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The usefulness of dermoscopy in canine pattern alopecia: a descriptive study

Zanna, G ; Roccabianca, P ; Zini, E ; Legnani, S ; Scarpella, F ; Arrighi, S ; Tosti, A

Abstract: **BACKGROUND:** Dermoscopic studies evaluating noninflammatory, nonpruritic progressive alopecia attributable to pattern alopecia are currently unavailable. **HYPOTHESIS/OBJECTIVES:** To evaluate the dermoscopic features observed in healthy skin of short coated dogs and compare these findings with those observed in dogs affected by pattern alopecia diagnosed by clinical and dermatopathological examination. **ANIMALS:** Thirty male and female, healthy, breed matched, young adult, short coated dogs (controls) and 30 male and female, young adult, short coated dogs affected by pattern alopecia. **METHODS:** Dermoscopy was performed with a Fotofinder II videodermoscope equipped with software that allowed the measurement of structures visualized in magnified images (20×-40×-70×). Skin biopsy samples were obtained from the thorax and evaluated dermoscopically for dermoscopic-histological correlation in affected dogs. **RESULTS:** Dermoscopic findings in canine pattern alopecia were hair shaft thinning, circle hairs and follicular keratin plugs; in the affected sun exposed areas there was a honeycomb-like pattern of pigmentation. Arborizing red lines reflecting vascularization were classified as a nonspecific finding because they were also common in healthy dogs. Dermoscopic features correlated with histology for selected hair follicle abnormalities. **CONCLUSIONS AND CLINICAL IMPORTANCE:** Although canine pattern alopecia is a visually striking disease, this study supports the value of dermoscopy for clinical examination and also opens promising perspectives for the identification of diagnostic dermoscopic patterns that may be useful for other skin disorders.

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1 **Dermoscopy in dogs: an absorbing perspective in evaluation of pattern** 2 **alopecia**

3
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25 Conflict of interest: No conflicts of interest have been declared.

26 27 **Abstract**

28
29 **Background** - No dermoscopic studies evaluating non-inflammatory, non-pruritic
30 progressive alopecia attributable to pattern alopecia are currently available.

31 **Hypothesis/objectives** - To evaluate the dermoscopic features observed in healthy
32 skin of short-coated dogs and compare these findings with those observed in dogs
33 affected by pattern alopecia diagnosed by clinical and dermatopathologic
34 examination.

35 **Animals** - Thirty healthy breed-matched young-adult short-coated dogs, both
36 females and males, were used as controls for the dermoscopic evaluation of 30
37 young-adult short-coated dogs of both genders affected by pattern alopecia.

38 **Methods** - Dermoscopy was performed with the Fotofinder II videodermoscope
39 equipped with software that allowed the measurement of structures visualized in
40 magnified images (20x-40x-70x). Skin biopsy samples were taken at sites evaluated
41 dermoscopically for dermoscopic-histological correlation in affected dogs.

42 **Results** - Dermoscopically, canine pattern alopecia was characterized by hair shaft
43 thinning, circle hairs, follicular keratin plugs, and, in the affected sun-exposed areas,
44 by honeycomb-like pattern pigmentation. Arborizing redlines reflecting
45 vascularization were classified as a non-specific finding because they are common
46 also in healthy dogs. Dermoscopic features correlated with histology for selected hair
47 follicle abnormalities.

48 **Conclusions and clinical importance** - Although canine pattern alopecia is a
49 visually striking disease, this study supports the value of dermoscopy for clinical

50 examination and opens promising perspectives for the identification of diagnostic
51 dermoscopic patterns that may be useful for other skin disorders as well.

52

53 Source of funding: ECVD-ESVD practioner grant 2013

54 I declare that the conflict of interests of each author are declared: Yes

55

56 **Introduction**

57

58 According to Stolz *et al*, skin surface microscopy dates back to 1663, when Johan
59 Kolhaus first looked at nail fold vessels with a microscope.¹ Nevertheless, it was only
60 at the end of the last century that several diagnostic methods were developed
61 utilizing surface microscopy. Today, the upcoming evidence for *in vivo* diagnosis is
62 represented by dermoscopy originally used to observe and diagnose pigmented skin
63 lesions such as melanocytic nevus and melanoma,²⁻⁴ and trichoscopy as hair and
64 scalp dermoscopy.⁵⁻⁸ This latter technique has been used to visualize normal hairs
65 and assess their number per follicular unit, to distinguish whether hair follicle
66 openings are normal, empty, fibrotic or containing biological material as
67 hyperkeratotic plugs, and to study the appearance of perifollicular epidermis and
68 cutaneous microvessels.⁹ Therefore, trichoscopy has proved relevant in the
69 differentiation of cicatricial from non-cicatricial alopecias. As a large group of
70 disorders characterized by permanent destruction of hair follicles, cicatricial alopecia
71 shows trichoscopic features such as loss of follicular ostia and presence of fibrous
72 tracts that mark extinct follicles.¹⁰ On the other hand, in all non-cicatricial alopecias
73 as alopecia areata and androgenetic alopecia (male and female pattern alopecia),
74 suggestive trichoscopic findings are represented by specific hair shaft and follicular
75 opening abnormalities.¹¹⁻¹³

76 Differently from background in humans, to date only a few studies on the application
77 of dermoscopy exist in veterinary medicine and mainly in feline dermatology.¹⁴⁻¹⁶

78 Moreover, except for an abstract regarding the dermoscopic features of 35 dogs with
79 juvenile-onset demodicosis and 35 breed- and age-matched dogs,¹⁷ the authors are
80 unaware, to the best of their knowledge, of any dermoscopic study on canine non-
81 inflammatory alopecia. Therefore, the purpose of this project was twofold. The first
82 aim was to evaluate dermoscopic features observed in short-coated healthy dogs
83 and compare these findings with those observed in short-coated dogs affected by
84 pattern alopecia diagnosed by clinical and dermatopathological examination. The
85 second aim was to assess whether dermoscopic findings correlated or agreed with
86 those observed at histopathology in order to generate dermoscopic criteria that would
87 be useful for the diagnosis of pattern alopecia.

88

89

90 **Material and methods**

91

92 *Study population*

93

94 A population of 30 healthy short-coated dogs was matched with 30 short-coated dogs
95 referred for non-inflammatory, non-pruritic progressive alopecia attributable to pattern
96 alopecia. Details about both groups are presented in Table 1. Dogs were owned by
97 amateur pet breeders or clients, and informed owner consent was obtained prior to

98 any procedure. Dogs were selected on the basis of the following criteria: (i) no other
99 clinical abnormalities at physical examination; (ii) except for pattern alopecia, no
100 evidence of additional skin lesions on dermatological examination; (iii) for intact
101 female dogs, not being pregnant or lactating; and (iv) normal complete blood count
102 and routine serum biochemical analysis.

103

104 *Dermoscopic examination*

105

106 A videodermoscope (Fotofinder® TeachScreen Systems software GmbH Bad
107 Birnbach, Germany) was used and six body sites including convex pinnae, periaural
108 area, ventral neck, thorax, abdomen and caudal thighs were selected. Alcohol
109 (Kodan® spray, Schulke & Mayr, Vienna, Austria) was applied as interface solution to
110 better observe surface and subsurface microscopic features.

111 In order to take a dermoscopic overview image of the selected cutaneous region,
112 images at 20-fold and 40-fold magnification were first observed. Then, as previously
113 reported by Rakowska *et al.*,¹¹ images at 70-fold magnification, which allows a high-
114 quality enlargement of 9 mm² of the skin area to the size of the computer screen,
115 were used for statistical purposes. An area of 3.14 mm² was calculated on the
116 selected 70-fold images by means of the FotoFinder® software, and dogs with
117 pattern alopecia and controls were compared for the following parameters: diameter
118 and total number of hair tufts next to follicular ostia per examined area, total number
119 of hairs per hair tuft plus the ratio between the number of secondary hairs/primary
120 hair, and diameter of both primary and secondary hairs in each hair tuft. Hair follicle
121 infundibula, perifollicular epidermis and vascular structures such as very small
122 capillaries were also observed.

123

124 *Dermoscopy vs. histopathology*

125

126 To contrast dermoscopy and histopathology, in 20 of the affected dogs a single skin
127 biopsy taken from the thoracic skin area previously circled with a marker during
128 dermoscopic examination was collected under local anaesthesia using a 4-mm skin
129 biopsy punch. The biopsies were fixed in 10% neutral buffered formalin, trimmed,
130 routinely processed, and paraffin embedded. Transverse serial sections (4 µm thick)
131 were obtained and stained with haematoxylin and eosin for histological examination.
132 Histological images were observed under an Olympus BX51 photomicroscope
133 equipped with an Olympus C-5060 Wide Zoom and DP software digital camera
134 (Olympus, Tokyo, Japan) for computer-assisted image acquisition and analysis. The
135 slides contained multiple transverse sections of the skin at different levels starting
136 from the panniculus and ending with the *stratum corneum*. For hair follicle number
137 assessment, transverse skin sections were examined at the level of the mid/lower
138 isthmus. The total number of follicular units per examined area and number of total
139 hairs per follicular unit were counted.

140 Other parameters included infundibular hyperkeratosis evaluated in the superficial
141 slides at the level of the infundubulus in cross section; vascularization scored in the
142 same slides used to examine infundibular hyperkeratosis; and pigment clumping
143 evaluated in overall sections and scored according to severity of clumping in bulbs
144 and hair shafts. All these findings were graded as - (absent), + (mild), ++ (moderate),
145 +++ (severe).

146

147 *Statistical analyses*

148
149 To assess whether dogs with and without pattern alopecia were correctly matched for
150 age and body weight, the Mann-Whitney test was used, and for sex and hair colour
151 the Fisher's exact test and $r \times c$ contingency table were used, respectively, within
152 each of the 3 breeds. For each breed investigated, dogs with pattern baldness and
153 controls were compared for the measured parameters on the six body selected
154 regions described above. The analysis was performed using the Mann-Whitney test
155 followed by Bonferroni correction. Furthermore, the same hair parameters were
156 compared between regions within each dog breed for those with and without pattern
157 alopecia, using the Friedman test followed by Dunn's multiple comparison. To assess
158 whether dermoscopic examination yielded similar results to histology, the
159 Spearman's rank correlation coefficient was calculated between the total number of
160 hair tufts next to follicular ostia per examined area based on the former method and
161 the total number of follicular units per examined area counted with the latter. The
162 same test was also used to verify whether the total number of hairs per hair tuft with
163 dermoscopy correlated with the total number of hairs per follicular unit identified with
164 histology. Significance was considered for $P < 0.05$. In addition, the Cohen's kappa
165 coefficient was used to assess whether there was agreement between the two
166 methods in the analysis of infundibular hyperkeratosis, vascularization, and pigment.
167 Kappa values < 0 indicated no agreement and 0-0.20 as slight, 0.21-0.40 as fair,
168 0.41-0.60 as moderate, 0.61-0.80 as substantial, and 0.81-1 as almost perfect
169 agreement. Software package was used for analysis (GraphPad Prism version 5.0,
170 GraphPad Software, La Jolla, CA, USA).

171 172 **Results**

173 174 *Group matching*

175
176 Population characteristics did not differ statistically in any of the 3 breeds between
177 dogs with pattern alopecia and controls, suggesting appropriate matching.

178 179 *Dermoscopic features*

180
181 On dermoscopy, normal values were considered: hair shafts grouped into follicular
182 units consisting of thick hairs emerging independently from their follicular ostia and
183 considered as primary hairs, and surrounded by a variable number of thinner hairs all
184 protruding through a common external orifice and considered as secondary hairs.
185 Observed were hair follicle openings that were not empty, fibrotic or filled with
186 material such as keratotic plugs; no scaling on perifollicular and interfollicular skin
187 surface; and thin arborizing red lines corresponding to vessels between follicular
188 units. In dogs with dilute hair colour, pinpoint black spots were also observed on
189 interfollicular skin surface.

190 In dogs affected by pattern alopecia, the most common dermoscopic findings were:
191 hair shaft thinning; scattered circle hairs; plugging of the follicular infundibulum with a
192 yellow-brown material, and on periaural and caudal thigh regions, a honeycomb-like
193 pigmented network. As in controls, pinpoint black spots on interfollicular skin surface
194 of dogs with dilute hair colour and thin arborizing vessels regularly distributed
195 between follicular units were also detected. All these findings are illustrated in Figure
196 1 (a-f).

197

198 *Histological findings*

199

200 In transverse histological sections taken from the thoracic region, hair follicles were
201 characterized by moderate to severe decrease in size (Figure 2a) without distortion
202 or irregularity of their contour or reduction of the overall number of adnexal units
203 (Figure 2b). Infundibular hyperkeratosis and melanin clumping were also variably
204 observed, whereas in some areas, vessels appeared more prominent but were not
205 increased in number.

206

207 *Dermoscopic parameters in dachshunds*

208

209 Comparing dachshunds with pattern alopecia and controls, the following significant
210 differences were documented: i) the median diameter of hair tufts next to follicular
211 ostia was smaller in those with pattern alopecia than controls in the convex pinnae
212 (0.05 mm; range 0.03-0.07 vs 0.08 mm; range: 0.06-0.09; $P < 0.001$), ventral neck
213 (0.07 mm; range 0.04-0.09 vs 0.08 mm; range: 0.07-0.11; $P < 0.01$), chest (0.06 mm;
214 range 0.05-0.08 vs 0.08 mm; range: 0.06-0.11; $P < 0.05$) and abdominal region (0.06
215 mm; range 0.05-0.09 vs 0.08 mm; range: 0.06-0.11; $P < 0.01$); and ii) the median
216 diameter of primary hairs was smaller in those with pattern alopecia in the ventral
217 neck (0.03 mm; range 0.02-0.04 vs 0.04 mm; range: 0.02-0.05; $P < 0.05$) and chest
218 (0.03 mm; range 0.01-0.04 vs 0.04 mm; range: 0.03-0.05; $P < 0.01$). No other
219 differences were documented between groups. In dachshunds with pattern alopecia
220 there was a significant difference in the ratio between the number of secondary
221 hairs/primary hair; in particular, the periaural region had a higher median ratio (7;
222 range: 4-14) than the abdominal region (5; range: 2-8; $P < 0.001$). No other
223 differences were documented for the hair tuft parameters in any region. In controls
224 there were significant differences in the diameter of hair tufts next to follicular ostia
225 and in the diameter of primary hairs; specifically, the periaural region had a smaller
226 median diameter of hair tufts located next to follicular ostia (0.07 mm; range: 0.04-
227 0.08) than the ventral neck (0.08 mm; range: 0.07-0.11; $P < 0.01$), the chest (0.08
228 mm; range: 0.06-0.11; $P < 0.01$) or abdominal region (0.08 mm; range: 0.06-0.11; P
229 < 0.05), while the periaural region had a smaller median diameter of primary hairs
230 (0.03 mm; range: 0.02-0.03) than either the ventral neck (0.04 mm; range: 0.02-0.05;
231 $P < 0.01$) or chest (0.04 mm; range: 0.03-0.05; $P < 0.01$). All these results are
232 summarized in Table 2.

233

234 *Dermoscopic parameters in Italian greyhounds*

235

236 Between Italian greyhounds with pattern alopecia and controls, the median diameter
237 of hair tufts next to follicular ostia was smaller in those with pattern alopecia (0.05
238 mm; range 0.04-0.07) than controls (0.07 mm; range: 0.07-0.08; $P < 0.01$) in the
239 ventral neck. No other differences were documented for the hair tuft parameters in
240 any region. In Italian greyhounds with pattern alopecia there were no significant
241 differences between the 6 regions for any of the 4 hair tuft parameters. Similarly, in
242 controls there were no significant differences. All these results are summarized in
243 Table 2.

244

245 *Dermoscopic parameters in miniature pinschers*

246

247 Between miniature pinschers with pattern alopecia and controls, the following
248 significant differences were documented: i) the median diameter of hair tufts next to
249 follicular ostia was smaller in those with pattern alopecia than controls in the convex
250 pinnae (0.05 mm; range 0.05-0.05 vs 0.08 mm; range: 0.06-0.10; $P < 0.001$), ventral
251 neck (0.05 mm; range 0.04-0.07 vs 0.08 mm; range: 0.07-0.08; $P < 0.05$) and caudal
252 thigh region (0.05 mm; range 0.05-0.06 vs 0.07 mm; range: 0.06-0.08; $P < 0.05$); ii)
253 the median diameter of secondary hairs was smaller in those with pattern alopecia
254 than controls in the convex pinnae (0.01 mm; range 0.01-0.01 vs 0.02 mm; range:
255 0.01-0.02; $P < 0.01$), ventral neck (0.01 mm; range 0.01-0.01 vs 0.02 mm; range:
256 0.02-0.02; $P < 0.001$) and chest region (0.01 mm; range 0.01-0.01 vs 0.02 mm;
257 range: 0.01-0.02; $P < 0.01$). No other differences were documented between groups.
258 Within pinschers with pattern alopecia there were no significant differences between
259 the 6 regions for any of the 4 hair tuft parameters. In contrast, in controls there were
260 significant differences in the ratio between the number of secondary hairs/primary
261 hair; in particular, the convex pinnae had a higher median ratio (9; range: 8-11) than
262 either the chest (5; range: 5-6; $P < 0.05$) or caudal thigh (5; range: 4-6; $P < 0.01$). All
263 these results are summarized in Table 2.

264 *Dermoscopy vs. histopathology*

266
267 Dermoscopic and histologic findings are presented in Table 3. A very strong positive
268 correlation was observed for the total number of hair tufts next to follicular ostia
269 based on dermoscopy and the total number of follicular units per examined area
270 counted with histology ($\rho=0.898$; 95% CI=0.750-0.961; $P < 0.001$), and the total
271 number count of hairs per hair tuft at dermoscopy and total number of hairs per
272 follicular unit identified at histology ($\rho=0.868$; 95% CI=0.683-0.948; $P < 0.001$)
273 (Figure 3). A fair agreement was observed between dermoscopy and histology for
274 the analysis of follicular hyperkeratosis ($\kappa=0.333$; 95% CI=0.013-0.679), with
275 only 12 of 20 (60%) agreements; a fair agreement was observed for the analysis of
276 vascularisation ($\kappa=0.200$; 95% CI=0.120-0.520), with only 9 of 20 (45%)
277 agreements; a moderate agreement was observed for the analysis of pigment
278 ($\kappa=0.294$; 95% CI=0.032-0.556), with only 11 of 20 (55%) agreements.

279 **Discussion**

281 In this study, dermoscopic findings in dogs affected by pattern alopecia have been
282 characterized for the first time, highlighting the value of dermoscopy as an adjunctive
283 technique for cutaneous clinical examination.
284 Canine pattern alopecia is a relatively common but poorly studied skin disorder
285 somehow similar to, but also clearly different from, human androgenetic alopecia.¹⁸
286 Fine hairs referred to as miniaturized hairs represent the hallmark clinical
287 presentation of this disorder. However, to the best of the authors' knowledge, *in vivo*
288 measurement of hair shaft thickness based on dermoscopy has not been performed
289 before. In this study, the first hair parameter dermoscopically measured was the
290 median hair tuft thickness diameter next to follicular ostia that was shown to be
291 smaller in all affected dogs compared with controls. This result is not surprising if we
292 consider that the relative thinning of hairs is the most striking feature of the disease.
293 Of note, however, differences between breeds and within the same breed were
294 detected, dependent on other hair parameters accounted for. For example, in
295 affected dachshunds the median ratio between the number of secondary

296 hairs/primary hair was shown to be higher in diseased animals than in controls in all
297 the skin regions evaluated. The periaural region demonstrated the largest number of
298 secondary hairs (7; range: 4-14). Moreover, within the group of dachshund controls
299 the periaural region was demonstrated as having the smallest median diameter of
300 primary hairs (0.03 mm; range: 0.02-0.03) indicating that thinning of hairs in this
301 region may be considered as a normal feature in this breed. In Italian greyhounds,
302 the ventral neck region was described as affected mainly by thinning hairs, and this
303 finding indicates the relevance of this region in distinguishing affected from healthy
304 dogs. In miniature pinschers, secondary hairs were smaller in affected dogs than in
305 controls, mostly in the convex pinnae, ventral neck and chest, whereas in controls,
306 the median ratio between the number of secondary hairs/primary hair was higher in
307 the convex pinnae (9; range: 8-11). All these results taken together reveal that hair
308 shaft thinning in canine pattern alopecia is a process that does not simultaneously
309 affect all hairs of all regions, and that great variability exists between and within
310 affected dog breeds. This variability may be the result of artificial selection pressure
311 for extremely fine haircoats sought by breeders who often attempt to manipulate the
312 appearance of a dog, thereby predisposing it to this presumptively genetic alopecia.¹⁹
313 In humans, androgenetic alopecia is considered an inherited condition caused by a
314 genetically determined hair follicle sensitivity to the effects of dihydrotestosterone,
315 with the result of a gradual shortening of anagen phase and a prolongation of
316 kenogen phase.²⁰⁻²⁴ Increased concentrations of both 5- α reductase isoenzyme and
317 androgen receptor have been detected in the balding scalp, suggesting that such
318 changes contribute to hair loss.²⁵

319 To date, the pathogenesis of canine pattern alopecia is not known and the
320 involvement of an abnormality in hair follicle hormonal receptor is still debated.¹⁹ The
321 alteration of the hair-cycle dynamics with an increase in the prevalence of kenogen
322 follicles has been demonstrated in some canine non-inflammatory alopecias but not
323 in pattern alopecia,²⁶ and the expression of 5- α -reductase genes has been evaluated
324 in only one study, performed in normal skin.²⁷

325 In order to provide both qualitative and quantitative diagnostic follicular information,
326 transverse sections of skin biopsy specimens were used in this study, as in human
327 literature.^{28,29} Some key information such as follicular counts was easily assessed,
328 and histological findings were shown to positively correlate with dermoscopic
329 calculations of hair parameters. However, accurate determination of growth stages of
330 the hair cycle was not possible on transverse sections due to the absence of the
331 entire length of the hair follicle including site, shape and depth of the hair inferior
332 portion and, specifically, of the bulb. Therefore longitudinal sections continue to
333 provide the best morphological and spatial information to assess specific growth
334 stages of hair cycle in dogs.²⁶ Additionally, in human beings hairs are mostly primary
335 while in normal and diseased skin of dogs it is difficult to determine the primary or
336 secondary origin of the follicle, especially without the hair shaft.

337 To detect other dermoscopic features that could differentiate diseased dogs from
338 controls, hair follicle openings, perifollicular and interfollicular skin surface, and
339 vascular structures were dermoscopically examined and evaluated in conjunction
340 with histological findings. Follicular ostia filled with light yellowish or brownish
341 material were mostly observed in the ventral regions of dogs affected by pattern
342 alopecia; this was histologically related to a variable amount of keratin filling the
343 follicular infundibulum. In humans, this dermoscopic finding, termed 'yellow dot',
344 represents sebum mixed with variable amounts of keratin secreted by normal, active
345 sebaceous glands through the miniaturized hair follicle.^{7,9,11} Therefore the result of

346 this process is the accumulation of yellow material at the top of the hair follicular
347 opening. Our hypothesis is that a similar mechanism may occur in canine pattern
348 alopecia.

349 Moreover, in some affected dogs, hairs with typical circular or spiraliform
350 arrangement were dermoscopically observed, but no histological change was
351 identified in relation to this dermoscopic feature. In humans, the pathogenesis of this
352 finding remains obscure, although some authors relate it to hairs with a small
353 diameter that renders it difficult to penetrate the *stratum corneum*. For this reason,
354 they grow in a circular tract and in a subcorneal location.³⁰ Based on this, our
355 dermoscopic finding may have an explanation, but further studies are needed to
356 better understand the pathogenesis of these hairs with this typical arrangement.
357 Variable infundibular melanin clumping in both healthy and affected dogs with dilute
358 hair colour was histologically detected and dermoscopically visualized as pinpoint
359 black spots on interfollicular skin surface.

360 Finally, a honeycomb-like hyperpigmentation pattern, characterized by hyperchromic
361 rings on the skin surface and resulting from solar exposure in thinning or completely
362 balding areas as demonstrated in humans,¹⁰ often coexisted as an additional feature
363 in the periaural and caudal thigh regions.

364 Cutaneous microvessels that arborize into thin red branches in a non-homogeneous
365 fashion were considered as non-specific dermoscopic findings because they are also
366 common in normal skin. Given that dermoscopy enables horizontal inspection of the
367 skin, vessels that run parallel to the skin surface are visualized as lines, while those
368 that run perpendicularly are generally viewed as dots, or even loops.^{31,32} However,
369 they are best evaluated when the pressure exerted by the dermoscope against the
370 skin is low. High outside pressure may indeed reduce blood flow in cutaneous
371 capillaries.¹⁰ In this study, the lack of dermoscopic visualization of cutaneous blood
372 vessels in some selected areas may have resulted from excessive pressure applied
373 to the skin. Translucent ultrasound gel that allows one to apply the lens against the
374 skin gently in order to better visualize blood vessels is expected in future studies.

375 In summary, the results of this study suggest that dermoscopy may provide a new,
376 relevant clinical perspective on hair disorders and offer the clinician a novel way in
377 which to uncover clinical aspects of cutaneous diseases.

378

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383

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465 **Figure legend**

466

467 **Figure 1.** Representative dermoscopic features in dogs affected by pattern alopecia.
468 (a) Diffuse hair thinning (20x). (b) Presence of hair circle (black arrows) between
469 miniaturized hairs (20x). (c) Hair circles (black arrow), thin arborizing vessels (red
470 arrow) and yellow-brown material around follicular ostia (blue arrow) (20x). (d)
471 Plugging of follicular infundibular with yellow-brown material (blue arrows) (70x).
472 Pinpoint black dots in a dog with diluted hair colour (black arrows) (70x).
473 Honeycomb-like pattern (black arrows) on the caudal thigh (70x).

474

475 **Figure 2.** Representative photomicrographs of hair follicle miniaturization in dogs
476 affected by pattern alopecia. (a) Decrease in size of hair follicular units without
477 distortion or irregularity of their contour or reduction of the overall numbers of adnexal
478 units. Scale bar represents 1000 μm . (b) Multiple thinner hair follicles at higher
479 magnification. Scale bar represent 200 μm .

480

481 **Figure 3.** Correlation of the number of hair tufts located next to follicular ostia based
482 on dermoscopy (x-axis) with number of follicular units counted with histology (y-axis).
483 The regression line is shown.

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