

1 **METASTATIC CANCER OF UNKNOWN PRIMARY IN 21 DOGS**

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3 F. Rossi,¹ L. Aresu,² M. Vignoli,¹ P. Buracco,³ G. Bettini,⁴ S. Ferro,² F. Gattino,³ F. Ghiani,⁵

4 R. Costantino,⁶ L. Ressel,⁷ E. Bellei,⁸ and L. Marconato.¹

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7 1. Centro Oncologico Veterinario, Sasso Marconi, Italy

8 2. Department of Comparative Biomedicine and Nutrition, University of Padova,
9 Legnaro, Italy

10 3. Department of Animal Pathology, Veterinary School, Grugliasco, Italy

11 4. Department of Veterinary Medical Sciences, University of Bologna, Italy

12 5. Studio Veterinario Associato Costa-Ghiani-Spallarossa, Genova, Italy

13 6. Clinica Veterinaria Orobica, Bergamo, Italy

14 7. Diagnostica Veterinaria di Laboratorio, Sasso Marconi, Italy

15 8. Centro Medico Veterinario, Finale Emilia, Italy

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23 **Corresponding author:**

24 Laura Marconato, DVM, DECVIM-CA (Oncology)

25 Centro Oncologico Veterinario

- 26 Via San Lorenzo 1-4
- 27 40037 Sasso Marconi, Bologna, Italy
- 28 lauramarconato@yahoo.it
- 29
- 30

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32 Abstract

33 The aim of this retrospective study was to describe clinical features, treatment and outcome of
34 21 dogs with metastatic cancer of unknown primary (MCUP), a biopsy-proven malignancy
35 being diagnosed at a metastatic stage, in which the anatomical origin of the primary tumor
36 cannot be detected. All dogs underwent total-body CT. Signalment, type and duration of
37 clinical signs, metastasis site, pathology results, treatment and outcome were recorded.
38 Carcinoma was the most common diagnosis (57,1%), followed by sarcoma, melanoma and
39 mast cell tumor. The median number of disease sites per dog was 2, with bones, lymph nodes,
40 lungs, and spleen being the most frequent metastatic locations. The median survival for all
41 dogs was 30 days. Overall, a primary site was not identified in 20 (95,2%) dogs. MCUP
42 encompasses a variety of different pathologic entities and harbors a poor prognosis.

43

44 Metastatic carcinoma of unknown primary (MCUP) is the seventh most frequently occurring
45 cancer and the fourth commonest cause of cancer-related death in people.¹ It refers to a
46 biopsy-proven malignancy in which the anatomical origin of the primary tumor cannot be
47 detected after a thorough patient history, careful physical examination, and extensive work-up
48 including laboratory testing, chest radiographs, endoscopy, abdominal ultrasound and/or
49 computed tomography (CT) of the head, chest, abdomen and pelvis and, in selected cases,
50 mammography.² Serum tumor markers are commonly of no help, since non-specific
51 elevations occur in the majority of MCUP patients.^{3,4} In people only 20% of primary sites are
52 identified by extensive diagnostic work-up before the patients die.⁵ In approximately 70% of
53 patients, the primary site cannot be identified even at necropsy.^{4,6} Typically, MCUP
54 progresses and spreads rapidly, with signs related to the metastatic site. Due to the lack of
55 consensus on diagnostic guidelines and optimal treatment, patients with MCUP have a poor
56 prognosis, with a median survival time of 6-12 months, thereby rendering this disease a
57 dilemma for oncologists.⁴ Standard management is based on empiric chemotherapy, including
58 several taxane/platinum regimens; nevertheless as the tumor recurs or progresses after first-
59 line chemotherapy, effective second-line treatments are not available.^{3,7,8}

60 To the authors' knowledge, there are no studies in veterinary medicine addressing MCUP. In
61 dogs, conventional techniques for evaluating primary tumor sites include laboratory testing,
62 radiographs, ultrasound, endoscopy, CT scan and/or magnetic resonance imaging (MRI),
63 depending on tumor site. Serum tumor markers being clinically useful in cancer diagnosis are
64 not available in veterinary medicine.⁹

65 Advanced diagnostic procedures in pathology, such as immunohistochemistry (IHC), may
66 enable diagnosis of the origin of primary tumors by biopsy of the metastasis in selected cases.
67 Some markers have been used in dogs to classify tumors according to their site of origin and
68 distinguish metastatic carcinomas, including thyroid transcription factor-1 (TTF-1) and

69 uroplakin III.^{10,11} Nevertheless, as IHC is not 100% specific and its interpretation may be
70 challenging, it is important to use markers for guidance in conjunction with the clinical
71 presentation and imaging studies.

72 The aim of this retrospective study was to describe clinical characteristics, treatment, and
73 outcome of dogs with MCUP.

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76 **MATERIALS AND METHODS**

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78 **Criteria for selection of cases**

79 Medical records of all contributing institutions were retrospectively searched to identify dogs
80 with a presumed diagnosis of MCUP. Determining whether the primary site is unknown or
81 whether it will be possible to detect it with further evaluation is difficult. In the present study,
82 members of the Italian Society of Veterinary Oncology (SIONCOV) were asked to look
83 through their records to identify MCUP cases. Once a possible case was identified, the
84 histological sample of the metastatic site was retrieved and reviewed in concert by skilled
85 pathologists with oncologic expertise (GB, SF, LR).

86 For the purposes of this study, dogs were considered to have MCUP if the following
87 diagnostic procedures did not reveal a primary tumor site: a detailed medical history;
88 complete physical examination; complete blood cell count and biochemistry; urinalysis;
89 histopathological review of biopsy material, and total body (TB) CT. Pathologic evaluation
90 included light microscopic evaluation in all cases. Poorly differentiated tumors had additional
91 immunohistochemical staining.

92 Dogs with tumors having the capability of arising at multiple sites simultaneously, such as
93 lymphoma or histiocytic sarcoma, were excluded from the study. In particular, to rule out

94 these tumors, additional immunohistochemical staining, including CD20, CD79, CD3 and
95 CD18, were performed whenever indicated.

96

97 **Procedures**

98 Data obtained from the medical records of dogs enrolled in this retrospective study included
99 signalment (i.e., age, sex, body weight, and breed), type and duration of clinical signs, results
100 of imaging, site of metastasis, pathology results, treatment, response to therapy, outcome and
101 necropsy data (if performed). Responses to treatment were defined according to the World
102 Health Organization criteria and were required to last for at least 28 days.

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105 **RESULTS**

106 Twenty-one dogs fulfilled the inclusion criteria. There were 2 Beagles, 2 German shepherd
107 dogs, 2 Labrador retriever, 2 Corso dogs, 2 mixed breeds, and one each of Schnauzer, Cocker
108 spaniel, Basset hound, Mongrel, Siberian husky, Boxer, American Staffordshire terrier,
109 Weimaraner, Rhodesian Ridgeback, West Highland white terrier, and Beauceron. There were
110 12 males (1 of which was castrated) and 9 females (7 of which were spayed). Median age at
111 presentation was 10 years (range, 7 to 15 years), whereas median weight was 24 kg (range, 6
112 to 42 kg). Patients' characteristics are listed in Table 1.

113 When considering clinical signs, 3 (14,3%) dogs were asymptomatic and their tumors were
114 diagnosed incidentally. Two of them had a painless enlargement of a peripheral lymph node
115 (mandibular: n=1, axillary: n=1), and the other dog developed multiple, painless,
116 subcutaneous nodules. Eighteen (85,7%) dogs showed clinical signs, such as dyspnea (n=7),
117 lameness (n=5), depression/lethargy/weakness (n=2), tenesmus (n=1), abdominal pain (n=1),

118 lethargy (n=1), and polyuria/polydipsia (n=1); in these dogs clinical signs had been present
119 for a median of 14 days (range, 3 to 150 days) prior to presentation.

120 All enrolled dogs underwent complete history, physical examination, CBC, chemistry profile,
121 urinalysis, and contrast-enhanced TBCT scan, showing metastatic disease without obvious
122 primary. The CT scan images were acquired with dogs in sternal recumbancy in order to
123 minimize lung collapse/hypostasis, which may hide small peripheral metastatic lesions. The
124 patients were scanned before and after the intravenous administration of a non ionic contrast
125 medium (Ioversol, Covedian, Milan, Italy) at the dose of 600-800 mgI/kg through a power
126 injector at a speed of 3 ml/sec. Different scanners were used. These included a single-slice CT
127 (GE, HiSpeed FX/I) in 3 dogs, and a 16-slice multidetector scanner (GE, BrightSpeed) in 18
128 dogs. The slice thickness was 1.25 to 3.0 mm, depending on the machine used. All images
129 were reviewed by two board-certified radiologists (FR, MV), who were unaware of the
130 histopathological diagnosis in the cases for which it was available. All cases included in this
131 study had multiple nodules of similar size and vascularization, located in different organs,
132 confirmed to be neoplastic in origin based on histopathological evaluation. An evaluation of
133 the CT did not suggest that any of these lesions could be the primary tumor. The metastatic
134 origin of the pulmonary lesions was supported by the finding of multiple nodules of soft
135 tissue density and similar size, growing in the pulmonary interstitium in an expansive way
136 and compressing the surrounding structures. CT features typically associated with primary
137 neoplasia, like the presence of a larger single mass or a focal area of soft tissue lung infiltrate,
138 associated with signs of local aggressiveness, were never observed in this group of
139 patients.¹² The sites of metastasis are listed in Table 1. The median number of disease sites per
140 dog was 2 (range, 1 to 11). Ten (47,6%) dogs had a single metastatic organ site, 3 (14,2%)
141 had 2, 2 (9,6%) had 4, 1 (4,8%) had 3, 1 (4,8%) had 5, 1 (4,8%) had 6, 1 (4,8%) had 7, and 1

142 (4,8%) had 11. Bones, lymph nodes, lungs, and spleen were the most frequent metastatic
143 locations (Table 1).

144 Pathologic samples were obtained by surgical excision (n=3) or core needle biopsy (n=18).

145 All biopsy specimens were of good quality allowing for accurate histological interpretation.

146 Twelve (57,1%) dogs were diagnosed with carcinoma (undifferentiated carcinoma: n=11;

147 squamous cell carcinoma: n=1), 7 (33,3%) with sarcoma (undifferentiated sarcoma: n=3;

148 fibrosarcoma: n=2; hemangiosarcoma: n=2), 1 (4,8%) with amelanotic melanoma and 1

149 (4,8%) with mast cell tumor. Immunohistochemical analysis to better characterize poorly

150 differentiated tumors was performed in 17 cases. Tests performed included cytokeratin,

151 vimentin, S-100 as standard panel, and PNL2 for amelanotic melanomas. Additional

152 pathologic evaluation (including TTF-1, Factor VIII and CD18) was individualized on the

153 basis of clinical and pathologic features. The pathological diagnoses of the dogs and details

154 on immunohistochemistry are reported in Table 1.

155 When considering histological type and metastatic pattern detected after TBCT, 8 out of 12

156 (66,7%) dogs with carcinoma had a single metastatic site. All dogs with sarcoma had multiple

157 metastatic sites, whereas both the dog with melanoma and the one with mast cell tumor also

158 had a single metastatic site.

159 Eleven (52,4%) dogs received no treatment and were euthanized shortly after diagnosis. Four

160 (19%) dogs received systemic chemotherapy (2 of which were treated with metronomic

161 chemotherapy), 2 (9,5%) dogs were treated with firocoxib (Previcox, Merial, Milanofiori,

162 Italy), 1 (4,8%) underwent surgery, 1 (4,8%) was treated with palliative radiation therapy and

163 immunotherapy with the canine melanoma vaccine (Oncept, Merial, Milanofiori, Italy), 1

164 (4,8%) dog was treated with surgery, chemotherapy and radiation therapy, and 1 (4,8%) dog

165 was treated with surgery and toceranib (Palladia, Pfizer, Italy).

166 The median survival time was 30 days for all dogs diagnosed with MCUP. Eighteen (85,7%)
167 dogs were dead at the end of the study for cancer-related causes after a median of 12 days
168 (range, 1 to 504 days). Seven of them underwent therapy; in five dogs the metastatic tumors
169 did not respond to any form of treatment, thereby being classified as progressive; 1 was
170 stable, and 1 obtained a partial remission before developing pulmonary metastases. The
171 median survival time for dogs undergoing any form of treatment was 80 days (range, 30 to
172 504 days). Necropsy was only performed in a single case; no primary tumor site was found.

173 Three dogs were still alive at data analysis closure, after 882, 101 and 80 days. Among these,
174 one dog had a metastatic amelanotic melanoma in the mandibular lymph node with no
175 evidence of a primary tumor based on physical examination and complete work-up, including
176 TBCT. Seven months after radiation therapy and immunotherapy, the dog developed a
177 melanoma in the ipsilateral footpad and was irradiated again, thereby obtaining a complete
178 response. It was hypothesized that the footpad was the primary site, possibly having remained
179 occult when the metastasis first appeared. The IHC staining pattern was similar between the
180 melanoma in the footpad and the lymph node.

181 The second dog had a metastatic carcinoma involving peripheral, intrathoracic and abdominal
182 lymph nodes, both adrenal glands, liver, pancreas, lungs, and muscles. At the time of writing,
183 the dog was still receiving daily firocoxib; however, the metastatic tumor was shown to be
184 progressive according to follow-up imaging.

185 The third dog had a carcinoma metastatic to the medial iliac lymph node. Due to
186 hypercalcemia, a clinically occult anal sac carcinoma was suspected, and the dog underwent
187 lymphadenectomy and bilateral anal saccullectomy. However, based on histopathology, both
188 anal sacs were morphologically normal, thereby prompting the diagnosis of MCUP metastatic
189 to the medial iliac lymph node. After surgery the hypercalcemia resolved, and at the time of

190 writing the dog is being treated with toceranib and is considered to be in complete remission
191 based on clinical and imaging features.

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193

194 **DISCUSSION**MCUP is perceived to be a very aggressive disease carrying a poor prognosis. It
195 refers to a biopsy-proven metastatic cancer in the absence of an identifiable primary tumor
196 despite a complete diagnostic work-up.¹ Although the biologic characteristics of MCUP
197 remain to be determined, some hypothesis have been postulated. The primary tumor may
198 remain diminutive, thereby escaping clinical detection, or it may undergo spontaneous
199 immune-mediated regression or dormancy after seeding the metastasis.¹² Alternatively, the
200 angiogenic incompetence of the primary tumor may lead to marked apoptosis and cell
201 turnover, resulting in a cancer that acquires a metastatic phenotype.¹³ Other explanations
202 include various theories, including stem cell and embryologic migration hypotheses.¹⁴
203 Nevertheless, all these theories cannot be clinically tested and remain speculative.

204 MCUP is not rare in people, representing 3-5% of all malignancies diagnosed in oncology
205 practice.⁴ There are no studies in veterinary oncology focusing on MCUP, therefore its
206 prevalence is unknown. In this case series, we described 21 dogs with MCUP for which
207 TBCT and tumor histology were available. Only dogs undergoing TBCT were included, as
208 conventional radiography and ultrasound may miss tumors located in the head and neck
209 region, pelvic cavity or intracardiac structures, thereby being inadequate for diagnosing
210 MCUP. It is unknown whether the use of more sophisticated imaging studies (such as, for
211 instance, MRI or PET, if available) would be beneficial and appropriate in these cases.
212 Indeed, the poor prognosis typically being associated with metastatic cancer raises issues of
213 cost effectiveness for intensive diagnostic work-up, which may be unrevealing and of unclear

214 benefit in terms of improving prognosis. Therefore, the list of investigations for MCUP is
215 difficult to define, and requires continual updating.

216 In this case series, in one single dog the primary site was suspected to be found antemortem,
217 being in accordance with most studies conducted in people.⁴⁻⁶ This dog had a metastatic
218 melanoma in the mandibular lymph node and 7 months later developed a melanoma in the
219 ipsilateral footpad, which was hypothesized to be the primary site. Although the mandibular
220 lymph node is not the draining lymph node for the footpad, nodal metastases may also occur
221 in a random process, with the second and third level lymph nodes being involved with
222 metastatic disease when compared with the nodes closest to the tumor. Beside the
223 identification of the neoplastic anatomical site of origin, great interest has been given to the
224 recognition of specific histological subtypes, as the chemotherapeutic regimens chosen to
225 treat MCUP cases in people depend not only on the site of primary origin, but also on the
226 cancer subtype.¹⁴ While epithelial histotypes are more frequently diagnosed,^{1,3,4} malignant
227 melanomas and sarcomas occasionally occur as apparent metastasis to lymph nodes or viscera
228 without a detectable or known primary lesion.¹⁵⁻²¹ IHC stains are an important complement to
229 light microscopy in the investigation of MCUP. Several panels of stains are recognized as
230 important in the diagnosis of specific subtypes of cancer by predicting with greater certainty
231 the likely tissue of origin of the malignancy.³ Several examples include GCDFP-15,
232 mammoglobin, oestrogen and progesterone receptors, in breast cancer, TTF-1 in
233 pulmonary carcinoma, HEPAR-1 in hepatocellular carcinoma, thyroglobulin/TTF-1 in
234 thyroid carcinoma placental alkaline phosphatase/OCT-4 in germ-cell tumors, CDX-2 in
235 colorectal cancer, and synaptophysin and chromogranin in neuroendocrine tumors
236 (OIEN). Because the morphological and immunohistochemical features are often not
237 characteristic, gene expression-based analysis are an emerging tool to help in identifying the
238 primary site and, possibly, selecting targeted treatment.²² Gene expression profiling is a new

239 frontier in veterinary oncology, and usually not routinely offered. In this retrospective series
240 of cases, we limited the pathological evaluation to morphology and, in selected cases,
241 immunohistochemistry. In agreement with the human counterpart, carcinomas were the most
242 frequently diagnosed tumors in this case series, followed by sarcoma, melanoma and mast cell
243 tumor. It must be stressed that a limited panel of IHC tests were used here, mainly due to
244 financial concern and to the lack of site-specific markers with high sensitivity and specificity,
245 thereby precluding the possibility to further characterize some of the tumors. Whether the use
246 of a large panel of antibodies is associated with clinical gain and change in management is not
247 known and cannot be recommended at the moment.

248 In human oncology, it appears that patients with MCUP have a limited life expectancy with a
249 median survival approximately of 6-12 months, and with fewer than 25% of patients
250 surviving beyond 1 year.⁴ The same holds true for dogs, as a median survival time of 30 days
251 was recorded here. This data is not unexpected, as proven metastatic cancer is typically
252 associated with a poor outcome, regardless of the recognition of the tumor's primary site.

253 In human patients, several clinical and biologic variables have been demonstrated to have
254 significant impact on survival, including performance status, weight loss, histological
255 subtype, presence of liver metastases, more than two metastatic sites, elevated levels of serum
256 alkaline phosphatase and lactate dehydrogenase, thereby allowing the inclusion of patients
257 into groups requiring specific guidelines that translate into prolonged survival.²³⁻²⁵ Favourable
258 subsets are usually treated with locoregional treatment or systemic platinum-based
259 chemotherapy, achieving responses and survival times that are similar to those of patients
260 with relevant known primary tumours.^{14,26} Conversely, patients in unfavourable subsets are
261 treated with empirical chemotherapy based on various combination regimens, but responses
262 and survival are generally poor.^{14, 26} Due to the small size of our population and the non
263 uniformity of treatment, prognostic factors were not identified in this work. More information

264 needs to accumulate to verify whether these data may provide useful diagnostic and
265 therapeutic information for dogs with MCUP as well.

266 The purpose of this study was to describe a collection of eclectic, previously unreported
267 cases; however, the retrospective nature of this study and the small population size represent
268 main limitations. Many questions still need to be answered: not only MCUP is a rare disease
269 entity, but it is also neglected, more over because of the lack of information and
270 understanding about the disease. The uncertainty which surrounds almost all aspects of care
271 for dogs with MCUP is most clearly seen when decisions need to be made about
272 investigations and treatment, as shown in this series of cases. Additionally, the consistently
273 poor prognosis is a further disadvantage when discussing options with the owners or when
274 trying to support research. Collaborative studies are warranted to improve the knowledge and,
275 possibly, the care of animals with MCUP.

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