n = 3; PLR > 216.2), and low PLR (n = 23; PLR ≤ 216.2).

3.10 | Prognostic factors

Significant prognostic factors are summarized in Table 5. Both renal disease (Figure 2) and NLR (Figure 3) were significantly prognostic for OST, PFS, and DFI. The NLR alone was significantly prognostic for SFR. None of the other factors held any prognostic significance with respect to OST, PFS, DFI, SFR, and TR.

4 | DISCUSSION

To the authors’ knowledge, ours is the first study to evaluate pulse dose melphalan and compare it to daily dose melphalan in dogs with MM. Our hypothesis was supported by the lack of significant difference in outcome and adverse event profiles between the pulse and daily dosing regimens, although small cohort size and high numbers of censored dogs could have impacted this result. Both protocols were associated with high response rates, a short TR and few dose-limiting adverse events. Previously reported negative prognostic factors including hypercalcemia and osteolytic lesions were not confirmed in our study, whereas renal disease and high NLR emerged as potential negative prognostic indicators.

Melphalan chemotherapy typically is well tolerated, with myelosuppression being the most common dose-limiting toxicity.1,2,39–42 In our study, both dosing regimens were well tolerated. Only 1 of 6 dogs that experienced grade 3 or 4 thrombocytopenia had BMA-confirmed evidence supporting melphalan as the likely cause, based on a lack of bone marrow plasmacytosis or other neoplasia. This dog received pulse-dose melphalan for 13 months. Conversely, 2 dogs in both cohorts that received ≥ 2 years of melphalan had no grade 3 or 4 thrombocytopenia events documented. Overall, both protocols were associated with few clinically relevant adverse events.

Three dogs were switched from daily to pulse dose scheduling, and responses after this switch varied from SD to CR. A possible reason for positive responses is the 2–3 fold increased dose intensity during the first 5 days of melphalan pulse dosing. Although the dose intensities between protocols were similar when compared over several months, this finding suggests that some dogs may benefit from high dose-intensity pulses of melphalan.

| Variable                | High NLR (n = 11)         | Low NLR (n = 15)         | P value |
|-------------------------|---------------------------|--------------------------|---------|
| Median OST (range)      | 330 days (117–930)        | 1198 days (70–1554)      | .008    |
| Median PFS (range)      | 227 days (64–928)         | 778 days (70–1406)       | .04     |
| Median DFI (range)      | 233 days (97–902)         | 778 days (9–1492)        | .002    |
| Median SFR (range)      | 259 days (97–1308)        | 1157 days (9–1492)       | .001    |

Renal disease (n = 9)  No renal disease (n = 29)

| Variable                | High NLR (n = 11)         | Low NLR (n = 15)         | P value |
|-------------------------|---------------------------|--------------------------|---------|
| Median OST (range)      | 330 days (103–1554)       | 1198 days (70–2262)      | .019    |
| Median PFS (range)      | 243 days (64–1406)        | 664 days (70–1481)       | .03     |
| Median DFI (range)      | 219 days (46–1492)        | 902 days (9–1308)        | .049    |

Abbreviations: DFI, disease free interval; NLR, neutrophil to lymphocyte ratio; OST, overall survival time; PFS progression free survival; SFR, survival from remission.
Contrary to previous findings, hypercalcemia and osteolytic lesions were not identified as negative prognostic indicators in our study. This disparity could be a consequence of our small sample size, high censoring, or improved pain management, with the latter allowing for longer treatment because of improved quality of life despite the presence of neoplastic bone disease. The importance of Bence-Jones proteinuria was not assessed in our study, because this test was performed in only 2 dogs. Few studies of humans with MM have evaluated the prognostic importance of hypercalcemia and bone involvement, and the results are inconsistent, similar to studies in dogs. Some studies have shown worse outcome associated with hypercalcemia and bone lesions (on magnetic resonance imaging and positron emission tomography/computed tomography) whereas other studies have not identified inferior outcomes associated with either factor.

In our study, renal disease was found to be significantly associated with shorter OST, PFS, and DFI. This finding is corroborated by studies in human patients with newly diagnosed MM with some studies relating prognosis to the severity of renal impairment and others showing a correlation between reversibility of renal impairment and improved overall survival. Other studies however refute the role of kidney disease as an independent prognostic factor when adjusted for MM stage. All dogs in our study had reversible azotemia consistent with AKI of various grades, although a component of underlying early chronic kidney disease could not be ruled out. Possible reasons for presentation with renal insufficiency include nephrotoxicity of monoclonal light chains, hypercalcemia, hypertension, dehydration, and use of nephrotoxic drugs. These findings along with those previously reported support renal impairment as a negative prognostic factor in dogs with MM, although studies stratifying dogs based on renal function are necessary for validation.

Increased NLR is a recently identified independent negative prognostic factor in people with MM, with a high NLR associated with shorter overall survival and progression- or event-free survival times. Similarly, we identified an association between increased NLR and shorter OST, PFS, DFI, and SFR in dogs. In neoplastic processes such as MM, an increased NLR may reflect a decreased antitumor immune response by lymphocytes with concurrent protumor activity by neutrophils, particularly an IL-6-mediated neutrophilia.

Abdominal ultrasonography with fine needle aspiration (FNA) cytology of intra-abdominal organs was performed in the majority of dogs in our study. This diagnostic combination helped confirm the diagnosis of MM in 11 dogs that had either a normal BMA result or no BMA performed, and provided additional staging information in 6 others. Five dogs with ultrasonographically normal-appearing spleen or liver had cytologic evidence of infiltration with neoplastic plasma cells, lending support to aspiration of normal-appearing visceral organs for complete staging. Collectively, these findings support the use of abdominal ultrasonography with FNA cytology as part of the initial diagnostic evaluation of dogs with MM, although visceral involvement was not associated with prognosis in our study.

Limitations of our study are attributable to its retrospective nature and include the lack of randomization as well as lack of standardization in staging tests, follow-up, response evaluation, and rescue treatment. Our study also included a relatively small sample size, which may be attributable to the relative rarity of MM in dogs. In conclusion, our findings suggest that dogs with MM being treated with melphalan in either the daily or pulse dose setting have a favorable prognosis with minimal chemotherapy-related toxicity. Renal disease and high NLR were found to be independent negative prognostic factors in our study population. Prospective, controlled, and randomized studies to confirm these results are warranted.

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CONFLICT OF INTEREST DECLARATION

The authors declare that they have no conflict of interest with the content of this article.

OFF-LABEL ANTIMICROBIAL DECLARATION

Authors declare no off-label use of antimicrobials.

INSTITUTIONAL ANIMAL CARE AND USE COMMITTEE (IACUC) OR OTHER APPROVAL DECLARATION

Authors declare no IACUC or other approval was needed.

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