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Title: The Utility of Diagnostic Tests for Immune-mediated Hemolytic Anemia

Short Title: Diagnosis of IMHA

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28 **Abstract**

29 **Background:** A definitive diagnosis of immune-mediated hemolytic anemia (IMHA) can be difficult to
30 make. However, it is critical to differentiate IMHA from other causes of anemia due to the impact on
31 prognosis and outcome for IMHA patients. Recently published American College of Veterinary Internal
32 Medicine recommendations for the diagnosis of IMHA should be followed to concurrently confirm
33 ongoing anemia, verify *in vivo* hemolysis, and detect anti-erythrocyte antibodies. The reliability of
34 immunologic IMHA tests varies depending on which test is used and how it is performed.

35 **Objectives:** Our aims were to determine which tests are currently used in veterinary medicine to
36 diagnose IMHA and to review the utility of assays that have historically been used to diagnose IMHA.

37 **Methods:** A short survey was designed to see which diagnostic tests for IMHA were currently being
38 used by veterinary practices. The survey was distributed *via* list-serves to veterinarians and veterinary
39 technologists. A literature review was performed to report the utility of diagnostic tests for the diagnosis
40 of IMHA.

41 **Results:** Survey respondents indicated variability in test protocols used to diagnose IMHA. Most
42 respondents perform saline agglutination or Coombs' tests to detect anti-erythrocyte antibodies.
43 Additional tests that can be used to support a diagnosis of IMHA are discussed in this review.

44 **Conclusions:** A standardized diagnostic approach should be followed to differentiate IMHA from other
45 causes of anemia. Test methodology can vary from one laboratory to another, and clinicians should be
46 familiar with the procedures used by their laboratory.

47

48 **Key Words:** Agglutination, Anti-erythrocyte antibody, Coombs', Diagnosis, Spherocytosis

49

50 **Introduction**

51 Immune-mediated hemolytic anemia (IMHA) is commonly diagnosed in canine patients with hemolytic
52 anemia. It is a less prevalent, but an equally important cause of hemolytic anemia in cats, horses, and
53 cattle.^{1,2} While there are certain clinicopathologic findings supportive of an IMHA diagnosis (e.g.,
54 peripheral blood spherocytosis, RBC agglutination, demonstration of immunoglobulins attached to RBC
55 membranes), the diagnosis of IMHA is often presumptive. Misdiagnosis of IMHA is problematic because
56 of important differences in treatment decisions and prognostic consequences for patients with IMHA as
57 compared with other causes of anemia.³ Additionally, it is critical to evaluate patients that have IMHA
58 for underlying diseases that drive the immune-mediated response. This article summarizes the results of

59 a short survey on the use of diagnostic tests for IMHA in veterinary medicine and reviews the utility of
60 diagnostic tests for IMHA.

61

62 **Diagnosis of IMHA**

63 In anemic patients, the mechanism causing the anemia (decreased RBC production, hemorrhage, or
64 hemolysis) needs to be determined. A CBC (including quantitative data and a qualitative blood film
65 review) provides fundamental information needed for the assessment of anemia. Anemia can be
66 classified as either regenerative or nonregenerative based on quantification of the reticulocytosis using
67 an absolute reticulocyte concentration or corrected reticulocyte percentage in the sample. If the anemia
68 is regenerative, blood loss or hemolytic anemia should be considered. Blood loss anemia can often be
69 differentiated from hemolytic anemia using clinical examination findings and additional
70 clinicopathologic data. Specifically, acute hemorrhage results in loss of serum proteins and,
71 consequently, decreased serum albumin and globulin concentrations. Although blood loss typically
72 results in a strongly regenerative anemia, chronic hemorrhage can cause a poorly regenerative anemia if
73 blood loss leads to iron depletion, since the resultant iron-restricted erythropoiesis impedes reticulocyte
74 production. It is important to assess patients for occult blood loss as gastrointestinal and urinary blood
75 loss can be easily missed. While the majority of IMHA patients present with a regenerative anemia, this
76 initial classification is not a certainty. IMHA can be nonregenerative or pre-regenerative if there has
77 been insufficient time for a clearly regenerative response to occur (anemia present < 5-7 days)⁴ or if
78 immune-mediated destruction of erythroid precursors is occurring (“precursor-targeted immune-
79 mediated anemia”).⁵

80

81 It is vital to document that true *in vivo* hemolysis is occurring in patients with IMHA. Accordingly, proper
82 collection of blood samples is critical to avoid *ex vivo* hemolysis, which can lead to false decreases in RBC
83 concentration, hematocrit (HCT), and packed cell volume (PCV); false increases in mean cell hemoglobin
84 concentration (MCHC); variable alterations in serum bilirubin concentrations (depending on the
85 methodology used); and can also interfere with several other serum chemistry values.⁶ Diagnostic test
86 results that support *in vivo* hemolysis are provided in **Table 1**.

87

88 To distinguish IMHA from other causes of anemia, an American College of Veterinary Internal Medicine
89 (ACVIM) consensus statement recommends documenting all three of the following diagnostic findings as
90 minimum criteria to diagnose IMHA in dogs:⁷

- 91 1) Decreased packed cell volume (PCV)
- 92 2) At least one of the following abnormalities supportive of hemolysis: erythrocyte ghost cells,
93 hyperbilirubinemia, bilirubinuria, icterus, hemoglobinemia, or hemoglobinuria
- 94 3) A positive saline agglutination test that persists with washing or at least two of the following:
95 a) A positive saline agglutination test without washing
96 b) ≥ 5 spherocytes per 1000 \times microscopic field
97 c) Detection of anti-erythrocyte antibodies by a Coombs' test or flow cytometry

98 Note that erythrocyte ghost cell numbers can markedly increase *ex vivo*, so examination of a freshly
99 made smear (at the time of blood collection) is recommended to assess this morphologic abnormality.⁶

100

101 A short survey (Appendix A) to determine how tests are being used to diagnose IMHA was sent to an
102 undetermined number of people in Australia, Canada, Europe, and the United States *via* list-serves and
103 email messages. Veterinary practitioners in both general practice and specialty hospitals, veterinary
104 clinical pathologists, and medical technologists and technicians in clinical pathology laboratories were
105 asked to reply. Respondents were not asked to self-identify. Ninety-four people completed at least the
106 first question of the survey.

107

108 The hematology parameters that > 75% of the respondents said were provided with a CBC are listed in
109 Figure 1. Seventy-two people (77%) indicated that, when needed, a saline agglutination test was
110 included with CBC results. An additional 13 respondents (14%) did not include a saline agglutination test
111 as part of a CB, but as an additional test. Seventy-two people (77%) recommended a test to detect anti-
112 erythrocyte antibodies (saline agglutination test, Coombs test, and/or flow cytometry). Additional tests
113 that were recommended by > 15 % of respondents included a biochemistry panel (66%), screening for
114 tick-borne diseases (60%), urinalysis (52%), and diagnostic imaging (34%).

115

116 Several tests have been used to aid in the diagnosis of IMHA, none of which are 100% sensitive and
117 specific. Differences in the protocols used to perform diagnostic tests for IMHA can dramatically affect
118 the performance of the test.^{8,9} Tests for IMHA must account for physical properties associated with
119 antibody-antigen interactions, one of which is temperature-dependent antibody-antigen binding.¹⁰

120 Therefore, the Coombs' test is often carried out at both 37°C and 4°C.¹¹ Additionally, inclusion criteria
121 for IMHA patients vary between references, which can alter the assessment of test performance. Most
122 studies evaluating diagnostic tests for canine IMHA list auto-agglutination or spherocytosis as inclusion

123 criteria. However, Reimer *et al.* (1999) concluded that 2/70 (3%) canine patients with IMHA did not have
124 auto-agglutination or spherocytosis, nor did they have polychromasia.¹²

125

126 Genetics

127 Several dog breeds have a higher incidence of IMHA than the general canine population including;
128 cocker spaniels,¹²⁻¹⁵ English springer spaniels,¹² miniature schnauzers,^{13,15} and old English sheepdogs¹⁶;
129 however, a specific gene association with IMHA has not be found. A few studies have associated the
130 presence of specific major histocompatibility complex alleles in dogs with the occurrence of immune-
131 mediated disease, but the predictive value of these associations remains uncertain.¹⁷⁻¹⁹ The frequency
132 of specific dog erythrocyte antigens found during blood typing was not significantly different in 33 dogs
133 with IMHA as compared to 1,014 dogs without IMHA.²⁰ Therefore, expectedly, genetic testing was not
134 recommended as an additional test for IMHA by survey respondents.

135

136 Non-specific Tests for Antibodies

137 Two proteins, staphylococcal protein A (SpA) and papain, have been used in several papers to document
138 increased immunoglobulin in serum samples. SpA is an immunoglobulin binding protein produced by
139 *Staphylococcus aureus* that has been shown to be more sensitive than indirect antiglobulin tests in dogs
140 with IMHA.²¹ Papain is an enzyme that digests antibodies into three 50 kDa segments comprised of one
141 fragment crystallizable (Fc) and two fragment antigen-binding (Fab) regions. When using a papain test to
142 detect anti-erythrocyte antibodies, false-positive results occur in up to 3/16 (19%) of dogs.²² One study
143 tested 23 papain positive anemic dogs with direct and indirect antiglobulin tests; 8/23 (35%) dogs were
144 positive using a direct test and none were positive using an indirect test.²³ Due to the non-specific
145 nature of SpA and papain reactions, they are no longer recommended as diagnostic tests for IMHA and
146 unsurprisingly were not listed as additional tests used by survey respondents.

147

148 Auto-agglutination

149 Red blood cell agglutination should raise suspicion for IMHA (**Fig. 2**). To determine whether
150 agglutination is likely related to the presence of antibody or complement on the surface of the red blood
151 cell (i.e., auto-agglutination), a saline agglutination test should be performed. It is important to note that
152 artifactual agglutination/rouleaux is not dispersed in all patients using a 1:1 dilution. However, if
153 agglutination persists after washing erythrocytes at least three times in isotonic or phosphate-buffered
154 saline, it is likely that the erythrocyte clumping is true agglutination rather than rouleaux formation. The

155 occurrence of auto-agglutination in dogs with IMHA varies from 42% to 86%.¹²⁻¹⁶ The number of patients
156 with a positive saline agglutination test is decreased when erythrocytes are washed extensively.^{24,25}
157 There is some evidence that the decrease in positive saline agglutination tests with washing is due to a
158 reduction in false-positive results,²⁵ but increased numbers of false-negative results should also be
159 considered.

160
161 Seventy respondents indicated whether they evaluated saline agglutination tests macroscopically or
162 microscopically: 34 looked for microscopic agglutination, 29 looked for macroscopic and microscopic
163 agglutination, and seven looked at the sample macroscopically only. Sixty-three people indicated how
164 they dilute blood samples for a saline agglutination test: 31 performed a 1:4 dilution, 24 performed a 1:1
165 dilution, and eight performed both a 1:1 and a 1:4 dilution of the sample. Of the 28 respondents who
166 indicated whether they wash RBCs for the saline agglutination test, 16 used unwashed RBCs, 11 used
167 washed RBCs, and one used both.

168
169 Erythron
170 Spherocytosis is often used as an inclusion criterion for studies of dogs with IMHA and was included in
171 CBC data by 90/94 (96%) of survey respondents (Fig. 1). However, IMHA also can be non-spherocytic, as
172 the occurrence of spherocytosis in dogs with IMHA ranges from 61% to 95%.^{12,13,15,16,24,26} Spherocytes are
173 generally only recognized in dogs (owing to the pronounced central pallor of their erythrocytes). In
174 contrast, spherocytes are rarely identified with certainty in cats, horses, and cattle since erythrocytes
175 from these animals lack central pallor on blood smears.

176
177 Spherocytosis and some other RBC membrane abnormalities can cause increased erythrocyte fragility.²⁷
178 The osmotic fragility of erythrocytes can be tested by diluting whole blood in progressively decreasing
179 concentrations of sodium chloride (NaCl), incubating the samples for 30 minutes at room-temperature,
180 recording absorbance of the samples at 540 nm, and then creating a data curve assuming that the
181 lowest NaCl concentration causes 100% hemolysis.²⁷ RBC hemolysis occurs as a result of a loss of
182 osmotic regulation and volume control, which is exacerbated in a number of RBC disorders. Increased
183 RBC hemolysis during osmotic fragility testing is commonly reported in spherocytic conditions (e.g.,
184 IMHA), but can be seen in spectrin deficiency,²⁸ hereditary stomatocytocytosis,²⁹ intestinal parasite-
185 associated microcytosis, *Babesia canis* infection,³⁰ or non-hemolytic samples that are lipemic.
186 Erythrocyte fragility testing is not commonly available to practitioners but can provide support for

187 ongoing hemolysis.³¹ One study observed increased erythrocyte fragility in 15/15 (100%) direct
188 antiglobulin test (DAT) positive and 4/12 (33%) DAT negative anemic dogs.²⁵ None (0/91) of the survey
189 respondents indicated that they recommended osmotic fragility testing for patients suspected of having
190 IMHA.

191
192 Most patients with IMHA have a moderate to marked regenerative anemia. A reticulocyte count was
193 listed as a component of a CBC by 77/94 (82%) of the survey respondents (Fig. 1). The occurrence of
194 reticulocytosis in dogs with IMHA ranges from 67% to 82%.^{13,16,26} IMHA patients often have increased
195 numbers of circulating nucleated RBCs and/or Howell-Jolly bodies. Interestingly, reticulocyte
196 hemoglobin (HGB) content was shown to be decreased in 5/14 (36%) dogs with IMHA suggesting that
197 iron-restricted erythropoiesis can be present in some canine IMHA patients.³²

198
199 A diagnosis of erythroid hyperplasia in bone marrow samples is a definitive indication of erythrocyte
200 regeneration. It is noteworthy that Weinkle *et al.* (2005) found 23/45 (51%) dogs with IMHA that
201 underwent bone marrow analysis had erythroid hyperplasia.¹³ Similarly, another study that analyzed
202 bone marrow samples from dogs with IMHA observed erythroid hyperplasia in 6/11 (55%) samples.¹⁵
203 Extramedullary hematopoiesis³³ and secondary myelodysplasia³⁴ also have been reported in dogs with
204 IMHA. Bone marrow evaluation is typically recommended for suspected IMHA patients that have a
205 nonregenerative anemia. Low numbers of survey respondents [6/91 (7%)] indicated that they
206 recommend bone marrow aspiration or biopsy to aid in the diagnosis of IMHA. Two of these
207 respondents specified that this recommendation was warranted in patients with persistent
208 nonregenerative anemia.

209
210 Nonregenerative anemia has been reported in 6/23 (26%) canine IMHA patients in one study²⁴ and 6/20
211 (30%) in another.¹⁵ In a retrospective analysis of dogs with nonregenerative anemia in which a bone
212 marrow sample was clinically indicated, 55/82 (67%) were determined to have IMHA (based on the
213 presence of either Coombs' positivity, auto-agglutination, or > 30% spherocytes).³⁵ In dogs with
214 nonregenerative IMHA, 38/55 (69%) had erythroid hyperplasia, and 17/55 (31%) showed incomplete
215 maturation of the erythroid line.³⁵ In another study of canine patients with a nonregenerative anemia
216 present for more than 5 days, 41/43 (95%) had erythroid hyperplasia in bone marrow samples, 23/43
217 (54%) had a spherocytosis, and 20/35 (57%) were positive by DAT.³⁶ To determine if dogs with a
218 nonregenerative anemia > 5 days duration have precursor-targeted immune-mediated anemia, bone

219 marrow should be evaluated for macrophage phagocytosis of erythroid precursors.⁵ Weiss (2008) also
220 evaluated 57 cats with nonregenerative anemia in which bone marrow analysis was indicated.
221 Approximately half of the cats 28/57 (49%) were determined to have IMHA (based upon measurement
222 of a HCT < 20% and either Coombs' positivity or auto-agglutination).³⁵ In cats with IMHA and
223 nonregenerative anemia, 24/28 (86%) had erythroid hyperplasia and 4/28 (14%) showed maturation
224 arrest of the erythroid line.³⁵ In another study, phagocytosis of erythroid precursors and abnormal
225 presence of stainable iron was documented in the bone marrow of cats with both primary and
226 secondary IMHA.³⁷

227
228 In cases of IMHA with RBC agglutination in the sample, many of the measured or calculated values of the
229 erythron are often erroneous [e.g., RBC concentration, mean cell volume (MCV), HCT, mean cell
230 hemoglobin (MCH), MCHC, red cell distribution width (RDW)] as impedance counters will count
231 erythrocyte clumps as single and large erythrocytes, which leads to a significant reduction in the
232 numbers of RBCs counted and an increase in the mean size of the cells.⁶ In cases of intravascular
233 hemolysis, HGB concentration is not clinically reliable, as it represents a combination of free (plasma)
234 and RBC HGB. If it is available, the determination of the cell hemoglobin concentration mean (CHCM)
235 using an advanced laser cell counter could help assess RBC HGB.

236
237 Leukon
238 Abnormalities in leukocytes are commonly observed in patients with IMHA. Leukocytosis was reported
239 in 43% to 99% of dogs with IMHA (WBC concentrations in these reports ranged from 5,300 cells/ μ L to
240 105,700 cells/ μ L).^{12,15,38,39} In one study, decreased survival time of dogs with IMHA was associated with
241 leukocytosis and lymphopenia,⁴⁰ while lymphocytosis was a positive prognostic factor in cats with
242 IMHA.³⁷ Neutrophil left shifts were noted in up to 16/20 (80%) of dogs with IMHA.¹⁵ One paper
243 observed decreased survival rates in dogs with IMHA with band neutrophil concentrations \geq 3000
244 cells/ μ L.¹³

245
246 IMHA patients with inflammatory leukograms typically have acute patterns such as a neutrophilia with a
247 left shift, lymphopenia, eosinopenia, and monocytosis.¹⁵ In severe inflammatory and erythroid
248 regenerative conditions, a leukoerythroblastic pattern can be observed with a highly acute inflammatory
249 leukogram and a high percentage of nucleated RBCs in different stages of maturation.¹⁵ Rubricytosis
250 causes a false increase in the automated WBC concentration that must be corrected mathematically

251 after the enumeration of nucleated RBCs by blood smear review. This further emphasizes the need for
252 blood smear examination, which can also help with the detection of neutrophil left-shifting and toxicity
253 that can be present in IMHA cases. Sixty-five/94 (69%) survey respondents indicated that blood smear
254 evaluation by a board-certified clinical pathologist was included in CBCs they performed or received (Fig.
255 1).

257 Serum Biochemistry and Urinalysis

258 Abnormalities in biochemical parameters have been associated with the clinical outcomes of IMHA
259 patients (**Table 2**), but not all studies report the same findings. An increase in total bilirubin
260 concentrations was observed in 60% to 100% of dogs with IMHA.^{12,15,26} It is important to note that
261 increased conjugated bilirubin can interfere with phosphorus measurements leading to
262 pseudohypophosphatemia in patients with IMHA.⁴¹ Although not linked to decreased survival, Klag et
263 al. (1993) observed hemoglobinemia and/or hemoglobinuria in 4/42 (10%) dogs with IMHA.²⁶ There are
264 also a few case studies of dogs with IMHA with biochemical abnormalities consistent with distal renal
265 tubular acidosis.⁴² In cats with IMHA, hyperglobulinemia is reported to be a positive prognostic factor.³⁷
266 Biochemistry profiles were recommended as additional tests for suspected IMHA patients by 60/91
267 (66%) of survey respondents.

268
269 Additional serum protein parameters have been reported to be altered in dogs with IMHA. For example,
270 cardiac troponin I was > 0.1 ng/mL in 20/27 (74%) dogs with IMHA (authors indicated that < 0.1 ng/mL is
271 expected in healthy dogs, but a true reference interval was not provided).⁴³ C-reactive protein was
272 increased in dogs with IMHA at presentation.⁴⁴⁻⁴⁶ Alpha-1 acid glycoprotein also was increased in dogs
273 with IMHA, while albumin can be decreased at presentation.⁴⁵ Increased serum concentrations of
274 several cytokines have been reported in dogs with IMHA (n = 20) as compared with six healthy dogs.⁴⁶
275 Interleukin-15 (IL-15), IL-18, granulocyte-monocyte colony stimulating factor, and monocyte
276 chemoattractant protein-1 concentrations were increased in animals with IMHA that died ≤ 30 days
277 after hospital admission.⁴⁶ Similarly, IL-2, IL-6, and tumor necrosis factor- α were present at higher
278 concentrations in dogs with primary IMHA (n = 19) when compared with dogs that had other
279 inflammatory diseases (n = 22) or healthy dogs (n = 32).⁴⁷ In question 4, none of the survey respondents
280 indicated that they recommended these protein assays to help diagnose patients with IMHA.

282 Thrombon and Coagulation

283 Thrombocytopenia is reported to occur in 29-70% of dogs with IMHA.^{12,14,15,26,38,39} In a study of 151 dogs,
284 a platelet concentration < 150,000 platelets/ μ L correlated with decreased survival rates.¹³ Also, a
285 decreased mean platelet component concentration was found in dogs with IMHA (n = 95) as compared
286 with healthy dogs (n = 95) or sick canine patients (n = 95)⁴⁸ which could indicate increased platelet
287 activation in IMHA patients.⁴⁹

288
289 Considerations for severe thrombocytopenia include a consumptive process [e.g., disseminated
290 intravascular coagulation (DIC), pulmonary thromboembolism (PTE)] or a concurrent immune-mediated
291 thrombocytopenia (IMT). In humans, concurrent IMHA and IMT have been termed Evan's Syndrome.
292 This disease process likely occurs in dogs; however, the presence of concurrent anti-erythrocyte and
293 anti-platelet antibodies has rarely been documented in veterinary patients.^{50,51} In 38 dogs with both
294 anemia and thrombocytopenia, 18/38 (47%) of patients were positive by DAT for anti-erythrocyte
295 antibodies.⁵² In a similar study of 21 dogs with concurrent anemia and thrombocytopenia, auto-
296 agglutination that persisted after washing was observed 6/21 (29%) dogs, and two of three dogs tested
297 by DAT were positive.⁵³

298
299 Several studies have assessed coagulation parameters in dogs with IMHA (**Table 3**). Importantly,
300 increased mortality was observed in dogs with IMHA that had thrombocytopenia, prolonged
301 prothrombin time (PT), prolonged activated partial thromboplastin time (APTT), decreased fibrinogen,
302 or DIC.⁴⁰ Reports using thromboelastography determined that 85-100% of dogs with IMHA were
303 hypercoagulable.⁵⁴⁻⁵⁶ Development of DIC is observed in between 10/31 (28%)¹⁴ and 9/20 (45%)¹⁵ dogs
304 with IMHA. One study reported that thromboemboli were found in 20/25 (80%) IMHA dogs at
305 necropsy.¹⁴ The analysis of coagulation parameters is typically recommended in IMHA patients that
306 have clinical signs of coagulopathy. One survey respondent indicated in Question 4 that they
307 recommended measurement of D-dimers in patients suspected of having IMHA.

308 309 Indirect Antiglobulin Tests

310 Indirect antiglobulin tests are not recommended in veterinary species due to low sensitivities and
311 specificities. When these studies were first evaluated for utility in canine patients with IMHA, the
312 sensitivity and specificity of an indirect antiglobulin test were 62.5% and 96.6%, respectively.⁵⁷ The DAT
313 performed by the same laboratory had a 83.3% sensitivity and 98.8% specificity.⁵⁷ None of the survey
314 participants recommended indirect antiglobulin tests in Question 4.

315

316 Direct Antiglobulin Tests (DATs)

317 Various methods for directly detecting RBC surface-bound anti-erythrocyte immunoglobulin (Ig) and
318 opsonizing complement protein (C3) are available. Of the 72 respondents who recommended a test to
319 diagnose IMHA with anti-erythrocyte antibodies, 66 recommended a Coombs' test, three recommended
320 flow cytometry and a Coombs' test, two recommended flow cytometry alone, and one recommended
321 flow cytometry and a Coombs' test at 4°C.

322

323 *Coombs' Tests*

324 Coombs' tests are often performed using a microtiter plate format. Additional methods include gel-
325 based microcolumn, immunochromatographic strip, and capillary DAT assays. Good agreement has been
326 reported between results of the Coombs' test and these methods.²⁵ False-positive DAT results have
327 been reported in anemic dogs. In theory, false-positive results could be due to technical difficulties (e.g.,
328 nonspecific absorption of the antibody, incomplete washing, contamination, assignment of an
329 inappropriate cut off) but patient factors are critical to consider. Dogs that recently received a
330 transfusion can have a positive DAT.^{8,58} Also, dogs with an autoimmune disease that are positive for
331 antinuclear antibodies have been reported to be DAT positive without conclusive evidence of IMHA.⁵⁹
332 Additionally, horses with equine infectious anemia can have a positive Coombs' test.⁶⁰

333

334 The Coombs' test uses species-specific antibodies to detect Ig and/or C3 bound to erythrocytes in a
335 patient blood sample. A positive test results in RBC agglutination. Sixty-five of the people surveyed
336 specified if they recommended a Coombs' test when auto-agglutination was observed, 50 people
337 recommended a Coombs' test if no auto-agglutination was seen (7 specified use of a microtiter plate,
338 and 2 recommended an immunochromatographic strip DAT assay). Fifteen people recommended a
339 Coombs' test with or without auto-agglutination (1 specified the use of a microtiter plate, and another
340 recommended the use of a strip DAT assay). The authors agree with the recent ACVIM consensus
341 statement,⁷ which indicates that a Coombs' test is unnecessary if true auto-agglutination that persists
342 after washing is present.

343

344 When a Coombs' test is warranted, polyvalent and monovalent test antibodies are available. These
345 antibodies are pre-adsorbed onto RBCs from healthy dogs before use in the test. False-negative results
346 can occur with either type of test antibody; therefore, including both polyvalent and monovalent

347 antibodies in a Coombs' test can be beneficial.¹¹ Including both polyvalent and monovalent antibodies
348 increased test performance in a study that reported a sensitivity of 82% and a specificity of 95% when
349 antibodies were combined.⁸

350
351 The antibody binding reaction is temperature-dependent, so it is recommended that testing is
352 performed at both 37°C and 4°C.¹¹ Thirty people specified that they ran a Coombs' test at 37°C, 11
353 respondents performed a Coombs' test at 37°C and 4°C, 1 respondent performed the test at 37°C and
354 room temperature, and (as mentioned above) one person performed flow cytometry plus a Coombs' test
355 at 4°C.

356
357 Also, prozone effects are commonly reported when agglutination is not observed at low serum dilutions
358 (i.e., high antibody concentrations) but is observed at higher serum dilutions (i.e., low antibody
359 concentrations). This is due to the presence of excess immunoglobulins that interfere with agglutination
360 induced by the interactions of the test antibodies with Ig and C3 on the RBCs.⁶¹ This improper test
361 antibody to anti-RBC Ig ratio leads to false-negative results if there are not enough serial serum dilutions
362 tested. A prozone effect was observed in 17/126 (13%) samples tested by Piek *et al.* (2012).⁶²

363
364 The reported performance of Coombs' tests vary, likely due to different samples, protocols, and test
365 reagents that are used at different laboratories. Important positive control samples for the Coombs' test
366 include Ig-coated and complement-coated canine RBCs, but these reagents are not readily available.⁹ It
367 has been reported that the use of whole blood in EDTA or acid citrate dextrose (ACD) yields similar
368 results; however, ACD anticoagulants were preferred in one study because of increased sample
369 hemolysis in EDTA.⁹ False-negative results that reduce the sensitivity, and negative predictive value
370 (NPV) of the Coombs' test can be caused by physical properties of the test antibodies (e.g., low antibody
371 affinity, an inappropriate antibody ratio, steric hindrance), poor technique (e.g., excessive washing,
372 delayed processing, assignment of an inappropriate cut-off), or patient factors (drug-dependent
373 reactions, blood transfusions, steroid administration).⁸

374
375 In dogs with IMHA, one study reported that a low percentage, 17/46 (37%), of patients had positive
376 Coombs' test results,¹² but other studies indicated that 77% of dogs with IMHA were positive.^{13,14} In a
377 small study of 12 dogs, the sensitivity, specificity, positive predictive value (PPV), and NPV of the
378 Coombs' test was 58%, 100%, 100%, and 62%, respectively.⁶³ Similarly, Quigley *et al.* (2001) calculated a

379 PPV of 100% and an NPV of 68%.⁶⁴ In a study of cats with IMHA, 2/89 (2%) healthy cats were reported
380 to have a strongly positive Coombs' test at 37°C.⁶⁵ Another study reported that 0/14 (0%) nonanemic
381 cats and 18/55 (33%) anemic cats were Coombs' positive.⁶⁶ Of the 18 Coombs' positive cats, 15 were
382 diagnosed with primary IMHA, two were feline leukemia virus positive (FeLV), and one had
383 cholangiohepatitis.⁶⁶ An older manuscript indicated a weak positive Coombs' test at 4°C in 9/20 (45%)
384 healthy cats and a positive Coombs' test at 4°C and 37°C in 16/20 (80%) anemic cats (12 of the Coombs'
385 positive anemic cats were FeLV positive).⁶⁷

386
387 It is expected that transfusion reactions can cause positive DAT results.^{58,68,69} Honeckman *et al.*
388 indicated that transfusions given 3 to 21 days before a Coombs' test could be particularly problematic.⁵⁸
389 However, only one study was found that reported DAT test results in seven dogs that had been given a
390 recent transfusion; results of two DAT test kits were reported.⁸ Samples were interpreted as truly
391 positive for five dogs (by at least one Coomb's test kit), falsely positive for one dog, and falsely negative
392 for one dog. This study did not specify the length of time between when the transfusion was given and
393 when the diagnostic testing was done. Interestingly, in humans, blood typing is recommended either
394 prior to transfusion or a minimum of 3 months after transfusion to avoid erroneous results.⁷⁰

395
396 Ninety-one of the 94 survey respondents indicated if they would interpret any diagnostic results with
397 caution following a transfusion. The timeframe of concern for people who interpreted results with more
398 caution varied (Fig. 3). It is evident from this small survey that there is uncertainty if recent transfusion
399 would cause false-positive results in CBC, agglutination, or DAT assays.

400 401 *Flow Cytometry Methods*

402 Flow cytometry can also be used to detect immunoglobulins bound to RBCs. One of the first evaluations
403 of flow cytometry as a diagnostic test for IMHA compared data from 12 dogs and three horses with
404 IMHA to 12 healthy animals from each respective species.⁶³ They reported low specificity of a goat anti-
405 equine IgG but 100% specificity of goat anti-equine IgG F(ab')₂ fragment in their assay.⁶³ In dogs, by
406 pooling anti-canine IgG, IgM, IgA, and C3 antibodies, the sensitivity of the test was 100%, specificity was
407 87.5%, PPV was 92%, and NPV was 100%.⁶³ There was no prozone effect with this assay.⁶³ Quigley *et al.*
408 (2001) reported a PPV of 100%, and a NPV of 93% when they used flow cytometry to evaluate 13 dogs
409 with IMHA and 13 healthy dogs.⁶⁴ In 2008, Morley *et al.* published an assessment of the utility of flow
410 cytometry to detect anti-erythrocyte antibodies in dogs.⁷¹ They found 26/147 (18%) anemic patients

411 had detectable anti-erythrocyte antibodies. This included 17/22 (77%) IMHA patients, 5/14 (36%) IMT
412 patients, and 3/71 (4%) cancer patients. However, 12/145 (8%) nonanemic dogs also were positive for
413 anti-erythrocyte antibodies, which included 3/5 (60%) patients with infectious disease and 5/81 (6%)
414 cancer patients, and the test had a PPV of 70% and NPV of 95%.

415

416 *Direct Enzyme-Linked Antiglobulin Tests (DELATs)*

417 Immunoglobulins and C3 bound to RBCs are detected by comparing the absorbance of patient samples
418 to healthy control animals in the DELAT. Early evaluation of a DELAT indicated that 1 mg/mL p-
419 nitrophenyl phosphate in carbonate buffer is the preferred substrate for the reaction, however false-
420 positive results were observed in 31/60 (52%) dogs tested.⁷² Another evaluation of DELAT performance
421 yielded comparable results with a Coombs' test; 12/23 (52%) samples were Coombs' positive, while
422 13/23 (57%) were DELAT positive.⁷³ To the authors' knowledge, this test for IMHA is no longer readily
423 available.

424

425 **Guidelines for Performing Diagnostic Tests for IMHA**

426 As mentioned previously, technical difficulties and test protocols can profoundly affect the performance
427 of diagnostic tests for IMHA. Tests performed at 4°C, room temperature, and 37°C can provide different
428 results. This document includes two example protocols for saline agglutination testing (Appendix B) and
429 an example of Coombs' testing (Appendix C). In different diagnostic laboratories, it is expected that
430 these protocols will be performed differently but that proper procedures will yield adequate results.

431

432 **Conclusions**

433 Making a definitive diagnosis of IMHA can be difficult due to the variability in patient presentation and
434 diagnostic test performance. Recommended tests for diagnosing IMHA in anemic patients include: 1) a
435 CBC with verified reticulocyte count, manual PCV with assessment of plasma color, and microscopic
436 examination of a (preferably fresh) blood smear, 2) serum chemistry profile, 3) urinalysis, 4) saline
437 agglutination test (preferably with washing), 5) Coombs' test or flow cytometric analysis if the saline
438 agglutination test is negative, 6) coagulation testing for thrombocytopenic patients and patients with
439 clinical signs of coagulopathy, and 7) tests to determine if any underlying disease is present (e.g., drug or
440 toxin exposure, infection, inflammation, neoplasia, other autoimmune diseases). Our survey proved that
441 clinicians choose different tests to diagnose IMHA and that laboratories perform tests differently.
442 Therefore, it is recommended that veterinarians contact a clinical pathologist or technicians or

443 technologist at the diagnostic laboratory they use to obtain details about the reliability of specific tests
444 being performed to diagnose IMHA.

445

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644

645 **Table 1.** Clinical pathology findings that support the diagnosis of hemolytic anemia.⁶

CBC/Blood smear exam	Serum Biochemistry	Urinalysis
Increased polychromasia	Hyperbilirubinemia	Hyperbilirubinuria
Reticulocytosis ± rubricytosis ± Howell-Jolly bodies	Hemoglobinemia ⁱ	Hemoglobinuria ⁱ
Macrocytosis & anisocytosis	Decreased haptoglobin concentration ⁱ	Increased urobilinogen
Decreased MCHC ^e or increased MCHC ⁱ	Decreased hemopexin concentration ⁱ	
Spherocytes ^e ; ghost cells ⁱ – immune-mediated damage or other cause		
Heinz bodies ^e ; eccentrocytes ⁱ – oxidative damage		
Hemoparasites – direct physical &/or immune-mediated damage		
Schistocytes ⁱ ; acanthocytes ⁱ ; keratocytes ⁱ – direct physical damage		

646 e = specifically associated with extravascular hemolysis

647 i = specifically associated with intravascular hemolysis

648 MCHC = mean cell hemoglobin concentration

649

650 **Table 2.** Serum biochemistry results associated with decreased survival in IMHA patients.

Abnormality	Species	References
Hyperbilirubinemia	Dog	12-14,40
	Cat	37
Hyperlactatemia	Dog	74
Increased alanine aminotransferase	Dog	40
Increased aspartate aminotransferase	Dog	40

Increased urea nitrogen	Dog	40
Increased alkaline phosphatase	Dog	12
Hypoalbuminemia (< 3.0 mg/dL)	Dog	13
Hypophosphatemia (< 3.5 mEq/L)	Dog	13
Increased creatine kinase (> 250 U/L)	Dog	13

651

652 **Table 3.** Evidence of coagulation abnormalities in dogs with IMHA.

Abnormality	Dogs with IMHA affected	References
Increased prothrombin time	10-28%	14,15
Increased activated partial thromboplastin time	45-47%	14,15
Increased fibrin degradation products	57-60%	14,15
Increased fibrinogen	17/20 (85%)	15
Increased D-dimers	16/20 (80%)	15
Decreased anti-thrombin III	10/20 (50%)	15
Increased Russell viper venom time	7/20 (35%)	15
Increased von Willebrand factor associated antigen	9/20 (45%)	15
Increased Kaolin clotting time	3/20 (15%)	15
Increased P-selectin*	15/20 (75%)	75
Hypercoagulability using thromboelastography	85-100%	54-56
Disseminated intravascular coagulation	28-45%	14,15
Thromboemboli found at necropsy	20/25 (80%)	14

653 * A second paper saw no increase in P-selectin.¹³

654 **Figure Legends:**

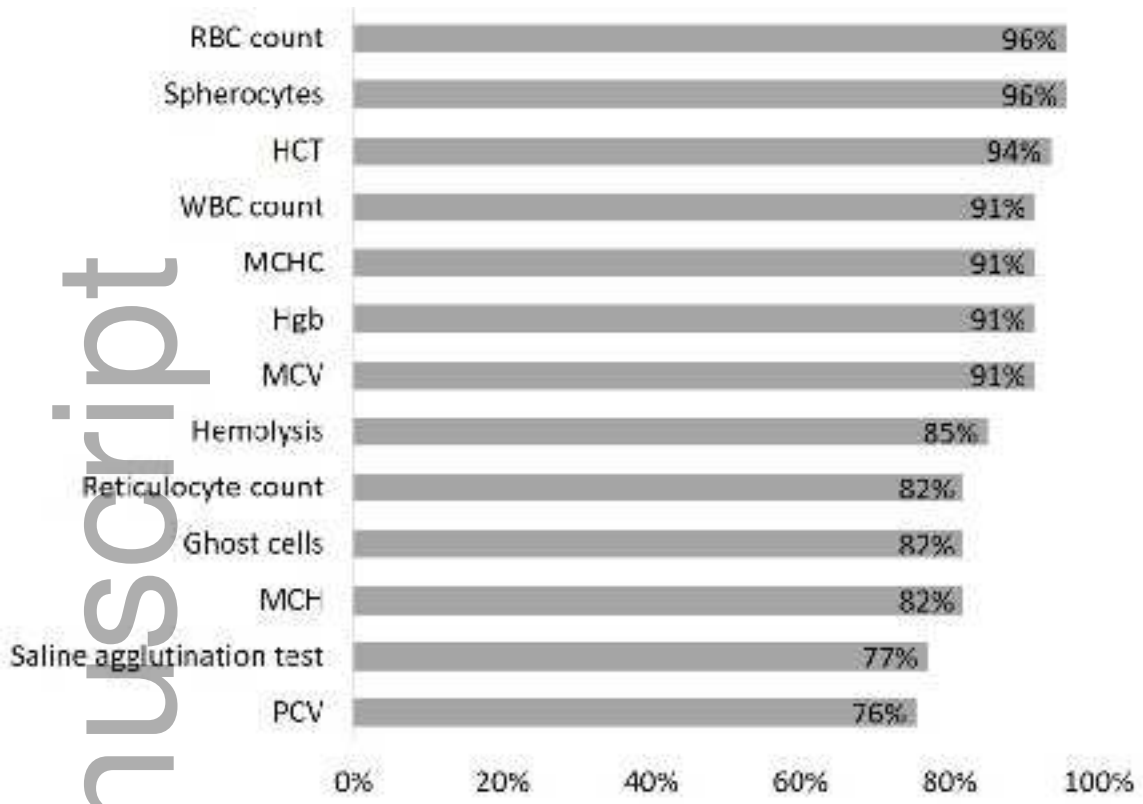
655 **Figure 1.** Proportions (above 75%) of hematologic parameters reported or received on the CBC results of
656 patients suspected of having IMHA by survey respondents (n=94).

657

658 **Figure 2.** Microscopic evidence of agglutination in a wet-mount saline agglutination test. Peripheral
659 blood in EDTA (0.15%) was diluted 1:4 in isotonic saline (0.9% NaCl). A drop of the mixture was placed
660 on a glass slide, and a coverslip was placed over the drop. Grape-like aggregates of erythrocytes can be
661 observed (unstained, 200× magnification).

662

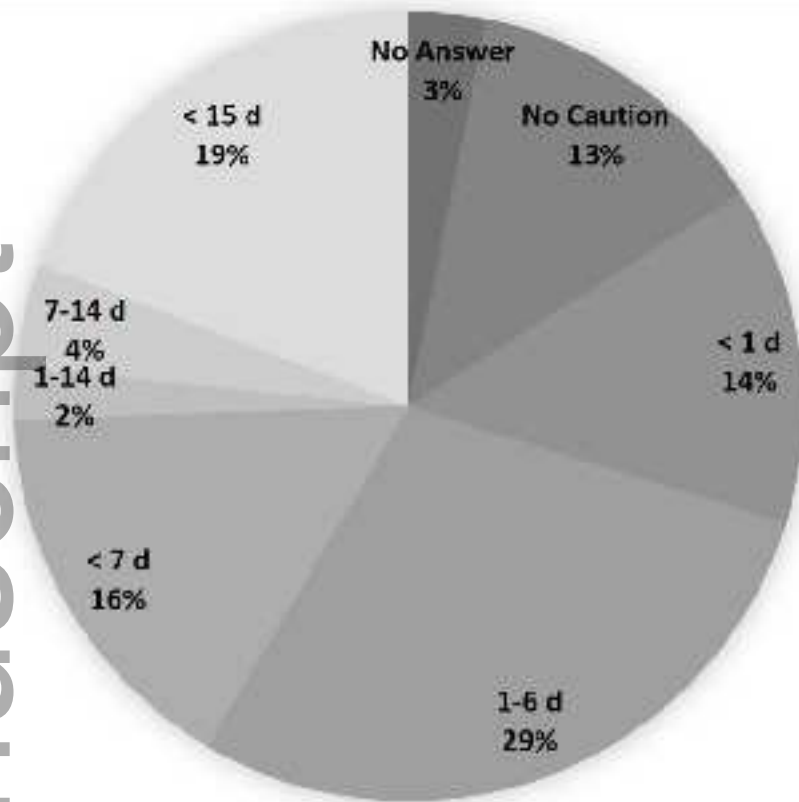
663 **Figure 3.** Survey responses indicating the post-transfusion time-frame in days (d) during which
664 respondents would cautiously interpret hematologic test results that support a diagnosis of IMHA
665 (n=94).



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