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AUTHORS: Anette Loeffler, David H. Lloyd

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1 **Review Article**

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3 **What has changed in canine pyoderma? A narrative review**

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5

6 Anette Loeffler \*, David H. Lloyd

7

8 *Department of Clinical Science and Services, Royal Veterinary College, University of London,*  
9 *United Kingdom*

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14 \* Corresponding author. Tel.: +44 1707 666333.

15 *E-mail address:* [aloeffler@rvc.ac.uk](mailto:aloeffler@rvc.ac.uk) (A. Loeffler).

16 **Abstract**

17 Canine pyoderma remains one of the main presentations in small animal practice and  
18 frequently leads to prescribing of systemic antimicrobials. A good foundation knowledge on  
19 pyoderma was established during the 1970s and 1980s when treatment of infection provided little  
20 challenge. However, our ability to treat canine pyoderma effectively is now limited substantially by  
21 the emergence of multidrug-resistant, methicillin-resistant staphylococci (MRS) and in some  
22 countries, by restrictions in antimicrobial prescribing for pets. The threat from rising antimicrobial  
23 resistance and the zoonotic potential of MRS add a new dimension of public health implications to  
24 the management of canine pyoderma and urge a revisit and the search for new best management  
25 strategies. This narrative review focusses on the impact of MRS on how we manage canine  
26 pyoderma, and how traditional treatment recommendations need to be updated in the interest of  
27 good antimicrobial stewardship. Background information on clinical characteristics, pathogens and  
28 appropriate clinical and microbiological diagnostic techniques are briefly reviewed in so far as they  
29 can support early identification of multidrug-resistant pathogens. We examine the potential of new  
30 approaches for the control and treatment of bacterial skin infections and highlight the role of owner  
31 education and hygiene. Pyoderma patients offer great opportunities for good antimicrobial  
32 stewardship by making use of the unique accessibility of the skin through cytology, bacterial culture  
33 and topical therapy. For long-term success and to limit the spread of multidrug-resistance, we need  
34 to focus on identification and correction of underlying diseases that trigger pyoderma in order to  
35 avoid repeated treatment.

36

37 *Keywords:* Antimicrobial resistance; Staphylococci; MRSA/MRSP; Cytology; Topical  
38 antimicrobial therapy

## 39 **Introduction**

40           Although good prevalence data for canine pyoderma are lacking, bacterial skin infections  
41 were the second most frequent cause for presentation to first opinion practice in a UK survey on  
42 canine skin problems (Hill et al., 2006). Rarely life-threatening, pyoderma substantially contributes  
43 to canine morbidity through associated pruritus or pain, and potentially widespread and severe  
44 inflammatory changes. Because pyoderma is always secondary to underlying disease, unless this is  
45 corrected, recurrence is likely requiring repeated therapy, and causing frustration and continuing  
46 expense.

47  
48           Indeed, pyoderma is one of the main presentations leading to antimicrobial prescription in  
49 small animal practice (Hughes et al., 2012). A recent UK first opinion practice survey showed that  
50 92% of 683 dogs with pyoderma, either suspected or confirmed, received systemic antibacterial  
51 therapy (Summers et al., 2014). With continuing emergence of methicillin-resistant staphylococci,  
52 mainly *S. aureus* (MRSA) and *S. pseudintermedius* (MRSP), it is necessary to reduce antimicrobial  
53 use as a principal driver of multidrug-resistance (MDR) and pyoderma provides excellent  
54 opportunities for good antimicrobial stewardship.

55  
56           In this narrative review, we focus **on how the emergence of MRSP, MRSA and other**  
57 **MDR zoonotic pathogens has changed our approach to the management of canine pyoderma,**  
58 **and on how** traditional treatment recommendations need to be adapted to deal with this increasing  
59 threat to antimicrobial effectiveness and to public health.

60

## 61 **Foundation knowledge and clinical disease**

### 62 *Aetiology and pathogenesis*

63           Since publication of the first comprehensive veterinary dermatology text books in the 1960s  
64 (Muller and Kirk, 1969), pyoderma has consistently featured as one of the major diseases affecting

65 canine skin. It has been suggested that this is partly a consequence of the comparatively thin and  
66 compact canine stratum corneum, of the paucity of intracellular emulsion in canine epidermis and  
67 of the lack of a sebum plug in the canine hair follicle (Lloyd and Garthwaite, 1982; Mason and  
68 Lloyd, 1993).

69

70 The critical question of why pyoderma, particularly superficial pyoderma, develops and  
71 frequently recurs, is still incompletely understood. The major role of primary underlying disease in  
72 its aetiology is supported by the observation that the predominant staphylococcal pathogens are  
73 colonisers of healthy dogs and that most staphylococcal skin infections involve ‘endogenous’  
74 strains, i.e. isolates genetically identical to those of the patient’s healthy cutaneous and mucosal  
75 microflora (van Eiff et al., 2001; Pinchbeck et al., 2006 & 2007).

76

77 Common underlying triggers such as ectoparasite infestations, allergic skin diseases and  
78 endocrinopathies have long been associated with pyoderma, with allergic disease likely the main  
79 driver for recurrent forms (Mason and Lloyd, 1989; Colombo et al., 2007; Bloom, 2014). More  
80 specific concepts of quorum sensing, of a minimum infective dose and most recently findings from  
81 microbiome studies showing significant changes in diversity and composition during atopic  
82 dermatitis have provided new insights on why infection with opportunistic bacteria may develop in  
83 skin (Lloyd, 2014; Pierezan et al., 2016; Rodrigues Hoffman et al., 2017). Immunological defects in  
84 innate and adaptive immunity were identified in deep pyoderma of German shepherd dogs  
85 presenting with widespread, highly inflammatory infections during the 1980s and 1990s (e.g.  
86 Wisselink et al., 1988; Chabanne et al., 1995; Shearer and Day, 1997) but could not be conclusively  
87 linked to the breed or the occurrence of pyoderma (Rosser, 2006). Fortunately, this devastating  
88 disease now seems to be rare, possibly following targeted breeding.

89

90           The gaps in our understanding remain frustrating but it is important to remember that, when  
91 underlying causes are not identified, use of the term “idiopathic pyoderma” does not represent a  
92 diagnosis. In such cases diagnostic investigations need to be continued as failure to eliminate or  
93 control underlying disease **or predisposing factors** will lead to recurrence.

94

#### 95 *Classification and diagnosis of pyoderma*

96           With its secondary aetiology and the need for responsible use of antimicrobials in mind, a  
97 diagnosis should always include i) recognition of suggestive skin lesions and likely depth of  
98 infection, ii) confirmation of bacterial infection through cytology and iii) identification of  
99 underlying primary disease.

100

101           Despite its prevalence, canine pyoderma is often misdiagnosed (Gortel, 2013) leading to  
102 inappropriate treatment. Recognition of suggestive skin lesions and their distribution is essential  
103 and requires careful inspection of the skin. Since the number of ways skin can react to insult is  
104 limited, classifications have been proposed to facilitate morphological diagnosis. The most widely  
105 used is based on depth of infection and distinguishes surface, superficial and deep pyoderma, all  
106 three associated with typical clinical presentations (Ihrke, 1987; White and Ihrke, 1987) (Fig. 1).

107

108           Surface pyoderma remains the least understood group. It includes frequently seen  
109 presentations such as acute moist dermatitis (“hot spots”, pyotraumatic dermatitis), fold pyoderma  
110 (intertrigo), and the more recently described microbial/bacterial overgrowth syndrome in which  
111 erythema is the only clinical sign but large numbers of bacteria on the inflamed skin can be  
112 demonstrated by cytology (Pin et al., 2006). Here, excessive multiplication of bacteria is confined  
113 to the skin surface and is seen as a minor player in the pathogenesis, triggered by a dominant  
114 inflammatory cause.

115

116 Superficial pyoderma involves invasion of the epidermis. Bacterial folliculitis extends into  
117 the follicular ostium and epidermal tissue and is likely the most frequent pyoderma type in dogs. It  
118 presents with papules, pustules and epidermal collarettes, typically on the ventral abdomen and  
119 medial thighs or on the trunk and often associated with areas of alopecia and varying degrees of  
120 pruritus; its interfollicular form (impetigo) occurs mostly in puppies. Coat type and immune-status  
121 can also influence clinical appearance as in the moth-eaten appearance of superficial pyoderma in  
122 short-coated breeds or in the large lesions (collarettes, pustules) associated with bullous impetigo or  
123 superficial spreading pyoderma in immune-compromised dogs (Bloom, 2014; Beco et al., 2013a).  
124 Mucocutaneous pyoderma is a disease of unknown aetiology. It primarily affects lips and perioral  
125 skin, with swelling, erythema and crusting which may lead to fissuring and erosion. It often  
126 responds slowly to therapy and can be confused with immune-mediated disease.

127

128 Deep pyoderma is less common but more serious, as its expansion into the dermis and  
129 proximity to blood vessels increases the risk of haematogenous spread and bacteraemia. It can be  
130 seen with any underlying trigger or acquired immuno-deficiency but is commonly associated with  
131 demodicosis (Kuznetsova et al., 2012; Mueller et al., 2012). Lesions include draining sinuses,  
132 fistulae, haemorrhagic crusts, nodules and varying degrees of erythema and swelling; pain is not  
133 infrequent. Common localised forms of deep pyoderma affect the head (chin acne, muzzle  
134 folliculitis and furunculosis) or limbs (interdigital nodules, callus pyoderma, acral lick granuloma).  
135 Nodular lesions quite often involve bacteria other than staphylococci and need to be differentiated  
136 from non-bacterial infected granulomas, sterile granulomatous disease, neoplasia and foreign body  
137 reactions by biopsy, special stains, macerated tissue culture and sometimes molecular techniques.

138

139 For initial diagnosis, cytology from slide or tape impressions, a frequent requirement of  
140 antimicrobial stewardship guidelines prior to antimicrobial prescription (e.g. BVA, 2015), is  
141 recommended to confirm bacterial involvement. Cytology of superficial pyoderma lesions is

142 reported to have 93% diagnostic sensitivity, based on presence of neutrophils and intracellular cocci  
143 (Udenberg et al., 2014) but, despite being rapid and inexpensive (Curtis, 2001), remains underused  
144 in general practice (Hill et al., 2006).

145

146 Bacterial culture, on the other hand, is of limited value in the initial diagnosis of pyoderma.  
147 It is likely to yield staphylococci from infected and non-infected skin (Doelle et al., 2015), and can  
148 therefore not distinguish infected from colonised skin. However, bacterial culture and antimicrobial  
149 susceptibility testing are essential for selection of systemic therapy after a diagnosis has been  
150 established. It is of note that sampling can be challenging, particularly in deep pyoderma for which  
151 surface swabs have been shown to predict relevant pathogens from deep infection in only about  
152 30% of cases (Shumaker et al., 2008) and submission of tissue (in saline, not formalin) obtained  
153 through biopsy is preferred.

154

#### 155 *Pathogens*

156 The predominant role of coagulase-positive staphylococci has been long recognised (Ihrke,  
157 1987). Originally all such infections were ascribed to *S. aureus*, but refinement of microbiological  
158 techniques allowed new species including *S. intermedius* and *S. pseudintermedius* to be described  
159 (Table 1). *S. pseudintermedius* is recognised to be most commonly involved, particularly in  
160 superficial pyoderma (Medleau et al., 1986; Shumaker et al., 2008). Other staphylococci, including  
161 *S. aureus*, *S. schleiferi* and *S. hyicus* may be involved in up to 10% of cases.

162

163 Staphylococci have an array of potential virulence factors but despite detailed investigation  
164 significant associations between specific virulence genes and disease have not yet been identified,  
165 shifting attention again to host factors that may facilitate infection (Bannoehr et al., 2012; Tanabe et  
166 al., 2013; Couto et al., 2015). However, biofilm production, which can promote resistance to host  
167 defence mechanisms and greatly enhance antimicrobial resistance, has been confirmed in many



168 isolates of *S. pseudintermedius* and other veterinary staphylococci (Götz, 2002; Hall-Stoodley et al.,  
169 2004).

170

171 Many other bacterial pathogens, including *Pseudomonas aeruginosa*, *Proteus spp.*,  
172 streptococci, *Burkholderia spp.* and *Escherichia coli* may be difficult to distinguish clinically  
173 (Rantala et al., 2004; Hillier et al., 2006; Cain et al., 2015; Tham et al., 2016). Isolation of  
174 coagulase-negative staphylococci, such as *S. lugdunensis* and *S. schleiferi subsp. schleiferi*, and of  
175 *Micrococcus spp.*, can also cause confusion in laboratories that are looking for coagulase-positive  
176 bacteria (Cain et al., 2011; Gobeli Brawand et al., 2016; Cotting et al., 2017). Surprisingly, the idea  
177 that coagulase-negative staphylococci are non-pathogenic persists even though they are the most  
178 common cause of nosocomial bacteraemia in human hospitals (von Eiff et al., 2002; Becker et al.,  
179 2014) and are increasingly reported in animal infections (e.g. Rook et al., 2012; Frank et al., 2008;  
180 Davis et al., 2013; Kern and Perreten, 2013; Ruzauskas et al., 2014).

181

## 182 **Emergence of multidrug-resistance**

183 Resistance to antimicrobials within bacterial populations is an ancient phenomenon, vital for  
184 bacterial survival (D'Costa et al., 2011; Perron et al., 2015). However, the accumulation of multiple  
185 resistance genes in bacterial pathogens, driven by overuse of antimicrobial drugs, has become a  
186 chilling threat to human and animal health (Gossens et al., 2005; Costelloe et al., 2010).

187

188 First concerns about MDR in canine pyoderma emerged twenty years ago when MRSA  
189 became recognised in sporadic skin and wound infections; later, the more epidemic spread of  
190 MRSP overtook it and now presents major challenges to our management of canine pyoderma. In  
191 addition, all key multidrug-resistant pathogens of relevance in human medicine, such as  
192 *Enterococcus faecium*, *Klebsiella spp.*, *Acinetobacter baumannii*, *Pseudomonas aeruginosa* and

193 *Enterobacter spp.* (Boucher et al., 2009), are now recognised to be associated with infection in pets  
194 (Grobbel et al. 2007; Kuzi et al., 2016; Abdel-Moein et al., 2017).

195

#### 196 *Meticillin-resistance in staphylococci*

197         Although methicillin is no longer available for clinical use, it still serves as a marker for  
198 broad resistance to all  $\beta$ -lactams (excepting some of the latest anti-staphylococcal molecules) and as  
199 an indicator of likely nosocomial epidemiology and additional multidrug-resistance. The genetic  
200 basis underpinning methicillin-resistance is the presence of the *mecA* gene held on a large mobile  
201 genetic element, the staphylococcal cassette chromosome (SCC). This is similar in all staphylococci  
202 and has been extensively studied for MRSA (Lindsay and Holden, 2006).

203

204         Since the first identification of MRSA from pets, isolates from pets and humans have been  
205 found genetically identical, providing indirect but good evidence that transmission between these  
206 hosts can occur in both directions (reviewed by McCarthy et al., 2012). MRSA was the first  
207 multidrug-resistant *Staphylococcus* to receive attention in animals when pets contaminated by  
208 human MRSA patients were shown to be involved in perpetuating human infection or recurrent  
209 outbreaks (Scott et al., 1988; Manian, 2003). Since then, sporadic infections, case series and  
210 outbreaks have been reported, typically involving skin and wound infections in dogs (Tomlin et al.,  
211 1999; Paterson et al., 2015; Morris et al., 2017). Most reports are from countries with a high MRSA  
212 prevalence in human hospitals, indicating a spill-over from humans; epidemic spread beyond clinic  
213 or kennel outbreaks has not been reported. Fortunately, the prognosis in such infections can be  
214 considered good, depending on underlying causes, as the great majority of these human hospital-  
215 associated MRSA remain susceptible to tetracyclines and potentiated sulphonamides and around  
216 50% to clindamycin. A less predictable prognosis needs to be considered for rare infections  
217 involving MRSA from human lineages that carry toxins such as Panton-Valentine-leucocidin

218 (Rankin et al., 2005; van Duikeren et al., 2005) and those associated with livestock-associated  
219 MRSA (Gómez-Sanz et al., 2013).

220

221 A much greater veterinary challenge is the emergence in dogs of MRSP, associated with  
222 even broader drug-resistance. Whole genome sequencing shows that only three genetic steps  
223 (acquisition of *mecA* on a SCC, acquisition of a large transposon (Tn5405-like element) carrying up  
224 to five resistance genes and genome point mutations for fluoroquinolone and sulphonamide  
225 resistance) are required for its rapid evolution to MDR, emphasising the important role of selection  
226 pressure (Loeffler et al., 2007; Perreten et al., 2010; Detwiler et al., 2014; McCarthy et al., 2015).

227

228 First reported from dogs in North America in the late 1990s (then MRSI), MRSP accounted  
229 for over 30% of staphylococcal isolates from American dogs within less than ten years (Gortel et  
230 al., 1999; Morris et al., 2006; Jones et al., 2007) and is now identified worldwide. An even higher  
231 prevalence was recently reported from China and Japan with nearly 50% and 70%, respectively  
232 (Feng et al., 2012; Kasai et al., 2016). In the UK, where MRSP was first recognised in 2009, the  
233 burden seems relatively low with rates below 5% of clinical *S. pseudintermedius* laboratory  
234 submissions reported in 2015 (Maluping et al., 2014; Beever et al., 2015). In contrast, studies from  
235 continental Europe, where MRSP had been identified three years earlier, prevalence was soon  
236 reported around 30% (Loeffler et al., 2007; DeLucia et al., 2011).

237

238 Substantial percentages of methicillin-resistance have also been reported in coagulase-  
239 variable *S. schleiferi* (subspecies *schleiferi* and subspecies *coagulans*), some from pyoderma but  
240 most from otitis (Cain et al., 2011).

241

242 *Clinical implications and early identification*

243 Clinically, MRS infections in animals are no different from infections involving less  
244 resistant staphylococci (Fig. 1) (Morris et al., 2017). In fact, early case-control studies showed that  
245 clinical outcome was no worse for MRSA and MRSP infections in pets compared to those  
246 involving their susceptible counterparts, provided that a safe antibacterial treatment option was  
247 available (Weese et al., 2012; Lehner et al., 2014). Finding treatment options may be troublesome  
248 and treatment has been shown to take longer than with susceptible staphylococci (Bryan et al.,  
249 2012). However, the bigger concern with MRS pyoderma is its potential for spreading these  
250 zoonotic, multidrug-resistant pathogens to other animals and people, and into the environment. For  
251 MRSP, the risk of zoonotic transmission is generally considered low as low carriage rates of *S.*  
252 *(pseud)intermedius* have been found in people regularly exposed to dogs (Havey et al. 1994;  
253 Goodacre et al., 1997; Han et al., 2016). As for all staphylococci though, the risk is increased for  
254 immune-compromised people and individual cases of zoonotic MRSP infection have been reported  
255 (Stegman et al., 2010; Somayaji et al., 2016; Lozano et al., 2017).

256

257 Transmission of staphylococci is supported by their ability to survive on dry surfaces and at  
258 healthy skin and mucosal carriage sites for many months, equipping them for nosocomial spread  
259 (Wagenvoort et al., 2000; Windahl et al. 2016). Early identification of MRS by clinicians is  
260 therefore crucial to limit outbreaks but it relies on awareness of risk factors. Risk factors for  
261 multidrug-resistant infection in human medicine are well documented and include frequent  
262 hospitalisation, length of stay in hospitals, surgical interventions and repeated antimicrobial therapy  
263 (Sadfar and Maki, 2002). Unsurprisingly, the same risk factors have been identified for MRSA and  
264 MRSP infections in dogs (Soares-Magalhães et al., 2010; Baker et al., 2012; Lehner et al. 2014;  
265 Weese et al., 2012).

266

267 *New focus on laboratory identification*

268           Before the emergence of MRS, species identity of a coagulase-positive *Staphylococcus* was  
269 probably of little importance to most clinicians. When MRS are involved, accurate differentiation  
270 between MRSA and MRSP is critical for further management as important epidemiological  
271 differences exist. Isolation of MRSA from a dog will prompt a focus on human health concerns and  
272 the need to inform the owner's human physician. In contrast, isolation of the dog-adapted MRSP  
273 should initiate all appropriate infection control measures recommended for veterinary nosocomial  
274 pathogens and advice on limiting contagion to other animals, while only a lower zoonotic risk needs  
275 to be considered (Morris et al., 2017).

276

277           Unfortunately, it has also become clear that identification through phenotypic assessment  
278 alone is more difficult than text books suggest as subtle morphological and biochemical variation  
279 occurs within bacterial populations (Pottumarthy et al., 2004; Sasaki et al., 2007; Geraghty et al.,  
280 2013; Bond and Loeffler, 2012). Similarly, recognition and accurate identification of coagulase-  
281 negative staphylococci is important as their pathogenic potential is increasingly recognised;  
282 reporting such isolates as 'consistent with microflora organisms' is no longer sufficient. Semi-  
283 automated and automated laboratory procedures help with speciation but are commonly set up for  
284 human bacterial pathogens and may lack precision for veterinary isolates. Currently, best accuracy  
285 in a diagnostic setting is achieved by Matrix Assisted Laser Desorption/Ionisation Time of Flight  
286 Mass Spectrometry (MALDI-TOF) which has been validated for many veterinary pathogens  
287 including the very similar SIG species (Decristophoris et al., 2011; Sauget et al., 2016; Somayaji et  
288 al., 2016).

289

290           Susceptibility testing is most often done by traditional disk diffusion with clinical  
291 breakpoints guiding predictions on clinical efficacy. Dilution testing and minimum inhibitory

292 concentrations (MICs) were rarely needed for the management of skin infections in the past but will  
293 be helpful in multidrug-resistant infections when a borderline MIC may still be overcome with high  
294 doses of an authorised antimicrobial rather than choosing a less safe drug, or when calculating  
295 dosages for treatment with an unauthorised agent. Resistance testing against methicillin, nowadays  
296 replaced by oxacillin, can also be misleading since *mecA*-independent mechanisms (other *mec*  
297 types, hyper-penicillinase producers, incomplete expression) can lead to inconsistent results (Morris  
298 et al., 2017). Confirmation of phenotypic methicillin-resistance by additional tests (molecular for  
299 *mecA* or by agglutination tests detecting an altered penicillin-binding protein encoded by *mec*) is  
300 desirable before MRS management decisions are initiated (Becker et al., 2014b).

301

### 302 **Management of canine pyoderma**

303 In the past, treatment of canine pyoderma was rarely challenging as *S. pseudintermedius*  
304 (formerly *S. intermedius*) was widely susceptible and broad-spectrum antibacterial agents such as  
305 cephalexin, potentiated amoxicillin and enrofloxacin became licensed for use in dogs during the  
306 1970s and 1980s, all with an indication for skin infection (Medleau et al., 1986; Kruse et al., 1996;  
307 Lloyd et al., 1996; Pellerin et al., 1998; Normand et al., 2000). It was already recognised that  
308 isolates from animals that had repeatedly received antimicrobials were likely to show more  
309 resistance (Noble and Kent, 1992; Holm et al., 2002) but empirical selection of drugs for systemic  
310 treatment of pyoderma was nearly always successful. This situation began to change around 20  
311 years ago when methicillin-resistant, multidrug-resistant staphylococci were recognised amongst  
312 canine clinical isolates and, in the UK, there is now evidence that resistance to most antimicrobial  
313 classes is gradually increasing (Beever et al., 2015). Based on recent data for small animal  
314 pathogens, this trend of increasing AMR is likely to continue worldwide (Ludwig et al., 2016).

315

316 *Topical therapy*

317           Topical antibacterial therapy has always been advised for surface infections and, in  
318 combination with systemic therapy, for superficial and deep pyoderma (Ihrke, 1987; Curtis, 1998 &  
319 1999). However, newer studies have provided good evidence that topical therapy can be effective as  
320 sole antibacterial treatment in superficial pyoderma, including cases with MRS (Murayama et al.,  
321 2010; Loeffler et al., 2011; Borio et al., 2015). In situations where pet and owner can be expected to  
322 be compliant and where clinicians are prepared to convince owners of its merits, topical treatment  
323 can help to reduce overall antimicrobial prescription.

324

325           A wide range of different formulations, such as shampoos, creams, gels and ointments, and  
326 more recently foams, is marketed for dogs and includes a variety of antibacterial agents; this can be  
327 confusing. A systematic review of topical therapy for canine bacterial skin infections concluded that  
328 while evidence from randomised controlled trials was sparse, good evidence supported the use of  
329 shampoos containing 2-3% chlorhexidine and, to a lesser extent, benzoyl peroxide (Mueller et al.,  
330 2012) and these continue to be the mainstay of topical therapy, at least for widespread disease.  
331 Localised infections can also be treated with creams or gels containing antibiotics such as fusidic  
332 acid, authorised for use in dogs in European countries and in Canada, or mupirocin ointment  
333 authorised in the USA for dogs but reserved for use in human medicine in most of Europe (Cobb et  
334 al., 2005; BNF, 2017).

335

336           While concern over resistance to topically used antibacterial agents exists, clinical treatment  
337 failure of topical anti-staphylococcal therapy has not been conclusively reported, to the authors'  
338 knowledge; MICs for staphylococci from animals have been consistently low and are likely to be  
339 substantially exceeded by achievable topical drug concentrations (Loeffler et al. 2008; Valentine et  
340 al., 2012; Clark et al., 2015). However, continual monitoring of resistance and clinical efficacy,

341 further evaluation of alternatives such as hypochlorite (bleach), Manuka honey, of potentially  
342 synergistic combinations and of anti-biofilm products will be critical (Walker et al., 2016).

343

344         Combination of topical treatment with systemic treatment is recommended whenever  
345 possible to potentially reduce the duration of systemic therapy, and in MRS infections, to reduce  
346 environmental contamination and risk of transmission to other hosts.

347

#### 348 *Systemic therapy*

349         Systemic therapy, required for deep pyoderma and for widespread or severe superficial  
350 infections, should follow the concept of ‘as little as possible but as much as necessary’ (RUMA  
351 2009). Efficacy depends predominantly on bacterial susceptibility but will also be determined by  
352 correct drug administration, appropriate dosing, owner compliance and clinical variables such as  
353 severity of infection and causative and concurrent diseases. Surprisingly, despite their universal use,  
354 evidence on efficacy of systemic antimicrobial agents is sparse as only few adequate studies  
355 documenting outcome exist (Summers et al., 2012).

356

357         While bacterial culture and susceptibility testing would be desirable for every patient and is  
358 never contraindicated in pyoderma, realistically, cost, perceived delay of effective treatment and  
359 clinical time pressure often motivate empirical drug selection. In countries with low MRS  
360 prevalence, empirical selection may still be effective for most superficial pyodermas. In high-MRS  
361 prevalence countries, this can no longer be considered reliable or cost-effective. Indeed, repeated  
362 testing may be required as antimicrobial therapy has been shown to promote acquisition of MRSP  
363 in dogs not previously MRSP-positive (Beck et al., 2012). Recent pyoderma guidelines further  
364 specify that culture and susceptibility testing is essential in all dogs with deep pyoderma, those with  
365 a history of MRS or with owners reporting MRS in themselves, and in dogs where appropriate  
366 empirical antibiotics has been ineffective. (Beco et al., 2013b; Hillier et al., 2014)



367

368           When prescribing antimicrobial drugs for dogs, it is important to remember that most are  
369 also used in human medicine, either as identical or related molecules, and that key agents for canine  
370 pyoderma are listed by the WHO as ‘critically important antimicrobials’ or ‘highly important for  
371 human medicine’ (WHO, 2011).

372

373           For non-MRSP pyoderma, most antimicrobials authorised for use in dogs would be effective  
374 if prescribed appropriately. Treatment recommendations have recently been detailed in two free  
375 access publications, one on pyoderma by a group of veterinary dermatologists (Beco et al., 2013b),  
376 the other on superficial bacterial folliculitis by the International Society for Companion Animal  
377 Infectious Disease (ISCAID)(Hillier et al., 2014). Briefly, antimicrobial drugs can be classified into  
378 first and second tier/line drugs, depending on the likelihood that they will be effective against  
379 staphylococci and their spectrum of activity against Gram-negative pathogens. First-tier drugs, such  
380 as **clindamycin**, first-generation cephalosporins, amoxicillin-clavulanate or **potentiated**  
381 **sulphonamides** may be chosen empirically in areas with a low prevalence of MRS. **Clindamycin,**  
382 **an antimicrobial with good efficacy against most staphylococci, can be considered as a**  
383 **responsible treatment choice due to its relatively narrow spectrum of activity. However,**  
384 **clinicians need to be familiar with their local *S. pseudintermedius* resistance pattern as**  
385 **differences in resistance have been recognised between countries and between isolates from**  
386 **first-time pyoderma versus those from recurrent pyoderma (Holm et al., 2002; Beever et al.,**  
387 **2015; Larsen et al., 2015).** Treatment with second-tier agents, such as for example  
388 fluoroquinolones, should always be based on bacterial culture and susceptibility results. Readers are  
389 referred to these guidelines for more detailed information on dose recommendations and adverse  
390 effects.

391

392 In MRS pyoderma, drugs predicted to be effective by *in vitro* testing are selected based on  
393 national licensing rules, their clinical and safety characteristics, dosing practicalities and cost, with  
394 no single drug shown to be better than another. Information specifically on MRS treatment is  
395 detailed in recently published open access Clinical Consensus Guidelines on MRS infections  
396 (Morris et al., 2017). Specifically on the interpretation of resistance testing, the guidelines point out  
397 that no representatives of  $\beta$ -lactam antibiotics should be used for MRS infections even if testing  
398 indicates susceptibility for individual agents of this class, that testing for inducible resistance to  
399 clindamycin is recommended for MRS to avoid treatment failure during therapy, that extrapolation  
400 of results for one type of tetracycline to another can be unreliable as resistance is mediated by a  
401 number of different genes and that resistance to one fluoroquinolone is likely to indicate resistance  
402 to others in MRSP; MIC determination may then help to inform treatment decisions (Kizerwetter-  
403 Świda et al., 2016).

404

405 When no susceptibilities to clinically relevant and authorised antimicrobials are reported,  
406 extended testing is required. Amikacin, rifampicin and chloramphenicol are most frequently  
407 mentioned for such infections (Frank and Loeffler, 2012; Papich, 2012) but their use should be  
408 preceded by appropriate dose calculations and toxicity monitoring, requires detailed owner  
409 education and compliance, and should include advice on infection control measures to limit spread  
410 (Morris et al., 2017). In the authors' opinion, glycopeptides, linezolid and potentially new  
411 compounds should be strictly reserved for use in humans. Some institutions may consider these  
412 under restriction-of-use protocols but this should rarely be necessary for pyoderma (Weese, 2008).

413

414 Recommendations on how long to treat pyoderma for remain controversial. Traditional  
415 advice, based on clinical expertise is three weeks or one week beyond clinical cure for superficial  
416 pyoderma, and four to eight weeks or two weeks beyond clinical cure for deep pyoderma (Ihrke,  
417 1987). In addition, many datasheets now recommend several weeks of therapy. In human medicine,

418 antibiotic courses are typically shorter but recently, even the advice to patients to complete a course  
419 after clinical signs have resolved has been questioned (Llewelyn et al., 2017). In the absence of  
420 better data, it is prudent to follow advice from ISCAID and adhere to the traditional  
421 recommendations but where shorter treatment is prescribed, plans for close monitoring of progress  
422 by veterinarian rather than owner should be made (Hillier et al., 2014). In addition, resolution of  
423 clinical signs will not signal the end of case management for MRS pyoderma as dogs can become  
424 carriers and carry the risk of contagion, including zoonotic transmission, and of self-re-infection.

425

#### 426 *Correction of primary triggers, follow-up and prevention*

427 After resolution of any type of pyoderma, prevention of recurrence is very important as  
428 multidrug-resistance may develop with repeated systemic treatment. Such prevention will depend  
429 on elimination or suppression of underlying triggers. A diagnosis of these triggers may not be a  
430 priority to owners compared to the urgency of resolving the pyoderma, and will present an extra  
431 challenge to communication during busy consultations. Most problematic, in the authors'  
432 experience, are those dogs that in the absence of pyoderma (i.e. when infection has been resolved)  
433 present either with no clinical signs suggestive of underlying triggers or with signs compatible with  
434 very mild allergic skin disease. In those cases, provided history and signalment are in line with  
435 allergic skin disease, empirical treatment with anti-inflammatory medication may help to prevent  
436 flares of bacterial infection. If successful, this approach can subsequently be optimised by further  
437 investigations into allergic skin disease (Olivry et al., 2015).

438

439 Importantly for MRS infections, once infection has resolved, animals will continue to  
440 harbour staphylococci on healthy skin and mucosae. For MRSP, a bacterium well adapted to dogs  
441 (Simou et al., 2005), carriage has been shown to continue for up to 11 months after infection has  
442 resolved (Windahl et al., 2012). Carriage and environmental contamination and the risk of  
443 subsequent self-re-infection have long been suspected as major contributors to the successful spread

444 of human MRSA and a similar epidemiology is suspected for MRSP in veterinary settings (Beck et  
445 al., 2012; Morris et al., 2017).

446

#### 447 **New approaches**

448 The growing problem of antimicrobial resistance and the lack of effective, new conventional  
449 antimicrobial drugs has promoted the development of different approaches to prevention and  
450 control of bacterial infections (Lloyd, 2012; Vale et al., 2016). Staphylococcal vaccines, either *S.*  
451 *aureus* lysates or autogenous bacterin preparations have been assessed in small studies and warrant  
452 further investigations (Glos and Mueller, 2006). Antimicrobial peptides, which are produced by the  
453 skin and function as a vital part of cutaneous antimicrobial defence, are now being exploited in  
454 veterinary products for dogs. Two promising approaches have been adopted. In the first, plant  
455 extracts promoting production of endogenous antimicrobial peptides by the treated skin have been  
456 incorporated in shampoos and ear cleaners (Marsella et al, 2013; Santoro et al, 2016). In the second,  
457 a synthetic peptide (Cabassi et al, 2013) has been incorporated in shampoo, foam and an ear  
458 treatment gel. A variety of other approaches are being investigated and developed (Lloyd, 2012) but  
459 have not yet led to the development of veterinary products (Table 2).

460

461 It is likely that at least some of these new approaches will prove successful, however we  
462 should not expect the development of agents which will allow us to ignore good drug stewardship  
463 and the adoption of rigorous hygiene.

464

#### 465 **Conclusions**

466 Canine pyoderma will need to be managed appropriately to reduce morbidity and to limit  
467 the spread of potentially MDR pathogens amongst pets and humans. However, the availability of  
468 effective and safe systemic antimicrobials will become - or already is in some countries -  
469 substantially limited, either by continued selection of antimicrobial resistance amongst pathogens or

470 by legislative restrictions on prescribing by veterinary surgeons. We will likely need to adapt our  
471 prescribing practices for all animal species and all affected organs in the future. For canine  
472 pyoderma though, the skin as the infected organ can be easily accessed for examination and  
473 treatment monitoring, rapid in-house tests and topical therapy. This provides unique opportunities  
474 to combine relatively small achievable adaptations in our pyoderma management with good  
475 antimicrobial stewardship and effective treatment outcomes. Comprehensive owner education and  
476 rigorous hygiene measures need to become an integral part of pyoderma management and will help  
477 to limit the spread of antimicrobial resistance and delay the end the Golden Age of Antibiotics  
478 (Gould, 2009).

479

#### 480 **Highlights**

- 481 1. Management of canine pyoderma is increasingly complicated by multidrug-resistant  
482 pathogens such as MRSP and empirical selection therapy is no longer reliable in areas with  
483 a high MRSP prevalence.
- 484 2. Use of in-house cytology can rapidly confirm bacterial infection and support responsible  
485 antimicrobial prescribing.
- 486 3. Topical therapy can be effective on its own in cases of superficial pyoderma, even in those  
487 involving MRSP.
- 488 4. Awareness of risk factors, contagious and zoonotic characteristics, laboratory requirements  
489 and necessary hygiene measures are critical in the management of MRSP pyoderma.
- 490 5. Diagnosis and treatment of underlying diseases needs to replace our reliance on  
491 antimicrobial therapy in dogs with recurrent pyoderma.

492

#### 493 **Conflict of interest statement**

494 The authors have no financial or personal relationship with other people or organisations  
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1046 **Table 1**  
 1047 The changing nomenclature of *Staphylococcus aureus*.  
 1048

Name	Year	Comments
<i>Staphylococcus pyogenes aureus</i>	1886 (Rosenbach)	Representing golden (rather than white) staphylococcal colonies
<i>S. aureus</i>	1951 (Shaw et al.)	Representing all coagulase positive staphylococci
Methicillin-resistant <i>S. aureus</i> (MRSA)	1961 (Jevons)	First recognition of methicillin resistance in <i>S. aureus</i>
<i>S. aureus</i>	1971 (Hajek and Marsálek)	Differentiation of animal species-related biotypes A-F
<i>S. intermedius</i>	1976 (Hajek)	Differentiated from <i>S. aureus</i> , representing biotypes E and F
<i>S.pseudintermedius</i>	2005 (Devriese et al.)	Differentiated from <i>S. intermedius</i> , representing biotype E.
<i>Staphylococcus intermedius</i> group (SIG)	2007 (Sasaki et al.)	Includes <i>S. intermedius</i> , <i>S. pseudintermedius</i> and <i>S. delphini</i> which are difficult to differentiate in routine laboratory testing. <i>S. intermedius</i> shown to be mainly associated with wild pigeons.*
Methicillin-resistant <i>S. pseudintermedius</i> (MRSP)	2007 (Sasaki et al.)	First recognition of methicillin resistance in <i>S. pseudintermedius</i>

1049 \*Canine SIG isolates are always considered as *S. pseudintermedius* (Bannoehr et al., 2007).  
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**Table 2**

New and alternative antimicrobial approaches.

Approach	Action
Efflux pump inhibitors	Suppress elimination of antimicrobial agents
Silencing of resistance and virulence genes	Antagonise function of specific genes
Quorum quenching	Agents suppressing virulence of pathogen
Probiotics and prebiotics	Provide or promote competitor bacteria
Microbial predation	Bacterial or fungal predators consume pathogen
Bacteriophages	Invade and destroy pathogen
Vaccines and immunoglobulins	Stimulate or passively provide immunity

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1056 **Figure legend**

1057

1058 Fig. 1. Examples of recurrent or chronic (> 3 months) pyoderma involving multidrug-resistant  
1059 bacteria. All cases had received repeated courses of systemic antimicrobials with initial  
1060 improvement. Pyoderma resolved when underlying triggering causes were diagnosed and treated in  
1061 combination with antibacterial therapy. (A) Acute moist dermatitis with methicillin-resistant  
1062 *Staphylococcus pseudintermedius* (MRSP) on the neck of a young atopic Saint Bernard. (B)  
1063 Purulent *Klebsiella* spp. infection complicating erosive pad lesions in a sterile granulomatous  
1064 disease. Both dogs were treated and remained in remission with topical antibacterial and systemic  
1065 anti-inflammatory treatment. (C) Recurrent superficial pyoderma with expanding epidermal  
1066 collarettes and focal crusts due to MRSA in a dog with early hyperadrenocorticism; infection  
1067 resolved with topical antibacterial washes alone when the endocrinopathy was treated. (D)  
1068 Widespread deep pyoderma involving *Pseudomonas aeruginosa* in a young Dalmatian dog with  
1069 juvenile-onset demodicosis; there was no evidence of pyoderma on cytology after 3 weeks of  
1070 systemic antibacterial and acaricidal therapy.

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A



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B



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C



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D