Background – Superficial bacterial folliculitis (SBF) is usually caused by *Staphylococcus pseudintermedius* and routinely treated with systemic antimicrobial agents. Infection is a consequence of reduced immunity associated with alterations of the skin barrier and underlying diseases that may be difficult to diagnose and resolve; thus, SBF is frequently recurrent and repeated treatment is necessary. The emergence of multiresistant bacteria, particularly meticillin-resistant *S. pseudintermedius* (MRSP), has focused attention on the need for optimal management of SBF.

Objectives – Provision of an internationally available resource guiding practitioners in the diagnosis, treatment and prevention of SBF.

Development of the guidelines – The guidelines were developed by the Antimicrobial Guidelines Working Group of the International Society for Companion Animal Infectious Diseases, with consultation and advice from diplomates of the American and European Colleges of Veterinary Dermatology. They describe optimal methods for the diagnosis and management of SBF, including isolation of the causative organism, antimicrobial susceptibility testing, selection of antimicrobial drugs, therapeutic protocols and advice on infection control. Guidance is given for topical and systemic modalities, including approaches suitable for MRSP. Systemic drugs are classified in three tiers. Tier one drugs are used when diagnosis is clear cut and risk factors for antimicrobial drug resistance are not present. Otherwise, tier two drugs are used and antimicrobial susceptibility tests are mandatory. Tier three includes drugs reserved for highly resistant infections; their use is strongly discouraged and, when necessary, they should be used in consultation with specialists.

Conclusions and clinical importance – Optimal management of SBF will improve antimicrobial use and reduce selection of MRSP and other multidrug-resistant bacteria affecting animal and human health.

Introduction

In dogs, superficial bacterial folliculitis (SBF) is the commonest form of canine pyoderma, which is in turn, the principal reason for antimicrobial use in small animal practice. As we face the problem of increasing antimicrobial resistance in both human and veterinary medicine, there is a pressing need for prudent and more focused use of antimicrobial drugs (AMDs). In the human field, adoption of guidelines for antimicrobial use at the hospital level has been shown to improve prescribing practices significantly, both alone and as part of broader antimicrobial stewardship programmes. Similar
benefits can be expected in the veterinary field, where there is a need for improved antimicrobial stewardship both in veterinary hospitals and in veterinary practice.

This document presents guidelines developed in 2011–2013 by the Antimicrobial Guidelines Working Group of the International Society for Companion Animal Infectious Disease (ISCAID). These guidelines were developed because of increasing concerns regarding widespread antimicrobial resistance in bacteria infecting dogs and cats. The members of the Group were Scott Weese (chair), Joseph Blondeau, Dawn Booth, Edward Breitschwerdt, Luca Guardabassi, Andrew Hillier, Michael Lappin, David Lloyd, Mark Papich, Shelley Rankin, Jane Sykes and John Turnidge. The group met in Miami (FL, USA) to develop the guidelines, then communicated by email and through telephone conferences to refine the wording of this document further. Input was also solicited from diplomates of the American College of Veterinary Dermatology (ACVD) and the European College of Veterinary Dermatology (ECVD). The guidelines are directed primarily at private small animal practitioners in primary care practice.

It should be noted that these guidelines are specific for SBF and apply only to dogs. Although the broad principles relating to AMD use in SBF are applicable to a variety of canine bacterial skin infections, significant differences exist amongst such infections that may be associated with the depth of the skin that is affected and the bacterial pathogens involved. These guidelines cannot be applied to other types of bacterial infections in canine skin without careful consideration. It is anticipated that guidelines for other bacterial skin infections in dogs will be developed in due course.

To the best of the authors’ knowledge, there is only one published peer-reviewed article that provides similar guidelines.7 Those guidelines differ from this document in that they are directed more generally at the treatment of skin and soft tissue infections in dogs and cats, they are directed at the use of systemic antibiotics only and do not address topical therapies, they suggest diagnosis and treatment of pyoderma according to an unpublished classification system based on the clinical appearance of lesions rather than the depth of the infection in the skin and they are authored by a group of European specialist dermatologists. Thus, apart from differences in content, we believe that our guidelines provide a different perspective from a broader international group of authors who also represent other pertinent areas of specialization in addition to dermatology.

Recommendations for the diagnosis of canine superficial bacterial folliculitis

The predominant pathogen that causes SBF is Staphylococcus pseudintermedius (previously known and referred to as Staphylococcus intermedius).8 Although dogs may carry or be colonized and infected by Staphylococcus aureus and by the coagulase-variable species Staphylococcus schleiferi,9,10 these are far less frequent pathogens in SBF. Coagulase-negative staphylococci (CoNS; such as Staphylococcus epidermidis and Staphylococcus xylosus) may rarely be cultured from lesions of SBF, usually in association with S. pseudintermedius. The clinical relevance of isolation of these species from SBF lesions is unclear. Other bacteria may, on rare occasions, cause lesions compatible with SBF. These include Streptococcus canis, Pseudomonas aeruginosa and other Gram-negative bacteria.11,12

Clinical signs

In practice, the diagnosis of most cases of SBF is based upon clinical signs and the presence of characteristic lesions; there is no evidence that these differ amongst infections caused by the different staphylococci. Common lesions of SBF are erythematous pappules (Figures 1 and 2) and pustules (Figures 2 and 3), typically associated with hair follicles (Figure 3). However, follicular involvement may be difficult to appreciate macroscopically. Crusts of variable thickness (Figure 4) are common lesions but are sometimes absent. Variable alopecia, erythema and hypo- or hyperpigmentation are often present. Multifocal to coalescing patches of alopecia providing a ‘moth-eaten’ appearance may be the only visible lesions in some short-coated breeds (Figure 5). Epidermal collarettes (Figure 6) and target lesions (annular areas of alopecia, scaling, erythema and hyperpigmentation; Figure 7) may be the most obvious lesions in some cases.

Cytology

Demonstration of coccic from lesionlal skin by cytology is a powerful adjunctive diagnostic test and is strongly encouraged for proper diagnosis. Appropriate techniques need to be used for both specimen collection and examination to optimize the value of this diagnostic procedure.13 Cytology is mandatory in the following circumstances: (i) typical lesions (pustules) are not present or scant and SBF is still suspected; (ii) typical lesions are present but there is a poor response to empirical antimicrobial therapy; or (iii) a bacterial culture is to be performed. This is because positive cytology in the face of a negative culture should prompt repeat culture rather than diagnosis of a sterile pustular disease.

Cytology is also essential for the diagnosis of co-infection with Malassezia pachydermatis (a frequent occurrence in dogs with SBF) or rod-shaped bacteria (a rare occurrence in dogs with SBF). The presence of coccoid bacteria in cytological specimens from typical lesions is highly supportive of bacterial infection; when associated with inflamatory cells and intracellular cocci from intact pustules, infection is confirmed. The absence or scarcity of bacteria and the absence of inflammatory cells or intracellular cocci do not rule out a bacterial infection. Inflammatory cells and phagocytes may be absent in dogs with underlying immunosuppressive diseases or those being treated with immunosuppressive agents, such as glucocorticoids.

Tests to rule out differential diagnoses

Superficial bacterial folliculitis should be distinguished from other inflammatory follicular diseases and is differentiated from dermatophytosis by dermatophyte culture (or Wood’s lamp evaluation or direct examination of hairs for spores) and from demodicosis by deep skin scrapings. Such testing is recommended, and is essential, when
Superficial bacterial folliculitis

Figure 1. Erythematous papules caused by superficial bacterial folliculitis. Note that the dog’s hair has been clipped for visualization of the papules.

Figure 2. Erythematous papules and a pustule (arrow) caused by superficial bacterial folliculitis.

Figure 3. Folliculocentric pustule caused by superficial bacterial folliculitis.

Figure 4. Erythematous papules and crusts on the ventral abdomen of a golden retriever caused by superficial bacterial folliculitis.

Figure 5. Patches of truncal alopecia on a short-haired dog caused by superficial bacterial folliculitis (so-called ‘short-haired dog pyoderma’).

Figure 6. An epidermal collarette caused by superficial bacterial folliculitis.

History and clinical findings are atypical of SBF or the disease is refractory to AMD treatment. Sterile pustular diseases (such as pemphigus foliaceus and sterile neutrophilic or eosinophilic pustulosis) are uncommon to rare and are differentiated on the basis of cytology (absence of bacteria, presence of acantholytic cells).
After initial treatment of SBF was limited to topical AMDs alone and the infections failed to resolve, it is acceptable either to perform bacterial culture and susceptibility testing or to institute empirical systemic AMDs. As systemic AMDs are suggested to be dispensed for a minimum of 3 weeks, it is important that veterinarians educate owners not to continue AMD therapy in the absence of improvement of SBF lesions during this time, or with the emergence of new lesions after 2 weeks of therapy, without veterinary advice.

Pustules are the preferred lesion for specimen collection, and a thorough search for pustules should be made. Clipping hair to facilitate examination of the skin surface and the use of a hand-held magnifying lens can be helpful in detecting pustules. In the absence of pustules, specimens may be obtained from beneath crusts (look for pus present under the crust), epidermal collarettes or papules. Specimen collection methods are summarized in Table 1. Immediate transport of the specimens to the laboratory is recommended, and transport medium should always be used (clinicians should consult with their laboratory if they are uncertain of how to transport their specimens). If delay in submission of specimens is unavoidable, advice on storage should be obtained from the relevant clinical microbiology laboratory.

To date, there are no published reports demonstrating that current use of AMDs has a significant effect on isolation of causative bacteria from dogs with persistent SBF; thus, it is acceptable to collect samples for bacterial culture and susceptibility testing from SBF lesions whenever indicated, regardless of the current use of topical or systemic AMDs.

Table 1. Sampling techniques for lesions of superficial bacterial folliculitis for bacterial culture and susceptibility testing

<table>
<thead>
<tr>
<th>Lesion</th>
<th>Sampling procedure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pustule</td>
<td>No surface disinfection. Clip hair with sterile scissors (avoid clippers). Lance pustule with sterile narrow-gauge needle. If purulent exudate is visible on the needle, apply to a sterile swab; if not, gently touch exudate expelled from pustule with sterile swab and place in transport medium or sterile container.</td>
</tr>
<tr>
<td>Crust</td>
<td>No surface disinfection. Use sterile forceps or a sterile needle to lift the edge of a crust. The presence of exudate under a crust indicates an ideal site for culture. Touch sterile swab to exposed skin surface and place in transport medium or sterile container.</td>
</tr>
<tr>
<td>Epidermal collarette</td>
<td>No surface disinfection. Clip hair with sterile scissors (avoid clippers). Roll sterile swab across border of collarette two or three times and place in transport medium or sterile container.</td>
</tr>
<tr>
<td>Papule*</td>
<td>Sampling by biopsy is probably more reliable. Provide local anaesthesia by subcutaneous injection of 2% lidocaine. Clip hair with sterile scissors or clippers. Clean skin surface by a single wipe with 70% alcohol. Using a sterile 3 or 4 mm punch and sterile surgical instruments, collect tissue sample and place in sterile container or transport medium. Suture biopsy site. Alternatively, papules may be prepared and disinfected as above, then sampled by insertion of a sterile needle and culture of emerging or expressed blood or exudate.</td>
</tr>
</tbody>
</table>

*There is no research to show which method is more appropriate.†This method of disinfection is suggested to kill any surface bacteria. However, there is no research to indicate the value or necessity for any disinfection of the skin surface prior to sampling of papules.
Where possible, laboratories should be used that observe protocols, including those published by the Clinical and Laboratory Standards Institute (CLSI), including material from the CLSI subcommittee on Veterinary Antimicrobial Susceptibility Testing (CLSI-VAST), or the European Committee on Antimicrobial Susceptibility Testing (EUCAST) and other internationally recognized public organizations.

The following AMDs should be tested with all staphylococcal isolates: erythromycin, clindamycin, tetracycline (for testing susceptibility to doxycycline), trimethoprim-sulfamethoxazole, gentamicin, cephalothin (or cefazolin, representing first generation cephalosporins), cefpodoxime (representing third generation cephalosporins), amoxicillin–clavulanate, oxacillin (mecillin) and enrofloxacin (for testing susceptibility to fluoroquinolones). Inclusion of other fluoroquinolones may be considered if enrofloxacin is not the fluoroquinolone drug of choice (CLSI breakpoints are available for difloxacin, enrofloxacin, marbofloxacin and orbifloxacin for dermal Staphylococcus spp.). If erythromycin resistance is determined in the presence of clindamycin susceptibility, the D-test should be performed (or molecular methods for detection of *erm* genes) to determine whether inducible clindamycin resistance is likely. Additional AMDs that may be important for infections with metcillin-resistant staphylococci (MRS) include amikacin, chloramphenicol, minocycline and rifampicin (rifampin). Consultation with a specialist is recommended when treatment with these drugs is being considered. Other antimicrobial drugs which clinicians intend to consider for therapy should also be included. However, regional and national restrictions relating to the use of specific drugs in animals should be observed.

Clinical microbiology laboratories must perform tests to differentiate coagulase-positive staphylococci from CoNS. *S. aureus* should be distinguished from other coagulase-positive staphylococci. This is important for two reasons: (i) the CLSI-determined breakpoints for oxacillin differ for animal species, such as those published by the Clinical and Laboratory Standards Institute (CLSI), including material from the CLSI subcommittee on Veterinary Antimicrobial Susceptibility Testing (CLSI-VAST), or the European Committee on Antimicrobial Susceptibility Testing (EUCAST) and other internationally recognized public organizations.

Recommendations for the treatment of canine superficial bacterial folliculitis

Veterinarians must consider the nature of the disease in each patient to determine the best mode of therapy. Traditional reliance on systemic AMDs and the expectation that empirical choices will always work are now being challenged by the growing frequency of MRS that are resistant to multiple classes of AMDs in addition to the β-lactams. The prevalence of MRS will vary in different localities, and it is important for veterinary practitioners to become familiar with typical local and regional resistance patterns so that they may be prepared to make appropriate selections of modes of treatment and AMDs.

Factors that impact therapy, in addition to antimicrobial resistance, include the severity and extent of lesions, patient factors (such as hair coat, temperament and environment), concurrent disease and the owner’s ability to perform topical or systemic therapy, all of which may affect the efficacy of the chosen therapy.

Owners’ compliance with instructions and completion of treatments is critical to the resolution of infection and prevention of recurrence. Clinicians should maintain contact with owners and support them as far as possible to promote effective compliance. When recurrence of SBF occurs, veterinarians should present owners with a diagnostic plan for evaluation of underlying primary disease (allergic dermatitis, endocrinopathy, etc.) and make it

| 1 | Note staphylococcal species isolated | *Staphylococcus aureus* is a human pathogen and therefore presents a higher public health risk. *Staphylococcus pseudointermedius* is the predominant pathogen in bacterial infections of canine skin. It is a rare cause of human infection but presents enhanced risk if metcillin resistant. Coagulase-negative staphylococci present a much lower level of risk but are often metcillin resistant. They are more likely to be involved in animals with reduced immunity and where implants are used. Low numbers of CoNS should be regarded as probable skin contaminants in patients that are not immunosuppressed, especially when isolated in mixed cultures. If quantitative information is not provided in the report, the laboratory should be consulted before initiating therapy against them. |
| 2 | Is the isolate reported as metcillin resistant? | Oxacillin is equivalent to metcillin and used as a marker of metcillin resistance. Oxacillin-resistant staphylococci are reported as ‘metcillin-resistant’ Metcillin (oxacillin)-resistant staphylococci are by convention resistant to all β-lactam AMDs (cephalosporins, penicillins, carbapenems and monobactams), regardless of occasional apparent *in vitro* susceptibility. Clinical microbiology laboratories must report these isolates as resistant to all β-lactam AMDs Metcillin-resistant staphylococci are commonly resistant to multiple antimicrobials in addition to the β-lactam AMDs, but this is not always the case. |
| 3 | Clinical disease status of patient and history of AMD use | Susceptibility results should always be interpreted in the context of the clinical disease and current and prior history of antimicrobial use in the patient, bearing in mind that susceptibility *in vitro* does not always parallel clinical response in infected animals. |

Table 2. Guidelines for interpretation of microbiology reports by clinicians

Abbreviations: AMD, antimicrobial drug; and CoNS, coagulase-negative staphylococci.

© 2014 ESVD and ACVD, Veterinary Dermatology 5
clear that this is the best means to control recurrence of SBF, reduce AMD use and reduce the likelihood of emergence of drug-resistant infections.

**Topical antimicrobial therapy**

Topical therapy of SBF is probably underused because of the perception that clients will find it more difficult to apply and that compliance may be poor. However, there are significant potential advantages for early and frequent use of the topical approach in this disease. These advantages include more rapid lesion resolution and a decrease in the duration of antimicrobial administration when combined with systemic AMD therapy, removal of organisms and debris from the skin surface, minimal adverse effects and greatly reduced exposure to AMDs of bystander organisms in other organ systems (reducing risk of inadvertent emergence of resistant strains). In addition, resistance to the high concentrations of antiseptics and AMDs used in topical products is very uncommon, and these agents are typically bactericidal to MRS. The emergence of highly multiresistant MRS with few or no options for systemic AMD therapy has provided a new stimulus for the topical approach, which is emerging as an important treatment for multidrug-resistant bacterial infections of the skin.

The benefits and importance of topical antimicrobial therapy and topical therapies that help to restore normal skin structure and function (promoting recovery and enhancing resistance to infection) are likely to emerge as significant options as systemic therapy becomes more limited.

In general, topical therapy is helpful in all patients with SBF. Topical therapy alone (without co-administration of systemic AMDs) is encouraged as a desirable and recommended approach to the treatment of SBF unless precluded by owner and/or patient factors. This is particularly true in the following circumstances: (i) localized lesions of SBF; (ii) early stages of generalized SBF when lesions are mild; and (iii) to help prevent recurrence of SBF while diagnostic procedures for primary underlying skin disease are pursued.

Table 3. Summary of topical antimicrobial treatment options for superficial bacterial folliculitis in the dog

<table>
<thead>
<tr>
<th>Application</th>
<th>Formulations</th>
<th>Agents and modes of use</th>
</tr>
</thead>
<tbody>
<tr>
<td>Extensive or generalized disease</td>
<td>Shampoos, lotions, sprays, rinses and conditioners</td>
<td>Antiseptics, including chlorhexidine (also in combination with miconazole), or benzoyl peroxide are preferred, although ethyl lactate, povidone iodine and triclosan may also provide benefit. Commonly used two or three times weekly until 7 days after lesions resolve and then weekly for prophylaxis.* Can also be used for more localized disease. For shampoos or conditioners that are rinsed from the skin, contact time of 10 min prior to rinsing is important.</td>
</tr>
<tr>
<td>Focal and localized infections</td>
<td>Gels, creams, ointments, lotions and wipes</td>
<td>Antiseptics, including a variety of hydroxyl acids (e.g. acetic, lactic and malic acids), benzoyl peroxide and silver sulfadiazine. Antimicrobial drugs, including novobiocin, pristinamycin, bacitracin, fusidic acid and mupirocin. Mupirocin and fusidic acid are used in human medicine for meticillin-resistant Staphylococcus aureus treatment and decolonization; resistance is increasingly reported. Reports indicate that resistance to topical therapy with these agents in meticillin-resistant staphylococci causing canine superficial bacterial folliculitis is very rare; however, it is recommended that they be reserved for targeted application in dogs with infections where culture and susceptibility indicate no other suitable antimicrobial drugs and where topical antiseptics have failed to resolve the infection.</td>
</tr>
</tbody>
</table>

*Extended treatment duration is based on clinical experience; further research is required to confirm the need for this.

Use of the agents listed should take account of local and regional restrictions on their use.
**Systemic antimicrobial therapy**

Selection of systemic AMDs is based on availability, safety, cost, local prevalence of resistant staphylococci and patient-specific factors (concurrent disease or drug administration, previous drug reactions, etc.). A recent systematic review found the evidence for efficacy of systemic AMDs for treatment of superficial pyoderma to be good for cefovecin, fair for amoxicillin–clavulanate, clindamycin, cefadroxil, trimethoprim–sulphamethoxazole and sulfadimethoxine–ormetoprim and insufficient for cefalexin, cefpodoxime, ibafloxacin, marbofloxacin and lincomycin.37 Despite the value of such reviews, the relative dearth of published studies, lack of standardization of methods for diagnosis and assessment of treatment outcome, as well as the absence of studies with many commonly used AMDs, prevent generation of comprehensive guidelines based solely on their findings.

Choices of suitable AMDs that may be selected for empirical therapy of SBF when risk factors for likelihood of AMD resistance are not present (see indications for bacterial culture above) are grouped as first tier drugs (Table 4). Those AMDs that may be chosen when first tier drugs and topical agents are not appropriate and when culture and susceptibility results indicate susceptibility are grouped as second tier drugs (Table 4). Third tier drugs are also listed, but their use is strongly discouraged and it is recommended that cases be referred for specialist consultation if such AMDs are being considered. Suggested doses for antimicrobial drugs for systemic treatment of superficial bacterial folliculitis in the dog are given in Table 5.

In principle, it would be ideal if veterinarians had available a selection of AMDs for empirical therapy that were narrow spectrum, labelled for treatment of SBF in the dog and to which a majority of S. pseudintermedius were still susceptible. Unfortunately, this is rarely possible because some commonly used AMDs do not have a veterinary label in some countries, few of the commonly used AMDs are narrow spectrum, many AMDs that are registered and approved for use in the treatment of SBF may be associated with the emergence of multidrug-resistant infections, and there is distinct geographical variability in susceptibility of S. pseudintermedius to many of the available AMDs.38,39

Members of this working group have been unable to reach consensus on how the cephalosporins, including cefalexin, cefadroxil, cefpodoxime and cefovecin, should be distributed as first or second tier AMDs. All are approved (in at least one global region) for use in the treatment of skin wounds and abscesses, or pyoderma, in dogs and have demonstrated efficacy in clinical studies; furthermore, a systematic review has shown fair to good evidence for the moderate to high efficacy of cefadroxil and cefovecin in the treatment of SBF.37,40–43 Simple consideration of clinical efficacy would support the inclusion of all these drugs as first tier AMDs. However, there is concern among some members of this panel about the potential selective effects of third generation AMDs for systemic antimicrobial treatment of serious MRSA infections in humans.43

---

**Table 4. Summary of systemic antimicrobial treatment options for superficial bacterial folliculitis in the dog**

<table>
<thead>
<tr>
<th>Category</th>
<th>When used</th>
<th>Suggested AMDs and comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>First tier</td>
<td>Primary choice empirical therapy of known or suspected SBF</td>
<td>Clindamycin or lincomycin First generation cephalosporins (e.g. cefalexin, cefadroxil), Amoxicillin–clavulanate</td>
</tr>
<tr>
<td>First or second tier</td>
<td>Additional choices only if local regional susceptibility of Staphylococcus pseudintermedius is known</td>
<td>Trimethoprim- and ometoprim-potentiated sulphonamides</td>
</tr>
<tr>
<td>Second tier</td>
<td>When empirical selection of first tier systemic AMD and topical therapy are not appropriate and when cultures indicate susceptibility</td>
<td>Doxycycline or minocycline Chloramphenicol Fluoroquinolones (such as enrofloxacin, marbofloxacin, orbifloxacin, pradofloxacin and ciprofloxacin) (should only be used when other feasible options are not available) Rifampicin, Commonly used in combination with another drug to which the causative organism is susceptible; however, this process may not reduce development of resistance in staphylococcal infection75 Aminoglycosides, including gentamicin and amikacin. See Table 5 for comments on nephrotoxicity and ototoxicity First tier AMD (clindamycin, lincomycin and potentiated sulphonamides) may also be considered when cultures indicate susceptibility</td>
</tr>
<tr>
<td>Third tier</td>
<td>When first and second tier are not appropriate and cultures indicate susceptibility</td>
<td>Linezolid, teicoplanin, vancomycin. Regardless of the fact that most (or all) MRSP are susceptible, the use of these three AMDs is strongly discouraged. These drugs can be considered reserved for the treatment of serious MRSA infections in humans.</td>
</tr>
</tbody>
</table>

Abbreviations: AMD, antimicrobial drug; ESBL, extended-spectrum b-lactamase; MRSA, meticillin-resistant Staphylococcus aureus; MRSP, meticillin-resistant Staphylococcus pseudintermedius; and SBF, superficial bacterial folliculitis. Use of the agents listed should take account of local and regional restrictions on their use.
cephalosporins (cefepoxide and cefovecin) on the Gram-negative microbiota, due to their broader spectrum of activity compared with first generation cephalosporins. Both drugs are marketed as extended-spectrum cephalosporins; in addition to approval for use in infections caused by *S. pseudintermedius*, cefepoxide is regarded as a broad-spectrum AMD and has been approved in the USA for use in the treatment of skin infections associated with *Escherichia coli* and *Proteus mirabilis*, whilst cefovecin has been approved in the Europe for use in the treatment of skin infections associated with *E. coli* and for urinary tract infection associated with *E. coli* and *Proteus*. Cefovecin is significantly more active against *E. coli*, *Klebsiella pneumoniae* and *Proteus* spp. compared with cefalexin and cefadroxil, and its *in vitro* activity against *E. coli* and *Proteus* spp. is comparable to that of cefpodoxime.44 Although cefovecin may be considered as a 'narrower-spectrum' drug due to the high-affinity protein binding (and subsequent low free plasma concentration), pharmacokinetic data provided by the manufacturer45 indicate that the free plasma concentration exceeds the MIC90 of *E. coli* for 2 days following injection and exceeds the MIC50 of *E. coli* for 6 days. Thus, concentrations can be sufficient to kill susceptible Gram-negative bacteria, as opposed to only Gram-positive bacteria, which are killed by lower drug concentrations.

### Table 5. Suggested doses for systemic antimicrobial drugs for treatment of superficial bacterial folliculitis in the dog

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dose</th>
<th>Comments</th>
</tr>
</thead>
</table>
| Amikacin                      | 15–30 mg/kg i.v., i.m. or s.c. once daily | Useful for treatment of multidrug-resistant organisms. Potentially nephrotoxic and ototoxic. Avoid in animals with renal insufficiency*
| Amoxicillin–clavulanate       | 12.5–25.0 mg/kg p.o. twice daily | Pharmacokinetic data are available to support the use in dogs with duration of 14 days. Repeat injection after 14 days in most cases if infection is not resolved and to meet the criterion for treatment to 7 days beyond resolution |
| Cefalexin, cefadroxil         | 15–30 mg/kg p.o. twice daily | Reserved for multidrug-resistant infections with few other options. Myelosuppression can occur, particularly with long-term therapy. Vomiting is frequently encountered. Avoid contact by humans because of rare idiosyncratic aplastic anaemia. Wearing of gloves by owners handling the drug is essential |
| Cefovecin                     | 8 mg/kg single s.c. injection | If there is erythromycin resistance with clindamycin susceptibility, the D-test should be performed (or molecular methods for detection of *erm* genes) to determine likelihood of clindamycin resistance |
| Cefpodoxime proxetil          | 5–10 mg/kg o.o. once daily | Potentially nephrotoxic. Avoid in animals with renal insufficiency* |
| Chloramphenicol               | 40–50 mg/kg p.o. three times a day | Pharmacokinetics and dose in dogs have not been evaluated, |
| Ciprofloxacin                 | 25 mg/kg p.o. once daily | If there is erythromycin resistance with clindamycin susceptibility, the D-test should be performed (or molecular methods for detection of *erm* genes) to determine likelihood of clindamycin resistance |
| Clindamycin                   | 5.5–10 mg/kg p.o. twice daily | Pharmacokinetics and dose in dogs have not been evaluated, |
| Doxycycline                   | 5 mg/kg p.o. twice daily or 10 mg/kg once daily | Potentially nephrotoxic. Avoid in animals with renal insufficiency* |
| Enrofloxacin                  | 5–20 mg/kg p.o. once daily | Pharmacokinetics and dose in dogs have not been evaluated, |
| Lincomycin                    | 15–25 mg/kg p.o. twice daily | Pharmacokinetics and dose in dogs have not been evaluated, |
| Gentamicin                    | 9–14 mg/kg i.v., i.m. or s.c. once daily | Potentially nephrotoxic. Avoid in animals with renal insufficiency* |
| Marbofloxacin                 | 2.75–5.5 mg/kg p.o. once daily | Pharmacokinetics and dose in dogs have not been evaluated, |
| Minocycline                   | 10 mg/kg p.o. twice daily | Pharmacokinetics and dose in dogs have not been evaluated, |
| Orbifloxacin                  | 7.5 mg/kg p.o. once daily | Pharmacokinetics and dose in dogs have not been evaluated, |
| Ometoprim–sulfadimethoxine    | 55 mg/kg on first day, then 27.5 mg/kg p.o. once daily | Pharmacokinetics and dose in dogs have not been evaluated, |
| Pradofloxacin                 | 3.0 mg/kg p.o. once daily | Pharmacokinetics and dose in dogs have not been evaluated, |
| Rifampicin                    | 5–10 mg/kg p.o. twice daily | Pharmacokinetics and dose in dogs have not been evaluated, |
| Trimethoprim–sulfadiazine or sulfoxamethoxazole | 15–30 mg/kg p.o. twice daily | Pharmacokinetics and dose in dogs have not been evaluated, |

*Abbreviations: i.m., intramuscular; i.v., intravenous; p.o., per os; and s.c., subcutaneous.

*See IRIS: International Renal Interest Society guidelines for prevention of aminoglycoside-induced acute kidney injury; www.iris-kidney.com

Use of the agents listed should take account of local and regional restrictions on their use.

© 2014 ESVD and ACVD, Veterinary Dermatology
concentrations. This raises concerns about possible selection of highly resistant extended-spectrum β-lactamase (ESBL)-producing *E. coli* by use of cefovecin. As for cefpodoxime, this extended-spectrum cephalosporin is administered as a prodrug, cefpodoxime proxetil, which is absorbed and de-esterified in the gastrointestinal tract to its active metabolite.46 Thus, it is questionable whether the active metabolite may reach sufficient concentrations in the large intestine to select for ESBL-producing bacteria. These concerns notwithstanding, at least one member of the panel was not convinced that there is sufficient published evidence indicating that cefovecin or cefpodoxime produce active concentrations in the intestinal lumen of dogs that are sufficient to affect the microbial population.

A few recent studies in dogs have identified antimicrobial drug use in general as a risk factor for the emergence of MRSP, and, at present, it is reasonable to assume that any cephalosporin or amoxicillin-clavulanic acid could select for MRSP. One small report has associated misuse of unspecified fluoroquinolones, macrolides and third-generation cephalosporins with persistence of MRSP colonization in a breeding kennel.47 The use of fluoroquinolones and extended-spectrum cephalosporins in humans,48–50 and of fluoroquinolones in dogs, is a known risk factor for selection of MRSA.51 Use of these AMDs is also a risk factor for selection of ESBL-producing *E. coli* in both humans and animals,52–55 and guidelines in human medicine recommend prudent use of these broad-spectrum agents to prevent spread of multidrug-resistant bacteria.56–58 These factors, along with the increasingly high prevalence of MRSP and ESBL-producing Enterobacteriaceae in dogs, support the promotion of precautionary principles and the limitation of extended-spectrum cephalosporins and fluoroquinolones as second tier AMDs. In accordance with this, the package insert for cefovecin in Europe specifies that ‘A sample of the lesion should be obtained for culture and susceptibility testing prior to beginning antimicrobial therapy’.45 and the technical monograph states, in addition, ‘It is prudent to reserve third generation cephalosporins for the treatment of clinical conditions, which have responded poorly, or are expected to respond poorly, to other classes of antimicrobials or first generation cephalosporins’.59

With regard to the fluoroquinolones, enrofloxacin, marbofloxacin, orbifloxacin and pradofloxacin are approved for use in dogs in some countries and have been shown to be effective for the treatment of superficial pyoderma. However, the use of this group of AMDs is a known risk factor for the emergence of MRSA in humans,48–50 and guidelines also recommend limited use of these agents.56–58

When recurrence of SBF occurs, careful consideration of culture and susceptibility testing is encouraged because previous exposure to AMDs is a risk factor for resistance24,25 and may be especially important in patients with previous MRSP infections or from households with other pets that have previously been diagnosed with an MRSP infection.26,27 Veterinarians should present a plan for evaluation of underlying primary disease to owners of dogs with recurrent infections. If culture is not performed on recurrence of the infection, the same AMD should be used that successfully resolved the previous infection. Most studies evaluating the efficacy of AMDs indicate that SBF infections are resolved after 3 weeks or more of systemic AMD treatment; rapid improvement over the first 1–2 weeks is typically observed, but resolution of all lesions and prevention of rapid recurrence of disease requires 3–6 weeks of treatment.17–22,28 Although there is no significant difference in the likelihood of resolution of MSSP after 3–4 weeks of systemic AMD treatment compared with MRSP infections, it has been reported that MRSP infections took longer to treat compared with MSSP infections.60

In a minority of patients, resolution of lesions may be achieved with 2 weeks of systemic AMDs. However, the assessment of complete resolution cannot be left to pet owners, and all patients should ideally be re-evaluated to ensure resolution of the infection. In particular, if attending veterinarians dispense <3 weeks of AMDs, they should anticipate and be confident that the patient will be presented for re-evaluation to determine whether additional antimicrobial therapy is indicated or the infection has resolved on completion of this period. Furthermore, patients with a history of recurrent SBF must be re-evaluated at the conclusion of AMD treatment.

In the absence of evidence to the contrary, continuation of treatment for at least 7 days beyond clinical resolution of lesions is recommended in all cases,14 because the inflammatory process and lesions will subside and become inapparent as the infection is eliminated. This extended duration of treatment is based on clinical experience. Further research is required to confirm the need for such additional therapy, whether a 7 day period is sufficient, and to determine methods that will confirm whether infection has been eliminated when clinical lesions have resolved. Concurrent glucocorticoid use during therapy of SBF is strongly discouraged because it may improve the clinical appearance of the lesions and result in premature discontinuation of AMD administration whilst also reducing the patient’s innate and adaptive immune response to infection.

Prevention of superficial bacterial folliculitis

The most effective measure to prevent recurrence is to identify and control the underlying primary disease. Protocols for the use of systemic AMDs to aid in the prevention of SBF, or to delay recurrence, have been published and advocated in public prior to the widespread emergence of MRS and have included pulse therapy (intermittent administration of therapeutic doses of AMD) and continuous use of subtherapeutic dosing.61,62 However, there is significant concern for the selection of resistance with these protocols. Accordingly, their use is strongly discouraged. The use of autogenous bacterins62,63 or commercial bacterial antigens64 is encouraged. However, very few studies of the efficacy and usefulness of these measures have been reported and further research is necessary. If pulse or subminimal AMD therapy is being considered for prevention of SBF, it is recommended that the patient be referred to a specialist for further evaluation and treatment.

Decolonization of carriage sites has been demonstrated to reduce the recurrence of MRSA infections in humans.65,66 Although recurrent MRSP infections are
common, there are currently no controlled studies in dogs that would indicate potential effective methods of decollo-
ination, nor the need for such procedures. Therefore, at
this time, routine decolonization of carriage sites of dogs
with recurrent MRSP infections is of questionable value
and not recommended.

Public health considerations
Staphylococci can be transferred in both directions between
animals and humans.71 Whilst the risk of infec-
tion with S. pseudintermedius and S. schleiferi is very
low in healthy humans, infections by pathogenic staphylo-
cocci acquired from pets have been documented.68,69
Such infections are a much greater hazard in the case of
MRS, particularly with MRSA.70

Precautions need to be taken to limit the possibility of
transfer of staphylococci from infected animals to owners
and veterinary staff in the clinic. Owners and veterinary
staff need to be aware of this potential hazard and
advised on measures to minimize the risk of transfer, par-
ticularly when susceptible individuals (elderly people,
those with lesions or diseases rendering them more sus-
sceptible to infection and those receiving immunsuppres-
sive therapy) are likely to come into contact with the
affected animals.

Infection control measures
Hygiene should be maintained rigorously in the clinic
when animals suspected of having staphylococcal infec-
tions are admitted. This should involve the development
and display of hygiene protocols specific for each clinic
environment. Staff should be trained to recognize risk fac-
tors for multiresistance and observe such protocols; com-
pliance should be monitored and enforced. Materials
likely to have been contaminated should be disinfected
after such animals are seen, and effective hand cleansing
with alcohol sanitizers must be carried out before and
after touching the animal. Owners of animals with sus-
pected staphylococcal infections should also be advised
of the importance of hygiene. Detailed recommendations
on hygiene in the clinic are beyond the scope of this arti-
cle, and readers are advised to refer to other published
material on this topic.71–73

Summary of recommendations
See Appendix 1.

References
1. Guardabassi L, Houser GA, Frank LA et al. Guidelines for antimicro-
ical use in dogs and cats. In: Guardabassi L, Jensen LB,
2. Edin Rantalai M, Haf I, Lillas A et al. Survey of condition-based
prescribing of antimicrobial drugs for dogs at a veterinary teach-
prior to admission to a veterinary teaching hospital. J Am Vet
guidelines to support judicious use of antibiotic therapy. J Clin
Pharm Ther 2010; 35: 71–78.
5. Mejilian TA, Prasad PA, Kogan A et al. Evaluation of an antimi-
icrobial stewardship program at a pediatric teaching hospital. Pediatr
6. Toth NR, Chambers RM, Davis SL. Implementation of a care
bundle for antimicrobial stewardship. Am J Health Syst Pharm
2010; 67: 746–749.
7. Beco L, Guaguere E, Lorente Méndez C et al. Suggested guide-
lines for using systemic antimicrobials in bacterial skin infec-
tions: part 2—antimicrobial choice, treatment regimens and
8. Devriese LA, Vancanneyt M, Baele M et al. Staphylococcus
pseudintermedius sp. nov., a coagulase-positive species from
222: 451–454.
10. Cain CL, Morris DO, O’Shea K et al. Genotypic relatedness and
phenotypic characterization of Staphylococcus schleiferi species
11. Fortin M, Higgins R. Mixed infection associated with a group B
12. Hillier A, Alcorn JR, Cole LK et al. Pyoderma caused by Pseu-
odomas aeruginosa in dogs: 20 cases. Vet Dermatol
13. Mendelsohn C, Rosenkranz W, Griffin CE. Practical cytology for
14. Miller WH, Griffin CE, Campbell KL, Muller & Kirk’s Small Animal
15. Toma S, Colombo S, Corneliani L et al. Efficacy and tolerability of
once-daily cephalixin in canine superficial pyoderma: an open
16. Miller WH, Griffin CE, Campbell KL, Muller & Kirk’s Small Animal
17. Frank LA, Kunkle GA. Comparison of the efficacy of cefadrox-
il and generic and proprietary co-amoxiclav in the treatment of
18. Harvey RG, Noble WC, Ferguson EA. A comparison of linco-
mycin hydrochloride and clindamycin hydrochloride in the
treatment of superficial pyoderma in dogs. Vet Rec
19. Littlewood JD, Lakhani KH, Paterson S et al. Clindamycin hydro-
chloride and clavulanate-amoxicillin in the treatment of canine
20. Lloyd DH, Carlotti DN, Koch HJ et al. Treatment of canine pyo-
derma with co-amoxiclav: a comparison of two dose rates. Vet
Rec 1997; 141: 439–441.
21. Messinger LM, Beale KM. A blinded comparison of the efficacy of
daily and twice daily trimethoprim-sulfadiazine and daily sulfa-
dimethoxine-ormetoprim in the treatment of canine pyoderma.
22. Bloom PB, Rossier EJ. Efficacy of once-daily clindamycin hydro-
chloride in the treatment of superficial bacterial pyoderma in
23. Soares Magalhães RJ, Loeffler A, Lindsay J et al. Risk fac-
tors for meticillin-resistant Staphylococcus aureus (MRSA)
infection in dogs and cats: a case-control study. Vet Res
2010; 41: 55.
Staphylococcus pseudintermedius among dogs admitted to a
25. Weese JS, Faires MC, Frank LA et al. Factors associated with
meticillin-resistant versus meticillin-susceptible Staphylococ-
cus pseudintermedius infection in dogs. J Vet Med Assoc
in-resistant Staphylococcus pseudintermedius (MRSP) from
skin and carriage sites of dogs after treatment of their meticil-
lin-resistant or meticillin-sensitive staphylococcal pyoderma.


© 2014 ESVAD and ACVD, Veterinary Dermatology


Appendix 1: Summary of guidelines for the diagnosis and antimicrobial therapy of canine superficial bacterial folliculitis

Superficial bacterial folliculitis in dogs is typically caused by *Staphylococcus pseudintermedius*.

**Diagnosis**: Initially based on clinical signs of papules, pustules, crusts, patchy alopecia or epidermal collarettes. Cytological demonstration of cocci and inflammatory cells is strongly encouraged to support the diagnosis. Bacterial culture and susceptibility testing is encouraged with recurrent infections and is essential when there is <50% reduction in lesions after 2 weeks of therapy, new acute lesions emerge after 2 weeks of therapy, infection has not resolved after 6 weeks of therapy, intracellular rods are detected on cytology or there is a history of prior multidrug-resistant infection. Pustules are the preferred lesion to culture, but crusts, epidermal collarettes and papules may also be sampled.

<table>
<thead>
<tr>
<th>Application</th>
<th>Formulations</th>
<th>Agents</th>
<th>Treatment recommendations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Extensive or generalized disease</td>
<td>Shampoos, lotions, rinses, sprays, conditioners</td>
<td>Antiseptics, including chlorhexidine (also with miconazole) and benzoyl peroxide, are preferred, but ethyl lactate, povidone iodine and triclosan may also be helpful</td>
<td>Two or three times weekly. Shampoos or conditioners: 10 min contact time prior to rinsing</td>
</tr>
<tr>
<td>Focal and localized infections</td>
<td>Gels, creams, ointments, lotions and wipes</td>
<td>Antiseptics, including hydroxyl acids (e.g. acetic, lactic and malic acids), benzoyl peroxide and silver sulfadiazine. Antimicrobial drugs, including novobiocin, pristinamycin, bacitracin, fusidic acid and mupirocin</td>
<td>Use one or two times daily</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Category</th>
<th>When used</th>
<th>Suggested antimicrobial drugs</th>
<th>Dosing</th>
</tr>
</thead>
<tbody>
<tr>
<td>Systemic antimicrobial therapy*†</td>
<td>Empirical therapy of known or suspected superficial bacterial folliculitis</td>
<td>First generation cephalosporins (e.g. cefalexin, cefadroxil)</td>
<td>15–30 mg/kg p.o. twice daily</td>
</tr>
<tr>
<td>First tier</td>
<td></td>
<td>Amoxicillin-clavulanate</td>
<td>12.5–25 mg/kg p.o. two to three times a day</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Clindamycin</td>
<td>5.5–10 mg/kg p.o. twice daily</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Lincomycin</td>
<td>15–25 mg/kg p.o. twice daily</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Trimethoprim–sulphonamides</td>
<td>15–30 mg/kg p.o. twice daily</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ormetoprim–sulphonamides</td>
<td>55 mg/kg on first day then 27.5 mg/kg p.o. once daily</td>
</tr>
<tr>
<td>First or second tier</td>
<td></td>
<td>Cefovecin</td>
<td>8 mg/kg s.c. once every 2 weeks</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Cefpodoxime</td>
<td>5–10 mg/kg p.o. once daily</td>
</tr>
<tr>
<td>Second tier</td>
<td>First tier systemic antimicrobial drug and topical therapy ineffective. Selection based on culture and susceptibility testing</td>
<td>Doxycycline</td>
<td>5 mg/kg p.o. twice daily; or 10 mg/kg p.o. once daily</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Minocycline</td>
<td>10 mg/kg p.o. twice daily</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Chloramphenicol Fluoroquinolones:</td>
<td>40–50 mg/kg p.o. three times a day</td>
</tr>
<tr>
<td></td>
<td></td>
<td>enrofloxacin</td>
<td>5–20 mg/kg once daily</td>
</tr>
<tr>
<td></td>
<td></td>
<td>marbofloxacin</td>
<td>2.75–5.5 mg/kg once daily</td>
</tr>
<tr>
<td></td>
<td></td>
<td>orbifloxacin</td>
<td>7.5 mg/kg p.o. once daily</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ciprofloxacin</td>
<td>25 mg/kg once daily</td>
</tr>
<tr>
<td></td>
<td></td>
<td>pradofloxacin</td>
<td>3 mg/kg p.o. once daily</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Rifampicin Aminoglycosides:</td>
<td>5–10 mg/kg p.o. twice daily</td>
</tr>
<tr>
<td></td>
<td></td>
<td>gentamicin</td>
<td>9–14 mg/kg i.v., i.m. or s.c. once daily</td>
</tr>
<tr>
<td></td>
<td></td>
<td>amikacin</td>
<td>15–30 mg/kg i.v., i.m. or s.c. once daily</td>
</tr>
<tr>
<td>Third tier</td>
<td>Vancomycin, teicoplanin and linezolid</td>
<td>Use strongly discouraged</td>
<td></td>
</tr>
</tbody>
</table>

*Therapy must be administered for at least 3 weeks or until 7 days beyond resolution of lesions.
†Use of the agents listed should take account of local and regional restrictions on their use.

Abbreviations: i.m., intramuscular; i.v., intravenous; p.o. per os; and s.c., subcutaneous.

© 2014 ESVD and ACVD, *Veterinary Dermatology*
Die superfizielle bakterielle Follikulitis (SBF) wird u. blicherweise von *Staphylococcus pseudintermedius* verursacht und routinemäßig mit systemischen Antibiotika behandelt. Eine Infektion ist die Konsequenz einer reduzierten Immunität, die mit Anderungen der Hautbarriere und zugrundeliegender Erkrankungen, deren Diagnose und Heilung manchmal schwierig sind, einhergeht; daher kehrt die SBF häufig wieder und eine Behandlung ist wiederholt nichtig. Durch das Aufkommen von multiresistenten Bakterien, vor allem Methicillin-resistentem *S. pseudintermedius* (MRSP), konzentriert sich die Aufmerksamkeit auf den Bedarf einer optimalen Behandlung der SBF.
Ziele – Bereitstellung einer international verfügbaren Quelle, die PraktikerInnen bei der Diagnose, der Behandlung und der Vorbeugung einer SBF unterstützt.


Schlussfolgerungen und klinische Bedeutung – Ein optimales Management von SBF wird die Verwendung von antimikrobiellen Wirkstoffen verbessern und die Selektion von MRSP und anderen multiresistenten Bakterien, die die tierische und die menschliche Gesundheit beeinträchtigen, reduzieren.

要旨 – 著者研究細菌性毛髪炎（SBF）は一般的に Staphylococcus pseudintermedius により生じ、通常全身性抗生物質を用いて治療される。感染は、表在性膿瘍と診断された場合、腫瘍の変化に関与した、皮膚の感染がその結果であり、そのため SBF は頻繁に再発し、繰り返した治療が必要となる。多剤耐性細菌の感染、特にウイルス性、S. pseudintermedius (MRSP) は SBF の治療の必要性に急がざるを得ない。目指す – SBF の診断、治療および予防に関して臨床家を指導する国際的に入手可能な資料の提供

ガイドラインの開発 – ガイドラインは Antimicrobial Guidelines Working Group of the International Society for Companion Animal Infectious Diseases により、American および European Colleges für Veterinary Dermatology の共同作成のもと、ガイドラインでは原因微生物の分離、抗生物質の感受性試験、抗生物質の選択、治療プロトコルならびに感染のコントロールについての助言を含む SBF の診断および管理に関する最善な方法を説明する。アドバイスとして MRSP に関しては適切なアプローチを含む耐性および感染の方法を示している。全身的な治療は 3つの段階に分類される。段階 1 の治療は診断が明確で抗菌薬の耐性に対するリスク因子がないときに使用する。一方で、段階 2 の薬剤は抗菌薬の感受性試験が必須として使用する。段階 3 は重篤な感染症の為に確実されていった薬剤を含み、それらの使用は極端に際して増加され、もし必要である場合は、臨床家の相談のもとに使用するべきである。

結論および臨床的な重要性 – SBF の適切な管理は抗菌薬の使用を改善し、動物やヒトの健康に影響を及ぼす MRSP および他の多重耐性血球の選択を減少する。

要約 – 表在性細菌性毛髪炎（SBF）は一般的に Staphylococcus pseudintermedius により生じ、通常全身性抗生物質を用いて治療される。感染は、皮膚の感染がその結果であり、そのため SBF は頻繁に再発し、繰り返した治療が必要となる。多剤耐性細菌の感染、特にウイルス性、S. pseudintermedius (MRSP) は SBF の治療の必要性に急がざるを得ない。