Virbac thanks everyone who has participated in the elaboration of this document for their invaluable collaboration.

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Pierre JASMIN, DVM

This book is a truly original manual of canine dermatology. Many dermatological guides or handbooks are available and may be pleasant to read and convey a good understanding of canine dermatoses. However, this book puts forward a quite novel and stimulating approach.

Pierre JASMIN, its author, has developed the concept of classifying the dermatoses in a special way which facilitates the clinical approach. He distinguishes three groups of cutaneous diseases: first, a group of infectious and parasitic, primary or secondary dermatoses which are considered initially. A second group containing the most common primary dermatoses is then presented and, finally, a third group of rare primary dermatoses is considered. His practical approach is thus applied sequentially from group 1 through to group 3. These three steps are modulated according to the age of the animal, enabling dermatoses of young dogs to be distinguished from those of adult dogs.

A diagnostic approach to otitis externa is proposed separately.

This original and efficient approach is complemented by summaries of the essential elements of every dermatological consultation: history-taking and the clinical examination. Finally, Dr. JASMIN reviews one of the most useful but least understood of the dermatological complementary examination procedures: cutaneous histopathology.

An important feature is the presentation of the common dermatoses as monographs in the middle section of the handbook. These diseases are reviewed in a concise and consistent manner covering definition and general information, aetiology and pathogenesis, epidemiology, clinical signs, diagnosis and, of course, treatment. Case follow-up and prognosis are considered at the end of each monograph.

This practical, rigorous and well illustrated document provides a fertile source for the student and the aspiring clinician, as well as the experienced practitioner already interested in dermatology. It provides rapid access to a wide range of knowledge and data, relevant to dermatological practice and in this way simplifies and facilitates the diagnostic approach to dermatological cases.

This book should find a place in the clinic of every canine veterinary surgeon positioned for ready reference as the problems of canine dermatology are encountered in daily practice.

Didier Noël CARLOTTI, DVM, Diplomate ECVD

Preface of the third edition

This edition has included new information on many dermatoses and recent advances on otitis externa. The practicability remains unchanged, of course, and the need for a 3rd edition demonstrates the usefulness of Dr Jasmin’s approach in every day practice. Both students and veterinary clinicians will have at their disposal an updated tool of great quality and readily usable.

Didier Noël CARLOTTI, DVM, Diplomate ECVD

Before speaking

Canine dermatology is, and will remain, a challenging field. It requires a very wide variety of knowledge ranging in many topics including: bacteriology, mycology, parasitology, immunology, allergology, endocrinology, oncology... But it is also this variety that makes it so engaging and enthralling.

And if it is true that therapeutic failure shows evidence, one should keep in mind that it is the same with clinical improvement. Dermatology may thus be frustrating at times, but it can also generate much satisfaction.

However complex this field is, keep in mind that a large number of cases may be achieved successfully: patient relief, owner satisfaction and gratification of the clinician. In order to accomplish this goal, it is necessary to follow a method, to master it, to refine it and to ultimately model it for itself.

May I by this handbook help you to achieve this. It is anyhow this ambitious intent that motivated me with passion all along its conception.

Pierre JASMIN, DVM

Clinical Handbook on Canine Dermatology
Presentation

The dermatological approach

Taking the history
Dermatological examination - Lesions
Differential diagnosis for groups of related dermatoses
  1/ General approach
  2/ Otitis externa
  3/ Dermatoses of adult dogs
  4/ Dermatoses of young dogs

Monographs of the major canine dermatoses

- Dermatophytosis
- Sarcoptic mange
- Dermatoses associated with infestations by other parasites
- Demodicosis and pyodermocosis (juvenile onset and adult onset)
- Pyoderma - General presentation
- Simple approach pyoderma
- Complex approach pyoderma and Bacterial overgrowth (BOG) syndrome
- Malassezia dermatitis and yeast overgrowth
- Flea Allergy Dermatitis (FAD)
- Adverse Food Reactions (AFR)(Food Allergy and Food Intolerance)
- Canine Atopic Dermatitis (CAD; Atopy)
- Iatrogenic and spontaneous Cushing’s syndromes (ICS and SCS)
- Hypothyroidism
- Otitis externa

Applications and use of skin biopsies

Elements of physiology and structure of healthy canine skin
Presentation

It will be important in the first instance for users to read the presentation and first part of the handbook in their entirety to gain a proper understanding of the aims and general philosophy of the dermatological approach proposed.

Concept of the document

This document is designed to be used as a reference aid for the diagnosis and management of canine dermatological cases on a daily basis. It is not an exhaustive bibliographic review, but should be used as a clinical handbook available for consultation at any time and presenting a rational approach to the diagnosis and management of canine dermatological cases. It aims to provide a practical and efficient pathway that, in most cases, will enable the practitioner to establish an accurate diagnosis and formulate an appropriate treatment or management plan.

The need for a clinical handbook

Why does Dermatology remain such a difficult area of veterinary medicine, even though skin lesions are readily observed and the skin is easily accessible for detailed examination and sampling? There are three principal reasons for this paradox.

1. The skin has a limited range of reaction patterns and so many different dermatoses have lesions which appear similar. Clinical signs are thus of limited diagnostic significance.

2. Although the skin lends itself to biopsy sampling the value of histopathology (even though rich in information) is quite variable amongst different skin diseases. Indeed, the diagnostic potential of histopathology is in many cases inversely proportional to the frequency of the dermatosis.

3. Several dermatoses may occur simultaneously on the same animal, presenting a complex picture. This may require sequential investigation, first of secondary dermatoses and then of the primary disease(s).
Solutions provided by the Clinical Handbook

1-Grouping of Dermatoses by Frequency and Sequence of Investigation.

Fortunately, most dermatological cases seen in practice are caused by a small number of specific diseases. In all of these conditions a clinical approach is essential and histopathology is seldom useful. **We can divide these conditions into two groups (Groups 1 and 2) based on the sequence of investigative procedures which is required.** Group 1 includes a small number of dermatoses associated with infectious or parasitic agents. This group is subdivided into 1a, uncommon primary dermatoses, and 1b, common secondary dermatoses. Group 2 is made up of a small number of commonly occuring primary dermatoses that are typically associated with the secondary conditions of group 1b.

The other dermatoses, which are uncommon, but which encompass a much larger range of diseases, are placed in **Group 3.** These conditions require a more sophisticated approach, in which histopathology plays a very important part. Investigation should include the use of skin biopsy samples. Biopsies are also required in all cases in which other diagnostic procedures have failed to provide a precise diagnosis.

<table>
<thead>
<tr>
<th>DIAGNOSTIC GROUPS OF RELATED DERMATOSES</th>
<th>SEQUENCE OF INVESTIGATION</th>
</tr>
</thead>
</table>
| **GROUP 1:** Specific infectious and parasitic dermatoses | **GROUP 1a:** Some primary dermatoses  
*Few and relatively uncommon.* | **Always as a first step** |
| **GROUP 1b:** Secondary dermatoses  
*Few but common.* | |
| **GROUP 2:** Commonest primary dermatoses  
*Few but very common.* | **Only as a second step** |
| **GROUP 3:** Rare primary dermatoses  
*Many different but rare dermatoses.* | **At any time if suspected** |

This approach must permit us to progress, in each case, toward the diagnosis of the encountered dermatosis, because it explains in a concrete and practical way which dermatoses have to be previously excluded (or confirmed) in a first step, to be able to carry on toward a precise diagnosis.
2-Separation of Animals into Two Categories: Young Dogs and Adults.

A second fundamental classification in this approach is the separation of cases into two age ranges, young dogs and adults, where the division between the two categories is puberty.

Why this separation? Sexual maturity is associated with the development of a mature endocrine system and immune system, changes which have major clinical consequences in dermatology. Dermatological conditions in dogs are actually fundamentally different depending on whether the dog is “mature” or not. This necessitates a specific dermatological approach for each of these two categories.

In dogs, there is considerable variation between breeds in the age of acquisition of puberty. One year is a good marker for medium-sized breeds but in small breeds puberty will tend to occur earlier; in large breeds it will be later. Exceptions to this pattern may exist amongst specific breeds and between individuals.

Courtesy of Pierre Jasmin
Content and Structure of the Clinical Handbook

Part 1 presents the general approach and specific procedures to follow when faced with a dermatological problem:

1. How to initiate the approach and take the history.
2. Dermatological examination, identification of the principal lesions and use of a lesion map.
3. Identifying the different diagnostic groups (1a, 1b, 2 and 3), assembly of a list of appropriate differential diagnoses corresponding to each group and the sequence of diagnostic procedures.
   - The concept is presented first as a table which specifies the sequence of diagnostic procedures according to their groups.
   - Specific dermatoses to be considered in the two age groups, adults and young dogs, are listed for each group.
   - The management of otitis externa, which is considered as a specific dermatological problem, is described separately in a table. A specific monograph is devoted to otitis and describes a clinical method of investigation.

Part 2 presents monographs in which each dermatosis or group of dermatoses is reviewed from a clinical and practical perspective. Only the useful key points essential to the proper management of the problem are described. The same general plan is followed for all the monographs, including definition, aetiology, pathogenesis, epidemiology, clinical signs, diagnosis, treatment, prognosis, follow-up and conclusions. Note that some agents mentioned under Treatment may not be licensed or available in your country.

Part 3 is devoted to a detailed analysis of the effective use of skin biopsies, enabling the reader to maximise the efficiency and diagnostic power of the histopathological examination.

Part 4 presents some elements on canine skin physiology, structure, epidermis renewal, follicular cycle and differences with human skin.

Philosophy of the Clinical Handbook

An analysis of cases encountered in practice led to the realisation that a diagnostic approach could be designed which would enable the small number of very common dermatoses to be efficiently diagnosed, using clinical procedures and differentiating the conditions suffered by young dogs and adults. Diagnosis of the many different but rarer dermatoses could be achieved with the aid of histopathological examination of skin biopsy specimens.

This approach has enabled us to provide a framework for diagnosis and management of the great majority of cases without neglecting the diagnostic approaches required for rarer conditions.

This document is the product of this analysis and has been assembled with the collaboration of some of the best-known specialists in veterinary dermatology. Virbac, faithful to its philosophy, hopes that this book will address both your needs and your expectations.
Part.1
The dermatological approach
Clinical Handbook on Canine Dermatology
A dermatological sign (Pruritus, Alopecia, Otitis...) can be:

**Taking the history**

A reason for consultation

Noticed at consultation

In every case plan sufficient time
to fill in the History Sheet and to initiate the approach.

Evaluate the time available to you and the owner. Allocate sufficient time to enable you to explain and initiate the approach.

You can give the history sheet to the owner and ask him/her to complete it in the waiting room.

Information on the sheet should be amplified by questioning the owner. You and the owner should be seated. Take the dog off the consultation table if it was first placed there.

Make a special appointment if necessary and give the owner the history sheet to fill in at home. Ask him/her to bring it back on the day of the appointment.
**History Sheet**

**General History**

1/ Age when acquired and origin (breeder, pet store...): .................................................................
2/ Geographical area of home, present and past (trips, kennels...):
3/ Way of life, environment (exercise, house, garden, flat...) and in-contact pets:
4/ Diet (current and previous):
5/ Reproduction (frequency of oestrus, pregnancy, behaviour of males...):
6/ Previous diseases and surgery:
7/ Unrelated previous dermatoses:
8/ Current non skin-related problems:
9/ Behavioural changes:

**Dermatological History**

1/ Chief complaint:
2/ Date of onset:
3/ Pruritus, severity and time of onset (before or after lesions):
4/ Specific clinical signs and related behaviour:
   • Itching of feet (licking, chewing):
   • Itching of head, lips, chin, peri-orbital area (scratching, rubbing):
   • Otitis, itching of ear pinnae (scratching, shaking):
   • Itching of anogenital area (rubbing, licking and chewing):
   • Ophthalmological signs, conjunctivitis and/or epiphora (ocular secretion):
   • Upper respiratory signs, nasal discharge and/or sneezing:
   • Lower respiratory signs, asthmatic problems:
5/ First cutaneous lesions observed and their development:
6/ Seasonal influences:
7/ Other affected animals (cohabiting, lineage) or human beings in contact:
8/ Presence of fleas and anti-parasitic treatments given:
9/ Shampoos and grooming:
10/ Owner’s view (personal opinion, remarks) on the dermatosis of his dog:
11/ Previous treatments (date, products, dosage and duration) and responses:

---

Clinical Handbook on Canine Dermatology
Dermatological examination - Lesions

Standard general examination

Including weight, temperature, pulse, auscultation, lymph nodes and abdominal palpation, ophthalmological examination...

Thorough dermatological examination

Search carefully for primary and secondary lesions with a lens if necessary.
Note all these lesions on the silhouette.
The entire skin and haircoat must be thoroughly investigated. This should be done first when the dog is standing up, starting from the face down to the tail, going through the back and the flanks. Then the dog is laid down to examine the ventral aspect and the limbs to their extremities.

Using the information provided by the history and the silhouette, and taking account of any abnormalities detected in the general examination, the list of differential diagnoses is developed and related to the defined disease group (see the table page 9 explaining the general approach to differential diagnosis for groups of related dermatoses).

These diseases are listed separately for the adult dog and the young dog in two different action plans. Adopt the plan appropriate to the age category of the case.*

The history and the lesion map are essential for the clinical approach but will also be found very useful in supporting interpretation of histopathological analysis and for the clinical follow-up.

* See the presentation of the handbook which defines the difference between young dog and adult dog.
## PRIMARY LESIONS

<table>
<thead>
<tr>
<th>Definition and Clinical Elements</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ERYTHEMA</strong></td>
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</table>

**PAPULES** | Raised, solid, circumscribed elevations of the skin, from 1 mm up to 1 cm in diameter, usually indicative of a cellular infiltrate. |

**PUSTULES** | **SUPERFICIAL PUSTULES AND FURUNCLES** | Raised, circumscribed, normally thin-walled lesions containing pus (typically 1 to 3 mm). If follicular, consider follicular infections involving bacteria (pyoderma), Demodex (demodicosis), dermatophytes (ringworm). If non-follicular - consider impetigo in young dogs, bullous autoimmune dermatoses in adults. Characteristic secondary lesions are epidermal collarettes. Pustules may extend by rupture of hair follicles to form furuncles. Dermal involvement causes vascular damage; blood is commonly present in the more abundant purulent discharge as in furunculosis, pyodemodicosis... |

**CRUSTED PAPULES** | Papules surmounted with a small central crust, may need examination with a hand lens. Commonly found in sarcoptic mange. |

**VESICLES BULLAE** | Small, raised, circumscribed, fluid-filled epidermal lesions. If larger than 1 cm are called bullae. |

**NODULES** | Raised, solid cutaneous masses greater than 1 cm in diameter. Caused by inflammatory infiltrates or neoplasia within the dermis or rarely, in the subcutis. |

**TUMOURS** | Solid masses involving any tissue in the skin or subcutis and usually caused by neoplasia or granuloma formation - usually circumscribed but may be diffuse. |

## SECONDARY LESIONS

<table>
<thead>
<tr>
<th>Definition and Clinical Elements</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SCALES</strong>*</td>
</tr>
</tbody>
</table>

**CRUSTS*** | Dried exudate adherent to the skin surface - may contain squames, blood. Often caused by rupture and desiccation of primary lesions containing pus or exudate, or following damage to the skin including self-trauma. |

**EROSIONS EXCORIATIONS** | Superficial lesions confined to the epidermis and which heal without scar formation (consider "hot spots"). Excoriations are self-inflicted and often linear - seen in pruritic dermatoses. |

**ULcers*** | Lesions in which the epidermis is breached, there is damage to dermal tissues and a haemorrhagic exudate. They are associated with dermal or subcutaneous pathology. Scarring occurs. Ulcers may be present in cellulitis. |

**EPIDERMAL COLLARETTES** | Scales arranged in a circular or arciform pattern and representing remnants of a pustule or vesicle or a zone of focal inflammation such as a papule. Strongly suggestive of superficial pyoderma. |

**FOLLICULAR CASTS*** | Squames and sebaceous debris which has been cast around the hairs within a follicle, often matting together several hairs. Consider follicular dermatoses such as demodicosis and dermatophytosis. |

**COMEDONES*** | Plugs of keratin and sebum, which dilate hair follicles, initially pale in colour they darken with time. This material can be expressed by digital pressure. Consider demodicosis and dermatophytosis, or endocrinopathy, particularly Cushing’s syndrome. |

**LICHENIFICATION HYPERPIGMENTATION*** | Chronic thickening and hardening of the skin with exaggeration of the normal markings - usually associated with prolonged and continual friction, and accompanied by excessive accumulation of melanin - hyperpigmentation. Cause may be endocrine (diffuse pattern) but is more often postinflammatory (latticework appearance) - a major sign of chronic inflammation. Alterations of hair colour may also occur commonly as a postinflammatory event. Consider atopic dermatitis or food allergy when there is dyscoloration of the paws due to continual licking. |

**ALOPECIA*** | Abnormal hair loss - may be partial or complete, circumscribed or diffuse, symmetrical or asymmetrical. Note the general pattern first and then follow close examination with a lens describe it clearly on the silhouette. Consider conditions altering hair development such as endocrine disorders (tend to be symmetrical), and those resulting in follicular damage, such as pyoderma, demodicosis and ringworm (asymmetrical and associated with areas or infection). |

*Lesions which may be defined as primary in some dermatoses.
Principal primary lesions
Clinical features

- Erythema
- Papules
- Non Follicular Pustules
- Furuncles
- Tumour (Mycosis fongoïde)

Erythema. Courtesy of: D.N. Carlotti
Papules. Courtesy of: D.N. Carlotti
Non Follicular Pustules. Courtesy of: D.N. Carlotti
Furuncles. Courtesy of: D.N. Carlotti
Tumour (Mycosis fongoïde). Courtesy of: D.N. Carlotti
Principal secondary lesions
Clinical features

Scales. Courtesy of D.N. Carlotti

Erosions. Courtesy of D.N. Carlotti

Follicular casts. Courtesy of D.N. Carlotti

Lichenification / Hyperpigmentation. Courtesy of D.N. Carlotti

Excoriations. Courtesy of D.N. Carlotti

Ulcers. Courtesy of D.N. Carlotti

Comedones. Courtesy of D.N. Carlotti

Alopecia. Courtesy of D.N. Carlotti
**Clinical Record**

Owner:  
Animal's name:  

<table>
<thead>
<tr>
<th>Date</th>
<th>Dermatological examination</th>
<th>Complementary examinations</th>
<th>Differentials</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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</table>

Clinical Handbook on Canine Dermatology
Differential diagnosis for groups of related dermatoses:

**DIAGNOSTIC GROUPS OF RELATED DERMATOSES**
(see lists of the specific dermatoses)

<table>
<thead>
<tr>
<th>GROUP 1: Specific infectious and parasitic dermatoses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Few and relatively uncommon.</td>
</tr>
<tr>
<td>GROUP 1a: Some primary dermatoses</td>
</tr>
<tr>
<td>Few but relatively uncommon.</td>
</tr>
<tr>
<td>GROUP 1b: Secondary dermatoses</td>
</tr>
<tr>
<td>Few but common.</td>
</tr>
<tr>
<td>(occurring secondary to the primary dermatoses, especially of group 2)</td>
</tr>
<tr>
<td><strong>Always as a first step</strong></td>
</tr>
<tr>
<td>• Eliminate these differentials definitively one by one, based on clinical criteria or on clinical pathology and/or therapeutic trials.</td>
</tr>
<tr>
<td>• More than one may be present at a time.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>GROUP 2: Commonest primary dermatoses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Few but very common.</td>
</tr>
<tr>
<td>(very often associated with secondary dermatoses)</td>
</tr>
<tr>
<td><strong>Only as a second step</strong></td>
</tr>
<tr>
<td>• When all the dermatoses of group 1 have been eliminated from the list of differentials.</td>
</tr>
<tr>
<td>• When some signs persist after successful treatment of secondary dermatoses.</td>
</tr>
<tr>
<td>• When secondary dermatoses, successfully treated, reoccur.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>GROUP 3: Rare primary dermatoses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Many different dermatoses but rare.</td>
</tr>
<tr>
<td><strong>At any time if suspected</strong></td>
</tr>
<tr>
<td>• When an uncommon lesional picture is seen.</td>
</tr>
<tr>
<td>• When sudden and/or severe lesion development occurs.</td>
</tr>
<tr>
<td>• Rarer but generally more severe.</td>
</tr>
</tbody>
</table>

**SEQUENCE OF INVESTIGATION**

**PRIORITY IN THE LIST OF DIFFERENTIALS**

**TREATMENT**

**USEFULNESS OF HISTOPATHOLOGY**

- Sometimes diagnostic.
- The clinical approach is essential.
- Rarely diagnostic. May help the clinician to re-orientate his/her approach. The clinical approach is essential.
- Diagnostic in the great majority of cases with good biopsies. Several to many skin biopsies are necessary.

"All the diagnosed dermatoses can be treated at the same time, but all the suspected dermatoses have to be managed successively!"
### Clinical Handbook on Canine Dermatology

#### Otitis Externa

**Adult dog**

<table>
<thead>
<tr>
<th>Type of Otitis Externa</th>
<th>Clinical Method of Investigation</th>
<th>Prevalence of Otitis and Associated Dermatoses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Otitis due to a Foreign Body (FBO)</td>
<td>Refer to the monograph</td>
<td>+</td>
</tr>
<tr>
<td>Parasitic Erythematoceruminous Otitis (PECO)</td>
<td></td>
<td>+</td>
</tr>
<tr>
<td>Non Parasitic Erythematoceruminous Otitis or Infectious Erythematoceruminous Otitis (NPECO/IECO)</td>
<td>This method provides an accurate diagnosis for all types of otitis</td>
<td>+++</td>
</tr>
<tr>
<td>Suppurative Otitis (SO)</td>
<td></td>
<td>+</td>
</tr>
</tbody>
</table>

**Young dog**

<table>
<thead>
<tr>
<th>Type of Otitis Externa</th>
<th>Clinical Method of Investigation</th>
<th>Prevalence of Otitis and Associated Dermatoses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Associated dermatoses</td>
<td>Integration of chronic or recurrent cases into the general dermatological approach</td>
<td>+++ (group 2)</td>
</tr>
</tbody>
</table>
## Sequence of investigation for the differential diagnosis of dermatoses in adult dogs

### ALWAYS AS A FIRST STEP

**Specific infectious and parasitic dermatoses (group 1)**

<table>
<thead>
<tr>
<th>Group 1a</th>
<th>Commonest primary dermatoses</th>
<th>Group 1b</th>
<th>Secondary dermatoses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sarcoptic mange</td>
<td>Folliculitis</td>
<td>Malassezia dermatitis</td>
<td>common and very often secondary to allergies</td>
</tr>
<tr>
<td>Other ectoparasitoses</td>
<td>Malassezia dermatitis</td>
<td>Bacterial overgrowth</td>
<td>less common and very often secondary to endocrine disorders</td>
</tr>
<tr>
<td>Intertrigo</td>
<td>Hot spot</td>
<td>(Pyo) Demodicosis</td>
<td></td>
</tr>
<tr>
<td>Idiopathic folliculitis</td>
<td>Deep pyoderma</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dermatophytosis</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### ONLY AS A SECOND STEP

**If no dermatosis of group 1 is diagnosed or if clinical signs persist after the successful treatment of a secondary dermatosis of group 1b**

**Commonest primary dermatoses (group 2)**

- If pruritus or if it persists after treatment of secondary dermatoses:
  - Investigate an ICS and/or wait for the withdrawal if there is no real urgency: 3/6 weeks if oral administration and 6/12 weeks if injectable administration.
- If no pruritus or if it has disappeared after treatment of secondary dermatoses:
  - Beware of the previous use of corticosteroids!

**Differential diagnosis of allergies**

- Flea Allergy Dermatitis (FAD)
- Adverse Food Reactions (AFR)
- Canine Atopic Dermatitis (CAD)
  - (Contact dermatitis by irritation or hypersensitivity)
  - (Other insect bite hypersensitivity)

**Differential diagnosis of endocrine disorders**

- Lateral Cushing's syndrome
- Spontaneous Cushing's syndrome
- Hypothyroidism
- Remember the euthyroid sick syndrome!
  - (Other endocrine disorders)

### AT ANY TIME IF SUSPECTED

**Rare primary dermatoses (group 3)**

- BULLOUS and NON BULLOUS AUTOIMMUNE DERMATOSES / NEOPLASIA / BULLOUS IMPETIGO
- LEISHMANIASIS (should be considered in group 1a in enzootic areas) / CUTANEOUS DRUG REACTIONS
- SEBACEOUS ADENITIS / STERILE NODULAR PANNICULITIS / HEPATOCUTANEOUS SYNDROME & MANY OTHER DERMATOSES
Sequence of investigation for the differential diagnosis of dermatoses in young dogs

**ALWAYS AS A FIRST STEP**

1. **Specific infectious and parasitic dermatoses (group 1)**
   - Some primary dermatoses (group 1a):
     - Sarcoptic mange
     - Other ectoparasitoses
     - Impetigo
     - Intertrigo
     - Folliculitis
   - Secondary dermatoses (group 1b):
     - Very common in adults, should be rare in young dogs as underlying primary dermatoses (allergies and endocrine disorders)
     - Should be rare too in young dogs (immature immune and endocrine systems)

2. **Commonest primary dermatoses (group 2)**
   - Very common in adults, should be rare in young dogs (immature immune and endocrine systems)

3. **At any time if suspected**
   - Rare primary dermatoses (group 3):
     - Genodermatoses / Juvenile cellulitis / Cutaneous drug reactions
     - Leishmaniasis (should be considered in group 1a in enzootic areas)
     - & many other dermatoses

---

**DIFFERENTIAL DIAGNOSIS OF ALLERGIES**

- Adverse Food Reactions (AFR)
- Flea Allergy Dermatitis (FAD)
  - (Contact dermatitis by irritation or hypersensitivity)
  - (Other hypersensitivity)

**DIFFERENTIAL DIAGNOSIS OF ENDOCRINE DISORDERS**

- Hypothyroidism
  - Remember the euthyroid sick syndrome!
  - (Other endocrine disorders)
Monographs of the major canine dermatoses
Infection of the hair, claw or stratum corneum by filamentous fungi belonging to the dermatophytes and normally involving species of the genera *Microsporum* and *Trichophyton*. Infection almost always affects the hair follicles leading to crusting and hair loss. *Microsporum persicolor* infects the stratum corneum and not the hair shaft. Affected animals usually recover spontaneously after a period of several weeks but chronic infection may occur. The disease is zoonotic. Dermatophytosis in dogs is not very common but clinical signs are highly variable and this disease is one of the major primary infectious dermatoses that have to be investigated as a first step in the diagnostic procedure.

### Aetiology / Pathogenesis

**Aetiology.**
- The dermatophytes are classified by their principal habitats into three groups. Geophilic species persist as saprophytes in soil whereas the zoophilic species live principally on animals (Table 1). Anthropophilic species are those which normally affect man.
- In dogs, species of the genera *Microsporum* and *Trichophyton* cause nearly all cases of dermatophytosis.

**Table 1. Principal Veterinary Dermatophyte Species, their Hosts and Habitat.**

<table>
<thead>
<tr>
<th>GENUS</th>
<th>SPECIES</th>
<th>HOSTS</th>
<th>HABITAT</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Microsporum</em></td>
<td><em>canis</em></td>
<td>cat, dog, horse, rodents, man</td>
<td>zoophilic</td>
</tr>
<tr>
<td></td>
<td><em>equinum</em></td>
<td>horse, cat, dog</td>
<td>zoophilic</td>
</tr>
<tr>
<td></td>
<td><em>gypseum</em></td>
<td>dog, cat, horse, man</td>
<td>geophilic</td>
</tr>
<tr>
<td></td>
<td><em>nanum</em></td>
<td>pig, man</td>
<td>zoophilic</td>
</tr>
<tr>
<td></td>
<td><em>persicolor</em></td>
<td>dog, cat, rodents</td>
<td>zoophilic</td>
</tr>
<tr>
<td><em>Trichophyton</em></td>
<td><em>equinum</em></td>
<td>horse, dog, cat, man</td>
<td>zoophilic</td>
</tr>
<tr>
<td></td>
<td><em>erinacei</em></td>
<td>hedgehog, dog, cat, man</td>
<td>zoophilic</td>
</tr>
<tr>
<td></td>
<td><em>mentagrophytes</em></td>
<td>cat, dog, rodents, man</td>
<td>zoophilic</td>
</tr>
<tr>
<td></td>
<td><em>verrucosum</em></td>
<td>cattle, sheep, man</td>
<td>zoophilic</td>
</tr>
</tbody>
</table>

- **Microsporum:**
  - *M. canis* is responsible for the great majority of canine infections.
  - As a geophilic species, *M. gypseum*, is influenced by soil conditions and the prevalence of infection with this dermatophyte varies in different regions.
  - *M. persicolor* is more difficult to diagnose as it infects the stratum corneum rather than the hair shaft.
  - Infection with *M. equinum* and *M. nanum* is rarely diagnosed.
- **Trichophyton:**
  - About 30% of dermatophyte infections are caused by *T. mentagrophytes.*
  - Infection with the other *Trichophyton* species is uncommon to rare.
**Pathogenesis.**

- **Dermatophyte infections** are acquired by **contact with dermatophyte arthrospores** derived from infected animals, soil or fomites.
- Following **adherence of the arthrospores** to cells of the *stratum corneum*, **germination** occurs with **production of hyphae**, which **invade the stratum corneum** aided by the secretion of keratinases.
- **Penetration of anagen hair shafts** then occurs with invasion extending **through the hair shaft** (endothrix invasion) as far as the new keratin at the base of the hair but not into the mitotically active hair matrix.
- Invasion ceases when hairs enter the telogen growth phase.
- Following **hyphal invasion** of the hair shaft, masses of infective **spherical arthrospores** are **formed** on the **surface of the hairs** (ectothrix production).
- Invasion provokes an **inflammatory response** and, under normal circumstances, this leads to resolution of the disease within 1-3 months. **Chronic infection** occurs when the host is unable to generate a curative immune response.

![Dermatophytosis: ringworm due to *Microsporum canis* (fungal folliculitis). Courtesy of: P. Jasmin](image-url)
Epidemiology

- **Infective forms** of dermatophytes are derived either from soil (*M. gypseum* infection) or from animals.
- Potential **animal sources** are listed in Table 1.
- **The commonest source of *M. canis* is infected cats**, whereas *Trichophyton* spp. are often acquired from wild rodent carriers. Hedgehogs are a source of *T. erinacei*.
- *M. canis* spores in infected premises can remain viable for over 1 year.
- In some countries, **hunting dogs** are predisposed to *M. gypseum* infections.
- **Terriers**, which like to investigate areas inhabited by wild rodents, are predisposed to *Trichophyton* spp. infections.
- *M. persicolor* is most often associated with wood mice and bank voles but canine infection may be acquired from domestic rabbits.
- **The disease is most often diagnosed in young dogs** (animals less than one year old).
- Older animals are less susceptible and, if previously exposed to dermatophytes, may have developed immunity.
- **Cats may develop a carrier status in the absence of clinically apparent lesions and represent an insidious source of contamination for dogs.**

![Cat affected by dermatophytosis. *Microsporum canis.* Courtesy of: D.N. Carlotti](image1)

![Dog affected by dermatophytosis. *Trichophyton mentagrophytes.* Courtesy of: D.N. Carlotti](image2)
Clinical signs

- Signs can be very variable and diagnostically misleading. For example: circular crusting lesions (as in human dermatophytosis) are caused by other common diseases e.g. superficial pyoderma; in contrast diffuse scaling and crusting with pruritus suggestive of allergy may be caused by dermatophytosis.

- Typical dermatophytosis.
  - Focal or multifocal areas of alopecia including broken hairs at the periphery, which extend centrifugally and may heal centrally.
  - Take care to differentiate these from the target lesions of superficial pyoderma.
  - Pruritus may be variable, typically low to absent.

- Other lesions.
  - Folliculitis, furunculosis and crusting, which may be well demarcated and can resemble autoimmune disease.
  - Scaling which may be quite extensive and may or may not have an erythematous margin.
  - Nodular, deep, inflammatory, suppurating lesions, also known as “kerion”.
  - In the healing phase, lesions may be smooth and shiny, particularly with Trichophyton infections.

- Onychomycosis: infection of the claws may occur with or without evidence of claw bed inflammation (paronychia) and onychodystrophy.

- Secondary bacterial infection, particularly with staphylococci, may occur.
**Diagnosis**

Clinical signs may not be diagnostic and thus **history**, indicating contact with infected animals or people, and **diagnostic tests** are of prime importance.

**History.**
- The disease is **more common in young dogs (animals less than 1 year old)**. Some dogs are more exposed (see epidemiology).
- There may be a history of **contact with pets**, particularly **cats**, or with **wild rodents**. **Humans** in contact may also have characteristic circular, alopecic, erythematous, scaling or crusting lesions.
- The disease is **generally progressive spreading from an initial site to other sites** or extending to form a single confluent lesion (typical of *Trichophyton*).

**Clinical elements.**
- **Focal alopecia**, **scaling lesions** with broken hairs at the periphery and **variable pruritus**.
- With *Trichophyton* lesions may be much more inflammatory with confluent progressive areas of **marked crusting and hair loss**.
- In *M. persicolar* infection, the hair shafts are not invaded and **broken hairs** are **not a feature** unless a consequence of self trauma.
- **Folliculitis, furunculosis, paronychia** and **onychodystrophy** may be present.

**Major elements of history and clinical signs.**
- History of **contact with infected animals**.
- **Ringworm lesions in humans, especially children**, in contact (typically *M. canis* infection).
- **Progressive, focal or multifocal alopecia with scaling**.

**Major differentials.**
- Owing to variable clinical signs, the possible differentials include a wide range of diseases (see page 3).
- Consider crusting, scaling and alopecic diseases including cases of **bacterial folliculitis** and **furunculosis**, **hypersensitivities**, **demodicosis**, **autoimmune crusting diseases**, follicular dystrophy/dysplasia and sterile eosinophilic pustulosis (Ofugi’s disease).
Complementary examinations.
- Wood’s lamp:
  - *M. canis* infected hairs will fluoresce with a characteristic apple green colour in about 50% of cases.
  - Other dermatophyte species do not induce this fluorescence.

**WOOD’S LAMP EXAMINATION:**
MAXIMISING EFFICACY

- Lamp should have a built-in lens to enable small broken hairs to be seen.
- Bulbs gradually lose their potency. Check function with known Wood’s Lamp positive hair from time to time.
- Allow the lamp to warm up for a few minutes before use.
- Do the examination in a dark room.
- Allow your eyes to become accustomed to the dark if possible.
- Look for apple-green fluorescence associated with the hairs.
- Collect fluorescent hairs for subsequent microscopic or cultural examination.

- **Direct microscopic examination:**
  - Collect crusted and broken hairs or Wood’s lamp positive material by hair plucking or skin scraping and suspend it in 10% KOH or in cotton blue-lactophenol, which stains fungal elements blue. Breaking up the crusts facilitates examination. Warm the slide gently to speed clearing with KOH. Apply a coverslip.
  - Examine at low power (total magnification c. X20 – X40) and look for hairs which are broken or fragmented, or which appear to have a thin coating of material over the surface.
  - Switch to high power (total magnification c. X400-500) and look for the spherical dermatophyte arthrospores on the surfaces of the hairs. You may also see hyphae within the hairs and corneocytes.

- **Cytology of smears:**
  - Sometimes dermatophyte infection is associated with purulent lesions (furuncles). Examination of stained smears (Diff-Quik, immersion: X1000) may indicate possible dermatophytosis (to see the procedure, refer to the monographs on pyodermas).
  - Microscopic observation shows non-degenerate neutrophils, macrophages and sometimes lymphocytes and plasma cells, usually without any bacteria.
Fungal culture:
- Isolation of the dermatophyte **confirms the diagnosis** and enables the **species** involved to be identified. This may aid **determination of the source** of the infection.
- Note that dermatophytes may sometimes contaminate the coat and thus compatible clinical signs should be present in addition to the presence of the pathogen before dermatophytosis can be confirmed.
- **Prior disinfection of the site to be sampled with 70% alcohol** may be useful in reducing growth of other organisms. Gently spray the sampling area.
- **Collect hairs** and/or crusts as described page 5.
- Send the collected material to a veterinary laboratory or culture the specimens yourself.
- Culture on **dermatophyte test medium (DTM)** is a very convenient method.
  - This medium contains inhibitors which suppress many but not all contaminants and the pH indicator, phenol red.
  - Gently apply the samples on the surface of the medium, in different areas and in particular around the periphery.
  - The DTM bottle should not be completely closed, as the growth of dermatophytes is aerobic.
  - **Incubate** at room temperature or, better at 25-30°C (above a radiator for example).
  - **Cultures** must be examined regularly, and saved for at least a month.
  - **Growth of flat white colonies producing a red colour** in the surrounding medium within 10-14 days is suggestive of a dermatophyte.
  - **Aspergillus** is an example of a contaminant which might grow and cause reddening with 14 days, however, its colonies are blue-green or greysish in colour and unlike those of the dermatophytes.
  - Alternatively, culture can be done on **Sabouraud’s dextrose agar**. However, this method requires a more complete knowledge of the appearance of dermatophyte cultures as there is no indicator to aid identification.
  - **Further tests can then be done to identify the species isolated**, either **within your practice**, if you have sufficient expertise, or the culture can be sent to a laboratory for identification.
The Mackenzie Brush Technique for sampling:
- This is a method which enables efficient screening of the skin and coat for the presence of dermatophytes and their arthrospores, especially when characteristic lesions are absent.
- It is a particularly effective method in searching for carriers, amongst in-contact cats for example. It is also suitable for screening the skin and coat for residual infection whilst monitoring the progress of treatment.
- The coat of the animal is brushed with a sterile brush and this is then pressed into the surface of the culture medium, which is incubated as above.
- Traditionally, a toothbrush has been used as these can be purchased packaged in an uncontaminated state. However, it is impossible to sample a thick coat efficiently with a toothbrush. Plastic grooming brushes with well-separated teeth are more efficient.

Biopsy:
- Biopsy samples of affected skin are an effective method of diagnosis.
- Samples are taken from the edges of the lesions and fixed in the normal way.
- Sometimes the pathologist will need to use special stains (e.g. PAS: Periodic Acid-Schiff) to demonstrate the dermatophyte elements.

Treatment

General remarks.
- It is important to note that many dermatophyte infections will resolve spontaneously within 3 months and thus recovery may not be directly related to the therapy used.
- Other animals within the home may also be infected at a clinical or subclinical level. These animals should be examined and treated if infection is detected.
- Infections with Trichophyton and cases of onychomycosis are often more difficult to treat successfully.

Topical treatment.
- Topical therapy is unlikely to be effective on its own in eliminating infection but it may help to reduce contamination of the environment and spread of infection to owners and other animals.
- A combination of topical and systemic therapy is best in most cases.
- Clipping:
  - The coat should be clipped around focal lesions for a distance of about 5 cm or, when the lesions are generalised or extensive, the whole coat should be clipped.
  - Clipping disrupts the skin surface and facilitates penetration by dermatophytes. Thus it is advisable to clip a few days after starting topical and systemic therapy, when the antifungal agents are established in the skin.
  - Clippers and premises must be disinfected after use and the removed hair should be autoclaved or incinerated.
  - Because the infection of hairs extends below the skin surface, it is wise to clip the coat again after a growth of about 1-2 cm has occurred to remove newly appearing hairs bearing arthrospores.
- Products:
  - Localised lesions can be treated with miconazole or clotrimazole creams and lotions applied daily.
  - More extensive infections can be treated with enilconazole washes at three-day intervals.
**Systemic treatment.**

- **Griseofulvin:**
  - It is teratogenic and so must not be handled by women of child-bearing age and should not be used on pregnant animals. Side effects including gastrointestinal and haematological changes sometimes occur.
  - Doses of 30–50 mg/kg daily should be effective but can be doubled if the lower dose is not effective.
  - The drug should be given with a fatty meal, e.g. add 2.5 – 5.0 ml of vegetable oil or of a veterinary EFA supplement, to aid absorption. EFA supplements also provides linoleic acid, which aids skin barrier function and improves coat condition.
  - Treatment will normally be needed for weeks to months and should be continued for at least two weeks after clinical recovery.

- **Azoles:**
  - Ketoconazole is employed at 10 mg/kg daily and itraconazole is used at 5 to 10 mg/kg daily. Itraconazole is registered for use in cats as a palatable liquid given at 5 mg/kg for 7 days on 3 occasions alternating with 7-day treatment free periods.
  - Side effects with ketoconazole include hepatotoxicity, adrenal and gonadal hormone suppression, anorexia and vomiting; it is contraindicated in pregnancy. Itraconazole is better tolerated.

**Additional treatment.**

- Dermatophytosis should be carefully investigated in all in-contact pets, especially in cats (which can be important asymptomatic carriers), and treated if necessary.

- **Environmental decontamination:**
  - In view of the extensive dispersal of infected hairs and squames from affected animals and the prolonged survival of arthrospores, decontamination should be attempted.
  - Thorough cleaning and the use of disinfectants where possible is advisable. Carpets and soft furnishings should be vacuum cleaned.
  - This procedure should be repeated on a number of occasions as the infection comes under control.

- **Owners’ lesions:**
  - If the owners have lesions, they should be advised to consult a dermatologist.
  - They are a potential source of reinfection for the animals.

**Comments.**

- It is important to involve the owner and to gain his/her co-operation, as dermatophytosis is sometimes difficult to resolve, especially where several animals in a household are involved.
- Concurrent topical therapy is an important aspect of the treatment.
- In older animals and where treatment does not seem effective, check for concurrent disease and possible causes of immunosuppression.
**Prognosis**

- Where only a single animal is involved the **prognosis is good**.
- In **groups of animals, particularly if cats** are involved, in **older animals** and in **onychomycosis**, the **prognosis should be guarded**.

**Follow-up**

- **Therapy must be continued after clinical recovery and negative culture results.**
- In Wood’s lamp positive cases, the progress of recovery can be followed using the lamp but ultimately should always be checked with cultures.
- Animals may be susceptible to **reinfection**.
- The owner should be warned to look for signs of recurrence.
- In difficult cases monitoring, e.g. with the Mackenzie brush technique during the months following recovery may be warranted.

**Conclusions**

- **Dermatophytosis** is a **very pleomorphic** disease and is easily missed when it appears other than the classical form.
- The possibility of a dermatophytosis is too frequently ruled out from the differential diagnosis when it appears in other than the classical form.
- **Fungal cultures should always be done if in doubt**.
- Treatment failures are often due to non-compliance with good treatment regimens.
- **Treatment** often needs to be **continued** for many weeks with **regular monitoring** of the progress of **response to therapy**.
Sarcoptic mange

Introduction / Definition

- Highly contagious parasitic dermatosis caused by the multiplication in the epidermis of an acarine mite of the species Sarcoptes scabiei var. canis (200 to 400 μm).
- Intensely pruritic.
- Quite common.

Aetiology / Pathogenesis

The parasite: Sarcoptes scabiei.

- Mite specific to dogs, although limited infection of cats is occasionally reported.
- Parasitic life cycle (egg-larva-nymph-adult): short, 2 to 3 weeks.
- The fertilised females on the skin surface move rapidly towards the warmer areas of the skin and burrow into the epidermis to lay eggs.
- Life expectancy of adult mites: 4 to 5 weeks.
- In the environment (off the host), the different parasitic stages survive for only short periods (2 to 6 days at 25°C). Nevertheless, at lower temperatures and high humidity, nymphs and females can survive for up to 3 weeks and may lead to reinfection.

Pathogenesis.

- The clinical signs may appear within one week after infection.
- They result from mechanical irritation and hypersensitivity.

Epidemiology

- The condition has remained a consistent problem over the years occurring with variable frequency.
- Major consideration in the differential diagnosis of pruritic dermatoses in dogs.
- Highly contagious infection but marked individual variation in disease expression with the possibility of asymptomatic carriers. Infection generally results from direct contact but sometimes by indirect contact, with the origin of the disease remaining obscure.
- Possible contagion to humans:
  - Appearance of pruritic papules on the trunk, arms and legs.
  - However, as the parasite is not well adapted to human skin, adults rapidly die and cannot reproduce; affected individuals normally recover spontaneously once the animal is successfully treated.
- Most commonly found in puppies and young dogs, especially those coming from pet stores or some breeders. However, it also occurs in adult dogs.
Clinical signs

- Symptoms and lesions may be more subtle and discreet than portrayed in the literature, particularly in well-groomed animals and/or when parasiticides are frequently used.
- There is severe and constant pruritus often leading to a rapid appearance of extensive excoriations.
- Primary lesions: erythematous papules, crusted papules. Typical primary lesions (to be scraped!) are crusted papules which represent the exact points where the fertilised females entered the epidermis. They appear either just prior to, or simultaneously with the development of increasing pruritus.
- Secondary lesions: crusts, excoriations, hyperpigmentation, lichenification.
- Associated dermatological findings: scaling and seborrhoeic problems, alopecia, pyotraumatic dermatitis, otitis externa affecting the margins of the ear flaps.
- Lesion distribution: the favourite habitats of the mites and thus the sites of the lesions are the margins of the ear pinnae and the bony prominences, especially elbows and hocks, and then the ventral portions of the chest and abdomen. When the disease spreads, the entire body may be involved, but the dorsal midline is always spared.

Affected margin of the ear pinnae. Courtesy of: D.N. Carlotti

Crusted papules on an elbow. Courtesy of: D.N. Carlotti
Clinical elements.

- **Pruritus**: the intensity is directly related to the number of parasites only at the onset. Very rapidly (3 to 4 weeks after infestation), hypersensitivity reactions lead to very marked pruritus, often with low numbers of parasites that are difficult to demonstrate even with the aid of multiple skin scrapings.
- Lesions and distribution pattern (see page 2).

Pinnal-pedal reflex.

- Highly suggestive, positive in more than 80% of cases and rare in other pruritic dermatoses.
- This test should be correctly and consistently performed.
**Differential diagnosis.**
- **Atopic dermatitis:** there is the possibility of cross-reactivity between Sarcoptes and *Dermatophagoides farinae* (Df), and positive skin tests to the latter may be found in cases of sarcoptic mange.
- **Food allergy.**
- **Folliculitis, Malassezia dermatitis,** bacterial overgrowth and rarely, atypical dermatophytosis and pemphigus foliaceus.

**Scrapings.**
- **From crusted papules (elbows and hocks).**
- **From the ear margins,** preferably in lesional areas.
- It is vital to note that:
  - This is the only way to obtain absolute confirmation of the diagnosis.
  - It is quite difficult to diagnose sarcoptic mange by skin scrapings.
  - It is important to emphasise this to the owners and inform them that treatment may be considered even if skin scrapings are negative.
  - **Failure to find any mites does not eliminate the diagnosis!**
- Following specific guidelines when looking for *Sarcoptes* greatly increases the chances of success: **The best lesion to scrape is a crusted papule,** which has resulted from penetration of a female into the epidermis. The mite and/or its eggs may be underneath!
  - **Look for these crusted papules first on the ear margins, elbows and hocks,** with a lens if necessary.
  - **“Much more time has to be spent searching for the area to be scraped than actually scraping”** (contrary to the case in demodicosis).
  - Scrape the ear margins, especially if no crusted papule has been found.
  - Scrape with a blunt scalpel blade. Put a few drops of liquid paraffin, mineral oil or lactophenol on a microscope slide, lubricate the scalpel blade and use that to soften the skin (this facilitates scrapings and the adherence of the scraped material to the blade and its transfer to the slide). Then scrape until capillary bleeding results. With the scalpel blade, macerate and spread the sampled material in the liquid that has been placed on the slide, then crush it under the coverslip.

**Microscopic observations:**
- **Scan under the 10X objective the entire sample** present under the coverslip.
- Every field must be examined carefully.
- **Focus on suspicious areas using the 40X objective,** if necessary.
- If the sample is quite thick, do not forget to focus up and down to ensure that the whole sample is checked. **Note the presence of adults** (200 to 400 μm), **of eggs or of faecal pellets.** Demonstration of any of these is diagnostic.

**Serology.**
- In some countries, tests for circulating IgG antibodies to *Sarcoptes* are available. They indicate recent infestation but may not give positive results until 6 weeks after infestation.

**Skin biopsies.**
- At best these are only suggestive, with parasites only rarely observed.
- The disease is characterised by hypersensitivity reactions that result in histopathological changes similar to those encountered in allergic dermatitis.
**Therapeutic trial** (see treatment).
- Must be undertaken rigorously, as if the diagnosis had been confirmed.
- Sometimes ambiguous as other parasitic dermatoses may respond.
- The evaluation of efficacy is based on the reduction in pruritus (be careful: see next paragraph!) and then of lesions. The response is generally rapid. No final conclusion can be made before the minimum recommended duration of treatment is achieved.
- An exacerbation of pruritus at the very beginning of treatment is frequently noted and a good indication of a positive diagnosis (this may result from the death of the parasites).
- The diagnosis is excluded with decreasing reliability with the use of a systemic injectable treatment, then a systemic treatment given orally, and finally topical therapy.
- Be sure to use products that are not out of date, and that have been properly stored.

**Topical treatment.**
- Selamectin (6mg/kg) and moxidectin (2.5mg/kg) are effective when applied as spot-on preparations at monthly intervals. Treatment for 2 to 3 months is advisable.
- Fipronil spray (0.25%) has been shown to be effective in 5-week old puppies when used on 3 occasions at 3-week intervals. It is useful in very young puppies and pregnant or nursing bitches.
- Acaricidal dips can also be used. These are usually given twice a week for a minimum of 6 applications (unless otherwise recommended in the directions for use).
- All parts of the skin must to be treated, with particular attention given to the lesions around the ears and eyes.
- Products: organophosphates (phosmet, 0.09%), lime sulfur (2 to 3%), amitraz (0.05%, 3 times at 15 day intervals). Resistance may occur.
- Clipping and pre-treatment with keratolytic and antiseborrhoeic shampoos are indicated.

**Systemic treatment.**
- Very effective.
- Ivermectin (250 to 400 μg/kg, 2 or 3 times at 10 or 15 day intervals) by subcutaneous injection (may be painful but, can be relieved by the addition of a local anaesthetic in the same syringe).
- Strictly contra-indicated in Collies, Shetlands, Bobtails, sheepdogs, and their crosses.
- As this is not an approved drug for this indication, it must be given with the consent of the owner and carefully supervised by the prescribing veterinarian.
- Milbemycin oxime (2 mg/kg, 3 times at 1 week intervals) orally. Again, this is not an approved drug.

**Additional therapy.**
- Keratolytic, antiseborrhoeic, antipruritic and emollient topical shampoos and lotions.
- Possible systemic corticosteroid treatment for the first week in cases with very intense pruritus (prednisolone, 0.5 to 1 mg/kg/day orally, 2 or 3 days): only when a definitive diagnosis has been made by scrapings.
Important remarks.

• An exacerbation of pruritus at the very beginning of the treatment is frequent and suggests the correct diagnosis and efficacy of the treatment (but it is useful to give the owners notice of this fact!). Be sure to treat the affected animal long enough after complete clinical recovery (one more systemic administration or two more topical applications).
• Treatment of in-contact dogs is strongly indicated.
• Treatment of the environment using an environmental parasiticide may be necessary.
• Change, or thoroughly cleanse the animals bedding.

Prognosis

Excellent response to the correct treatment and a rapid cure.

Follow-up

• When relapses occur, it suggests that either the therapy was too short, or that there has been reinfection from the environment or another infected animal.
• Repeated recurrences will necessitate a complete and thorough epidemiological investigation.

Conclusions

• The possible confusion with allergic dermatitis, together with the difficulty and the time sometimes needed for the differential diagnosis of allergies, make it vital to exclude this condition prior to the work up for the other pruritic dermatoses.
• “When in doubt, treat!”
Dermatoses associated with infestations by other parasites: Fleas, Lice, Cheyletiella, Neotrombicula and Straelensia

Introduction / Definition

- Infestations with ectoparasites other than Sarcoptes, including insects (fleas, lice) and mites (Cheyletiella, Neotrombicula and Straelensia) are described.
- Infestation may be subclinical for all of these except Neotrombicula and Straelensia.
- Clinical disease is associated with variable degrees of pruritus.
- Frequency of occurrence of ectoparasitic infestations by decreasing order: fleas, Sarcoptes, Neotrombicula, Cheyletiella and lice. Infestation with Straelensia is rare but is reported in France and the Iberian Peninsula.

Aetiology / Pathogenesis

- Aetiology.
  - Flea infestation. Normally caused by Ctenocephalides felis but Ctenocephalides canis also infests dogs in certain areas and is more common in southern Europe. Infestation with species from man (Pulex irritans, P. simulans) and wildlife (e.g. Archeopsylla erinacei from hedgehogs and Spilopsyllus cuniculi from rabbits) also occurs. Only the adults live on the dog. Eggs are laid after the female has fed on blood and fall into the environment where they develop through larval and pupal stages to give rise to new adults, which seek a new mammalian host. Most of the life cycle is spent off the host. Typically it lasts 3-4 weeks but can take up to 6 months.
- **Louse infestation (pediculosis).** Dogs may be infested with the sucking louse, *Linognathus setosus*, which ingests blood via its piercing mouthparts. It is found particularly on the long ears of breeds such as the spaniel, basset and Afghan hounds. The biting louse of dogs, *Trichodectes canis*, which feeds on epidermal debris and is more active, generally causes more irritation. It is chiefly found on the head, neck and tail and tends to cause problems particularly in puppies. It can act as an intermediate host for the tapeworm, *Dipylidium caninum*. Lice spend their entire life cycle on their host. Eggs laid by the female are cemented to the hairs at one end and give rise to nymphal stages and then adults. Louse life cycles take 20-40 days.

- **Cheyletiella infestation (cheyletiellosis).** Normally caused by *Cheyletiella yasguri*. Other species may also infest the dog, particularly *C. parasitivorax* and *C. blakei*, typically found in rabbits and cats respectively. Owners may also develop transient infestation. The mites live in “pseudotunnels” within cellular debris at the skin surface and feed on tissue fluid with their piercing mouthparts. Eggs are attached to the hairs with fine filaments and are smaller than those of lice. Nymphs emerge from the eggs and eventually develop into adults. The life cycle entirely spent on the host, takes about 21 days.

- **Neotrombicula infestation (trombiculidosis).** Infestation by the larval stage of the free-living mite, *Neotrombicula autumnalis* which emerges from eggs laid in damp soil by the female in late summer and autumn and climbs upwards, typically on grass stems, and awaits the arrival of a host to which it attaches. The larva feeds on host fluids via piercing mouthparts for 3 days or more and then falls off to continue its life cycle in the environment.

- **Straelensia cynotis infestation (trombiculidosis).** As with *Neotrombicula* infestation is by free living larvae in later summer and autumn in dogs in rural environments. The larvae invade the hair follicles and after a prolonged feeding cycle may emerge and return to the environment.

**Pathogenesis.**

These infestations **commonly affect young dogs** (especially from breeding colonies) but also occur in adult dogs.

- Cutaneous irritation is generally associated with hypersensitivity reactions to the bites of these parasites (see monograph on flea allergy dermatitis). Where there is a state of anergy or low-grade sensitivity, infestation may be subclinical.
- In flea, louse and *Neotrombicula* infestations, crusted papules and excoriations associated with self-trauma may occur. Infestation with *Neotrombicula* normally causes intense irritation. In cheyletiellosis, abundant scaling is usually present. *Straelensia* infestation is associated with development of nodular lesions, often with a purulent dermatitis, folliculitis, furunculosis and crusts; pruritus is uncommon but can be intense.
- Anaemia may be caused by heavy infestations of fleas or lice, particularly in young or small dogs.
Clinical signs

- **Fleas.** In the absence of flea allergy, minor irritation but anaemia and accumulation of flea dirt in the coat in heavy infestations. See also monograph on flea allergy dermatitis.
- **Lice.** These insects are unable to tolerate high temperatures at the skin surface and populations generally fall in the summer. Infestations tend to be greater in winter. Transfer between animals occurs with close contact and infestation is promoted by poor management, crowding and debility. Lice are quite host specific and so transmission does not occur to dogs from other mammalian species. The entire life cycle is spent on the host.
- **Cheyletiella.** Transfer between animals may occur via contact or from the environment. Adults can survive off the host for up to 10 days in a cool environment but the nymphal stages survive no longer than 48 hours. The entire life cycle is spent on the host. Sequestration of the mites in the nasal cavity has been reported and may enable survival and continuing infestation despite use of topical parasiticides.
- **Neotrombicula.** Larvae are picked up from vegetation growing in moist, cool environments in the late summer and autumn and feed for a few days before falling off. The remainder of the life cycle occurs in the environment.
- **Straelensia.** Larvae are acquired in rural environments, especially woodlands, by hunting dogs. Spontaneous remission can occur but may take up to 12 months.

Epidemiology

- **Fleas.** See monograph on flea allergy dermatitis.
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Clinical elements.

- **Fleas.** See monograph on flea allergy dermatitis.
- **Lice.** Crowding and poor management, including malnutrition, are usually important factors. Dogs may have been brought from breeding establishments with endemic infestation or may have visited infested premises.
- **Cheyletiella.** Direct or indirect contact with infested dogs, cats, or rabbits. Papular lesions may be present on the owners, particularly the arms, around the waist and on the thighs. These normally resolve when the infestation in the dog has been cleared.
- **Neotrombicula.** Occurrence of marked, focal pruritus of the head, ears, feet or ventrum occurring in the late summer or autumn in dogs that have visited infested areas of ground. Infestation may extend into the winter in warmer climates. Often there is local knowledge of such areas and owners may have had a similar problem in previous years.
- **Straelensia.** Appearance of alopecic nodules and papules in late summer and autumn in dogs living in rural and wooded environments, particularly hunting dogs, in Southern France and the Iberian Peninsula.

Central elements of history and clinical signs.

- **Fleas.** See monograph on flea allergy dermatitis.
- **Lice.** Inapparent infestation also occurs.
- **Cheyletiella.** Direct or indirect contact with infested dogs, cats, or rabbits. Papular lesions may be present on the owners, particularly the arms, around the waist and on the thighs. These normally resolve when the infestation in the dog has been cleared.
- **Neotrombicula.** Occurrence of marked, focal pruritus of the head, ears, feet or ventrum occurring in the late summer or autumn in dogs that have visited infested areas of ground. Infestation may extend into the winter in warmer climates. Often there is local knowledge of such areas and owners may have had a similar problem in previous years.
- **Straelensia.** Appearance of alopecic nodules and papules in late summer and autumn in dogs living in rural and wooded environments, particularly hunting dogs, in Southern France and the Iberian Peninsula.
### Treatment

#### Topical treatment.
- **Fleas.** See monograph on flea allergy dermatitis.
- **Lice.** Shampoo the dog with a good cleansing shampoo and then apply an insecticide of your choice. Lice are readily killed by insecticides. Selamectin is effective against biting lice.
- **Cheyletiella.** Four weekly treatments with a parasiticide effective against Cheyletiella will usually be curative. Most insecticides and acaricides are effective. Selamectin is reported to be effective as a spot-on. All dogs and cats in contact should be treated. Heavy-coated dogs should be closely clipped to ensure effective penetration of the parasiticidal agent. An antiseborrhoic shampoo can also be useful. Environmental treatment is advisable (see below). Where topical treatment is unsuccessful or cannot be effectively applied (aggressive animals, large numbers affected, owners cannot cope, sequestration of the mites within the nose suspected) systemic therapy may be necessary.
- **Neotrombicula.** One or two applications of an insecticidal shampoo are normally effective. Animals should be kept away from the infested area until the Neotrombicula larval season has ended. Combined permethrin and pyriproxyfen is reported to be effective as a spray or a spot-on.
- **Straelensia.** Topical therapy alone is not effective but fipronil spray every 15 days combined with systemic therapy may be useful. Antimicrobial shampoos may help to control crusting and secondary infection.

#### Differential diagnosis.
- **Fleas.** See monograph on flea allergy. Where there is no allergy, causes of anaemia (including lice).
- **Lice.** Ectoparasitic infestations including fleas, Cheyletiella, Neotrombicula, Otodectes and Sarcoptes, and other causes of debility and anaemia.
- **Cheyletiella.** Ectoparasitic infestations including fleas, Neotrombicula, Otodectes and Sarcoptes, and keratinisation disorders.
- **Neotrombicula.** Ectoparasitic infestations including fleas, Cheyletiella, Otodectes and Sarcoptes.
- **Straelensia.** Superficial and deep pyoderma; demodicosis.

#### Complementary examinations.
- **Coat brushings.** This method is very efficient. The objective is to have a maximum of cutaneous debris (hairs, scales...) fall down onto a piece of paper, by brushing vigorously the back of the animal. Collected debris is first observed by eye (it is then possible to see some scales “moving”) and then dropped onto a slide with lactophenol or liquid paraffin and examined under a coverslip at low magnification. This technique is effective for searching for Cheyletiella present on the skin surface, but also allows the detection of flea faeces (wet paper test), or lice and their eggs attached to the hairs (lice are more mobile and are about 1-2 mm in length).
- **Hair plucks.** Mount in paraffin and examine as above for eggs and lice or mites.
- **Skin scrapings.** Remove hair (scissors), apply paraffin to the skin and scrape in areas with scaling or crusting and adjacent skin. Mount scrapings and examine as above.
- **Tape strips.** Use transparent adhesive tape. Apply to the affected area. Useful for trapping lice and collecting hair to check for the presence of eggs and lice or mites. Also useful for Neotrombicula. After sampling, press the adhesive side of the tape onto a glass slide and examine in the microscope as above.
- **Biopsy.** Straelensia infestation is best diagnosed by histopathology; skin scrapings are usually negative.
**Systemic treatment.**

- **Fleas.** See monograph on flea allergy dermatitis.
- **Lice.** Systemic insecticidal treatment is not required.
- **Cheyletiella.** Where topical therapy is not effective or is not feasible, ivermectin (300 μg/kg subcutaneously) on two or three occasions at two week intervals may be used. See monograph on sarcoptic mange, and especially contra-indications.
- **Neotrombicula.** Systemic insecticidal treatment is not required.
- **Anti-inflammatory therapy.** In all these diseases use of glucocorticoids at anti-inflammatory doses may be useful to control pruritus and reduce inflammation initially. To be avoided if possible.
- **Antibiotic therapy.** Where there is secondary pyoderma, treatment with an antibiotic effective against *Staphylococcus pseudintemedius* should be instituted. See monographs on pyoderma.
- **Straelensia.** Subcutaneous ivermectin (300 μg/kg) and fipronil spray every 15 days, together with oral cephalosporins are reported to be effective in achieving complete cure or preventing complications until remission. Spontaneous remission may occur after up to 12 months or more.

**Additional treatment.**

- **Treatment of the environment.** Environmental cleaning, particularly the use of a powerful vacuum cleaner, followed by the use of a parasiticide which persists in the environment for two weeks or more is advisable in cheyletiellosis and may be useful in infestation with lice where many animals are affected. See monograph on flea allergy for anti-flea measures.
- **Better management.** Parasitic infestation, particularly with lice, is likely to be more severe in animals that are not well groomed, are malnourished or in poor bodily condition. Management may need to be improved to aid recovery and reduce the likelihood of recurrence, especially in louse infestation.

**Comments.**

- Lack of success is likely to occur most often because of failure of owners to properly and fully prescribed treatments. Clear explanation of what is required and maintenance of good communication with owners until the parasites have been eliminated is very important.

**Prognosis**

- Provided that treatments are properly applied, the prognosis is good. In trombiculidiasis, owners should be warned to avoid infested areas in late summer and autumn.
- Flea allergy is much more difficult to treat successfully (see monograph on flea allergy dermatitis).

**Follow-up**

- It is important to re-check the animals to ensure that the parasites have been eliminated. If in doubt, extend the period of treatment.
- Occasionally, lack of response may be due to resistance to the agent used and a new course of treatment with a different product may be necessary.

**Conclusions**

- These parasitic infestations require careful evaluation and treatment. Good communication with the owners and careful application of the treatments are the keys to success.
- **Their frequency should not be underestimated.**
- **Flea infestation** of course is frequent and must be vigorously treated (even in absence of allergic signs).
- **Cheyletiellosis is underestimated and must be investigated.**
- Trombiculidosis due to *Neotrombicula* is more often recognised, because of its seasonal influence, and also because it is more easily detected and often recognised by the owners, but it is rarer than cheyletiellosis.
- Lice infestation is quite rare in many countries but it may be commoner in others.
Demodicosis and pyodemodicosis (juvenile onset and adult onset)

Introduction / Definition

- **Demodicosis** is a relatively common disease that affects predominantly pure-bred dogs.
- It is associated with proliferation of the mites *Demodex canis*, which are normal inhabitants of the hair follicle, and sometimes of the sebaceous glands.
- **Secondary bacterial infection** of the hair follicles often occurs, and rupture of the hair follicle wall may lead to the presence of free mites in the dermis, and a severe pyogenic infection.
- Classically, **two forms** of demodicosis are described, namely **localised** and **generalised**. Cases of the former usually self-cure, whereas **generalised demodicosis** is a severe disease requiring aggressive therapy. However, the distinction frequently made that they are completely different diseases is an oversimplification, as overlapping spectra occur. Also, it should be remembered that every case of generalised demodicosis was once localised.
- **In addition** to localised and generalised demodicosis, **the following forms are recognised**:
  - **Juvenile onset demodicosis** (onset prior to puberty) and adult onset demodicosis:
    - **Juvenile onset** is by far the most common, and although a serious disease, offers a better prognosis than does adult onset.
    - **Adult onset** is usually associated with severe internal disease and is often very difficult to control.
  - **Squamous demodicosis** (with no secondary infection) and **pustular demodicosis** (pyodemodicosis) where there is an associated pyoderma. Treatment of the latter involves appropriate antibiotics in addition to parasiticidal therapy.
  - **Pododemodicosis** is a demodicosis affecting the feet, generally pustular and often very refractory to treatment.

Aetiology / Pathogenesis

- **The parasite.**
  - The condition is associated with the mite *Demodex canis*.
  - These mites reside in the hair follicles and sometimes the sebaceous glands where they live on sebum and cellular debris.
  - Four stages are seen, the diamond shaped egg, the 6-legged larvae, and the 8-legged nymphal form which develops into the adult.
  - Their habitat is restricted to the skin, but may be found in the ear canal. In severe demodicosis all stages may be found in the lymphoid tissues and in many other internal organs. It is assumed that they reach these sites by simple drainage via the lymphatics.
Heredity.
• There is absolutely no doubt that the tendency to develop generalised demodicosis is inherited. Although the precise mode has yet to be elucidated, it is clear that the condition can be transmitted both by the dam and the sire.
• Thus at this point, the recommendation is that neither the sire nor the dam of litters that have resulted in generalised demodicosis should be used again for breeding.
• It is difficult to be as certain in the case of localised demodicosis, although lines have certainly been encountered within breeding kennels that have a higher than expected incidence.

Proliferation of the mites.
• As already indicated, *Demodex canis* is a normal inhabitant of canine skin.
• The puppy acquires the mites within the first few hours of suckling, and only animals reared artificially are free of mites.
• The factors that cause proliferation are probably multiple and complex. However, much interest has focussed on the immunological factors.
  • There appears to be a high incidence of demodicosis in malnourished puppies, and also in association with severe internal parasitism.
  • Adult onset generalised demodicosis is usually associated with a serious internal disease and is not infrequently seen with multicentric lymphoma, hyperadrenocorticism and hypothyroidism. It may also develop secondarily to chronic corticosteroid or immunosuppressive therapy.
  • However, in some cases, no underlying disease can be identified.
• Data from several studies are certainly consistent with demodicosis being associated with a T cell immunodeficiency.
  • But it is not always clear as to whether this defect is the cause or the result of the disease.
  • In particular, it fails to explain what triggers the initial mite proliferation.
  • It has been hypothesised that this results from an antigen-specific T cell defect.
  • In support of this, it has been shown that normal dogs show positive delayed intradermal skin tests to crude *Demodex* antigen, whereas animals with demodicosis react negatively.
  • Clearly, such tests must be repeated with purified antigens before conclusive deductions are made.
• In addition, there is some evidence in favour of hormonal involvement.
• Although many dog with Demodex suffer from a "euthyroid sick syndrome", with low T4 levels, but a normal TSH response, occasional cases are truly hypothyroid, and will not respond to parasiticidal therapy unless supplemented with thyroid hormone.
• Intact bitches with generalised demodicosis that have been successfully treated, will frequently relapse when they come into oestrus.
• In conclusion, although immunological factors are certainly relevant, there are probably a number of different and interrelated factors that can lead to proliferation of the mites.
**Epidemiology**

The epidemiology can be considered in terms of three differing types:

- **Juvenile onset of localised demodicosis.**
  - Peak age of onset is 3-6 months.
  - Although some lines of differing breeds may be predisposed, localised demodicosis is generally thought of as being a relatively common disease, but without any inherited predisposition.

- **Juvenile onset of generalised demodicosis.**
  - Again, onset is usually from 3-6 months and it is rare for this to commence after puberty.
  - Fortunately, this disease is less common.
  - It usually commences as localised demodicosis, of which some 10% of cases may progress to the generalised form.
  - Here, there is a clear inherited predisposition. Breeds commonly quoted as being predisposed include: Old English Sheepdog, German shepherd, Great Dane, Chinese shar pei, Collie, Doberman pinscher, English bulldog, Boxer, Afghan hound, Dalmatian, Dachshund, Beagle...
  - This predisposition can be transmitted both by the dam and the sire.

- **Development of pustular demodicosis (pyodemodicosis).**
  - The following sequence of events occurs:
    - Proliferation of mites within the hair follicle interferes with the normal clearing mechanisms of the follicle and predisposes to infection with *Staphylococcus pseudintermedius*.
    - Subsequent rupture of the hair follicle leads to a bacterial furunculosis and free mites in the dermis initiate a foreign body reaction.
    - Free keratin from the hair shaft represents further foreign body material.
  - In long-standing cases the infection can go deeper and cellulitis develops.
    This is particularly serious when it involves the interdigital area, leading to severe pododemodecosis.
Clinical signs

Demodicosis may have any of the following appearances:
- Areas of predilection are the face, particularly the periocular area and commissures of the lips, and the forelegs.
- Focal, demarcated areas of alopecia accompanied by a fine scale.
- Such areas may be erythematous.
- Areas of follicular plugging are often seen and may be the only sign when the disease involves the hairless areas.
- Focal, or generalised areas of seborrhoea manifested by excessive scale and a waxy surface lipid film.
- Most cases of demodicosis are non-pruritic unless there is a secondary pyoderma. However, rarely, some localised and some more generalised cases may be quite pruritic, markedly erythematous and occasionally appear clinically compatible with atopic dermatitis.
- Some cases, usually those with marked follicular plugging, develop a bacterial folliculitis and show pustules which tend not to progress to the development of marked epidermal collarettes.
- Deep pyoderma, either furunculosis or cellulitis is often a sequel and indeed this may be the only presenting sign.
- Very rarely, ulceration may develop, especially on the face and mucocutaneous areas which may mimic autoimmune diseases.
- Caution: all cases of deep pyoderma, especially if present in an unusual site or in an unusual breed, must be carefully evaluated for demodicosis (deep pyoderma = scrapings!).

Adult onset of generalised demodicosis.
- This is far less common, and is usually associated with a severe internal disease (e.g. lymphoreticular neoplasia, endocrine disorders, or the use of corticosteroids or immunosuppressive drugs).
- Sometimes, demodicosis appears prior to overt signs of internal disease, and so a continuing search must be made for such diseases during the course of treatment.
**Diagnosis**

**History.**
- Juvenile onset localised demodicosis is the most common.
- Although *Demodex canis* is acquired by puppies within the first few hours of suckling it is a normal inhabitant of canine skin and demodicosis is not a contagious disease.
- Adult onset generalised demodicosis is far less common and is usually associated with an identifiable internal disease, which may be of a serious nature. In some cases the onset of demodicosis precedes the overt development of the underlying disease.

**Clinical elements.**
- Major lesions are focal to generalised areas of alopecia and seborrhoea.
- With a predilection of development on the face and forelegs.
- Areas of follicular plugging are often seen.
- Demodicosis is usually non-pruritic except when there is a secondary pyoderma.
- Secondary pyoderma (folliculitis, furunculosis or cellulitis) is not infrequent.

**Skin scrapings.**
- The diagnosis is most often made by demonstration of the mites by skin scraping.
- *Demodex* mites are exceedingly rare in normal skin, and the presence of more than one mite in a scraping, and particularly the presence of young forms and eggs, enable confirmation of the diagnosis.
- The performance of the skin scraping:
  1. Some 3-5 sites are selected for skin scraping.
  2. The hair, if present, is clipped.
  3. The skin is gently squeezed between thumb and forefinger to force the mites more superficially in the hair follicle.
  4. The skin is moistened with liquid paraffin or mineral oil.
  5. Some is also placed on the slide.
  6. The skin is then scraped using a blunted scalpel blade until capillary bleeding is observed.
  7. The material is then transferred to the slide.
  8. The entire slide is scanned using the 10X objective.
  9. Focus on suspicious areas using the 40X objective if necessary.
  10. The proportion of live and dead mites, of adult and young forms and of eggs should be recorded.
- Note that mites are more readily demonstrated in demodicosis than in Sarcoptic mange and failure to demonstrate mites by skin scraping eliminates the diagnosis. The only exception to this is in cases of pododemodecosis, when the mites may be very hard to find.
Other diagnostic aids.

- Examination of hair "pluckings": as the mites are deep in the hair follicle, they are often readily observed around the hair bulb of hairs plucked from the periphery of lesions. However this method is less sensitive than is skin scraping.
- Examination of pustular exudate: material squeezed from areas of deep pyoderma frequently reveal many mites free in the exudate.
- Skin biopsies:
  - Biopsy samples of affected skin are an effective method of diagnosis, but are unnecessary for routine cases.
  - Sometimes the mites may be very deep, especially when associated with a deep pyoderma such as in pododemodicosis and not readily demonstrated by skin scrapings.
  - In some breeds, such as Chinese shar pei, the skin is sometimes difficult to scrape.
  - If the clinical index of suspicion is high such cases should be biopsied if negative skin scrapings result.
  - In demodicosis, contrary to the case in Sarcoptic mange, mites are easily seen during histopathological examination of skin biopsies.

Major differentials.

- The localised form is readily confused clinically with dermatophytosis.
- Lesions of demodicosis also resemble those of leishmaniasis and occasionally, the two diseases may coexist.
- The pustular form resembles any other form of deep pyoderma.
- Other deep, exudative diseases that may be confused with demodicosis are nodular panniculitis and (in areas of the world where they exist) the intermediate and deep mycoses such as sporotrichosis.
Treatment

**Localised demodicosis.**
- This is usually a self-limiting disease that cures spontaneously.
- Treatment with parasiticides is usually not warranted.
- In general it can be stated that:
  - Cases that have very few mites are probably curing themselves and treatment is not warranted.
  - Where many mites are seen and especially young forms and eggs treatment should be given.
- However, always remember that some 10% of cases of localised demodicosis go on to become generalised. Therefore, whether or not treatment is given a careful follow-up is necessary.

**Generalised demodicosis.**
- Aggressive acaricidal therapy is necessary for this is a potentially fatal disease.
  - Amitraz is used at 250 parts per million (ppm) applied every 2 weeks in the USA, but at 500 ppm weekly in most other countries.
  - It should not be used in Chihuahuas, and is used at lower concentrations in debilitated animals.
  - Other considerations in the use of amitraz are:
    - long and medium coated dogs should be clipped and the hair kept short during treatment,
    - any crusts should be removed by warm water soaks,
    - prior shampooing with a sulphur/salicylic acid or benzoyl peroxide shampoo is helpful,
    - skin scrapings should be taken every 2-3 weeks to monitor the response in terms of the live/dead mite ratio, and the number of eggs and juvenile forms,
    - therapy should be continued for at least 2 weeks or 2 dips after mites were last demonstrated.
  - Amitraz is also marketed as a spot-on product in association with metaflumizone with a label claim for demodecosis.
  - Milbemycin oxime in doses of 1-2 mg/kg for 8 weeks is generally effective, and licensed in some countries.
  - A combination of 10% imidacloprid and 2.5% moxidectin has recently been licensed in most countries. It is applied topically in spot form at monthly intervals for up to four months. Recent studies have shown that it is safe and more effective when applied weekly.
- **Antibiotic therapy.**
  - Concomitant antibiotics should be used in cases of pustular demodicosis.
  - Generally, an antibiotic that is appropriate for *Staphylococcus pseudintermedius* is used, irrespective of whether gram-negative organisms are also isolated.
  - In cases with severe secondary infections, culture and sensitivity are mandatory, and if the animal is severely debilitated, an antibiotic that will encompass both Gram-positives and Gram-negatives may be necessary and indeed life-saving.
  - Antibiotic therapy should be continued until the mite population is well-controlled.
- **Further supportive therapy.**
  - Remember that cases of adult-onset demodicosis are likely to be associated with an underlying disease process or with drug administration.
  - Any contributing disease must, therefore, be treated.
  - All intact bitches must be subjected to ovariohysterectomy, or recurrence will be very likely.
**Follow-up**

**General remarks.**
- Follow up involves a full clinical examination and multiple skin scrapings from affected areas.
  - Live mites are readily identified by their movement, even under oil.
  - The proportion of live/dead mites, and the proportion of eggs, young forms and adults are recorded.
- **All cases**, both localised and generalised, should be re-checked initially at intervals of 2 weeks and assessed as above.
  - Cases of localised demodicosis in which it was elected not to give treatment initially, should be immediately subjected to parasiticidal therapy if they show signs of spreading, or if a greater proportion of live mites, eggs and young forms are found.
- **Cases under long term treatment can be checked monthly**, and if the proportion of live/dead mites is not diminishing, then consideration should be given to the use of an alternative therapy.

**The approach to recurrent and/or resistant cases.**
- **Recurrent cases:**
  - Be sure that the treatment has been correctly applied.
  - If using amitraz at 250 ppm, and on every other week, consider using it at 500 ppm and weekly.
  - If there is an inadequate response to amitraz, consider using imidacloprid/moxidectin, or vice versa.
  - Be sure that the entire mite population has, as far as can be ascertained, been eliminated.
  - **Be sure that any underlying disease has been appropriately and successfully treated.**
  - In cases that tend to recur, consider using prophylactic dips with amitraz every week, every other week, or maybe monthly according to the response.
  - Alternatively, consider longer term therapy with imidacloprid/moxidectin.
- **Resistant cases:**
  - Before proceeding to use unlicensed products, extra-label use of amitraz by using it at 1000 ppm applied to half the body only, or spot-treatment of resistant areas such as pododemodicosis on a daily or alternate day basis should be considered.
  - **Other parasiticides** that are unlicensed, but that have been shown to be effective are:
    - **oral ivermectin** at 400-600 μg/kg daily for up to 8 months, as required. **Collies of all types, old English sheepdogs, other herding dogs and their crosses, and any other dogs with the MDR 1 gene mutation should never receive ivermectin**, which is not licensed for this use in any country of the world. Toxicity has been recorded even in dogs lacking this mutation.
    - **milbemycin oxime** has been used **orally** at doses of 0.5-2.0mg/kg. Investigators recommend starting with 0.5mg/kg, and increasing the dose each month if there is not a good response in terms of mite counts. This drug appears to be safe in collies,
    - **moxidectin** has also been used **orally** at 200-400 μg/kg daily for up to 6 months. In contrast to ivermectin, milbemycin oxime and moxidectin appear to be well-tolerated in all breeds.

**Prognosis**

**The prognosis for juvenile onset localised demodicosis is very good.**

**The prognosis for juvenile onset generalised demodicosis is always guarded.**
- **Recurrence** even over one year after apparently successful therapy may occur.
- Essentially, the case of severe, generalised demodicosis can never be considered as cured.
- **In particular, pododemodicosis has an especially guarded prognosis.**
- However, upon recurrence, reinstitution of therapy may still be successful.

**The prognosis for adult onset generalised demodicosis is especially guarded.**
- In that it is often associated with serious underlying disease.
- However, if the underlying disease is successfully controlled, then the demodicosis should also be controlled.

**Conclusions**

Demodicosis is often a frustrating disease to manage and one where the precise aetiological factors are poorly understood.

Owner education is vital in obtaining co-operation in the therapeutic approaches which may be expensive and are often long-term.

In adult dogs, generalised demodicosis is a very serious disease and is usually secondary to an underlying disease process. Both the demodicosis and the underlying disease must be treated.

In young dogs generalised demodicosis is still a very serious disease with a guarded prognosis, but there is usually an absence of underlying disease.

Juvenile localised demodicosis is usually a benign and self-limiting disease, although all such cases should be carefully monitored for any tendency to develop generalised disease.
Pyoderma - General presentation

Introduction / Definition

- Canine pyoderma is a **pyogenic bacterial infection** of the dog’s skin.
- Pyodermas are **very frequent, more or less pruritic** dermatoses of dogs.
- They should always be considered as a possible differential diagnosis at primary presentation of canine dermatoses.
- Correct **diagnosis** is of **major importance**, especially in the case of pruritic skin diseases, as pyoderma can **mimic a great number of dermatoses and above all be secondary to them**.

Key points and classification

- According to the origin and/or immune status of the dog, they can be **superficial or deep, localised or generalised**.
  - A recurrent pyoderma is a pyoderma that responds well to appropriate therapy (topical and systemic antibiotic) but that relapses regularly, which implies repeated treatments.
  - A resistant pyoderma is a pyoderma that does not respond to a number of usually effective therapies, particularly classical antibiotic treatments and thus necessitates aggressive treatment, including with particular antibiotics.
- Pyoderma has different causes and/or characteristics depending upon whether it is found in **young or adult dogs**. Even though the first fundamental division of the general dermatological approach proposed is done according to the age of the animal, for practicality and the purposes of clarity, pyodermas will be differently but closely classified as **“simple approach pyodermas”** and as **“complex approach pyodermas”**.
- **Terminology.**
  - Simple approach pyodermas may be **primary** or **idiopathic**, or **related** to causes that need to be investigated simultaneously in the first instance. There can be evident causes (folded skin in intertrigo) or other dermatoses, mostly ectoparasitic infestations (demodicosis and Sarcoptic mange in particular).
  - Complex approach pyodermas are **secondary** to underlying causes (mostly allergies and endocrine disorders) that need to be explored in the second instance, after treating the pyoderma when it interferes with any further diagnosis.
- **In the young dog.**
  - Only simple approach pyodermas are present; they are **superficial**: intertrigo, folliculitis and **impetigo** (exceptionally a juvenile pyodemodicosis may be deep).
- **In the adult dog.**
  - Simple approach pyodermas are **rarely present**. In most cases, complex approach pyoderma is present secondary to allergic dermatitis or endocrine disorders.
• Simple approach pyoderma are superficial: intertrigo, idiopathic folliculitis (in short-haired breeds or due to so-called bacterial hypersensitivity) and folliculitis related to ectoparasitic infestation (except for demodicosis that is secondary in adult dogs; pyodemodicosis then has a complex approach).

• Complex approach pyoderma are secondary and chiefly very frequent (directly related to the frequency of the underlying cause). They can be:
  - superficial (folliculitis): usually secondary to allergic dermatitis, very frequent.
  - deep (furunculosis and cellulitis including pyodemodicosis): generally secondary to endocrine disorders, much lower frequency.
  - “pseudo-pyoderma” (pyotraumatic dermatitis or “hot spots”) which has a “semi-complex approach” as they are generally secondary to a flea allergy dermatitis - relatively frequent.

• NB: some rare cases of localised deep pyoderma (e.g., pododermatitis) are idiopathic and then of simple approach.

⇒ Simple approach pyoderma therefore represent the majority of pyoderma found in young dogs and in some rare cases, adults. They are chiefly superficial.

⇒ Complex approach pyoderma follow the occurrence of underlying dermatoses and concern only adult dogs. They are more frequently superficial and secondary to allergic dermatitis.

⇒ Table summarising the different classifications:

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<thead>
<tr>
<th>AGE CATEGORY</th>
<th>YOUNG DOG</th>
<th>ADULT DOG</th>
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<tbody>
<tr>
<td>TYPES OF PYODERMAS</td>
<td>SUPERFICIAL PYODERMAS</td>
<td>DEEP PYODERMAS</td>
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<td>SIMPLE APPROACH PYODERMAS</td>
<td>Intertrigo</td>
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<td>Folliculitis</td>
<td>(Pyodemodicosis)</td>
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<td>Impetigo</td>
<td>(Canine acne)</td>
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<tr>
<td>COMPLEX APPROACH PYODERMAS</td>
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Typography indicating the relative importance of these pyoderma: very frequent, frequent, not frequent, rare.

Because of their specific nature, some pyoderma are classified in the third group of the diagnostic approach:
* Juvenile cellulitis is a rare pseudo-pyoderma affecting puppies of only few months old, which must however be thoroughly investigated.
** Impetigo is rarely seen in adults but it is usually severe and secondary to trauma or debilitating underlying diseases (Cushing’s syndrome...).
Canine pyoderma is due to the multiplication of bacteria in the epidermis and its appendages (hair follicles in folliculitis, or sweat glands in hidradenitis) in the case of superficial pyoderma, and with invasion of the dermis in the case of deep pyoderma.

The bacteria.
- In most cases, the sole agent involved is a coagulase-positive staphylococcus specific to the dog – *Staphylococcus pseudintermedius*.
- In other rare cases, other species of pathogenic staphylococci can be found - *Staphylococcus aureus* or *Staphylococcus hyicus* or *Staphylococcus schleiferi* (*var schleiferi* or *coagulans*).
- Other bacteria can be involved, particularly gram-negative bacteria (*Pseudomonas aeruginosa*, *Proteus mirabilis*, *Escherichia coli*...), but their development is usually secondary to the outgrowth of *Staphylococcus pseudintermedius*.
- Therefore, the latter will always have to be the principle target of the treatment, whatever the results of bacterial culture.

Pathogenesis.
In the dog, the *statum corneum* is far thinner and more compact than that of any other species, and there is a paucity of intercellular emulsion. Furthermore, the hair follicle infundibulum of the dog is open, lacking a sebum plug.
- *Staphylococcus pseudintermedius* belongs to the resident flora of the dog's skin and mucosa.
- It becomes pathogenic when the equilibrium of the cutaneous ecosystem is disturbed by modifications in the surface micro-environment. Pathogenic staphylococci can easily colonize an inflammed and excoriated, seborrhoeic skin. In fact, inflammed skin has accelerated epidermal proliferation and desquamation and is more humid and warmer. These alterations of the skin surface micro-environment promote the multiplication of bacteria. Furthermore, self-trauma and excoriations due to pruritus further degrade the epidermal defences and allow inoculation of bacteria into the skin.
- This explains why pyodermas are essentially secondary to underlying dermatoses. They are therefore very frequent in adult dogs, which group is more susceptible to skin diseases and in particular, allergies. Allergic skin disease, notably atopic dermatitis, is thus probably one of the most common causes of canine pyoderma and particularly of recurrent pyoderma.
In addition, it has been demonstrated that dogs affected with primary seborrhoea have markedly higher cutaneous bacterial counts than normal dogs, with a flora composed primarily of coagulase-positive staphylococci. Any seborrhoeic skin disease (including allergic skin disease and endocrinopathies) is therefore a possible cause of canine pyoderma. If the keratoseborrhoeic condition is not controlled, the pyodema can become recurrent.
- A vicious circle can then take place, staphylococci secreting virulent factors aggravating the cutaneous lesions and therefore the bacterial overgrowth. Antibacterial therapy is hence the key component in general therapy.

Pyodermas are not contagious neither to animals nor to humans. Nevertheless, routine hygienic rules must be respected when the animal presents with suppurative lesions.

The aetiology of recurrent pyoderma includes:
- Anatomic defect i.e. excessive folds that cause intertrigo.
- Allergic skin disease, generating skin inflammation and consequently skin infection, mostly superficial (folliculitis) but also deep in severe and poorly treated cases.
- Keratinisation disorders.
- Immunological deficiencies.
- Inappropriate treatments:
  - Poor selection of antibiotics.
  - Too short antibiotic therapy.
  - Too low dosage of antibiotic therapy.
  - Simultaneous use of contra-indicated drugs, particularly glucocorticoids that are never indicated in pyoderma, even in case of severe pruritus, because of their rebound effect; in addition, the pruritus due to pyoderma disappears very quickly after the initiation of specific therapy.

The main cause of resistant pyoderma is infection with resistant strains, including MRSP (and eventually Methicillin Resistant *Staphylococcus aureus* – MRSA – or Methicillin Resistant *Staphylococcus schleiferi* – MRSS) or *Pseudomonas* spp.
Diagnostic methods

Clinical suspicion is fundamental and may justify by itself the initiation of systemic antibacterial therapy, especially where there are pustules.

Cytology.

- An excellent method to support the clinical diagnosis, providing there are good data indicating the presence of pyoderma or other possible causes. It is also reliable and allows rapid performance and interpretation (at the patient’s side) and relatively inexpensive.

- The cellular types observed in pyodermas are the following:
  - Microscopic observation of pus is substantiated by the presence of neutrophils which can be degenerate (also called “impaired”) or non-degenerate. The degenerate neutrophils indicate an active immune status. They do not stain much and appear bulky, with hypersegmented and pyknotic nuclei while the non-degenerate neutrophils appear healthy and well stained (dark purple).
  - Presence of bacteria, generally cocci (and more rarely rods).
  - Bacteria may be extracellular (next to the neutrophils) or intracellular. In the latter case a few bacteria are gathered in vacuoles in a neutrophil appearing degenerate: this a sign of engulfment (phagocytosis) which indicates an immune response by the host and is suggestive of an active disease status, mainly if that phagocytosis is seen inside a lesion (pustule) and not on the surface of the skin; the engulfed bacteria can be considered as pathogens and therefore a diagnosis of a pyoderma can be made.
  - Macrophages and/or lymphoplasmocytes may be present, supporting a more specific immune response which indicates a more chronic and/or severe condition.
  - Red blood cells may be present, indicating capillary lesions and therefore a deeper involvement.
  - Most cellular elements are easily observed, however, signs of phagocytosis, which represent a fundamental diagnostic feature, must be explored. They are easily found - observation should be focused on accumulations of cocci.
  - The quantity of all the elements observed must be assessed (0 to +++) in order to evaluate the intensity of the condition and to help the follow-up. As an example, the decrease of phagocytosis observed during a therapy may be a sign of the efficacy of that therapy.

- Cytologic findings may also allow consideration of other dermatoses, for example:
  - The lack of bacteria in the presence of many non-degenerate neutrophils may suggest a sterile pustulosis. When occurring with acantholytic epidermal cells (large cells with a single, polygonal aspect) and especially with healthy neutrophils, an auto-immune dermatosis may be suspected - most commonly pemphigus foliaceous).
  - The presence of macrophages and/or lymphoplasmocytes combined with the absence of bacteria may indicate a dermatophytosis.
  - Eosinophils are found in pustular reactions to arthropod stings, etc.
• Practical guidelines to sampling and taking direct impression smears:
  • In the cases of fold pyoderma (or even “hot spots”), perform an impression smear by pressing the slide against the lesion, the edge (or a corner) is sufficient for sampling small areas.
  • In the cases of impetigo, folliculitis and furunculosis, look for an intact purulent lesion (pustule or furuncle), open it gently with the corner of the slide or a needle pinching the skin between two fingers in order to express the fluid, and apply the slide on to several areas of the pus.
  • In cases of cellulitis, take samples under the crust, selecting crust revealing pus and apply the slide to several areas.

• Slide preparation and fast-acting dye (Diff-Quik type):
  • Note on the slide where the sample has been placed (write the nature and the location of the sample). This will be the examination side.
  • Let the samples dry for 1 to 2 minutes in open air.
  • Immerse the slides in the three different dyes, following the advised contact times (generally 30 seconds each).
  • Rinse the slides thoroughly with clear water and blot them carefully with a paper towel, examination side upwards, until the water is completely absorbed.
  • A hair-dryer may be used (gently) for final drying.

• Microscopic examination:
  • First locate evenly spread and fully stained areas and place them under the microscope, examination side upwards!
  • Under low (10X objective) magnification look for the areas with most cells particularly the neutrophils that appear as agglutinated “Ω”.
  • Then change to the 40X objective.
  • Without modifying the depth setting, place the slide between two objective lenses to apply a few drops of immersion oil on the chosen area. Turn to the high power objective (100X immersion) adjusting the focus.
  • If the image stays blurred, the slide may have been accidentally turned upside down, and the reading side may be on the other side!
  • Scan the sample over a large area for a few minutes and look for characteristic images, such as cocci phagocytosis, eventually looking in other areas of the sample (adding some immersion oil if necessary).

• The images commonly found for each type of pyoderma are described in the chapter concerning the specific diagnosis.
**Skin biopsies.**

- May reveal characteristic images of pyoderma.
- Skin biopsies are required when support of the clinical diagnosis is needed, particularly if therapy has failed in spite of a strong suspicion (histopathology will then either confirm the diagnosis and support the continuation of an antibacterial therapy, or give proof or suggestion of another disease).

**Culture and Sensitivity testing.**

- **This test is not a diagnostic procedure by itself in dermatology**, but it can confirm the presumption, revealing the presence of one (or several) pathogen(s) inside a lesion.
- It should above all be considered as a **guide to treatment**.
- It normally allows the identification of the bacteria present in the pus, and the testing of their sensitivity:
  - with the conditions that the causal bacteria has been sampled (and not a normal skin resident species), that it survived during transportation (use the appropriate medium), that it is not inhibited by contaminating species and there is agreement with a laboratory that routinely isolates *Staphylococcus pseudintermedius*.
- Choose antibiotics that need to be included in the sensitivity test and be sure that the laboratory employs adapted discs/systems and refers to validated critical concentrations, especially for *Staphylococcus pseudintermedius*.
- These tests must be kept for difficult cases, e.g. therapy failure, cases refractory to treatment (resistant pyoderma), multiple relapses on animal treated several times (recurrent pyoderma), severe deep pyoderma...

Obviously, the identification of a MRSP (Methicillin Resistant *Staphylococcus pseudintermedius*) or *Pseudomonas* spp. is mandatory since these bacteria can be considered as a cause of pyoderma resistant to a well conducted antibiotic treatment. It is likely that the increase of number of resistant strains will lead to a more frequent use of bacterial culturing and sensitivity testing.
General therapy

Classically, pyoderma therapy requires antiseptic topical therapy combined with systemic antibiotics. This association hastens the healing process and allows a faster recovery, compared with systemic treatment alone.

Antibiotics.

- Systemic antibiotics are necessary in most pyoderma cases.
- The initial prescription must be for a minimum of two weeks plus one to two additional weeks after clinical recovery.
- The antibiotic can be renewed for courses of two to three weeks if necessary.
- Appropriate antibiotics for skin infections:
  - must be active on staphylococci and resistant to their enzymatic inactivators (e.g. *Staphylococcus pseudintermedius* secretes penicillinas),
  - must have an appropriate kinetics and good cutaneous penetration,
  - must be active in pus and reactive tissues (particularly in deep pyoderma),
  - must be easily taken orally to facilitate long term treatment by the owner,
  - because of duration of the treatment it must be well tolerated, particularly in young dogs,
  - its clinical efficacy must have been proven in recent trials.
- Useful antibiotics in dermatology are the following:
  - Macrolides and lincosamides, but resistance is increasing and with crossing between these two families.
  - Diaminopyrimidines potentiated sulfonamides. Possibilities of development of polyarthritis (principally in Dobermans) and keratoconjunctivitis sicca in long term treatments.
  - Penicillins resistant to penicillinas: Amoxicillin + clavulanic acid (clavulanic acid is an essential inhibitor of penicillinas, amoxicillin alone is not efficacious in pyoderma treatments) and oxacillin (infrequently used; may induce vomiting and levels of resistance are increasing).
  - Cephalosporins (*cephalexin* and *cefadroxil*). *Cephalexin* is a molecule that has been used for many years in veterinary medicine. Thanks to its excellent clinical efficacy, it remains a first choice antibiotic in veterinary dermatology.
  - Fluoroquinolones, should not be used in young dogs (especially in big breeds) because of their toxic effect on cartilage. As resistances increase quickly, they should not be used as first intention antibiotics, particularly in superficial pyoderma.

The best recent clinical results have been obtained with clindamycin, fluoroquinolones and *cephalexin*. Always choose veterinary specialities for which efficacy has been proven in practice.
**Cortisone therapy.**

- Inadequate and indiscriminate use of corticosteroids can produce **difficulties in diagnosis and treatment**. Except for special cases (pseudo-pyoderms such as juvenile cellulitis), the use of corticosteroids is **strictly contraindicated in pyoderma**, even in the presence of pruritus because of “rebound effect” - a severe relapse.
- As a factor of immunodeficiency it favours the outbreak of pyoderma, in particular deep pyoderma (with a possible development of iatrogenic Cushing's syndrome in the worst cases).
- Furthermore, diminishing the pruritus in a symptomatic manner does not allow evaluating the component due to the pyoderma itself (which will disappear with the healing of the lesions) and the component due to an eventual underlying pruritic dermatosis.
- The pruritus due to a pyoderma disappears rapidly (in a few days) following successful antibacterial treatment.

**Topical therapy.**

- Antiseptic topical products include shampoos, ointments and lotions (used as adjuncts to antibiotics or for symptomatic relief). In case of resistant pyoderma, they become a major part of the treatment.
- Shampoos (especially antiseptic) are very useful, at least at the beginning of the treatment. They allow the mechanical removal of debris, pus, crusts and bacteria (also eliminated by the antiseptic action). They contain various antiseptics: chlorhexidine, benzoyl peroxide, ethyl lactate or piroctone olamine. Products containing sugars (glycoteknology) can decrease the adherence of staphylococci and pseudomonas.
- Ointments, creams and gels containing fucidic acid or benzoyl peroxide are useful for treating localized lesions of pyoderma. Chlorhexidine can be used in lotions and sprays, as an antibacterial agent.
- The use of **topical therapy** (in particular antiseptic shampoos) is of benefit in all pyodermas.

**Therapy duration and follow-up.**

- **Antibiotic therapy in dermatology is protracted!** The recovery time for canine pyoderma is totally different from almost all other bacterial infections. **The owner must be made aware of this peculiarity** of canine dermatology.
- Whatever the case, a **minimum of three weeks** to initiate the treatment must be prescribed.
- The first re-visit should be planned after this time; the owner must of course be aware of the need to come back before if the condition deteriorates (e.g. non-effective treatment, or worse, a cutaneous drug reaction due to the molecule prescribed).
- **At that visit, prescribe a further two weeks of treatment if it has not completely recovered for at least one week.**
- In case of resistance or recurrence, several cultures and sensitivity testings may be done during the follow-up.
Simple approach pyoderma
All young dog pyodermas and rare cases in the adult.

Young dogs: Intertrigo - Folliculitis - Impetigo
Adult dogs: Intertrigo - Idiopathic folliculitis

Introduction / Definition

- The simple approach pyoderma can be primary or idiopathic, or related to underlying causes that need to be investigated in the first instance. These can be self-evident causes (folded skin in intertrigo) or other skin diseases, mostly ectoparasite infestations (particularly demodicosis and sarcoptic mange).

- Therefore these pyodermas do not have identifiable underlying causes or they have causes that can be diagnosed simultaneously during the first step.

- They concern mainly superficial pyodermas: intertrigo (or “fold pyoderma”), folliculitis and impetigo.

- Intertrigo occurs in young as well as in adult dogs and is due to anatomical defects (exaggerated skin folds).

- In the young dog folliculitis and juvenile impetigo are often found; most frequently idiopathic, but they can also be linked to parasitic infestation (especially sarcoptic mange). In some rare cases of juvenile demodicosis, a pyodemodicosis may develop with presence of a furunculosis and/or a cellulitis (rarely a folliculitis), both demodicosis and deep pyoderma must be then treated simultaneously (always perform skin scrapings when deep pyoderma occurs, especially in the very rare cases in young dogs!). See the monograph on demodicosis.

- In the adult dog idiopathic folliculitis (short-coated breed pyoderma, so-called bacterial hypersensitivity and/or superficial spreading pyoderma) or folliculitis linked to parasitic infestation (mostly Sarcoptes scabiei because when Demodex are involved, the approach becomes “complex” as their presence in adults requires the exploration of an underlying cause). Impetigo in adults is a rare and severe condition that can be easily confused with a bullous auto-immune dermatitis and is almost always due to severe underlying causes. It can then be classified in group 3 of the diagnostic approach (information is given in the chapter on complex approach pyodermas).

- Relatively frequent in young dogs, the simple approach pyoderma is seldom found in adults, and is essentially idiopathic folliculitis in short-coated breeds.

Aetiology / Pathogenesis

- Intertrigo.
  - Some animals are anatomically or “physiologically” (obesity, lactation...) predisposed to have an exceptionally folded skin that creates a warm, humid and dark environment favourable to bacterial colonisation and overgrowth. An excess of sebum, tears, saliva or urine in some folds are also aggravating factors.
  - The infection is due to cutaneous micro-trauma caused by continuous friction.

- Folliculitis.
  - Development of micro-abscesses (pustules) centred on hair follicles by bacterial multiplication and invasion.
**Impetigo.**
- Development of micro-abscesses (pustules) in or under the epidermal *stratum corneum* by bacterial multiplication and invasion.

**Epidemiology**

**Intertrigo.**
- Concerns the young dog as well as the adult, without any particular distinction.
- The infection develops on skin surfaces that are in intimate contact with each other. Folds are therefore predilection sites of infection.
- Dogs with very marked folds are more susceptible. The presence of folds may be due to hereditary or acquired anatomical defects.
- According to the site of the infection, there are different types of fold pyodermas: labial, facial, vulval, caudal, ventral (between pendulous mammary glands), obese fold pyodermas, and/or more or less generalised in other parts of the body in very “folded” breeds.

**Folliculitis and impetigo.**
- In young dogs, juvenile folliculitis and impetigo:
  - Impetigo and folliculitis are non-contagious and most often idiopathic. They can be linked to an ectoparasite infestation and both may be present in the same animal.
  - Impetigo is seen almost exclusively in young dogs. It is sometimes associated with a viral infection or intestinal parasitism and is favoured by poor conditions and poor nutrition.
  - These pyodermas evolve generally just before (or during) puberty: hormonal imbalances in an immature endocrine system (or at maturation) may be implicated.
  - Relapses can be frequent until adulthood, but generally stop spontaneously after puberty.
- In adults dogs:
  - Idiopathic folliculitis is non-contagious. Relapses may be frequent.
  - Impetigo is seldom seen in adults, in which it is a secondary pyoderma (see next chapter).

**Clinical signs**

**General remarks.**
- Presence of pruritus, erythema (even though not always seen and sometimes discreet in impetigo), papules and/or pustules must automatically lead to a suspicion of pyoderma (chiefly superficial in the young dog).

**Intertrigo.**
- “Unfold” the suspected folds (it may lead to an interesting discovery during a routine examination!).
- Erythema, exudate and nauseating odour may be discovered in the folds.
- Pruritus of variable intensity may be present.
Folliculitis.

- More or less pruritic, sometimes intensely, with scratching responsible for modification of lesion.
- Primary lesions: marked erythematous reaction (inflammation generally intense), relatively small follicular papules and pustules (therefore centred on hair follicles, that need to be observed with a lens!).
- Secondary lesions: follicular pustules are quite fragile and transient: they result in crusts and epidermal collarettes after rupture. Due to the location of the pustules, significant hair loss (alopecia) may result. It is frequently the major sign, and sometimes the only sign in cases of short-coated breed folliculitis (coat with a “moth-eaten” look): look for pustules through the hair-coat with a hand lens, usually visible on the ventral aspect; this will lead to your diagnosis!
- In cases of folliculitis due to “bacterial hypersensitivity”, the erythema and epidermal collarettes are more pronounced and extended (target lesions), and the pruritus may be very severe.

Impetigo.

- Pruritus is generally not severe (if intense, it usually indicates follicular involvement).
- Primary lesions: erythema (generally less marked than in folliculitis), non-follicular papules and pustules (therefore randomly distributed, irrespective of hair follicles), commonly located on the axilla and the abdomen (easily visible in this area). Follicular pustules present in folliculitis are not as easily recognised as the non-follicular pustules found in impetigo.
- Secondary lesions: impetigo pustules are very fragile and transient; after rupture, a yellow exudate dries into honey-coloured crusts.
**Diagnosis**

**History.**

- **General remarks:**
  - Pruritus and lesions appear at the same time.

- **Intertrigo:**
  - Labial: **bad smell** from the animal’s mouth.
  - Facial: brachycephalic breeds with skin folds between the muzzle and the eyes; frequently associated with corneal ulceration.
  - Vulval and on the limbs: due to obesity.
  - Caudal: in breeds with corkscrew tail, folds at the tail base.
  - Mammary (ventral midline): lactation.

- **Folliculitis:**
  - Young dog.
  - Short-coated breeds.
  - Pruritus varies but can be severe. Folliculitis is a major differential diagnosis in young dogs with pruritus.

- **Impetigo:**
  - Abdominal and axillary lesions in a young dog.

**Clinical elements and distribution pattern of skin lesions (see pages 2 and 3).**

- **Intertrigo:**
  - Folds involvement, moist and greasy aspect with a bad odour.

- **Folliculitis:**
  - In young dogs: evolves preferentially on the ventral aspect of the body (axilla, groin and abdomen), sometimes generalised, lesions are more papular than pustular.
  - In adult dogs (short-coated breed folliculitis): lesions on the head, the back, the chest and the limbs, but can sometimes be extended to the whole body with a “moth-eaten” look:
    - multiple small areas, with varying degrees at alopecia and small groups of hair tufting together and rising above the skin’s surface with a different orientation.
    - careful examination reveals the characteristic lesions, papules and pustules.

- **Impetigo:**
  - Lesions are essentially abdominal and axillary (variable pruritus and erythema, sometimes subtle).

**Major differentials.**

- **Intertrigo:**
  - Especially if labial and vulval: demodicosis, candidiasis, auto-immune skin disease at the onset (mucocutaneous junction affected) and neoplasia.

- **Folliculitis and impetigo:**
  - Sarcoptic mange with presence of papules:
    - non follicular in sarcoptic mange,
    - follicular in folliculitis,
    - also non follicular in impetigo, but generally distinctly associated with pustules.
  - Demodicosis, dermatophytosis and exceptionally sterile pustular dermatoses (pemphigus, sterile eosinophilic pustulosis: Ofugi’s disease).
Cytologic findings characteristic of each type of pyoderma.

- **General remarks:**
  - Direct smear is a valuable aid to the diagnosis of pyoderma.
  - Refer to the chapter “general diagnostic means in pyodermas”.

- **In intertrigo, bacterial colonisation will mostly be observed:**
  - Numerous **healthy neutrophils**.
  - Extracellular cocci (or possibly some rods).
  - Impaired neutrophils, sometimes some are in the phagocytic state.

- **In folliculitis and impetigo, images of bacterial invasion:**
  - Numerous **impaired neutrophils**.
  - Cocci (rarely some rods):
    - more or less abundant in an extracellular position,
    - but mostly in an intracellular position (in impaired neutrophils in phagocytic state), **evidence of phagocytosis** is variable in frequency but should be looked for since it is always present!
  - Eosinophils are sometimes present.

**Skin biopsies.**

- Histopathology is not a diagnostic aid routinely performed in the diagnosis of simple approach pyoderma. However, it may rule out or confirm its presence.
- Very beneficial in **atypical cases**, particularly when pustular lesions are not found in spite of a thorough examination with a hand lens (in those cases, biopsies of simple papular lesions will certainly lead to the diagnosis).
- In all cases, **perform biopsies on several primary lesions**: pustules (keeping them intact!) and papules, but also any other suspect lesion, in particular when the presence of pyoderma is doubtful.

**Therapeutic trial.**

- Refer to “Therapy” and the chapter “General therapy of pyodermas”.
- Empirical systemic antibiotic therapy is required.
- It is perfectly conceivable when pyodermas are superficial, on a first intention basis.
- Prescribe a treatment for courses of 3 weeks (2 courses minimum + 1 additional week after clinical recovery) and monitor the patient:
  - continue the treatment if healing is evident (at least 50% of the lesions and/or the pruritus has disappeared),
  - occasionally, efficacy of the treatment may be questionable, mostly if improvement is transient, or if only discrete healing is present; however treatments rarely fail when proper antibacterial therapy is used (proper choice of the antibiotic, dosage and duration). This may be due to the onset of a bacterial resistance.
- If there is no demonstrated bacterial resistance, cast doubts on the bacterial involvement and challenge the diagnosis if no improvement is seen, or if the case worsens.
Treatment

Refer to the chapter “General therapy of pyodermas”.

Intertrigo.

- **Topical therapy:**
  - The use of antiseptic and/or antiseborrheic shampoos is necessary at least when initiating the treatment in order to physically remove most of the debris, pus, scales, crusts and bacteria. They must be used once daily, at least during the first week, then twice a week until lesions begin to heal, and then routinely to prevent recurrence.
  - Various antibacterial or antiseptic topical products can also be used in addition to the shampoos, and in maintenance and prevention. Exceptionally (intense inflammation), an antibacterial lotion with corticosteroids may be prescribed for a few days.

- **Systemic antibiotic therapy:**
  - Generally not necessary, may be prescribed only in addition to the topical therapy which remains essential.

- **Adjunctive therapy:**
  - Corrective surgery could be performed in some cases.
  - Weight loss is of benefit in obese fold pyoderma.

Folliculitis and impetigo.

- **Topical therapy:**
  - Antiseptic shampoos:
    - very useful as an adjunct to systemic therapy,
    - hastens healing and allows a faster recovery,
    - may be sufficient in young dogs, particularly in cases of impetigo.
  - Antiseptic lotions:
    - appropriate after the use of shampoos in particular for preventing relapses.

- **Systemic therapy:**
  - Systemic antibiotic therapy (usually empirical), by courses of 3 weeks, renewable until resolution, including 1 week after lesion healing:
    - beneficial in the young dog for a faster recovery,
    - mandatory in the adult dog.
  - Corticosteroid therapy: contraindicated.
Important remarks.
• Intertrigo:
  • As long as predisposing factors are present (folds), relapses are frequent.
  • A maintenance prophylactic topical therapy is usually necessary.
• Folliculitis and impetigo:
  • In young dogs:
    • they can clear spontaneously without any treatment (but not in adults!),
    • relapses can be frequent, without any particular reason. However, they respond well to therapy and above all disappear spontaneously at puberty (persuade the owners to wait!),
    • a sole topical therapy is sometimes sufficient and may prevent or limit relapses,
    • pruritus ceases when the lesions disappear! (As compared to folliculitis secondary to allergic dermatitis in adults).
  • In adult dogs:
    • systemic therapy is mandatory,
    • topical therapy is very useful in addition to systemic therapy or as a maintenance therapy to prevent recurrence.

Prognosis
• Intertrigo.
  • Variable in the long term if predisposing factors (folds) are not controlled.
  • However, appropriate topical therapy is normally effective.
• Impetigo and folliculitis.
  • In young dogs:
    • Good with appropriate systemic therapy.
    • Good in the long term, often clears completely at puberty.
  • In adult dogs:
    • The prognosis is relatively good in short-coated breeds if folliculitis is treated properly with systemic antibiotic therapy, but a permanent follow-up is necessary in the long term.
    • NB: impetigo in adults (rare) has a poorer prognosis because the underlying cause is frequently serious.

Follow-up
• During the treatment.
  • Follow-up by courses of 3 weeks if systemic therapy has been prescribed.
  • Renew the antibiotic therapy by courses of 2 weeks until lesion healing, normally 3 to 8 weeks, including a week after clinical recovery.
  • Preferably monitor after 1 week for intertrigo and at the beginning for juvenile folliculitis and impetigo if only a topical therapy is prescribed.
• Recurrence follow-up after recovery.
  • Simple approach pyoderma relapse!
  • Start another systemic therapy if necessary as soon as the lesions reappear, based on culture and sensitivity testing.
  • Limit relapses by prescribing permanent topical therapy - this will be accomplished as often as possible:
    • With antiseptic shampoos, perhaps once to twice a week or eventually every 15 days if sufficient - needs close co-operation of owners.
    • With the use of antiseptic lotions, easy to use, in particular when they are available in pump spray bottles.
**Conclusions**

- **Pyoderma** is a major consideration in the differential diagnosis of skin diseases in young dogs.

- **Particular cases:**
  - A rare deep pyoderma in the young dog is called “canine acne” and clinically appears as furunculosis on the chin. The only feature in common with human acne is its chronic character and resistance to therapy. It seems to be related to keratinisation defects.
  - In adult dogs, "nasal furunculosis" is idiopathic and "pressure points cellulitis" is due to repeated trauma. Those rare cases of deep pyoderma have a simple approach. They require prolonged systemic antibiotic therapy coupled with local treatment.

- **A folliculitis** can be an idiopathic primary folliculitis, as well as a secondary folliculitis. The symptoms resolve in the former after treatment and the follow-up is focused on treating the relapses. When the latter clears, other lesions and/or pruritus generally persist; follow-up includes treating the relapses as well as exploring an underlying cause (e.g. an allergic dermatitis). Refer to the following chapter.

- **Giving a 3 week antibiotic course** followed by thorough and efficient monitoring is a prerequisite above all else when a pyoderma is suspected.

- **“When in doubt, treat!”**
Complex approach pyoderma
Secondary to underlying dermatoses and only in adults.

Superficial pyodermas: Folliculitis - Impetigo
Deep pyodermas: Furunculosis - Cellulitis
Pseudo-pyodermas: Pyotraumatic dermatitis ("Hot Spot")

Introduction / Definition

⇒ Complex approach pyoderma is a skin infection for which an underlying cause needs to be explored.
⇒ It can also be defined as a secondary pyoderma from an aetiology and pathogenesis point of view.
⇒ It can be a superficial pyoderma (folliculitis or very rarely impetigo), a deep pyoderma (furunculosis and cellulitis) or a pseudo-pyoderma (pyotraumatic dermatitis or "hot spot").
⇒ Secondary pyoderma (or complex approach pyoderma) concerns only the adult dog.
⇒ Impetigo is primary in young dogs and secondary in adult dogs, but because of its singularity, its seriousness, its possible confusion with a bullous auto-immune skin disease and its constant relation to a severe underlying disease, it is classified in group 3 of the dermatological approach and as such is not detailed here (only a few basics will be given).
⇒ Secondary pyoderma is very frequent, especially folliculitis.
⇒ The most frequent underlying skin diseases are classified in two major groups:
  • Allergies, which lead essentially to folliculitis or "hot spots", especially atopy in the former and flea bite allergy in the latter.
  • Endocrine disorders, mostly responsible for the occurrence of deep pyoderma.

Aetiology / Pathogenesis

⇒ Introduction.
  • Any physical or functional skin modification may result in secondary bacterial infection, superficial at first and deep when there is an immunodeficiency component.
  • Bacterial multiplication is either confined to the epidermis and its appendages in superficial pyoderma (hair follicles in folliculitis, or even sweat glands in hidradenitis) or progresses over the basement membrane and invades the dermis in deep pyoderma. When infection is present in “hot spots”, it is rather superficial and always of minor importance.
  • The initial pathogenic agent is generally Staphylococcus pseudintermedius. Even though other pathogens may develop taking advantage of the lesions created by the staphylococci (especially Gram negatives - particularly in deep pyoderma), their pathogenic role is not clearly established. For this reason, the outcome of bacteriological cultures (especially isolation of Staphylococcus pseudintermedius) must always be considered in the context of the overall clinical investigation.
**Secondary folliculitis.**
- Development of micro-abscesses (pustules) in the hair follicles by **bacterial** multiplication and **invasion** (in impetigo the micro-abscesses develop in or under the epidermal *stratum corneum*).

- In rare cases, a “bacterial hypersensitivity” may occur and seems responsible for diffuse lesions (“Superficial Spreading Pyoderma”) with very severe inflammation. A similar situation may also be found in idiopathic folliculitis (see previous chapter).

**Deep pyoderma.**
- In furunculosis, the hair follicle ruptures, resulting in intradermal micro-abscesses (pustules) centred on a follicular residue called a furuncle. Most of them follow folliculitis (which is generally transient).

- Cellulitis occurs when furuncles spread outward. Hence, the dermis (or even the hypodermis) is totally invaded by the infectious process that continues to extend.
Evolution of superficial pyoderma into deep pyoderma is possible and may be due to:
- Trauma by pressure, licking, chewing or scratching.
- Follicular lesions as in demodicosis.
- Spontaneous immunodeficiency status (endocrine disorders) or most frequently iatrogenic in origin (corticosteroid therapy).

NB: demodicosis is frequently related to deep pyoderma lesions. Pyodemodicosis is one of the most severe canine skin diseases; its occurrence follows a generalised demodicosis and involves lesions of cellulitis. That is why skin scraping is the first line of approach to these lesions!
- More specific deep pyodermas occur rarely and have a simple approach: “canine acne” (juvenile furunculosis of the chin), “nasal furunculosis” and “pressure points cellulitis”.

“Hot spots”.
- Localised erosive lesion following self-inflicted trauma.
- Occurrence of “hot spots” confirms the presence of a pruritic condition.
- Primary or secondary lesions of vasculitis could be present.
- The warm environment due to a very dense undercoat may be a predisposing factor.
- The role of self trauma is much more important in perpetuating the lesions than the initial infection.
- Hot spots should be differentiated from pyotraumatic folliculitis, a true pyoderma (intact pustules of folliculitis/furunculosis surround the oozing and suppurative area).

Epidemiology

- Secondary pyodermas are non-contagious and generated by underlying diseases.
- Folliculitis and “hot spots” are very frequent, deep pyoderma is much rarer!
- Secondary folliculitis.
  - Accounts for a very large proportion of canine dermatological cases.
  - Occurs preferentially in dogs affected with allergic dermatitis, in particular atopic dermatitis.
- Deep pyoderma.
  - It is much rarer, but also more severe.
  - It occurs generally in dogs with immunodeficiency.
  - Occurrence in the feet (especially if symmetrical) may have an allergic origin (atopy, food allergy).
  - Secondary furunculosis, more or less generalised, is the most common. More localised deep pyodermas (canine acne, nasal furunculosis and pressure points cellulitis) generally have a simple approach. Furunculosis can progress in cellulitis lesions.
  - An idiopathic furunculosis-cellulitis complex is seen in German Shepherds. It must be assessed as idiopathic only after having explored all possible underlying causes (refer to the conclusion).
- “Hot spots”.
  - Pyotraumatic dermatitis is a frequent canine skin disease.
  - It is secondary to pruritic skin diseases, especially allergies and more particularly flea allergy dermatitis.
  - It may occur after a self-inflicted trauma, but only in certain individuals and without any rational explanation.
Clinical signs

**General remarks.**
- Clinical signs are extremely variable according to the type of pyoderma.
- **NB:** Impetigo is seldom seen in adult dogs. The pustular pattern could be due to repeated micro-traumas (e.g. thorny bushes during hunting...). Numerous pustules are found on the most exposed areas (ears, neck, shoulders, axilla, limbs, abdomen...). A severe bullous condition occurs sometimes, particularly in severe immunodeficiency (Cushing’s disease...).

**Secondary folliculitis.**
- Primary lesions: **marked erythematous reaction** (inflammation generally intense), relatively small **follicular papules and pustules** (therefore **centred on hair follicles that need to be observed with a lens**).
- Secondary lesions: follicular pustules are quite fragile and transient, they result in **crusts** and **epidermal collarettes** (frequently generalised) after rupture. Due to the location of the pustules, a significant **hair loss (alopecia)** may result. It is frequently a major sign, especially in short-coated breeds (coat with a “moth-eaten” look). Always **explore thoroughly all the haircoat with a hand lens to locate the pustule** - this will lead to your diagnosis!
- **More or less pruritic**, sometimes **intensely**, with scratching responsible for lesion modification. **NB:** when pruritus occurs, it is absolutely necessary to treat the folliculitis first in order to assess if the underlying dermatosis is pruritic or not.

**Deep pyoderma.**
- **Pruritus** may be present but is usually **not severe**.
- Presence of **pain** is more **common**, as well as systemic signs.
- **Furunculosis lesions:** large prominent pustules, **erythema, cutaneous thickening, pus with blood, alopecia** and **scars**. These lesions can evolve in more extended cellulitis lesions: **crusts, ulcers, fistulas, suppuration and necrosis**. The former lesions can also be observed at first, without a visible furunculosis stage.
- **Cellulitis can be localised or generalised, more severe and frequently associated with hyperthermia, generalised lymphadenopathy and debilitation.**
- An exclusively pedal form called “interdigital pyoderma complex” can be observed. Generally secondary and very uncomfortable for the dog, its therapy and diagnosis are a challenge for the clinician. Both sides of the web are affected, and the foot is oedematous. The pads are usually intact but perionyxis and onyxis may be observed.
“Hot spots”.

- **Pruritus** is initially very intense, causing the dog to lick and bite itself incessantly. This self-inflicted trauma is responsible for most clinical signs. The pruritus is soon followed by pain.
- The lesion is frequently unique and well demarcated, very inflamed, erythematous and exudative. It appears suddenly and extends very quickly (in a few hours). Cutaneous thickening follows and the skin may be covered by a layer of pus due to bacterial colonisation (but not invasion).

**Diagnosis**

**History.**

- **General remarks:**
  - Remember that folliculitis is very frequent.
  - Previous antibacterial therapy does not exclude this suspicion:
    - first, because of the specific requirements of antibacterial therapy in dermatology (especially concerning the choice of an adequate drug and the aspect of a sufficient duration of treatment), the previous treatment(s) may not have been properly conducted (most frequently),
    - second, because even after well conducted therapy (at least three weeks with an established efficient antibiotic, clinical monitoring and continuation for one or two weeks after clinical recovery), relapses are typical and sometimes occur suddenly since these dermatoses are secondary.
• **Folliculitis:**
  - Pruritus (scratching) may be severe. Folliculitis is a major differential in the case of generalised pruritus in a dog.
  - Pruritus and lesions may appear simultaneously. More frequently, in cases of underlying allergic dermatitis, the pruritus appears prior to the lesions. In between:
    - pruritus can first be localised and then generalised when the lesions appear.
    - it can also be manifest in different ways before and after the lesions occur: extremity chewing and/or licking, rubbing the face (e.g. underlying atopy) followed by body scratching when the lesions occur (e.g. folliculitis).
  - Partial corticosteroid response (the pruritus, and sometimes the lesions, regress) followed by a severe relapse (rebound effect).
  - Eventually, only a partial response to well conducted antibacterial therapy but coupled with corticosteroid therapy.
  - Good response to well conducted antibacterial therapy (relapses are normal if the cause is not controlled!).
  - Improvement but no resolution when the antibiotic course is too short (less than three weeks), and failure when the antibiotic used is not adequate (see choice of antibiotics).

• **Deep pyoderma:**
  - The lesions appear gradually and oozing is noted by the owner.
  - Pruritus or pain appears simultaneously or after the lesions.
  - Frequent debilitation and lameness when the pedal area is affected (sometimes exclusively).
  - Previous corticosteroid treatments (particularly with long-acting injectables).

• **“Hot spots”**:  
  - Usually a sole lesion, appearing suddenly with initial severe pruritus followed by marked pain.
  - Constant licking/biting and lesion extension.
  - Possible previous occurrences with spontaneous recovery.

► **Clinical elements and distribution pattern of skin lesions (see pages 4 and 5).**

  - **Folliculitis:**
    - Generally pruritus resulting in scratching of the body.
    - More papular than pustular lesions, found on the ventral area of the body (axillae and abdomen), the head, the back, the thorax and the limbs, but the whole body may be affected.

  - **Deep pyoderma:**
    - A local or generalised lymph node reaction is commonly observed with a debilitating condition.
    - Lesions of furunculosis can extend to the flanks and back and become generalised.
    - Cellulitis lesions tend to occur on the rump and the thighs. Localised cellulitis tends to be secondary to furunculosis whether the generalised form is due to demodicosis or immunodeficiency (e.g. endocrine disorders).
    - An exclusively pedal pattern of furunculosis and cellulitis is sometimes observed (“interdigital pyoderma complex”), and causes lameness. In this area, a very transient folliculitis stage may exist but irritation, digital friction and self-trauma are all factors favouring rapid development into deep pyoderma.
• “Hot spots”:
  • Intense pruritus followed by frequent marked pain.
  • Spreading and localised lesion, usually on the dorsal area (particularly in case of underlying flea allergy dermatitis), or on the thighs.

► **Major differentials.**
  • Folliculitis:
    • Skin diseases that can result in follicular pustules: demodicosis and ringworm (other than bacteria, *Demodex* and dermatophytes are the two other invaders of the hair follicle). Cytological examination may give some clues (refer to corresponding monographs).
    • Other pustular dermatoses (may not be follicular): bullous auto-immune dermatoses (in particular pemphigus foliaceous) and exceptionally subcorneal pustular dermatosis and eosinophilic pustulosis. Cytological examination is a fundamental diagnostic step for these dermatoses.
  • Deep pyoderma:
    • Consider pyodemicosis! Furunculosis and cellulitis = skin scrapings! Especially when the feet are affected.
    • Auto-immune dermatosis, neoplasia and when the pedal area is affected: foreign bodies, leishmaniasis, trombiculidiasis, *Pelodera* dermatitis, hookworm dermatitis, dermatophytosis (sometimes with onyxis) etc.
  • “Hot spots”:
    • Pyotraumatic folliculitis and eventually, dermatophytosis, localised demodicosis, neoplasia and calcinosis cutis.

► **Cytologic findings characteristic of each type of pyoderma.**
  • General remarks:
    • Direct smear test is of paramount importance in the diagnosis of pyoderma.
    • Refer to the chapter describing general diagnostic methods of pyoderma.
  • Folliculitis cytologic findings are images of bacterial invasion:
    • Numerous impaired neutrophils.
    • Coci (rarely rods):
      • more or less abundant in an extracellular position,
      • but mostly in an intracellular position (impaired neutrophils in state of phagocytosis),
      • phagocytic pictures vary in number but should be looked for since they are always present!
    • Eosinophils are sometimes present.
  • In deep pyoderma:
    • It is possible and necessary to verify the presence of an infectious process, considering the following:
      • it is usually easy to sample pus from a furuncle (preferably), under crusts, or from a fistula,
      • the condition is severe and the differential diagnosis is complex,
      • it is not reasonable to perform a therapeutic trial.
    • Images of bacterial invasion, i.e. altered neutrophils and Coci, similar to those found in folliculitis will be observed.
• But micro-organisms and phagocytic pictures are rarer, but they must however be looked for!
• In addition, the presence of macrophages and/or lymphoplasmocytes suggest a more chronic infection (pyogranulomatous reaction).
• Red blood cells are frequently present indicating a deep injury with vascular lesions.
• Epithelial cells (as in superficial pemphigus!) are not unusual.

• “Hot spots” are characterised by images of bacterial colonisation:
  • Microbial agents found are numerous Cocci (sometimes rods) in an extracellular position. Their pathogenic role is doubtful.
  • Some healthy neutrophils.
  • More or less numerous degenerate neutrophils in a state of phagocytosis (which has not an important significance since it is not observed inside a cutaneous lesion).

Skin biopsies.
• Histopathology is not a diagnostic aid routinely performed in the diagnosis of complex approach pyoderma. However, it may rule out or confirm its presence.
• Very beneficial in atypical cases:
  • In folliculitis, when pustular lesions are not found in spite of a thorough examination with a hand lens (in those cases, biopsies of simple papular lesions will certainly lead to the diagnosis).
  • They should be done with no hesitation when deep pyoderma is suspected, particularly in the pedal area (biopsies are sometimes difficult to perform, but they provide consistent diagnostic data).
• In all cases, perform biopsies on several primary lesions: pustules, furuncles (keeping them intact!) and papules, but also any other lesion like ulcers (inside, around and on the margins), in particular when the presence of pyoderma is uncertain.

Therapeutic trial.
• Folliculitis:
  • Refer to “Therapy” and the chapter “General therapy of pyodermas”.
  • Empirical systemic antibiotic therapy is required and may evaluate the importance of the bacterial involvement in the condition.
  • It is perfectly conceivable when pyodermas are superficial.
  • However, resistant pyodermas become common. Cultures and sensitivity testing should be used in suspicion of such cases.
  • Prescribe a treatment for a minimum of 3 weeks and monitor the patient:
    • pursue the treatment if healing is evident (at least 50% of the lesions and/or the pruritus has disappeared),
    • cast doubts on bacterial involvement and challenge the diagnosis if no improvement is seen, or if the case worsens,
    • occasionally, efficacy of the treatment may be questionable, mostly if improvement is transient, or if only discrete healing is present. However, treatments rarely fail when proper antibacterial therapy is used (proper choice of antibiotic, dosage and duration).
  • Make your assessment exclusively on the regression of papules, pustules and epidermal collarettes, since erythema and pruritus may persist if the underlying dermatosis is pruritic.
• Deep pyoderma:
  • A therapeutic trial is not conceivable when deep pyoderma is suspected.
  • Establishing an acute diagnosis and ruling out the other differentials are necessary (history, clinical features, cytology, bacteriology may be sufficient, histopathology may be useful).
• “Hot spots”:
  • Since the diagnosis is relatively simple, it is not really a “trial”.

Clinical Handbook on Canine Dermatology
Treatment

➤ Refer to the chapter “General therapy of pyodermas”

➤ Folliculitis.
   • Topical therapy:
     • Antiseptic shampoos:
       • very useful as an adjunct to systemic therapy,
       • hastens healing and allows a faster recovery.
     • Antiseptic lotions:
       • useful after the use of shampoos in particular for preventing relapses.
   • Systemic therapy:
     • Imperative systemic antibiotic therapy (usually empirical but culture and sensitivity testing should be done in case of suspicion of resistance), by courses of three weeks, renewable until resolution, including one to two additional weeks after clinical recovery.
     • Corticosteroid therapy: contraindicated.

➤ Deep pyoderma.
   • Topical therapy:
     • Antiseptic shampoos:
       • very useful as an adjunct to systemic therapy,
       • very beneficial in removing the majority of pus, debris, crusts along with bacteria,
       • allows active oxygenation of the tissues essential to healing,
       • hastens healing therefore allows a faster recovery and above all provides immediate relief to the patient.
     • Antiseptic lotions:
       • useful between shampoo applications to maintain local antiseptic pressure.
   • Systemic therapy:
     • Imperative prolonged systemic antibiotic therapy most often based on sensitivity testing (refer to the general advice concerning culture and sensitivity at the beginning of the monograph), by courses of three weeks, renewable until resolution, including two to four additional weeks after clinical recovery.
     • Corticosteroid therapy: strictly contraindicated.
   • Adjunctive therapy:
     • Clipping the lesional areas is of extreme benefit and greatly contributes to lesion healing, particularly in the case of cellulitis. It must be done under general anaesthesia (close clipping down to the skin, even if the lesions bleed).
     • It is worth bathing the patient afterwards (with an antiseptic shampoo). Owners should be warned in advance of the procedure - to return a dog with multiple bleeding ulcers or showing much larger lesions than originally seen can be quite traumatic for pet owners.
     • Draining of potential abscesses (always in addition to a local antiseptic therapy and systemic antibiotic therapy) may be performed in interdigital pyoderma.
     • In some cases of recurrent idiopathic cellulitis, immunotherapy with staphylococcal extracts or autogenous vaccines (preferably with a Staphylococcus pseudintermedius strain).
**"Hot spots".**

- **Topical therapy:**
  - Local disinfecting with **antiseptic shampoos** is useful.
  - **Various** antiseptic and **topical corticosteroid products** may be used as a sole treatment.
  - Resolution should be obtained in a few days. Alopecia persists longer!

- **Systemic therapy:**
  - Antibiotic: generally not necessary.
  - Corticosteroid: generally not necessary but can be useful if the inflammation or the pain is intense (prednisolone 0.5-1 mg/kg/day, 3 to 5 days). To be avoided if a differential diagnosis of allergies is undertaken.

**Important remarks.**

- **General remarks:**
  - Prescribe a minimum duration of treatment of three weeks plus one to four additional weeks after clinical recovery and monitor the patient:
    - the therapy must be pursued if healing is obvious
    - efficacy of the treatment must be questioned if improvement is temporary, or if only discrete healing is present.
  - In **true pyoderma**, **topical therapy** is **essential** and **systemic therapy** mandatory.

- **Folliculitis:**
  - Pruritus is usually present but may be due, at least in part, to the underlying dermatosis.
  - **Pruritus ceases** when the lesions disappear **if the underlying dermatosis is non-pruritic and persists if it is pruritic (most frequently)**. The “remaining” pruritus can then appear differently:
    - the pruritus can first be generalised and then localised (face, ears, feet) when the lesions disappear,
    - therefore, it can appear differently before and after the lesions disappear: prevailing body scratching followed by extremity chewing and/or licking, rubbing the face when the lesions disappear (e.g. underlying atopy).

  - Beware of previous use of **corticosteroids**. Even if used long before, they could interfere with this assessment.

- **Deep pyoderma:**
  - If the patient’s condition is causing concern, particularly if pruritus is absent, undertake an immediate investigation into an underlying immunodeficiency.
  - **Beware of the euthyroid sick syndrome.** In this type of disease (or any other serious disease) low levels of thyroxin (basal T4) are frequently observed, which might lead to a suspicion that the patient is suffering from hypothyroidism when it is in fact perfectly euthyroid. Hypothyroidism can be an underlying cause of deep pyoderma, but this is not as common as endocrine assay results might suggest.

- **“Hot spots”:**
  - The treatment should be aimed at preventing self-trauma and to break the vicious circle lesions-pruritus-trauma. An effective treatment leads to rapid lesion healing.
  - It is then mandatory to prevent any hypersensitivity reactions, largely suspected to be the cause of the lesions (especially from vasculitis). Among the **underlying causes** of pyotraumatic dermatitis, flea allergy dermatitis is the commonest, followed more rarely by atopy, food allergy or parasitic infestation (in particular sarcoptic mange).
Conclusions

- Secondary folliculitis is as common as deep pyoderma is rare!

- Pyoderma, in particular folliculitis, is a major consideration in the differential diagnosis of skin diseases in dogs.

- Treatment of the secondary pyoderma is a prerequisite at first, and effective dermatological diagnosis will be established by reassessing the patient after recovery.

- The most frequent case is a folliculitis - recovery will allow a further differential diagnosis of allergies. An appropriate antibiotic therapy of three weeks, including one or two weeks after clinical recovery, followed by a thorough and efficient monitoring with regard to history is a prerequisite above all suspicion of folliculitis.

“When in doubt, treat!”

- Managing a deep pyoderma involving a differential diagnosis of endocrine disorders (or broadly of immunodeficiency) is rare. However, these cases are more severe and depend upon a precise diagnosis.
Importance of the follow-up and owner education in complex approach cases

• A regular follow-up of the case is very important.
• A patient affected with pyoderma must not be “dropped off” its antibiotic treatment without any follow-up. Success is due as much to a good follow-up as to the treatment!
• Explain to the client that this treatment is only a first step: even though it is mandatory and will relieve the discomfort, symptoms due to an unavoidable underlying dermatosis may persist and/or the pyoderma has a high chance of recurring. Exploring the cause will certainly be necessary if we want to control the dermatological problem of the patient in the long term.
• In the case of folliculitis, any decision for further investigation must be taken with the owner’s co-operation, and when the patient’s discomfort is obvious or when very frequent antibacterial treatment is needed.
• In the case of a deep pyoderma, explain that the underlying cause must be explored immediately because of the severity of the condition and unavoidable recurrence. Chiefly, an immunodeficiency must be explored: first as an endocrine origin (Cushing’s syndrome, hypothyroidism…) then consider other more unusual possible causes (auto-immune dermatosis, leishmaniasis, ehrlichiosis…). If no cause is found, this will allow us to declare the pyoderma as “idiopathic”, then prolonged and repeated antibacterial treatments must be prescribed. Immunotherapy must still confirm its efficacy.
• Indicate as well that corticosteroids are contraindicated, first for medical reasons, and then strictly if any further dermatological work up needs to be done.
• Corticosteroids act as “pro-infectious” factors and are responsible for “rebound” effects.
• Eventually, if necessary, and always out of pyoderma periods, a corticosteroid may be prescribed to help control an allergic dermatitis underlying a folliculitis: mostly atopy as flea allergy dermatitis and food allergy are quite easy to diagnose and control by other means (refer to the corresponding monographs). Prescribe then the lowest dosage possible and monitor rigorously and regularly for microbial over-infections such as pyoderma, Malassezia dermatitis and bacterial overgrowths (Staphylococcus pseudintermedius and Malassezia pachydermatis). Stop corticosteroids and initiate another anti-infectious treatment as soon as a cutaneous infection recurs. Always try to discontinue the use of corticosteroids; inform, educate and convince the owner about that.
• Moreover, pruritus due to pyoderma decreases very quickly (in a few days) after the initiation of the antibiotic treatment.
• Keep also in mind that controlling the bacterial problem may lead to a sufficient reduction in general pruritus, thus allowing a delay in the necessity for controlling the eventual underlying pruritic dermatosis!
Bacterial overgrowth (BOG) syndrome
(Staphylococcus pseudintermedius)
Only in adults, idiopathic or secondary, generally to an allergic dermatitis.

Introduction / Definition

➤ Until recently BOG was not believed to be a major dermatological problem, now it is being considered as an entirely separate dermatological entity: the “BOG syndrome”.
➤ A bacterial overgrowth is a surface bacterial proliferation that appears without any lesions of pyoderma. Generally Staphylococcus pseudintermedius is involved.

Aetiology / Pathogenesis

➤ A hyperproliferation of Staphylococcus pseudintermedius all over the body surface of dogs (skin and mucosae) can cause a syndrome exhibiting "superficial lesions" (see "clinical signs") and pruritus, without formation of pyoderma lesions, called bacterial overgrowth (BOG) syndrome.
➤ Clinically, a main feature of BOG syndrome is the absence of papules, pustules, epidermal collarettes and crusts which can be explained by the location of the bacteria at the surface of the skin, not in the hair follicles, and the absence of folliculitis.
➤ Lesions are mainly ventral, probably because moisture is the principal factor affecting cutaneous microbial populations.
➤ Anti-staphylococcal IgE levels of affected dogs are usually low, except maybe in "true idiopathic cases" where a real bacterial allergy could also explain the dermatitis. This suggests that staphylococcal hypersensitivity is usually not the pathogenic process of BOG syndrome. However, some bacterial factors such as a variety of toxins and enzyme secretions may cause an irritation and hypersensitivity reaction leading to erythema and pruritus.
Epidemiology

- There does not seem to be any sex or breed predisposition.
- BOG syndrome may affect young adults and adults.

Clinical signs

- **Pruritus and offensive odour are usually severe** and may be the principal chief complaint.
- Lesions are erythema, lichenification, hyperpigmentation and excoriations associated with alopecia and localised or generalised greasy keratoseborrheic disorder.
- Presence of a bilateral erythematoceruminous *otitis externa* is very frequent.
- Lesions are mainly ventral, particularly on the axillary and inguinal regions. However, this condition may occur in all the body areas and may be confused with an atopic dermatitis.
- It is the case that the BOG syndrome can be secondary to an underlying allergic skin disease, notably atopic dermatitis. The approach of such cases will need two steps.
**Diagnosis**

- **Cytology is an important diagnostic aid:** the microscopic observation of numerous bacteria present on the surface of the skin, combined with erythema and pruritus (generally intense) suggests a true bacterial proliferation that requires a treatment.
- In many cases, Cocci are found to be adherent to corneocytes.
- By the same sampling procedure (surface direct impression smear or better with a "tape strip test"), cytology may also reveal the presence of Malassezia pachydermatis.
- **This syndrome should be systematically included in the differential diagnosis of pruritic dermatitis** and cytology is the complementary examination of choice, it is therefore mandatory in the presence of pruritus in a dog, particularly in case of suspicion of atopic dermatitis. It can quickly reveal or confirm the presence of either a bacterial or yeast overgrowth, or both!
- **The diagnosis can then be pursued** immediately if these pathogens are absent, or after having eliminated them from the skin of the animal if they are present.

**Treatment**

- **Oral antibiotic courses of three weeks** (at least two courses may be necessary), including one or two weeks after clinical recovery, in conjunction with a topical antiseptic therapy (shampoo and/or lotion) is usually efficient to eliminate cutaneous bacterial overgrowth and associated clinical signs.
- **Cephalexin has been proven to also reduce mucosal staphylococci hyperproliferation**, also present in BOG cases, and suspected to be a potential "reservoir" for skin proliferation. This may help in the BOG syndrome control.
- In some cases, treatment may alleviate the pruritus tremendously, and this might explain why some cases of atopic dermatitis seems to be antibiotic responsive.

**Prognosis**

- **Prognosis is usually good:** BOG may however sometimes need a long course of antibiotic treatment and antiseptic shampoo.
- It may also frequently reoccur, until the underlying dermatosis is controlled.
Follow-up

- A regular follow-up is mandatory.
- An underlying allergic skin disease is usually present and a differential diagnosis of allergies may be then undertaken when BOG syndrome is controlled (bacteriological recovery but some remaining clinical signs).
- When BOG and associated clinical signs totally disappear, the investigation of an underlying cause becomes difficult, and this bacterial overgrowth will be labelled as “idiopathic”. This is less frequent and may be secondary to an immunologic abnormality or an underlying defect in skin integrity promoting multiplication of bacteria.

Conclusions

- The BOG syndrome (Staphylococcus pseudintermedius overgrowth) is a superficial cutaneous disorder without any characteristic primary lesions, and may then mimic and worsen an allergic skin disease.
- Secondary BOG syndrome is, at least potentially, frequent!
- It is a major differential in the diagnosis of pruritic dermatoses in dogs.
- The most frequent case is a BOG syndrome where recovery leads to the need for carrying out a differential diagnosis of allergies.
- The underlying dermatosis and relapses need to be controlled.
- Staphylococcus pseudintermedius and Malassezia pachydermatis may have similar and concomitant roles.

Importance of the follow-up and owner education

- A regular follow-up of the case is very important.
- A patient affected with BOG must not be “dropped off” its antibiotic treatment without any follow-up. Success is due as much to a good follow-up as to the treatment!
- Explain to the client that this treatment is only a first step; even though it is mandatory and will relieve the discomfort, symptoms due to an underlying dermatosis may persist and/or the BOG has a high chance of recurring. Exploring the cause will certainly be necessary if we want to control the dermatological problem of the patient in the long term.
- In the case of BOG, any decision for further investigation must be taken with the owner’s cooperation, and when the patient’s discomfort is obvious or when very frequent antibacterial treatments are needed.
- If no cause is found, this will allow us to declare the BOG as “idiopathic”, then prolonged and repeated antibacterial treatments must be prescribed.
- Topical antiseptic therapy is very important to contribute to the prevention of resistances to antibiotics used systemically.
- Pruritus due to BOG may decrease quickly as soon as the BOG starts to be controlled.
- Keep also in mind that controlling the bacterial overgrowth may lead to a sufficient reduction in general pruritus, thus allowing a delay in the necessity for controlling the eventual underlying pruritic dermatosis!
Malassezia dermatitis and yeast overgrowth

Introduction / Definition

- Malassezia dermatitis is a fungal skin infection.
- The pathogenic agent, Malassezia pachydermatis (Pityrosporum canis in the past) is a lipophilic yeast belonging to the cutaneous microflora of the normal dog, along with Staphylococcus pseudintermedius (principal pathogenic agent of canine pyoderma).
- In certain conditions, related to cutaneous and/or immune-mediated factors, Malassezia may proliferate within the stratum corneum.
- The subsequent yeast overgrowth may then acquire a real pathogenic capacity and initiate a true dermatitis.
- Malassezia dermatitis is relatively frequent in dogs, and is usually very pruritic.
- Its diagnosis is of major importance, particularly in the differential diagnosis of pruritic dermatoses, as it can mimic many dermatoses and mostly be secondary to them.
- It must therefore always be suspected in an adult dog.
- Since Malassezia are frequently associated to other dermatoses (which are also often pruritic) and since the subsequent lesional aspect is far from specific, the component of the clinical signs due to the yeast can only be determined after an efficient antifungal treatment.
- Consequently, “Malassezia dermatitis” designation should be only used when the presence of yeast is effectively responsible for the clinical signs, and therefore when the latter disappear after treatment (meaning after a “therapeutic trial”). The remaining clinical signs are then due to an underlying dermatosis, or if there are no symptoms left, the Malassezia dermatitis is idiopathic.
- However, and for this reason amongst others, any yeast overgrowth must be eliminated whether an actual Malassezia dermatitis is present or not.

Aetiology / Pathogenesis

- Malassezia pachydermatis.
  - The Malassezia genus includes unicellular lipophilic yeasts, i.e. their growth is favoured by the presence of lipids.
    - Several species (e.g. Malassezia furfur, M. sympodialis, M. globosa, M. obtusa, M. restricta, M. sloofiae) are lipid-dependent, which means that this development requires the presence of long-chain fatty acids.
    - Only one species, Malassezia pachydermatis, is lipophilic but not lipid-dependent; it contents itself with short-chain fatty acids and readily grows on usual mediums.
    - The identification of species is difficult and is based on molecular biology techniques.
    - M. pachydermatis is characterised by a monopolar budding from a broad base. This type of reproduction leads to its characteristic “peanut-shaped” appearance under the microscope.
    - These yeasts are small (2-7μm) and they are better observed under high power magnification (100X objective).
- Malassezia pachydermatis belongs to the resident microflora of the dog's skin.
  - Approximately 50% of normal dogs carry this yeast in their haircoat and skin.
  - The perianal region seems to be the preferential site for Malassezia portage and appears as a dispersion area, as is the case for staphylococci.
  - The external ear canal does not seem to be a preferential portage area (nevertheless, Malassezia otitis is frequently observed).
  - The most frequent cutaneous locations in the normal dog appear to be the lips and limb extremities.
Pathogenesis.

• Normal canine response to yeast presence on the skin surface:
  • non specific response mechanism (neutrophil phagocytosis),
  • specific cell-mediated response mechanism: antigen presentation by Langerhans’ cells, T cell induction, lymphokine production which stimulates macrophagic phagocytosis and basal keratinocyte multiplication.

• Consequences: yeast elimination, either directly by destruction, or indirectly by the shedding mechanic action.

• Some pathogenic alterations favour the development of the normally saprophytic Malassezia pachydermatis into a pathogenic status able to multiply and proliferate:
  • disruption of the cutaneous ecological equilibrium,
  • modifications of the capacity and mechanism of host defence.

• Predisposing factors enhancing the multiplication of Malassezia pachydermatis: excessive production and/or modification of the nature of the sebum and/or cerumen, excessive humidity, disruption of the epidermal barrier, presence of skin folds.

• These modifications are mainly due to structural alterations generated by primary causes.

• This explains why Malassezia dermatitis is essentially secondary to other (so called) underlying dermatoses and/or associated with other secondary dermatoses.

• It is thus frequent in adult dogs which may be affected by many primary dermatosis, essentially allergies and endocrine disorders.

• Amongst the underlying or associated dermatoses, the following are frequently observed:
  • skin allergies and most particularly atopic dermatitis,
  • endocrine disorders and most particularly hypothyroidism,
  • iatrogenic origin, most particularly corticosteroids treatment and especially when it leads to iatrogenic Cushing’s syndrome,
  • parasitic infestation and in particular demodicosis,
  • pyoderma,
  • keratoseborrhoeic disorders of various origin.

• These conditions are responsible for skin alterations that generate Malassezia proliferation, but the latter still aggravates the cutaneous problems:
  • by producing lipases that modify the surface lipid film,
  • by speeding up the “epidermal turn-over”,
  • by the development of an immediate hypersensitivity to Malassezia pachydermatis which then becomes an allergen as shown for example by: positive skin testing with Malassezia extracts and high levels of IgG and IgE in atopic dogs with or without Malassezia dermatitis in comparison with non atopic affected dogs and healthy dogs.

• All these modifications contribute in diminishing the skin integrity and aggravating the lesional status and are therefore responsible for the appearance or exacerbation of pruritus, which can become intense.

• A true vicious circle can develop; an antifungal therapy then becomes the first key point of the dermatological treatment.
Malassezia dermatitis is not contagious to other animals or humans. It affects adult dogs of all ages. There is no sex predisposition (neutered animals could be predisposed). Certain breeds seem predisposed to develop Malassezia dermatitis: West Highland white terrier, Basset hound, English setter, Shih Tzu, Dachshund, Cocker spaniel, American Cocker, Poodle, German shepherd, Collie, Shetland sheepdog, Jack Russell terrier, Silky terrier, Australian terrier, Springer spaniel and Shar pei. In temperate countries of the Northern Hemisphere, Malassezia dermatitis often begins during the summer, a period of the year also favourable to the expression of allergic dermatitis. Thereafter, it may persist all winter.

Epidemiology

- Pruritus is always present and is usually intense. For this reason, Malassezia dermatitis must be unconditionally treated at first to be able to evaluate if the underlying dermatosis is pruritic and to what degree.
- Primary lesions: generalised or localised erythema, papules and erythematous maculae.
- Secondary lesions: greasy seborrhoea, crusts, diffuse alopecia, hyperpigmentation and lichenification when chronic.
- Lesions are preferentially localised on the ventral aspect of the body (neck, axillae, abdomen, and inguinal area) and the perianal region.
- Other areas also frequently affected are the face (ears, lips, muzzle) and the limbs (medial thighs and extremities).
- An offensive odour of rancid grease is also frequently present.
- Erythematoceruminous otitis externa is frequently associated (refer to the Otitis Monograph).

Clinical signs

- Malassezia dermatitis. Lesional aspect of the chin and neck. Courtesy of: D.N. Carlotti
- Malassezia dermatitis. Axilla lesional aspect. Courtesy of: D.N. Carlotti
- Malassezia dermatitis. Lesional aspect of the groin and abdomen. Courtesy of: D.N. Carlotti
- Malassezia dermatitis. Lesional aspect of a limb. Courtesy of: H. Koch
- Malassezia dermatitis. Lesional aspect of the extremities. Courtesy of: D.N. Carlotti
Diagnosis

**General remarks.**
- The diagnosis is based on **history**, **clinical examination**, **Malassezia** observation on the cutaneous lesions and on the **response** to a specific antifungal **therapy**.
- Yeast identification may be obtained by various means.
- **Clinical examination and cytology are essential** and justify by themselves the initiation of antifungal therapy.

**History.**
- **All breeds may be affected**, but certain breeds seem predisposed (see above).
- **Constant pruritus** (scratching), sometimes **intense**. Malassezia dermatitis is a major differential in case of generalised pruritus in a dog.
- **Pruritus and lesions** generally **appear simultaneously or the lesions are preceded by the pruritus** when the patient is affected by an underlying allergic dermatitis: very frequent case!
- **Partial response to corticosteroids** (pruritus and sometimes lesion) followed by a **severe relapse**.

**Clinical signs and distribution pattern** (see page 3).
- **Generalised pruritus**.
- **Pruritus localisation mimics atopic dermatitis** since the distribution is similar and since both conditions are frequently associated.
- **Erythema and keratoseborrhoeic disorders** on the ventral aspect of the body, the neck, the face, the perianal region and the limbs. The **whole body may also be affected**.

**Cytology.**
- **General remarks:**
  - **Cytology is reliable**, allows **rapid** performance and interpretation (it needs a few minutes at the patient’s side) and **inexpensive**!
  - Various sampling techniques exist:
    - direct impression smear,
    - cellophane tape test (tape strip test),
    - smear from a skin scraping,
    - swab smear.
  - The first two methods must be preferred, the swab smear being reserved for auricular sampling or when cutaneous exudate is important.
• Direct impression smear performance and fast-acting staining (Diff-Quik type):
  • A slide is placed and rubbed against the lesional areas. The edge (or even a corner) of the slide may be sufficient if the area to sample is small (interdigital area for example).
  • Perform several smears in different lesional areas.
  • Note the side where the sample has been placed. This will be the examination side.
  • Let the samples dry for 1 to 2 minutes in open air.
  • Immerse the slides in the three different solutions used for classic haematological fast-acting staining, following the advised contact times (generally 30 seconds each).
  • Rinse the slides thoroughly with clear water and blot them carefully with an absorbent paper, examination side upwards, until the water is completely absorbed.
  • A hair-dryer may be used (gently) for final drying.

• Technique for a cellophane tape test and fast-acting staining (Diff-Quik type):
  • A clear cellophane tape is pressed firmly on the lesions.
  • Sample several times in different lesional areas.
  • Make a loop with the piece of cellophane tape by fixing it at the end of a slide, the adhesive side carrying the material sampled outwards.
  • Immerse the different pieces of tape in the three different solutions used for classic haematological fast-acting staining, following the advised contact times (generally 30 seconds each).
  • Rinse the pieces of adhesive tape thoroughly with clear water. Eliminate excess water with an absorbent paper.
  • A hair-dryer may be used (gently) for final drying.
  • Unroll the adhesive tape and stick it on the slide.
**Microscopic observations:**

- Locate macroscopically well stained areas and centre them under the microscope lens.
- Take care in positioning the slide on the right side: material sampled (for a direct impression smear) or cellophane, upward.
- Look for the areas with most cells under low (10X objective) magnification, particularly epidermal cells debris, where Malassezia tend to cluster.
- Then locate the areas likely to be best interpreted under the 40X objective.
- Without modifying the depth setting, place the slide between two objective lenses and apply a few drops of immersion oil on the chosen area. Turn to the high power objective (100X immersion) adjusting the focus.
- If the image of a direct impression smear stays blurred, the slide may have been accidentally turned upside down, and the reading side may be on the other side!
- Scan the sample over a large area for a few minutes.
- Quote semi-quantitatively (scale from 0 to ++++) the numbers of Malassezia present.

**Interpretation:**

- The minimum number of yeasts allowing a diagnosis is not known; in fact it may be variable from one dog to another and from one site to another in the same.
  - The simple fact of observing yeast (*Malassezia pachydermatis*), whatever their number, must be taken into account for an animal clinically affected with a dermatological disorder:
    - even if the implication of Malassezia in the dermatitis has not been demonstrated, it represents a potential “danger”;
    - and even if it is true that the higher the number of yeast present, the higher the probability that they have a pathogenic role in the dermatitis, a small number may cause important clinical consequences in case of hypersensitivity to the yeast.
- The treatment could simply be adapted to the number of yeasts present, and/or to the gravity of the clinical signs.
- However, the ultimate proof of the implication of *Malassezia* is a positive response to therapy.

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**Microscopic view of a sample on cellophane tape. Malassezia pachydermatis. (Diff-Quik stain, x1000). Courtesy of: D. Pin**

**Microscopic view of a sample on cellophane tape. Malassezia pachydermatis. (Diff-Quik stain, x1000). Courtesy of: D. Pin**

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**Microscopic view of a direct impression smear. Malassezia pachydermatis. (Diff-Quik stain, x1000). Courtesy of: D. N. Carlotti**

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**Microscopic view of a direct impression smear. Malassezia pachydermatis. (Diff-Quik stain, x1000). Courtesy of: D. N. Carlotti**
• Fungal culture and sensitivity:
  - This test may be useful in rare, specific cases. Its realisation has no actual additional benefit compared to cytology, being more complex and results are delayed.
  - Several methods are used for collecting material to culture: collected hairs, swab, Mackenzie Brush Technique...
  - The material may be placed on Sabouraud medium supplemented with antibiotics and olive oil.
  - Sensitivity testing is generally useless, since Malassezia is normally sensitive to classic antifungal and antiseptic agents (see “therapy”).
  - These tests should rather be reserved for difficult cases: unsuccessful therapy, multiple relapses when patients are treated several times...
  - Keep in mind that Malassezia belongs to the normal microflora of the dog’s skin: as for any opportunistic agent, it is the number of colonies that should be taken into account.

• Histopathology:
  - Skin biopsies are usually not necessary in the diagnostic work-up of Malassezia dermatitis and the sensitivity of this test is chiefly inferior to that of cytology.
  - May be performed to confirm the presence of yeast, to appreciate their implication in the dermatosis, but never to exclude their presence or their implication.
    - Yeasts can be seen virtually at the skin surface (in particular in samples stained with Periodic Acid-Schiff: PAS), but their absence does not exclude the possibility of their presence on the animal, even of a high population.
    - The presence of a histopathological pattern evoking a true Malassezia dermatitis may lead to conjecture regarding its implication in the dermatosis.

"Therapeutic trial".
  - Its goal is not to give evidence of yeast overgrowth, but of its implication or not in the clinical signs observed, i.e. the presence or absence of a true Malassezia dermatitis.
  - This "trial" must therefore always be preceded by the observation of Malassezia "overgrowth" on cytology, so that an allowance can be made for the interpretation of "overgrowth".
  - A systemic antifungal therapy will be preferred. Nevertheless, the use of efficient topical therapy may confirm the diagnosis in maintaining remission (see “treatment”).

Differential diagnosis.
  - The differential diagnosis is complex since the clinical signs of this pruritic dermatitis are not specific, and the principal primary lesion, the erythema, may be observed in all dermatitis!
  - Moreover, Malassezia dermatitis is associated and/or aggravated by most dermatoses belonging to the differential diagnosis.
  - However, this dermatitis is a major differential to be ruled out or treated, particularly as a secondary dermatosis, before investigating primary causes (essentially allergies and particularly atopic dermatitis).
  - The differential diagnosis includes (refer to corresponding monographs):
    - Hypersensitivity skin conditions (atopy, FAD, food allergy, or even contact dermatitis) which are also the most frequent primary causes.
    - Parasitic dermatoses (sarcoptic mange, and more rarely, infestations by other parasites).
    - Any other cause of keratoseborrhoeic disorders combined with a dermatitis.
Treatment

General remarks
- The therapy of Malassezia dermatitis is based on the sole use of topical therapy or in combination with a systemic antifungal treatment, depending on the gravity of the clinical signs, the importance of the overgrowth, the desired potency of the treatment and the potential secondary effects.
- The combination ensures a faster improvement and healing and this confirms the diagnosis (see above). Furthermore the maintenance of remission with the sole topical therapy is also helpful in confirming the diagnosis. However, when confronted by tolerance and economic reasons, a sole topical treatment correctly applied may be justified and sufficient.

Antifungal systemic treatment.
- Products:
  - The systemic treatment of Malassezia dermatitis is essentially based on ketoconazole.
  - The dosage for dogs is 5 to 10 mg/kg SID taken at the beginning of meals.
  - This drug is well tolerated, but biochemical parameters must be monitored when treatments are prolonged. Indeed, an increased level of serum ALT precedes the onset of clinical signs of intolerance (anorexia, vomiting) due to liver toxicity.
  - Itraconazole may be used at 5 to 20 mg/kg every 24 to 48 hours.
  - Griseofulvin is not active against Malassezia.
- Evolution and duration:
  - The initial prescription must last a minimum of three weeks.
  - Pruritus diminishes in the first week and the lesions start healing during the second.
  - A treatment of a few weeks is generally necessary and must be continued after clinical recovery (7 to 10 days).

Corticosteroid therapy.
- Corticosteroids are contra-indicated in Malassezia dermatitis, even in the presence of pruritus, for there is a risk of “rebound” effects, i.e. of severe relapse.
- They represent a factor of immunodeficiency favouring the onset of skin infections (with a risk of developing an iatrogenic Cushing’s syndrome).
- Furthermore, eliminating the pruritus symptomatically does not allow evaluation of the component due to Malassezia overgrowth and the component due to an underlying pruritic dermatosis. Non implication of Malassezia in the dermatitis and/or absence of underlying dermatosis and/or presence of a non-pruritic underlying dermatosis may be falsely concluded.
- Inappropriate use of corticosteroids may lead to therapeutic difficulties on one hand and diagnostic misunderstandings on the other.
**Topical therapy.**
- Topical treatment is essential, and may be sufficient by itself. In any case, it is extremely beneficial when combined with a systemic treatment.
- Available topics are mainly shampoos and lotions (which can be used in addition to or after shampooing).
- These topical agents include antiseptic or antifungal agents (chlorhexidine, miconazole, econazole, enilconazole, ketoconazole...).
- The use of antiseptic or antifungal shampoos, 2 or 3 times a week for two weeks and then every week until healing, first enables the mechanical removal of a large quantity of debris and yeast and then directly counteracts the microbial proliferation.
- The use of antiseptic lotions, as often as possible in combination with shampoos reinforces the efficiency of topical therapy and may allow reducing the frequency of shampoos.
- Antifungal lotions are also available for use after an antiseborrheic shampoo for example.
- Regular use of antiseptic shampoos and/or lotions may effectively prevent relapses.

**Course follow-up.**
- Assess the animal for the first time preferably after 3 weeks (efficacy of the treatment may not be properly assessed before 15 days).
- Then assess the animal at least every 3 to 4 weeks.
- Assess by cytology at each consultation the decrease of cutaneous yeast counts.
- The remission of clinical signs may not be obtained if the underlying dermatosis is severe, therefore the efficacy of the treatment will be assessed by yeast counting.
- Interpretation of clinical signs improvement:

If clinical signs disappear or regress after yeast overgrowth clears, it was indeed a Malassezia dermatitis.
- If the clinical signs disappear completely, the Malassezia dermatitis is “idiopathic” (rare).
- If they simply regress, an underlying dermatosis must be investigated (frequent);
  - If pruritus ceases, the underlying dermatosis is not pruritic (rare case).
  - If pruritus persists, the underlying dermatosis is pruritic (very frequent case).
- Beware of previous use of corticosteroids which may come to confuse this analysis.

If the clinical signs persist once the yeast overgrowth has cleared up, there was no Malassezia dermatitis.
- The microbial population must be always controlled in any case.
- Further dermatological work-up needs to be done.

**Prognosis**
- The prognosis is usually good with an appropriate therapy.
- Nevertheless, Malassezia dermatitis secondary to an underlying dermatosis requires a permanent long-term follow-up.
- Control of the underlying dermatosis may sometimes be difficult (atopy) and relapses may be frequent.

**Follow-up**
- Frequent follow-up is necessary and remains the key point to therapeutic success.
- Cytological examination at each consultation also represents a key feature.
- Treatment and recovery of a secondary Malassezia dermatitis are of course necessary as a first intention but they are only a step in the management of the dermatological case.
- Investigation of an associated or underlying dermatosis, follow-up and prevention of relapses after healing are indispensable.
Importance of the follow-up and owner education

• A regular follow-up of the case is very important.
• Explain to the client that this treatment is only a first step, certainly essential and will relieve the discomfort. Yet, specify that symptoms due to a possible underlying dermatosis may persist and/or that the dermatitis has a high chance of recurring and that exploring the cause will certainly be necessary in order to control the dermatological problem in the long term.
• The decision for further investigation must be taken with the owner’s co-operation when the patient’s discomfort becomes obvious or when treatments become too frequent.
• Indicate that corticosteroids are contra-indicated, first for medical reasons, and then strictly if any further dermatological work-up needs to be done.
• Corticosteroids act as “pro-infectious” factors and are responsible for “rebound” effects.
• Eventually, if necessary, and always out of infectious periods, corticosteroids may be prescribed to help control an underlying allergic dermatitis: mostly atopy as flea allergy dermatitis and food allergy are easily controlled by other means (refer to corresponding monographs). Prescribe the “lowest” dosage possible and monitor rigorously and regularly for microbial infection or overgrowth: pyoderma, Malassezia dermatitis, bacterial overgrowth or both (concurrent Staphylococcus pseudeintermedius and Malassezia pachydermatis overgrowth). Stop corticosteroids and renew anti-microbial therapy as soon as a skin infection recurs. Always try to discontinue the use of corticosteroids: inform, educate and convince the owners about that.
• Moreover, pruritus due to Malassezia dermatitis decreases quickly (in a few days) after initiating the treatment.
• Keep also in mind that controlling the fungal problem may lead to a sufficient reduction in general pruritus, thus allowing a delay in the necessity for controlling the eventual underlying pruritic dermatosis!
Flea Allergy Dermatitis (FAD)

Introduction / Definition

- FAD is a pruritic dermatitis arising from a hypersensitivity to the bites of fleas (most often *Ctenocephalides felis*).
- Very frequent, it is one of the commonest causes of pruritus in dogs.
- FAD is one of the three major allergic skin diseases that have to be considered in the differential diagnosis of pruritic dermatoses, the others being canine atopic dermatitis (CAD) and adverse food reactions (AFR): food allergy and food intolerance.

Aetiology / Pathogenesis

- The flea.
  - Fleas are extremely common in temperate and warmer climates, with varying levels of infestation of pets and their environment, closely linked to ambient temperature and humidity.
  - A clear understanding of the life cycle and environmental requirements is essential in order to develop flea control strategies and obtain owner compliance, especially when overt signs of infestation are lacking.
Clinical Handbook on Canine Dermatology

Part 2

Flea Allergy Dermatitis (FAD) • 2

➤ The development of hypersensitivity.
- Factors that determine whether or not a dog becomes allergic to fleas include:
  - Exposure to fleas early in life tends to reduce the likelihood that a dog will become allergic to fleas.
  - *Intermittent exposure* to the bites of fleas *favours* the development of hypersensitivity, whereas continual exposure tends to be protective, and leads to immunological tolerance.
  - *Dogs* that are atopic, and suffer from atopic dermatitis (CAD) are predisposed to the development of flea allergy.

➤ The pathogenesis in a hypersensitive dog.
- Both immediate and delayed hypersensitivity are involved, and can be visualised following intradermal skin testing when a reaction at both 15 to 20mins and 48hrs is seen.
- The sequence of events that occurs following the bite of a flea in a hypersensitive individual are as follows:
  - The *flea saliva* contains a number of protein antigens to which the patient will have been sensitised by prior exposure.
  - *Hypersensitive dogs* can develop their disease via a number of immunopathogenic pathways, which are:
    - the majority of sensitive dogs develop *IgE antibodies* (immediate, or type I hypersensitivity), and thus mast cell-derived mediators are involved,
    - in addition, some dogs manifest cutaneous *basophil hypersensitivity* with an influx of basophils armed with IgE antibody into the site of the application of allergen,
    - most dogs also have *cell-mediated, delayed, or type IV hypersensitivity*. Injection of the allergen causes an influx of lymphocytes and macrophages and a variety of interleukins are involved. In a certain number of cases (probably 15 to 30%), delayed hypersensitivity alone exists.
The incidence and epidemiology of this condition is inextricably linked to the environmental requirements of the cause which is the flea. Types of fleas involved are:

- *Ctenocephalides felis* is the major species involved world-wide.
- In some areas of some countries, *Ctenocephalides canis* is implicated. The latter is generally less well-adapted, and less hardy.
- Infestations with flea species more usually associated with human infestations, namely *Pulex irritans* and *Pulex simulans* are occasionally found.
- Sometimes, fleas whose natural hosts are poultry and wildlife are implicated. Such infestations are usually readily controlled.

The environmental conditions most suitable for the reproduction and survival of the flea are:

- Moderate temperatures of 18-30°C
- High relative humidity of 70-80%
- They do not survive in colder climates, or in hot climates with low humidity.

Flea infestation without allergy is actually much more common than is FAD.

FAD is non contagious but flea infestation is!

The major features of the biology of *Ctenocephalides felis* and the pre-adult components of the life cycle are as follows:

- Adult fleas are host-dependant blood-sucking ectoparasites.
- They may attack humans in heavily infested environments where more suitable hosts are scarce or absent.
- They can leave their host and survive in a standard house-hold environment but will die quickly unless a new host is found. They should then be considered permanent rather than transient parasites.
- Under optimum conditions, adults can live for long periods without feeding and 1 to 2 weeks under household conditions, but a blood meal is necessary for reproduction, which takes place on the host.
- Feeding fleas can survive up to 100 days on their host.
- When well fed, a female can lay up to 50 eggs/day and up to 2000 during her entire lifetime.
- Eggs are laid on the host and fall into the environment for development. Females lay preferentially at night and therefore the animal's sleeping areas will be most heavily infested and require special attention in the control programme.
- Eggs hatch to larvae (3 larval stages).
- Larvae are very active, negatively phototactic and positively geotactic. After hatching they migrate downwards and away from light. They will then be found preferentially for example in cracks in the flooring, between floor planks and deep in the pile of carpets. This knowledge is of major importance when treating the environment. It is not necessary, for example, to treat all the tiled floor of a very bright kitchen, but one must be careful to treat cracks and crevasses and underneath and behind furniture.
- Very little food is necessary for the development of the larvae which live on blood-containing faecal pellets that fall from the host.
- The third larval stage pupates into an oval sticky cocoon.
This stage is the most resistant to both insect growth regulators (IGRs) and to adulticides. Thus, environmental control alone may not be sufficient, and FAD affected pets must be protected from the newly emerged adult fleas by treatment with adulticides, at least at the beginning, and when any new environmental contamination occurs.

- Each cocoon contains a pre-emerged adult which can lie dormant for long periods, up to 3 or 4 months.
- In the presence of appropriate stimuli (vibration, increased carbon dioxide and heat), adult fleas emerge from cocoons. These stimuli can, of course, be provided by the presence of an animal. This explains for example why a house where an infested animal has lived, even if uninhabited for a long time, may remain an important source of infestation for a newly arrived animal.
- The life-cycle is typically completed in around 4-5 weeks. It may be accelerated to 3 weeks or less, or slowed to 6 months, depending upon environmental conditions.

Approximate percentages of the different parasitic stages present in the household.

<table>
<thead>
<tr>
<th>Stage</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adults</td>
<td>5%</td>
</tr>
<tr>
<td>Pupae</td>
<td>15%</td>
</tr>
<tr>
<td>Larvae</td>
<td>30%</td>
</tr>
<tr>
<td>Eggs</td>
<td>50%</td>
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</table>

Lifetages of Ctenocephalides felis felis. Courtesy of: B. Blagburn, Auburn University, AL, USA
**Ctenocephalides felis** life cycle in the environment and on the hosts

Adult fleas may survive 1 to 3 months on the host. They may represent a very low percentage of all the parasitic stages which occupy the house!

**Sources of infestation**

**Reservoirs of fleas (host-environment life cycle)**

Scanning electron micrograph of *Ctenocephalides felis* adult. Courtesy of: Royal Veterinary College, London

Scanning electron micrograph of *Ctenocephalides felis* egg. Courtesy of: Royal Veterinary College, London

Scanning electron micrograph of *Ctenocephalides felis* larva. Courtesy of: Royal Veterinary College, London

Scanning electron micrograph of *Ctenocephalides felis* pupa. Courtesy of: Royal Veterinary College, London
Clinical signs

- FAD is still sometimes called "summer eczema!"
- The primary eruption is an erythematous papule, which may develop a crust before disappearing in 2-4 days.
- The distribution involves mainly the lower back and posterior and inner thighs. These are also the sites most favoured by the fleas.
- Secondary lesions resulting from self-trauma due to the intense pruritus.
- Changes seen in chronic cases include seborrhoea, crusting, alopecia and eventually lichenification.

Diagnosis

- History.
  - Age of onset: young adult dogs (1 to 6 years) in 75% of cases.
  - The history should document exposure to fleas.
  - However, in exquisitely sensitive animals, only one or two fleas may be present, and may escape notice.
  - The owner also may notice the presence of flea faeces, small comma-shaped dark material which leaves a red stain behind when rubbed on moistened white paper.
  - Demonstration of fleas or flea excreta supports the diagnosis but absence of those elements do not exclude it: in one study up to 2/3 of confirmed FAD affected animals showed no evidence of fleas, and 1/3 showed no evidence of fleas or of flea faeces!
  - However, a careful examination for both fleas and flea faeces must always be undertaken.
  - Fleas and flea faeces are more readily demonstrable with the use of a flea comb. Also, the use during consultation of a quick knock-down flea spray (e.g. permethrin) will kill many fleas, which then will fall onto the examination table. A brushing may help render them directly visible to both the owner and the veterinarian. This may convince those who state that their dog never has any fleas!
  - If faecal pellets only are found, demonstrate to the owners that the findings are not merely dirt by rubbing the flea faeces on moistened white paper. They may then be convinced at least that fleas have recently been on the pet.
• The area of pruritus is also an important consideration.
• Previous parasiticidal treatments may not exclude the diagnosis depending on the products used and the rigour of application.
• There is usually a good response to systemic glucocorticoids, followed by a relapse unless the fleas are controlled at the same time.

• Seasonal influence:
  • More likely to appear in favourable seasons (e.g. late spring, summer and early autumn in temperate climates).
  • May however be present during the whole year in favourable climates, with the constituent parts of the life cycle surviving indoors in the winter.
  • Clinical signs may also persist year-round, even in the absence of fleas, if the FAD is associated with a concomitant disease (e.g. atopic dermatitis due to house dust mites).

► Clinical elements.
Key clinical features that are supportive of the diagnosis are:

• Characteristic distribution.
• Evidence of a papular and crusting primary eruption, accompanied by a variety of secondary changes.
• The presence of fleas and/or flea faeces, although absence of these may not exclude the diagnosis.

► Major differentials.
• The major differential is atopic dermatitis, although it must be remembered that the two conditions may co-exist in the same patient:
  • In one study 4/5 of dogs with FAD were also affected by atopic dermatitis!
  • Also, 1/3 of dogs with atopic dermatitis were also affected by FAD!
• Key distinguishing features of atopic dermatitis:
  • The distribution is facial and ventral.
  • There is usually a concomitant otitis externa.
  • Refer to the monograph on atopic dermatitis.
• Care must also be taken to exclude other parasitic dermatoses including infestation with Cheyletiella, Trombicula and lice.
• Depending upon the geographic location, FAD may well be by far the most common allergic skin disease in small animal practice, and must be confirmed or excluded with certainty during the differential diagnosis of allergies, once the common parasitic and microbial skin diseases have been excluded (dermatoses of groups 1a and 1b).

➡ Diagnostic tests.
• Confirmation of the diagnosis of FAD should be made.
• The “gold standard test” remains the intradermal skin test with a positive immediate (10-20 minutes) or delayed (24-48 hours) reaction to flea allergens. This test can be performed in conjunction with intradermal injections of other allergens to detect hypersensitivity in atopic dogs, but may also be performed alone with a positive (histamine) and negative (diluent) control.
  • Refer to the monograph on atopic dermatitis for the method of determination.
  • In the event of a negative result at 10-20 minutes, it is important to check the site at 48 hrs for any delayed reaction. The site must therefore be carefully marked with an indelible marker. It should be noted that in contrast to the urticarial immediate reaction, the delayed reaction may be evidenced by local erythema, palpable thickening or a crusted papule.
• **In vitro assays** for allergen-specific IgE are available but they are less reliable, both in terms of sensitivity and specificity. They can be performed using either monoclonal or polyclonal anti-canine IgE, or with the Fcε receptor assay. However, some 15-30% of cases of FAD will show delayed hypersensitivity only and will thus have a true positive delayed skin test, but a negative in vitro test.

• It must be remembered finally that the demonstration of hypersensitivity does not alone justify a diagnosis of FAD. All it means is that the patient is hypersensitive to flea extracts, but it does not necessarily mean that the dermatitis observed is the result of FAD or of FAD alone. **Response to parasiticidal therapy is then of major significance.**

**Therapeutic trial.**

• The only true way to confirm the diagnosis is by a therapeutic trial.
• For full details, see the section on flea control.
• The aim is to eliminate the flea bites over a 2-3 week period to allow resolution of the clinical signs.
• Main considerations:
  • All pets should be treated.
  • If there is a heavy environmental contamination, this must also be treated with an adulticide and an IGR.
  • A spray formulation is likely to give the quickest response for on-animal use.
  • Ideally a product with a quick killing action should be used.
  • Permethrin deserves special consideration for such a trial as it fulfils the above criteria, and to a greater extent than do some newer products, but should not be used on cats.
  • Symptomatic, anti-inflammatory therapy must not be used (or at best for only 2-3 days if the patient is severely affected) as it will interfere with the conclusions.
• Conclusions:
  • A positive response (successful therapeutic trial): if the clinical signs resolve, the diagnosis may be confirmed since the positive predictive value (PPV) is very high.
  • A negative response (unsuccessful therapeutic trial): the dog may not be affected by FAD, but the negative predictive value (NPV) is not as high as the PPV.
    • a negative response may be due to the fact that the patient is still being bitten,
    • increase this NPV using permethrin and following an adequate protocol (see below) to be able to conclude that actually the dog is not suffering from FAD.
  • The therapeutic trial may have been effective, but leading to a partial response:
    • pursue (or reinforce) the treatment if specific clinical signs of FAD remain,
    • the dog may have concomitant atopic dermatitis (and/or an adverse food reaction) if non specific signs remain, control of the FAD is then still necessary, and the necessity of initiating a control of the atopic dermatitis should be evaluated afterwards.
The recommended protocol is summarised below:
- 1, 2 and 3 are mandatory.
- 4 is highly recommended.
- 5 considerably increases the NPV.
- The long term treatment may be adapted in each case (depending on the occurrence of clinical signs in the affected animal and of the environmental parasitic infestation).

<table>
<thead>
<tr>
<th>Days of treatment and/or consultation</th>
<th>D₀</th>
<th>D₇</th>
<th>D₁₄</th>
<th>D₁₅ to D₂₁</th>
<th>Long term</th>
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<tbody>
<tr>
<td>1. Taking the history and clinical examination</td>
<td></td>
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<tr>
<td>2. Treatment of the affected dog with an efficient product (adulticide + IGR)</td>
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<td>+</td>
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<tr>
<td>3. Treatment of the in-contact pets with IGRs</td>
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<td>-</td>
<td>-</td>
<td></td>
<td>+</td>
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<tr>
<td>4. Treatment of the in-contact pets with adulticides</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td></td>
<td>+ / -</td>
</tr>
<tr>
<td>5. Perfect environmental treatment</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td></td>
<td>-</td>
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</table>

**Treatment**

**General remarks.**
- The only way to avoid the clinical signs (symptoms and lesions) of FAD in affected dogs is to eliminate the source of allergenic challenge i.e. prevent all flea bites by eliminating the fleas.
- Resolution will then occur within 15 days.
- Prevention of access of allergen to the animal is the best way to control any allergy. And, if most of the time, this is unfortunately not possible in atopy, it is possible in FAD, so it should be the goal.
- FAD can be perfectly controlled with the correct use of parasiticides.
- See “flea control” and “therapeutic trial” sections.

**Symptomatic and anti-inflammatory.**
- General remarks.
  - Symptomatic anti-pruritic therapy should be used with caution, remembering that it will no longer be possible to assess the efficacy of the anti-parasite strategy.
  - If they are required, it is preferable to use short-term oral corticosteroids than long-acting injectable forms which may induce serious side effects with repeated use.
  - Reassure the owner that pruritus will rapidly decrease with a successful parasiticidal therapy.
  - However, symptomatic antipruritic therapy should not be used in the case of a therapeutic trial.
- Topical therapy:
  - Topical therapy is not generally employed in the therapy of FAD, except in relation to flea control (see below).
  - The use of shampoos and/or lotions may be appropriate in some cases.
• Systemic therapy:
  • Corticosteroids:
    • Effectively control the inflammatory response.
    • Preferably, oral products rather than long-acting injectables.
    • May be sometimes helpful at the beginning, but do not replace an appropriate parasiticidal treatment (which would be rapidly effective alone!).
    • Their use does not allow correct evaluation of the “flea control” efficacy.
  • Other anti-inflammatory agents:
    • Antihistamines are generally ineffective in FAD.
    • Essential Fatty Acids, in sufficient doses may help, especially if there is concomitant atopic dermatitis.
  • Antibiotics:
    • Any secondary bacterial infection is usually minimal, and does not require systemic treatment.
    • Occasionally, however, animals are encountered where the acquisition of a flea infestation initiates a generalised pyoderma. In such cases, a 2 week course of a suitable antibiotic is indicated (see complex approach pyoderma monographs).
• Additional therapy:
  • Immunotherapy with the currently available products is not effective.

⇒ **Flea control.**

  • General remarks:
    • Although less rigorous protocols may be adequate to keep flea infestation controlled in non-allergic animals, the final objective in a case of FAD is to avoid any flea from biting the affected animal during its life.
    • Total elimination of the flea population within the household is the best way.
    • There are three distinct but related means of achieving the final objective:
      • eliminate the flea population from each animal in the household (the affected animal and all the in-contact pets),
      • protect them from continued infestation from adult fleas from the environment,
      • eliminate from the environment the reservoir of eggs, larvae and pupae.
    • Three principles must be considered when selecting the optimal method for flea control.
      • careful consideration of the environmental conditions and of any in-contact animals must precede the formulation of the treatment regime,
      • the goal is to eliminate the flea population as rapidly as possible, and with products that have repellent and/or anti-feeding properties to treat the affected animal,
      • use of a combination of products, so called "integrated flea control", is likely both to be more effective and also to limit the development of insecticidal resistance.

• Informing the owner and constructing the flea control programme:
  • After the diagnosis is established, the most cost-effective flea control has to be chosen, bearing in mind any special circumstances pertaining to the pets, the environment and the owner.
  • The most appropriate programme must be formulated aiming at all in-contact pets and the environment (both inside and outside if appropriate) where they spend time.
  • A major factor in achieving success is client education, taking the time to explain essential features of flea biology, as they affect the selected strategy, and of the properties of the products selected and the rationale behind their use.
The owners must be convinced that the most effective control programme is not likely to be the cheapest, but that time and money will be saved in the long run by adopting an effective strategy:

- The least treatment acceptable is the application of efficient and quick acting topical parasiticidals on the affected animal at appropriate intervals.
- The efficacy will be greatly increased by an efficient environmental treatment, even if done only once at the beginning.
- After this discussion, explain that the animal’s treatment might be reduced later (especially in winter...) and that if clinical signs of FAD reappear, this is because of ineffective parasite control, which will then have to be reinforced, first by re-initiating topical treatment of the affected animal and second by re-evaluating the treatment of the in-contact pets and of the environment.

ENVIRONMENTAL TREATMENT

- Adulticides and IGRs applied in each habitat where the affected dog goes (even once a year), in each room, using foggers, sprays on pump-sprays to treat where foggers do not.
- In warmer climates contamination of the outside environment may be a significant source of reinestation, and an adulticide and an IGR should be applied to shady areas where there is plenty of organic debris. The IGR chosen should be resistant to ultraviolet light (e.g. pyriproxyfen—as opposed to methoprene which may be inactivated by ultraviolet light).
- Do not forget all the areas frequented less commonly by the affected animal, including the car, the garage...
- Affected animals and all in-contact pets may be “used” to assist in the environmental treatment, giving them topical or systemic IGR (pyriproxyfen and lufenuron for example). Then, if a flea “accidentally” takes a blood meal on such treated animals, it will not reproduce thus preventing environmental contamination.

TREATMENT OF IN-CONTACT PETS

- Treatment with IGRs and/or adulticides.
- Topical applications are preferable when possible. IGRs may be sufficient in long-term control.

TOPICAL TREATMENT OF AFFECTED ANIMALS

- The best way to protect an affected dog from FAD is topical applications with locally acting products containing adulticides preferably in association with an IGR.
- Maintenance of continual and effective anti-parasite pressure is recommended.
Adulticides and IGRs in Flea Control

Example of an aggressive first treatment

**Reservoirs of fleas (host-environment life cycle)**

- Newly emerged hungry adult fleas
- Adult flea responsible for the FAD clinical signs
- Adulticides and IGRs in Flea Control
- Example of an aggressive first treatment

**ANIMAL TREATMENT:**
SPRAYS, SPOT-ON or LINE-ON for ADULTICIDE (preferably SPRAY for the FAD affected DOG) & IGRs (preferably TOPICAL)

**ENVIRONMENTAL TREATMENT:**
SPRAYS & FOGGERS

**NOTE:** IGRs given to animals contribute to the environmental treatment
Adulticide and IGRs in Flea Control

Example of maintenance treatment

Possible presence of newly arrived adult fleas (from the outside cat for example)

Adult flea which could have been responsible of FAD clinical signs!

ANIMAL TREATMENT:
SPRAYS, SPOT-ON or LINE-ON for ADULTICIDE
(preferably SPRAY for the FAD affected DOG)
& IGRs (preferably TOPICAL)

NOTE: IGRs given to animals participate to the environmental treatment

Reservoirs of fleas (host-environment life cycle)
Sources of infestation
**Types of products available.**
- Products used can be divided into **two types**, namely the **insecticides**, which kill the flea, and **insect growth regulators (IGRs)** which act on other stages of the life-cycle. The latter can be applied to the environment or to animals.
- **Insecticides**: amongst the older products, organophosphates and carbamates are still very effective. Pyrethroids, such as permethrin, are still potent insecticides and the latter product has been shown to have an **anti-feeding as well as a quick killing action.** A number of new products have become available in recent years, including fipronil, imidacloprid and selamectin.
- **Insect growth regulators (IGRs)**: these consist of the chitin synthesis inhibitors (lufenuron) and the juvenile hormone analogues which are ovicidal and larvicidal. Methoprene and fenoxycarb were the early products, but the recently developed **pyriproxyfen** is far more potent than either. Also, it has recently been shown to have insecticidal activity. Pyriproxyfen may be used alone for environmental treatment or on in-contact pets. It has been shown to act against newly emerged adults fleas, limiting their ability to take their first blood meal (which has to be taken very quickly otherwise fleas rapidly die).
- **Effective flea control involves use of a combination of insecticidal and growth regulating products.**

**Topical parasiticidal therapy.**
- **Shampoos:**
  - Shampoos are generally only effective in mild infestations and where there is little environmental contamination. However, they may form part of an approach using combinations of products.
  - Pyrethroids or carbamates are usually employed
- **Sprays:**
  - These are still popular ways by which to apply pyrethroids and also fipronil.
  - They are **often used in therapeutic trials.**
- **Collars:**
  - Although collars are often used for controlling flea infestation, more aggressive therapy is usually required for the treatment of FAD. Organophosphates and pyrethroids are often employed in collars.
- **Spot and line application:**
  - This is a new and **very popular way** in which to apply insecticides. Following application the products quickly disperse throughout the skin surface.
  - Permethrin, fipronil, imidacloprid, selamectin, metaflumizone and pyriprole are insecticides often used in this way.
  - The potent insect growth regulator pyriproxyfen is also available in this form alone, or admixed with permethrin as a potent integrated approach.

**Oral parasiticidal therapy.**
- The older oral products were of limited efficacy in FAD, as the flea has to bite before it gains access to the product (as is the case for systemically acting topical products).
- Two recent products may be effective in this formulation, namely nitenpyram (very short acting) and spinosad.
  - Also, two IGR can be administered orally: tablets containing lufenuron, and **pyriproxyfen** in a registered pet-food.
**Prognosis**

- The prognosis for FAD is good so long as a complete therapeutic strategy is involved which takes into account all in-contact animals that could provide a reservoir, as well as the environment.

**Follow-up**

- After the initial treatment, it is important to evaluate the clinical situation 2-3 weeks later. If the treatment is only partially successful, a reassessment of all aspects, and especially the environmental situation, is important.
- It must be remembered that once the animal is allergic, it is likely to remain so indefinitely.
- Therefore it is imperative that a long-term prevention strategy is developed for the future and regularly reassessed.

**Conclusions**

- FAD is still a very common skin disease in temperate climates.
- The only effective means of curing and preventing FAD remains the protection of the affected animal from flea bites.
- Newer products and the use of an integrated approach employing both an insecticide and an insect growth regulator have now made what used to be a very difficult problem, readily controllable.
- However, the key to success is still to develop a programme tailored to the individual situation and including all in-contact animals.
- **Combination therapy** using an insecticide and an insect growth regulator tremendously enhances the speed of success and the efficacy of long term prevention.
- Management of FAD is not really complex in itself, but it has to be remembered that it may be associated with secondary cutaneous infections and/or other allergic dermatitis, especially with atopic dermatitis (increasing the risk of cutaneous infections).

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Photomicrograph of *Ctenocephalides felis* egg.  
Courtesy of: P. Bourdeau

Photomicrograph of *Ctenocephalides felis* larva.  
Courtesy of: P. Bourdeau
Adverse Food Reactions (AFR) (Food Allergy and Food Intolerance)

Introduction / Definition

- Adverse food reactions can involve either immunologic or non-immunologic mechanisms. The former are called food allergy, and the latter are food intolerance. The relative incidence of each in veterinary medicine is unknown, and as the clinical signs are probably indistinguishable, they will be considered as one entity.
- Typically, AFR leads to a non seasonal pruritus.
- AFR may coexist with Canine Atopic Dermatitis (CAD), and Flea Allergy Dermatitis (FAD) can, of course, be superimposed.
- AFR is one of the three major allergic skin diseases that have to be considered in the differential diagnosis of pruritic dermatoses, the others being FAD and CAD, clinical signs of AFR being similar to the latter in a certain number of cases.
- During this differential diagnosis of allergies, AFR takes a particular place: after exclusion of FAD, even though AFR is not so common, it should always be investigated and excluded (or controlled) before initiating any treatment of CAD (even if the dog showed a positive intradermal test).

Aetiology / Pathogenesis

- The allergens.
  - Most dog owners believe that their dog cannot be affected by AFR because the dog has been on the same diet for years!
  - The offending antigen is usually a basic food ingredient of the diet, theoretically containing proteins (of unknown nature), and possible also carbohydrates.
  - Sensitisation to an antigen often follows a long refractory period, up to two years or more, before clinical manifestations become evident.
  - Major food ingredients involved in allergic reactions include meat (beef, pork, lamb...), fish, egg, milk and dairy products, wheat, rice, soybean, maize and other cereals.
  - In most cases, pruritus is due to one allergen or possibly two, but some dogs are multisensitive.
  - It has also been shown that some food antigens cross-react. Thus most animals reacting to beef (probably the major food allergen in dogs) will also react to lamb and milk.
  - Commercial canned or dry pet food (which may contain some or many of these ingredients) can as readily cause AFRs as do home prepared diets.
  - Involvement of these ingredients varies between regions of the world.
  - Several studies on dogs affected by AFR showed that 3/4 of cases were due to beef, dairy products and wheat.
  - Adverse reactions in dogs to corn, rice, pork and fish are rarely reported.
  - Contrary to what occurs in man, additives or preservatives are not considered to be a common cause of AFRs in dogs but it may be possible.
- Pathogenesis.
  - The specific immunological mechanisms involved in food allergy are not well understood.
  - Type I (immediate anaphylactic response), type II, type III (Arthus phenomenon) and type IV (cell-mediated reactions) have all been hypothesised.
  - It is likely that animals that respond quickly to a diet trial and relapse upon challenge within 1-2 days are IgE-mediated.
  - Dogs affected by AFR may be predisposed to secondary staphylococcal and/or Malassezia infections and/or proliferation.
Epidemiology

- AFR is not an uncommon skin disease.
- Estimates of its incidence range from 1 to 5% of all dermatological cases and 5 to 30% of allergic dermatoses.
- The following features are highly characteristic:
  - Age of onset has been reported to be very variable, from a few months (more probably due to dietary intolerance) to more than 10 years (almost certainly involving real allergic reactions). Onset of pruritus late in the animal’s life (> 7yrs) may be a compatible sign. However, in general, AFR has an earlier age of onset that does CAD, with almost one half of cases commencing prior to one year of age.
  - Breeds predisposed to AFRs include those predisposed to CAD and in addition, German shepherds, pugs and Rhodesian ridgebacks.

Clinical signs

- Perennial pruritus is the most common manifestation of AFR.
  - The degree of pruritus is variable but often intense and relatively constant from the time of onset and sometimes quite refractory to corticosteroid therapy.
  - Pruritus and lesions can be generalised but the distribution is predominantly facial and ventral (axillary and inguinal), with scratching of the trunk, rubbing of the face (lips, chin…) and chewing of the feet frequently seen. Sometimes it can be rather focal, and involve a distribution unusual for CAD, such as the distal part of all the limbs.
  - Erythema and a papular eruption may occur and are the most common primary lesions. Other primary cutaneous manifestations may include urticaria and angio-oedema (particularly in very young dogs).
  - In most cases secondary lesions develop and include excoriations, crusts, lichenification, hyperpigmentation and keratoseborrhoeic changes.
  - Secondary dermatoses may occur, such as bacterial overgrowth, folliculitis, Malassezia dermatitis, bacterial overgrowth, pyotraumatic dermatitis (hot-spot) and bacterial pododermatitis.
  - After treatment of secondary dermatoses, typically the patient remains pruritic in the areas of predilection for AFR.
  - A primarily erythematous bilateral otitis externa may be observed, with a rapid occurrence of a secondary infection involving bacteria and/or the yeast Malassezia (see the monograph on otitis).
  - Gastrointestinal signs, such as vomiting and diarrhoea, are reported in 10 to 15% of cases, and are sometimes the only apparent clinical signs of food allergy. Colitis (in some cases associated with perianal fistulas) has been reported. In some cases, increased frequency of defecation, and/or flatulence, can be an indication of a mild gastrointestinal disorder that can point to AFR. Thus concomitant gastrointestinal and dermatological signs are highly suggestive of an AFR.
  - Neurological signs (seizures) are rarely seen in dogs, but many owners report that their animal seems to “feel better” after initiation of the elimination diet.
  - It is not known if canine “asthma”, which is very uncommon, may sometimes be associated with AFRs.
  - It is important to re-emphasise that similar clinical signs can result from other diseases (CAD, FAD, Malassezia dermatitis, idiopathic bacterial folliculitis, Sarcoptic mange…). These must therefore be excluded and the secondary complications treated before AFR can be implicated in the disease process.
**Diagnosis**

► **General history.**

- Sudden or progressive onset of a mild to severe pruritus, sometimes late in the animal’s life.
- Onset of pruritic disease at <6 months of age, with or without urticaria.
- **Perennial pruritus** with no seasonal influence.
- The delay between ingestion of the offending food and onset of signs may depend on which type of hypersensitivity response predominates and on the particular antigens involved. However, most affected dogs have a constant pruritus with no obvious relationship to the diet.
- **Pruritus refractory to corticosteroid therapy,** particularly in chronic cases (this is compatible with AFR and might be an important point in the diagnostic criterion in the differential diagnosis of allergies). Early cases, however, sometimes respond quite well to corticosteroids.
- Because of a possible **poor corticosteroid response** to pruritus in chronic cases, dogs with food allergy are more likely to present with secondary **iatrogenic Cushing’s syndrome.**

► **Dietary history.**

- The dietary history should be carefully reviewed with the owner with two main objectives:
  - first, to identify ingredients commonly associated with AFR,
  - second, to determine what ingredients that have never been eaten by the dog should be used for the elimination diet.
- This history should include commercial foods, treats, biscuits, supplements, chewable medication, chew toys and also access to other sources of food: e.g. various leftovers from human meals, commercial cat food...

► **Clinical elements.**

- Remember that clinical signs may be similar or identical to atopic dermatitis.
- Pruritus of the face (rubbing) and of distal extremities (licking, chewing).
- Distal, facial and ventral to generalised distribution of lesions.
- Bilateral pododermatitis.
- Otitis externa.
- Secondary pyoderma and/or bacterial overgrowth.
Major differentials.

- FAD (Flea Allergy Dermatitis):
  - The major differential in flea-endemic areas is FAD.
  - The two diseases may co-exist in the same patient but FAD is more frequently associated with atopic dermatitis.
  - Key distinguishing features of FAD are (see monograph on FAD):
    - There is always a primary eruption, which is papular and crusting.
    - The distribution involves the lower back and posterior and inner thighs.
    - There is no obvious association with otitis externa.

- CAD (Canine Atopic Dermatitis):
  - It should be emphasised that clinical features of CAD can be identical to those of AFR. Differential diagnosis is thus very important.
  - Thus most dermatologists prefer to look for AFR prior to exploring further a diagnosis of CAD (see monograph on atopic dermatitis).
  - As has been noted in the case of FAD, atopic dermatitis and AFR often exist in the same patient.
  - Key distinguishing features of CAD are (see monograph on atopic dermatitis):
    - Peak-age of atopic dermatitis is between 1 and 3 years old.
    - Primary papular eruption might be less frequent than in AFR.
    - There is often an associated otitis externa, as in the case with AFR.

- Is the following case a dog affected by AFR or atopic dermatitis?

- The above pictures show a dog affected by food allergy (diagnosed by elimination diet and provocation tests) which perfectly mimics atopic dermatitis.
- At that time, clinical signs were not due to aeroallergen hypersensitivity revealed by the IDT.
- This emphasises first that a positive IDT is not diagnostic for atopic dermatitis, and second that it is necessary to perform an elimination diet test before trying to control any atopic dermatitis, even in the face of a positive IDT.
• **Parasitic Dermatoses:**
  - As is the case in FAD, **care must also be taken to exclude the pruritic parasitic dermatoses.**
  - These include **Sarcoptic mange** (see monograph on Sarcoptic mange) and infestation with **Cheyletiella, Trombicula** and lice (see monograph on dermatoses associated with infestations by other parasites).

• **Allergic Contact Dermatitis (ACD):**
  - Food allergy may also be confused with allergic contact dermatitis, which is **very rare in dogs.**
  - Key features of allergic contact dermatitis are:
    - There is always a primary eruption which is macular (erythema) and papular.
    - The eruption is restricted to contact areas where the hair is absent or thin.

> **Diagnostic tests.**

• **General remarks:**
  - The use of **in vitro tests** for food allergen-specific IgE and/or IgG is controversial. It has been shown that although normal dogs have IgG antibodies to food antigens, and that some also have IgE antibodies, the frequency and level of these antibodies is significantly greater in dogs with confirmed AFR than in normal dogs. So although serology cannot be used to make the diagnosis, it may be helpful in identifying the suitable composition of the elimination diet – particularly in relation to cross-reacting antigens.
  - **Histopathology** is not diagnostic but may be **suggestive of cutaneous hypersensitivity.**
  - The only definitive way to confirm the diagnosis of AFR is to perform a prolonged elimination diet trial ("hypoallergenic" test diet).

• **The purpose of this trial is to evaluate the improvement of clinical signs and especially pruritus.**
  - In order to appreciate the effect of the trial diet, **no other therapy is permitted,** neither systemic nor topical, especially corticosteroids! (However if the dog is severely pruritic, a short course of short-acting corticosteroids may be used for symptomatic relief early in the dietary trial)
  - Furthermore, **all other primary dermatoses** (including FAD) and **secondary dermatoses** (especially cutaneous infections) have to be excluded or treated before and during the diet trial.
  - In a dog affected by AFR, clinical signs will gradually decrease, over 3-8 weeks once the offending allergen has been eliminated from the diet.

• **Practical guideline for the elimination diet test:**
  - **Constraints and contents of the diet:**
    - The elimination diet must **exclude all former diet components that have been previously ingested by the animal:** see the dietary history.
    - If serology has been undertaken, foods should be chosen to which no IgE antibody, and at best low levels of IgG antibody, are shown.
    - Although commercial "hypoallergenic" diets are available (both limited antigen and hydrolysed protein), it is **better** to perform the test with a homemade diet. The **positive predictive values are comparable** with both diets but the negative one is much higher with the homemade diet.
    - If owners are really reluctant to cook an elimination diet, commercial **“hypoallergenic diets”** should be considered. However, owners have to be advised that it is not the best way and that **food allergy may remain questionable in case of a negative response.**
    - The elimination diet is usually composed of a **single protein source** and a **single carbohydrate source,** a meat and a vegetable.
    - The dog must “**never**” have previously eaten the included foods. In case of difficulties in finding foods that the dog has never eaten before, it can be considered that the animal might not be sensitised to a food component ingested less than **once per month** previously.
    - Note that **cross-reaction between beef and lamb** has been recently demonstrated. If the dog has been fed with one, do not use the other in the elimination diet.
    - In Europe, elimination diets commonly include pork, horse meat, or boiled fish with boiled potatoes, turnips, tomatoes or spinach. Advise the owner **not to use commercial tins.**
    - Even though a hypoallergenic diet may not be adequate for long term nutrition, gastrointestinal problems are rare. In the case of constipation or diarrhoea, the best course of action is to vary the respective quantities of the meat or the vegetable.
Absolutely no other food is permitted during this period:
- Likewise, no salt, butter, sauce, supplement, vitamins, treats, rawhide strips, or even chewable toys, especially bones, are allowed. If cooked, food has to be boiled (without salt), broiled (without oil or butter) or cooked in a micro-wave oven.
- Be careful that the dog has no access to another pet food, e.g. cat food.
- Dogs are frequently given titbits; owners are often reluctant to stop this. If this is the case (ask the owner), it is easy instead to give a piece of meat which is included in the elimination diet, especially prepared and stocked in the fridge.
- All people who might be in contact with the affected dog (even once!), e.g. all members of the family (especially children), friends, guests... have to be aware of the “treatment” of the dog and clearly advised that they must not give anything to the dog.
- Tell the owner that if an “accident” occurs during the test (e.g. a piece of bread dropped on the floor and eaten by the dog), they have to advise you of this event and that the test should be re-started.
- The animal should be controlled during the trial, to avoid any incidental ingestion of foods other than the permitted ones.

Duration:
- Improvement of clinical signs is usually gradual and may take a long time to become evident.
- Some dogs respond within 3-4 weeks, but if not, the diet must be given for a minimum of 8 weeks, and prolonged beyond (up to 12 weeks if necessary), in case of partial amelioration after 8 weeks.
- Conducting an elimination diet is not easy, the basic condition is that the owner has to be convinced of its importance. But when the owner agrees to perform it, do not hesitate to prolong it for as long as necessary.

Examples of three different elimination diet results:
• Provocation test:
  - In case of positive response to the elimination diet, the diagnosis must be confirmed by reintroduction of the former diet components, one by one at 10 to 15 day intervals. This is because it has been shown that in 20-30% of cases the improvement following dietary change is coincidental, and not a result of the dietary change itself. These cases will not relapse upon challenge.
  - Recrudescence of the clinical signs may occur within 2-3 days, but sometimes is delayed up to 14 days.
  - This allows the determination of the specific allergenic agent but also confirms the diagnosis.
  - Previous foods may be reintroduced:
    - in addition to the elimination diet (with only a decrease in the amount given), especially in case of the use of a commercial diet,
    - or instead of one of the components of a homemade elimination diet (e.g. beef meat instead of chicken meat).
  - When a food component causes the recurrence of clinical signs, stop it immediately, and feed the dog again with the previous foods that caused no signs. This should resolve the clinical signs induced by the food challenge (within a few days to a few weeks).

• Examples of two different provocation test results:

1. Homemade elimination diet with chicken.
   - Beef instead of chicken.
   - Elimination of beef, and chicken again instead.
   - Reintroduction of the former homemade diet with chicken instead of beef.
   - AFR to beef is diagnosed (frequent cases).

2. Commercial "hypoallergenic" diet.
   - + Beef
   - + Rice.
   - + Wheat.
   - + Pork.
   - Elimination of pork.
   - Reintroduction of the former commercial diet with which the dog was previously fed. It did not contain any pork! But beside its diet, the dog was fed with pork from time to time.
   - AFR to pork is diagnosed.

• Conclusions:
  - When the offending component has been found, it must never be given to the dog again (although some dogs may tolerate a new exposure later in life, after a long and strict avoidance).
  - If a commercial food is suspected, it may be impossible to determine the exact allergenic agent, so changing to a homemade diet may be the best solution. As commercial dog foods contain many common foodstuffs and additives, changing from one to another is not recommended.
  - AFR often occurs in association with CAD, and in some regions also with FAD.
  - Dogs affected by concomitant AFR and CAD may show a significant but not total improvement of their clinical signs during a dietary elimination trial. These patients must then be worked up for concomitant CAD.
  - One of the most important sources of errors is owner compliance. To increase compliance, it may be pertinent to ask the owner to precisely notice the date, foods given and clinical sign modifications during the whole dietary elimination trial and provocation test.
Clinical management

**General remarks.**
- Contrary to atopic dermatitis, food allergy can usually be perfectly controlled.
- Avoidance of the allergen(s) by omitting the offending food(s) identified during the test diet is the key to a successful “treatment”.
- Diagnosis and clinical recovery are then concomitant.
- An accurate identification of the offending allergen is necessary.
- Once the diagnosis has been made, feeding a commercial hypoallergenic diet – either limited antigen or a hydrolysed protein diet, is preferable to feeding a home prepared diet for maintenance. Although the hydrolysed protein diets should be the ultimate limited antigen diet, unfortunately not every animal is satisfactorily controlled by such diets.
- A tolerable home-prepared diet is an alternative, and can usually be achieved, but must be balanced with vitamin, mineral and essential fatty acid supplements.
- Secondary complications such as bacterial and/or Malassezia infection (including otitis externa) have to be treated before and during the elimination diet, but should not recur since food hypersensitivity is controlled.
- Any associated FAD has to be managed by flea control.

**Topical treatment.**
- Shampoos and lotions formulated to help control allergic skin diseases are useful (see Atopic Dermatitis Monograph).
- Otitis externa should be treated with an antibiotic, anti-yeast and anti-inflammatory otic preparation.
- An antiseptic ear cleanser may be useful for regular ear cleaning.

**Systemic treatment.**
- Systemic treatments and especially corticosteroids are usually unnecessary, bearing in mind that food allergy may not respond well to corticosteroid therapy.
- However, essential fatty acids may improve the skin and haircoat quality.

Prognosis

The prognosis for food allergy is good because the offending allergen can be eliminated. Rarely, dogs will become sensitised to the new diet, necessitating further investigations and dietary manipulations. Very rarely, dogs may become sensitised to new diets so quickly that they have to be maintained on anti-inflammatory or immunosuppressive therapy.

Follow-up

All dogs with confirmed AFR should be checked periodically to ensure that they are still asymptomatic.

Conclusions

Food allergy (or AFR) is not a common skin disease but is not rare. An elimination diet should be instituted when appropriate, i.e. where there is suspicion of an AFR, and also in cases of CAD to ensure that there is not also a contribution from an AFR. Even if the understanding of this skin disease is not complete at the present time, diagnosis and long term management of dogs with AFR is achievable.
Canine Atopic Dermatitis (CAD; Atopy)

Introduction / Definition

- Canine atopic dermatitis (CAD) is defined as: “A genetically predisposed inflammatory and pruritic skin disease with characteristic clinical features associated with IgE antibodies most commonly to environmental allergens.”
- Very frequent, atopy affects from 3-15% of the canine population and in some studies up to 50% of dermatological cases involve atopic dogs.
- Clinical signs are variable and are frequently complicated by concomitant flea allergy dermatitis (FAD), adverse food reactions (AFRs) and/or associated secondary infections (especially staphylococcal folliculitis).
- CAD is one of the three major allergic skin diseases that has to be considered in the differential diagnosis of pruritic dermatoses, the others being FAD and AFRs.
- The clinical diagnosis is facilitated by eliminating the possibility of other diseases, and treating or controlling any concomitant diseases: remaining clinical signs should then be carefully assessed for compatibility with those associated with CAD.
- Atopy, atopic disease and canine atopic dermatitis (CAD) are all acceptable terms for the same disease, with the latter preferred by the International Task Force on Canine Atopic Dermatitis (ITFCAD).

Aetiology / Pathogenesis

- The allergens.
  - Formerly called “inhalant allergens”, the current preferred name is “aeroallergens”, as they are believed to gain access to the body via the percutaneous route.
  - The majority of CAD cases result from hypersensitivity to house dust mites, leading to a non seasonal dermatitis (although the severity of the signs may vary with changes in the environmental allergenic load).
  - Less commonly, pollen allergies are involved, which may typically lead to a seasonal dermatitis.
  - Involvement of other allergens (moulds, human or animal dander...) are less common.
  - The list of allergens that should be included in intradermal test (IDT) protocols must be adapted for each different area, in each different country and in each different continent!
  - However, throughout the world, the major allergens involved in CAD are house dust mites, and especially Dermatophagoides farinae (Df), and to a lesser extent Dermatophagoides pteronyssinus (Dp). This leads to a non seasonal pruritus, although its intensity may vary due to the activity of the dust mites, and also if there are additional seasonal allergens involved.
*Pathogenesis.*

- The symptoms that affect atopic dogs are directly linked to an **inherited dysfunction of the immune system and** also to **defects in cutaneous barrier function that facilitate access of allergens and predispose them to bacterial and yeast proliferation and infections.**
- The following steps are believed to be important in the pathogenesis:
  - Access of allergen is via the **percutaneous** route. **Alteration of epidermal function barrier in atopic dogs facilitates this penetration.**
  - Allergen capture is aided by epidermal Langerhans’ cells which are armed with allergen-specific IgE.
  - The pathogenesis also involves **interaction of allergen with IgE antibody attached to mast cells.**
  - A variety of mediators are involved including histamine, leukotrienes and proteases from mast cells, and probably also interleukins from keratinocytes.
  - Staphylococci attach more readily to the skin of atopic dogs, and this leads to the development of a bacterial overgrowth and/or secondary pyoderma which is usually a folliculitis. Atopic dogs also have impaired cell-mediated immunity which favours the persistence of the infection, and they may develop IgE anti-staphylococcal antibodies which add to the allergenic load.
  - Furthermore, there is often proliferation of the yeast *Malassezia pachydermatis* with the development of IgE antibodies against soluble antigens further compounding the disease process.
  - Chronically affected atopic dogs frequently develop severe seborrheic changes.
Epidemiology

- With the exception of flea allergy dermatitis in flea-endemic areas, CAD is the most common allergic skin disease, and one of the most common pruritic dermatoses.
- The following features are highly characteristic:
  - The peak age of onset is between one and three years of age. It is uncommon for it to commence at less than 6 months of age and very rare at more than 7 years of age.
  - Although it can affect any breed, including mixed-bred dogs, there is a greater than expected incidence in terriers, setters, golden and Labrador retrievers, Dalmatians, English bulldogs, boxers, bichon frisés, Lhasa apsos, English springer spaniels and the Chinese shar pei.
  - Most cases suffer from perennial pruritus that may be subject to seasonal exacerbation in severity. However, a minority are affected for a limited time each year if their sensitivities are restricted to seasonal pollen allergens.

Clinical signs

- The degree of pruritus ranges from mild to intense.
- The distribution of pruritus and lesions is predominantly facial and ventral, with scratching of the trunk, rubbing of the face (lips, chin...) and chewing of the feet frequently seen.
- In the early case, there may be little in the way of primary eruption, but by the time of presentation there are usually erythematous macules and papules. However, some 60% of cases suffer from a secondary pyoderma which is usually a folliculitis. Treatment with antibiotics alone will clear the pyoderma, but typically, the patient remains pruritic with the pruritus then restricted to the sites of predilection for atopic dermatitis. In occasional cases, the pruritus ceases when the pyoderma is controlled, which seems to represent a subclinical atopic state. The pyoderma, and the pruritus generally return shortly after the cessation of antibiotic therapy.
- Evidence of self-trauma: a variety of skin changes may result, commonly including excoriations and occasionally acral lick granulomas and episodes of acute moist dermatitis (hot-spots).
- An otitis externa accompanies some 80% of cases (!), and occasionally it may be the sole presenting sign. Characteristically, the otitis commences by affecting the inner surface of the pinna and the vertical ear canal, but in longstanding cases the horizontal ear canal also becomes involved with a secondary infection involving bacteria and/or the yeast Malassezia (refer to the Otitis Monograph).
- Seborrhoeic skin changes are commonly seen in chronic cases with a greasiness of the skin and hair coat and marked scaling accompanied by varying degrees of alopecia.
- It is important to re-emphasise that similar clinical signs can result from other diseases. These must therefore be excluded, and the secondary complications controlled, before the clinical diagnosis of CAD can be made.
Diagnosis

**Historical and clinical diagnostic criteria.**

- Those **criteria** (key historical and clinical features) provide **consistency** in the diagnosis of atopic dermatitis. **They should be documented after ruling out or controlling other primary dermatoses and secondary cutaneous infections.**
- Considering the key criteria, the **greater the number** that are satisfied, the more **specific** will be the diagnosis (but lower will be the diagnostic sensitivity).
- In practice, the best compromise for sensitivity and specificity is where three to four key criteria are satisfied.
- **Presence of other relevant criteria supports the diagnosis.**

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**KEY CRITERIA**

- Onset of clinical signs between 1 and 3 years old.
- Gradual onset of pruritus, especially on the face (rubbing) and on the extremities (licking, chewing).
- Corticosteroid responsive pruritus (at least at the beginning). Discoloration of the hair on the extremities may be a very significant sign.
- Bilateral front feet pododermatitis.
- Otitis externa.
- Inflammation of the internal surface of the pinnae.
- Facial erythema (around eyes, lips and of the chin).
- Positive IDT (or serological test for allergen-specific IgE).

**OTHER RELEVANT CRITERIA**

- Breed predisposition and/or family history.
- Chronic or chronically relapsing dermatitis for more than two years.
- Secondary cutaneous infections, especially bacterial folliculitis.
- Bilateral conjunctivitis and/or epiphora.
- Lesions (lichenification) of the flexural surface of the tarsus.
- Seasonal aggravation of symptoms.
- Variation in severity with environmental changes.
- In some cases, there is exacerbation when in contact with grasses (may also result in contact dermatitis).

**IMPORTANT EXCLUSIONS ARE:** Absence of dorso-lumbar involvement (as is seen commonly in FAD). Absence of involvement of the ear margins (as is commonly seen in scabies).

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**Part 2: Canine Atopic Dermatitis (CAD; Atopy)**

**KEY CRITERIA**

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- Bilateral front feet pododermatitis.
- Otitis externa.
- Inflammation of the internal surface of the pinnae.
- Facial erythema (around eyes, lips and of the chin).
- Positive IDT (or serological test for allergen-specific IgE).

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**IMPORTANT EXCLUSIONS ARE:** Absence of dorso-lumbar involvement (as is seen commonly in FAD). Absence of involvement of the ear margins (as is commonly seen in scabies).
Part.2

Canine Atopic Dermatitis (CAD; Atopy) • 5

Clinical Handbook on Canine Dermatology

**Major differentials.**

- Flea allergy dermatitis (FAD):
  - The major differential in flea-endemic areas is FAD.
  - **However, the two diseases often co-exist** in the same patient, and most atopic dogs will develop FAD if exposed to fleas.
  - Key distinguishing features of FAD are (see monograph on FAD):
    - There is always a primary eruption, which is papular and crusting.
    - The **distribution** involves the **lower back** and posterior and inner thighs.
    - There is **no obvious association with otitis externa**.

- Adverse food reactions (AFR):
  - **More difficult is the distinction from** AFR **whose clinical features** can be **similar** to those of CAD.
  - Thus most dermatologists prefer to eliminate the possibility of an AFR by feeding a hypoallergenic diet for 8 weeks prior to exploring further a diagnosis of CAD (see monograph on adverse food reactions).
  - Adverse food reactions and canine atopic dermatitis can exist in the same patient.
  - Key distinguishing features of AFR are (see monograph on adverse food reactions):
    - AFR can occur at any age, with some 50% of cases commencing prior to 1 year of age.
    - There might be more often a primary papular eruption than in CAD.
    - There is often an associated otitis externa as in the case with CAD.

- Parasitic Dermatoses:
  - As is the case in FAD, **care must also be taken** to exclude the pruritic parasitic dermatoses.
  - These include **Sarcoptic mange** (see monograph on Sarcoptic mange) and infestation with **Cheyletiella, Trombicula** and lice (see monograph on dermatoses associated with infestations by other parasites).

- Allergic Contact Dermatitis (ACD):
  - Atopic dermatitis may also be confused with allergic contact dermatitis, which is **very rare in dogs**.
  - Key features of allergic contact dermatitis are:
    - There is always a primary eruption which is macular (erythema) and papular.
    - The eruption is restricted to contact areas where the hair is absent or thin.

**Diagnostic tests.**

- Confirmation of the clinical diagnosis of CAD should be made:
  - The **“Gold Standard test” remains the intradermal test** with a positive immediate (10-20 minutes) reaction to environmental allergens. Positive (histamine) and negative (diluent) controls are always included and the possible role of concomitant FAD can be assessed by including flea antigen.
  - **In vitro** assay tests for allergen-specific IgE are also available, but some may suffer oversensitivity, with many false positives and thus a lack of specificity, and others from a lack of sensitivity, and they may thus be less reliable. The selection of a reliable laboratory is thus essential. They can be performed either using monoclonal or polyclonal antisera specific for canine IgE, or alternatively using the cloned α chain of the Fcε receptor (the mast cell receptor that binds IgE).
  - Cellular techniques (“basophil degranulation tests” or “heterologous passive transfer”) are useful laboratory tests in the diagnosis of allergies mediated by immediate hypersensitivity, but not practical for routine use.
  - It is critical to remember that a positive intradermal skin test or a positive **in vitro** test for IgE does not of itself justify a diagnosis, as many normal animals may also be positive. The test results must be interpreted in the light of a carefully taken history and carefully observed clinical signs.
  - These tests have two purposes; one is to **support the clinical diagnosis** and the other is to **permit the selection of allergens for immunotherapy**.
  - In a small minority of cases whose clinical signs are typical for CAD, negative serology and negative IDTs may be obtained. Obviously it is possible that the allergens responsible for the disease were not included in the tests, or assays. However it has become clear that this represents an important subset of CAD, and the International Task Force on Canine Atopic Dermatitis (ITFCAD) has proposed the use of the term “Canine atopic-like dermatitis” for such cases.
Practical guideline for intradermal skin tests.

1. Corticosteroids interfere with the skin test and the patient should be off therapy for 3 weeks (in the case of oral products) or at least 6 weeks after the effect has worn off in the case of long-acting injectables. Paradoxically, the pruritus recurs well before a valid positive skin test can be obtained. If there is doubt, it is wise to inject just the positive control alone and if there is no response, or just a poor response, the animal should be reassessed in a month. Remember also that topical, ophthalmic and otic corticosteroid containing preparations may also interfere. Similarly, antihistamines will interfere for up to 10 days after the last dose.

2. Excessive stress may also inhibit the intradermal reactions (maybe due to endogenous release of cortisol).

3. Most dermatologists routinely use sedation with e.g. xylazine. In some cases sedation may not be necessary, and suitable restraint may be obtained with the assistance of the owner and/or veterinary nurse. It is important, however, not to use any sedative with antihistamine properties.

4. The animal is positioned in lateral recumbancy.

5. The injections must be performed on the animal’s thorax. An area is clipped (use equipment in perfect condition) around 15 cm x 10 cm, that is as far as possible free from any dermatitic changes. This area is sufficient for up to 60 allergens.

6. For esthetic reasons, it may be wise to use an area that is lowest and closest to the anterior limb.

7. Mark the injection sites with an indelible pen.

8. The animal is ready. The allergens should be stored at 4°C (degrees Celsius) in glass vials and returned to the refrigerator immediately as they are unstable at room temperature. A small amount only is thus placed in the plastic, 1ml numbered syringes.

9. Note the time of the commencement of the test.
Perform the injections above and below the pen marks. Stretch slightly the skin site between two fingers and puncture the skin in a tangential manner, the bevel upwards in order to penetrate into the dermis. The amount to inject is about 0.05 ml. The intention is to create a small intradermal “bleb”, big enough to be visible but not too much. The most important point is to inject the same amount of each allergen.

If the “bleb” does not appear or disappears quickly, it may be because the injection has been administered subcutaneously: start again. If liquid leaks on the skin surface, remove it immediately with absorbent paper to avoid contaminating adjacent injection sites.

The injection sites may bleed a little. Swab any such sites gently with a paper towel at the end of the test.

The times at which the test is started and completed must be noted. With a little training, 20 to 40 injections can be given in 5 to 10 minutes. After the injections are completed, the animal is maintained in lateral recumbency, and must not be allowed to traumatize the injection sites.

The results are read in batches, each 15-20 minutes after the injections were administered.

A positive reaction manifests as an erythematous wheal. The presence of erythema is important as well as the elevation above the skin surface. The tests are best read in the dark, using incident light.

Grade each site and record the result (refer to the grading methods).

For the interpretation of the flea allergen results, refer to the FAD monograph.

Do not hesitate to ask the owner to repeat the test if equivocal results are obtained, or if the histamine reaction is poor. An interval of 2-4 weeks is appropriate, and, of course, corticosteroids should not be administered in the interim.

It should be remembered that an animal with Sarcoptic mange may show positive reactions to house dust mites because of cross-reactions.
List of major European allergens to be included in the screen.

1/ Phleum pratense, Timothy grass
2/ Salix, Willow
3/ Quercus robur, Oak
4/ Betula verrucosa, Birchtree
5/ Robinia pseudoacacia, Robinia
6/ Artemisia vulgaris, Mugwort
7/ Plantago lanceolata, Plantain
8/ Urtica dioica, Nettle
9/ Taraxacum officinale, Dandelion
10/ Graminaceae
11/ Bird feathers
12/ Cat dander
13/ Human dander
14/ Dermatophagoides pteronyssinus, Dp
15/ Dermatophagoides farinæ, Df
16/ Cotton
17/ Moulds
18/ Ctenocephalides felis, Flea
19/ Diluent
20/ Histamine

Results and interpretation.

• A possible grading method for intradermal skin tests reactions.
  • A grade of up to 20 is assigned to each reaction.
  • Measure the diameter of the reactions and allow 1 point per mm (refer to the largest width if the reaction is not circular).
  • Add from 1 to 4 points according to the intensity of the erythema and elevation above the skin.
  • If the grade is over 20 (rare cases) assign a grade of 20 (maximal grade).
  • An allergen reaction will ordinarily be considered as positive if its grade R is above the mean of the positive control grade (P) and the negative control grade (N) (which are respectively the histamine and diluent reactions):

\[
R_n \text{ positive if } R_n > \frac{P + N}{2}
\]

• Use another system or your own if you prefer, but always use the same.

• Another classical method is the use of a 1-4 scale, where 1 is just discernibly greater that the negative control and 4 approximates to the size of the histamine wheal.

• Example of the above skin test results:
  • Df: 10 mm reaction and 3 points for the erythema, \( R_{Df} = 13 \).
  • Histamine: 14 mm reaction and 4 points for the erythema, \( P = 18 \).
  • Diluent: 1 mm reaction and 0 points for the erythema, \( N = 1 \).
  • No other reaction has been noted.
  • \( R_{Df} = \frac{P + N}{2} = \frac{18 + 1}{2} = 9,5 \)
  • Conclusion: The skin test is positive to Df (\textit{Dermatophagoides farinæ}) and the animal is hypersensitive to house dust mites.
Treatment

General remarks.
- Atopic dermatitis cannot usually be cured but only controlled.
- It is one of the most difficult skin disorders to control as, most of the time, dogs cannot be protected from allergen exposure (especially in the case of hypersensitivity to house dust mites).
- However, quality of life can be considerably improved and atopic dermatitis much easier to control when secondary complications such as bacterial and/or Malassezia infection or overgrowth are effectively treated. Also, of course, vigorous treatment of any concomitant otitis externa and any associated FAD (or AFR), must be undertaken.

Allergen avoidance.
- This is theoretically the best treatment.
- But usually, avoidance of allergens is not possible or practical. However, it may be in some cases an important aspect of the management.
- An accurate identification of the offending allergen is first necessary.
- In very rare cases the complete elimination of allergens may be possible (birds in case of allergy to feathers, some plants responsible for allergy to their pollens...).
- In most cases, the aim will be to achieve a decrease in exposure. For example, in the case of hypersensitivity to house dust mites, the exposure may be decreased if pets stay longer outside and/or away from the bedrooms, which have the highest concentration of dust mites.
- If in doubt, special kits are available for home use by which the relative dust mite concentration of different areas can be assessed.
- The use of a miticide in the environment may also help, and recent studies have shown the considerable clinical benefit can be derived. However this is not immediate, as the allergens can persist in the environment for some months. Pyriproxyfen has also been shown to be useful in reducing mite proliferation and the combination with an adulticide such as permethrin, can be very effective.

Topical treatment.
- Topical products which are effective against the major pathogenic factors will be very useful: i.e. if they are able to help eliminate allergens from the skin surface, to help restore the epidermal barrier and to help control the inflammatory process and any secondary cutaneous infections.
- Shampoos:
  - Shampoos are helpful as an aid to the control of atopic dermatitis. Dependant on the stage of the disease, different types may be of value.
  - A shampoo with good cleansing power will assist in removing allergen, and thus limit allergen penetration.
  - A shampoo containing immunomodulating factors such as mono-oligosaccharides may provide a soothing effect, acting directly against some immune mechanisms.
  - A shampoo containing EFA (such as linoleic acid) may help to restore the epidermal barrier.
  - Where cytological examination reveals proliferation of Malassezia and/or staphylococci, antimicrobial shampoos are very helpful.
  - A shampoo with antimicrobial activity may also help to prevent secondary microbial overgrowths.
  - Antipruritic shampoos, such as those containing colloidal oatmeal will provide soothing effects.
  - Attention must also be given to the skin barrier function directly, and shampoos containing ceramides will assist in this.
Lotions:
- Lotions with the same properties described above for the shampoos will reinforce the effects of the shampoos and facilitate treatment.

Other topicals:
- Other topicals are often indicated, again, dependent upon the presenting signs.
- Spot-on preparations containing ceramides have been shown to be effective in restoring skin barrier function and should form part of the therapeutic approach.
- If there is only a limited area of involvement, topical corticosteroid preparations may be preferable to systemic administration. Particularly indicated is hydrocortisone aceponate which is not absorbed, and has fewer side effects than do other topical steroids.
- An antiseptic ear cleanser may be useful for regular ear cleaning.
- An antibiotic, anti-yeast and anti-inflammatory otic preparation for any concomitant otitis externa.
- Astringents and antiseptic solutions applied to any acute moist dermatitis (hot-spot), possibly more frequent in FAD.

Systemic treatments.
A number of systemic drugs are used in the control of atopic dermatitis. These include:

Corticosteroids:
- Corticosteroids are generally highly effective in relieving the pruritus. However, oral, short-acting products only should be employed, and, ideally, using an alternate-day regimen.
- The effect of corticosteroids is only transient.

Ciclosporin:
- When dosed at 5mg/kg this is as effective as corticosteroids, although with a slower onset of action.

Antihistamines:
- These may sometimes give partial, or very occasionally complete relief. Although hydroxyzine (2mg/kg TID) is often favoured, some cases respond better to one antihistamine than another; and this is quite unpredictable.
- Try one after another for courses of 10 to 15 days.

Essential fatty acids (EFAs):
- These may have significant anti-inflammatory action and assist in restoring the barrier function of the skin if used in a formulation providing sufficient quantities of both n-6 and n-3 EFAs with a ratio close to 5:1.
- There is synergistic activity with both antihistamines and with corticosteroids, and so sometimes combination therapy is employed.

Antibiotics:
- Where there is significant secondary bacterial folliculitis, appropriate antibiotic therapy for renewable courses of 2 weeks plus 1 to 2 additional weeks after clinical recovery should be used.
- Antibiotics may also be helpful in cases where there is bacterial overgrowth, with or without a hypersensitivity to the organism involved.
- The condition of some animals is dramatically improved with antibiotic treatment.
- See Complex Approach Pyoderma and Bacterial Overgrowth Monographs.
• **Immunotherapy:**
  - **Immunotherapy** or hyposensitisation with allergen solutions is the cornerstone of the therapy in atopic dermatitis.
  - However, it may not be appropriate for cases in which:
    - effective control is achieved with medical therapy other than corticosteroids,
    - there is a short season of involvement that is readily controlled with low dose corticosteroids...
  - All cases should be treated with immunotherapy after undertaking intradermal tests (or *in vitro* tests for allergen-specific IgE).
  - It is usual to limit the vaccine to 10 allergens, being sure to include house dust mite, even if there is only a weak positive reaction.
  - **Response to immunotherapy may be quite brisk (within 3 months) or delayed for up to 9 months.** Animals failing to respond by then are unlikely to respond with further therapy and reassessment is necessary.
  - **Concomitant medical therapy** can be used *during the course of immunotherapy* and in most cases the doses required can be *reduced* with time as the immunotherapy treatment starts to become effective.
  - It is extremely important to monitor frequently and to introduce effective therapy for any secondary complications if and when they arise.

**Conclusions.**
- It is important to re-emphasise that the *continuous control of secondary infections* is the first step!
- The control of CAD itself generally requires *combination therapy*.
- The basis of the therapeutic approach is **hyposensitisation**, together with concomitant medical therapy including EFAs and the frequent use of topical products (shampoos and lotions).
- With good management, the necessity for using corticosteroids will be considerably reduced.
Conclusions

- Atopic dermatitis is a common skin disease.
- Following the proposed dermatological approach allows the diagnosis of CAD if clinical signs remain after having ruled out (or controlled), parasitic dermatoses, secondary cutaneous infections and other allergies (FAD and AFR).
- Successful management is dependant on a thorough understanding of the pathogenesis and of the potential complications, and on a willingness to modify the therapy in the light of a changing situation.

Follow-up

- Frequent follow-ups, allowing one to be alert for the clinical situation to change are the key to success in managing atopic dermatitis.
- Bacterial folliculitis and bacterial or Malassezia proliferation can recur.
- Also, the frequency and dose of immunotherapy may need to be adjusted in the light of the response. Some cases improve with each injection, and then relapse after a few days. More frequent, lower dose therapy may be helpful. Also, some studies have shown that when the usual full dose immunotherapy is ineffective, lower doses may paradoxically achieve better results.
- The main challenge is to follow the clinical evolution of the plethora of clinical signs resulting from atopic dermatitis itself, and from the associated secondary complications. Then, a thorough historical and clinical assessment is needed at each consultation time with the aim of objectively following the response to therapy.

Prognosis

- The prognosis for atopic dermatitis is quite good.
- However, the percentage of cases that are “cured” and require no further therapy is small.
- Most cases require lifelong control with immunotherapy, supplemented from time to time with other medical therapy.
Iatrogenic and spontaneous Cushing’s syndromes (ICS and SCS)

Introduction / Definition

- Systemic disease as a result of chronic glucocorticoid excess, often with cutaneous involvement.
- Two spontaneous types of Cushing’s syndrome:
  - pituitary-dependent hyperadrenocorticism,
  - hyperadrenocorticism caused by adrenocortical tumours.
- In addition: Iatrogenic hypercorticism.
- Cushing’s syndrome is a common disorder in dogs.

Aetiology / Pathogenesis

- Regulation of glucocorticoids secretion.
  - Cortisol is secreted and released by the adrenal glands.
  - The amount of secretion depends directly on the plasma concentration of adrenocorticocorticotropic hormone (ACTH) which is synthesised in the anterior lobe of the pituitary gland from the precursor molecule pro-opiomelanocortin (POMC).
  - The ACTH synthesis is regulated by the hypothalamus and the central nervous system via neurotransmitters that cause the release of corticotropin releasing hormone (CRH) and arginine vasopressin (AVP).
  - The ACTH secretion is also regulated by stress factors (increased production) and exogenous glucocorticoids (negative feedback leading to the inhibition of ACTH secretion).
  - The blood cortisol levels vary by a factor of 9 to 10 during the day in healthy dogs and normal levels range between 1-6μg/dl (27-165 nmol/l). Thus a single blood test for cortisol is not diagnostic for either hyper- or hypoadrenocorticism.
**Pituitary-dependent hyperadrenocorticism (PDH).**
- 85% of spontaneous Cushing's cases.
- Caused by changes ranging from hyperplasia to neoplasia of the pituitary gland.
- 70% occurring in the anterior lobe (mainly microadenoma).
- 30% occurring in the pars intermedia (often macroadenoma).
- Consequences:
  - increased ACTH production,
  - stimulation of both adrenal glands,
  - often bilateral hyperplasia,
  - increased cortisol production, decreased endogenous ACTH production.

**Adrenal-dependent hyperadrenocorticism (ADH).**
- ∼15% of spontaneous Cushing's cases.
- Caused by a tumour of one (rarely both) adrenocortical glands.
- Adenoma or adenocarcinoma.
- Consequences:
  - atrophy of contralateral adrenal gland,
  - increased cortisol production.
**Iatrogenic hypercorticism.**
- Caused by inappropriate chronic use of glucocorticoid therapy, systemic or in some cases topicals, particularly the more potent glucocorticoids.
- Consequences:
  - elevated blood level of circulating glucocorticoids,
  - reduction of endogenous corticoid production,
  - often atrophy of both adrenal glands,
  - secondary adrenocortical insufficiency with reduced production of both glucocorticoids and mineralocorticoids which may lead to a life-threatening Addisonian crisis when the glucocorticoid administration is stopped.

**Consequences on the cortisol level in plasma.**

**Epidemiology**

**Breed and age predilections for spontaneous hyperadrenocorticism (SCS).**
- Poodles, Yorkshire terriers and dachshunds are more commonly affected, but Cushing’s syndrome is also seen in many other breeds.
- The disease is frequently seen between the ages of 6 to 9 years.

**Breed predilections for iatrogenic hypercorticism (ICS).**
- Tolerance to glucocorticoid therapy is very variable amongst individuals and breeds, and depends on the duration and level of administration.
- Small dogs and certain breeds (such as Boxers) show a higher risk of developing iatrogenic hypercorticism.

**Clinical signs**

Chronic excessive glucocorticoid blood levels result in a varying spectrum of both cutaneous and systemic signs.

**Skin and hair.**

<table>
<thead>
<tr>
<th>COMMONLY OBSERVED</th>
<th>LESS COMMONLY OBSERVED</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hair loss (alopecia) mainly affecting the trunk, mostly bilateral.</td>
<td>Easy epilation of hairs.</td>
</tr>
<tr>
<td>Head and legs are usually unaffected.</td>
<td>Secondary infection and/or colonisation of the skin by bacteria and/or Malassezia with or without pruritus.</td>
</tr>
<tr>
<td>Regrowth of hair reduced or absent following clipping.</td>
<td>Demodicosis.</td>
</tr>
<tr>
<td>Decreased thickness and elasticity of the skin.</td>
<td>Calcinoisis cuts, often with focal inflammation and pruritus.</td>
</tr>
<tr>
<td>Telangiectasis, prominent blood vessels.</td>
<td></td>
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<tr>
<td>Comedones.</td>
<td></td>
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<tr>
<td>Dry skin.</td>
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<tr>
<td>Hyperpigmentation.</td>
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</tbody>
</table>
Non dermatological signs.

<table>
<thead>
<tr>
<th>COMMONLY OBSERVED</th>
<th>LESS COMMONLY OBSERVED</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polydipsia / polyuria.</td>
<td>(Recurrent) urinary tract infections (cystitis).</td>
</tr>
<tr>
<td>Polyphagia</td>
<td>Acute pancreatitis.</td>
</tr>
<tr>
<td>Increased lipogenesis leading to obesity, abdominal enlargement, hepatomegaly.</td>
<td>Pulmonary embolism.</td>
</tr>
<tr>
<td>Panting, exercise intolerance.</td>
<td>Testicular atrophy, aspermatogenesis.</td>
</tr>
<tr>
<td>Muscle weakness and atrophy (particularly involving the temporal and gastrocnemius muscles).</td>
<td>Prolonged intervals between oestrus cycles.</td>
</tr>
<tr>
<td>Anoestrus.</td>
<td>Congestive heart failure.</td>
</tr>
<tr>
<td>Calcification, particularly of blood vessels, lung and adrenal glands.</td>
<td>Glycosuria, diabetes mellitus (“steroid diabetes”), often insulin resistant.</td>
</tr>
<tr>
<td>Poor wound healing.</td>
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</tbody>
</table>
**Diagnosis**

**History.**
It is very important to obtain accurate details of all corticosteroid preparations administered to assist in the distinction between spontaneous and iatrogenic Cushing’s syndrome. The diagnosis of the latter may be possible without recourse to exhaustive tests, but just based upon the history and clinical signs.

**Clinical signs.**
Very variable, see previous paragraphs.

**Diagnostic tests:**
- **Haematology and blood chemistry:** may be a useful first indicator in the differential diagnosis of endocrine disorders alongside differential blood cell counts (especially lymphocytes, eosinophils), cholesterol, AP, AST, ALT, glucose and basal free T4.
- **Hormone assays and function tests.**
  - Also helpful in some cases:
    - **Diagnostic imaging:** X ray, ultrasonography, computertomography, NMR (Nuclear Magnetic Resonance).
    - **Skin biopsies.**

**Haematology / Blood chemistry.**

### MOST COMMON ABNORMALITIES AND THEIR FREQUENCY IN DOGS WITH HYPERCORTICISM

<table>
<thead>
<tr>
<th></th>
<th>Common</th>
<th>Less common</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>80%</td>
<td>70%</td>
</tr>
<tr>
<td>AP ↑↑ (86%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leucocytosis (84%) with Eosinopaenia (0 to 1% d’eosinophils) (84%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AST, ALT ↑↑ (50%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glucose ↑↑ (45%)</td>
<td></td>
<td></td>
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<tr>
<td>Phosphorus ↓ (38%)</td>
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</table>

- Classically, dogs with Cushing’s syndrome, from whatever cause, show a leucocytosis, lymphopenia, eosinopenia, a markedly elevated AP, and moderately elevated AST, ALT, cholesterol and glucose.
- Such tests are used for a first screening although they do not enable a definitive diagnosis.
- Low free T4 serum levels do not necessarily indicate hypothyroidism in patients with Cushing’s disease. They may just be a secondary effect caused by elevated cortisol levels (euthyroid sick syndrome).

**Hormone assays and function tests.**
- **Please note:**
  - For the evaluation of cortisol serum or plasma levels the blood needs to be centrifuged within 30 minutes after collection.
  - Serum, plasma and urine samples for the evaluation of cortisol levels need to be kept in the refrigerator until mailing.
  - Serum samples for the evaluation of ACTH need to be shipped on dry ice to the laboratory immediately after collection.
  - Procedures are given below but also follow any procedures recommended by your own laboratory.
**ACTH stimulation test.**
- First test in case of high suspicion of ICS or to eliminate the hypothesis before investigating a SCS.
- Test principle: cortisol determination immediately before and after stimulation with ACTH.
- Indication: evaluation of the adrenocortical reserve in cases of suspected iatrogenic hypercorticism (ICS), in cases of suspected spontaneous hyperadrenocorticism (SCS) and also in cases of suspected hypoadrenocorticism.
- Performance: a fasting blood sample is taken in the morning, immediately before and 90 minutes after the intravenous injection of 0.25 mg synthetic ACTH.
- Interpretation:
  - ICS and hypoadrenocorticism: cortisol basal level normal or low normal (1-6 μg/dl / 27-160 nmol/l) only mild or no increase after stimulation.
  - SCS: variable, but sometimes high cortisol basal level (10 μg/dl / 270 nmol/l) or more than 3 times higher than basal level after stimulation and reaching > 20 μg/dl (550 nmol/l).

**Determination of Urinary Cortisol Creatinine Ratio (UCCR).**
- This test is a good first screening test in case of low suspicion and to eliminate an hypothesis.
- It is also easier to conduct, as the samples are taken “at home” by the owner.
- UCCR has a high negative predictive value (NPV) for SCS:
  - If the ratio is low, it is very unlikely that the dog is affected.
  - The hypothesis may then be excluded.
  - The positive predictive value (PPV) is lower. A high UCCR may be due to endogenous cortisol secretion due to stress of the animal. To increase this PPV, tell the owner to try to avoid any stress of the dog.
  - If the ratio is high, to confirm the diagnosis, a dexamethasone suppression test should then be performed.
- UCCR has also a high negative predictive value (NPV) for ICS:
  - If the ratio is high, it is very unlikely that the dog is affected.
  - The hypothesis may then be excluded.
  - If the ratio is low, to confirm the diagnosis, the ACTH stimulation test should then be performed.
  - The ratio is derived by dividing the urine cortisol level in nmol/l by the creatinine level in nmol/l. Reported reference ranges vary between laboratories, with a normal value ranging from < 10 x10^-6 to < 60 x10^-6 and so your individual laboratory should be consulted for interpretation.
**Low Dose Dexamethasone Suppression Test (LDDST).**

- **Test principle:** cortisol determination in the blood before and some time after the administration of a low dose of exogenous corticosteroids. In healthy dogs the negative feedback mechanism causes a decrease in ACTH secretion and thus a suppression of cortisol production (<1μg/dl or 27nmol/l) after 3 or 4 hours and 8 hours.

- **Indication:** suspicion of spontaneous hyperadrenocorticism (SCS).

- **Procedure:** blood is taken in the morning immediately before and, 3-4 and 8 hours after the i.v. injection of 0.01 mg/kg dexamethasone.

- **Interpretation:** in dogs with hyperadrenocorticism, levels either remain high or are suppressed after 4 hours and then increased again after 8 hours.
  - If the value at 4 hours is at least 50% lower than the basal one and increases again after 8 hours, PDH is very likely.
  - If levels remain high, it indicates a SCS and more likely an adrenal tumour but it may be a dexamethasone-resistant pituitary tumour (often macroadenoma of the pars intermedia which is under dopamine control). Then a HDDST or ACTH measurement will permit differentiation between PDH and ADH.

**High Dose Dexamethasone Suppression Test (HDDST).**

- **Test principle:** cortisol determination in the blood immediately before and after the administration of a high dose of dexamethasone. The ACTH production in dogs with PDH can be suppressed by high doses of a potent exogenous corticosteroids, leading to a decreased cortisol production. In adrenal adenomas or adenocarcinomas the autonomous hypersecretion of cortisol is not influenced by high doses of exogenous corticosteroids.

- **Indication:** suspicion of adrenal hyperadrenocorticism (ADH).

- **Performance:** blood is taken in the morning immediately before and 4 hours and 8 hours after the intravenous injection of 0.1 mg/kg dexamethasone.

- **Interpretation:**
  - If no suppression, ADH is very likely.
  - Suppression of 50% or more from basal cortisol at sample 2 and 3 may indicate a pituitary tumour (PDH).
  - Definitive confirmation of the diagnosis can be achieved by ACTH measurement, which will be low in adrenal tumors, or by ultrasonography of the adrenal glands and computertomography of the pituitary.
Part 2

Iatrogenic and spontaneous Cushing’s syndromes (ICS and SCS) • 8

Always consider haematology and blood chemistry tests first, in the differential diagnosis of endocrine disorders.

FIRST SCREENING HORMONAL TESTS

If, high suspicion of ICS due to history, or, no precise idea if ICS or SCS:

ACTH stimulation test, then diagnosis of ICS or investigation of SCS.

If, no precise idea if ICS or SCS and wish of surely eliminating ICS or SCS:

Determination of UCCR, then investigation of ICS or SCS.

SECOND STEP HORMONAL TESTS

If, hypothesis of ICS excluded, and, suspicion of SCS due to history, clinical signs and first screening laboratory tests:

To exclude the hypothesis (low suspicion), choose a test with a high NPV:

Urinary Cortisol Creatinine Ratio (UCCR) alone.

To confirm the hypothesis (high suspicion), choose a test with a high PPV:

Low Dose Dexamethasone Suppression Test (LDDST).

Urinary Cortisol Creatinine Ratio (UCCR) with High Dose Dexamethasone Suppression Test (HDDST) orally.

To precisely differentiate between PDH and ADH:

study the results of low dose dexamethasone suppression test (LDDST), and/or High Dose Dexamethasone Suppression Test (HDDST) I.V., and/or measurement of endogenous ACTH, and/or diagnostic imaging.

Measurement of endogenous ACTH.

Very useful but no routine laboratory test.

Ask a specialist how and where to perform this test.

The endogenous ACTH levels allow the differentiation between PDH and ADH:

> 120 pg/ml: PDH
< 30 pg/ml: ADH

Summary: diagnosis of Cushing’s Syndrome (CS) iatrogenic (ICS) and spontaneous (SCS).

HAEMATOLOGY / BLOOD CHEMISTRY

UCCR with HDDST.

Indication: suspicion of spontaneous hyperadrenocorticism.

This test is done on 3 consecutive days and should be performed at home since the stress of hospitalisation can influence the results.

Test principle: cortisol determination in urine, twice before and once after the administration of a high dose of exogenous corticosteroids.

Performance: the first (fasting!) urine in the morning is collected at day 1 and 2 and kept cool. After the second sample is taken, 0.1 mg/kg dexamethasone is given orally every 8 hours. On day 3 the first morning urine is collected and the samples mailed to the laboratory to determine the Cortisol/Creatinine (C/C) ratio.

This procedure also gives information about the origin of hyperadrenocorticism, pituitary or adrenal (PDH or ADH) because it combines a test for basal adrenocortical function and a dynamic test for the differentiation between primary and secondary hyperadrenocorticism.

Interpretation (the values given below are for illustration – be sure to obtain normal values from your laboratory):

Day 1 and 2

Hyperadrenocorticism: C/C ratio > 25 x10^-6
Normadrenocorticism: C/C ratio <15 x10^-6
Questionable: C/C ratio 15-25 x10^-6

Day 3 (in addition to increased C/C ratio on samples 1 and 2)

ADH or dexamethasone resistant PDH: C/C ratio of sample 3, less than 50% lower than mean of samples 1 and 2.

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ADH or dexamethasone resistant PDH: C/C ratio of sample 3, less than 50% lower than mean of samples 1 and 2.

PDH: C/C ratio of sample 3, more than 50% lower than mean of samples 1 and 2.
**Schematic representation of the diagnostic possibilities depending on hypotheses and chosen tests.**

**Legends**

- **Affections:**
  - N = Normal dog
  - CS = Cushing’s Syndrome (Hypercorticism)
  - ICS = Iatrogenic Cushing’s Syndrome
  - SCS = Spontaneous Cushing’s Syndrome
  - PDH = Pituitary Dependant Hyperadrenocorticism
  - ADH = Adrenal Dependant Hyperadrenocorticism

- **Tests:**
  - LDDST = Low Dose Dexamethasone Suppression Test
  - HDDST = High Dose Dexamethasone Suppression Test
  - UCCR = Determination of Urinary Cortisol Creatinine Ratio
  - ACTH-S-T = ACTH Stimulation Test

**ACTH-S-T, LDDST and HDDST**

![Diagram showing diagnostic possibilities for ACTH-S-T, LDDST, and HDDST with timelines and test outcomes.](image-url)
ACTH-S-T, HDDST and LDDST

ICS
- CS ? / N ?

ACTH-S-T

SCS (ADH) ? / N ?

HDDST

ADH
- T0 T0+1h30
- T0 T0+4h T0+8h

PDH ? N ?
- T0 T0+1h30
- T0 T0+4h T0+8h

LDDST

N
- T0 T0+4h T0+8h

PDH
- T0 T0+4h T0+8h

UCCR with HDDST orally

Do not forget the use of UCCR + HDDST orally which provides useful information and is easier to conduct (no need for hospitalisation) compared to other tests using blood samples.
ACTH-S-T, and UCCR alone

ICS

SCS ? / N ?

ACTH-S-T

SCS ? N ?

UCCR alone

N

SCE

D1

Other tests to differentiate SCS / N

UCCR as a first screening test

no SCS ? or no ICS ?

UCCR

SCS

ICS

D1

Other tests to differentiate N / PDH / ADH

ACTH-S-T to differentiate N / ICS

Clinical Handbook on Canine Dermatology
Treatment

PDH

**Hypophysectomy.**
- This therapeutic procedure is *directed against the primary lesion in PDH* and leads to the *complete destruction of the pituitary gland*.
- Today, computer tomography or MRI (Magnetic Resonance Imaging) is recommended before surgery to allow accurate determination of the size and localisation of the gland.
- The trans-sphenoidal microsurgical hypophysectomy technique should only be performed by an *experienced surgeon*.
- In PDH, surgery shows a lower recurrence rate than the conservative (medical) therapy.
- *Diabetes insipidus*, permanent central or secondary *hypothyroidism* and *keratoconjunctivitis sicca* are the most common *sequelae* and must be controlled periodically.
- A lifelong L-thyroxine replacement *therapy* is mandatory.

**Chemical destruction of the adrenal cortices with o,p’DDD.**
- There are two main protocols – one aims to selectively destroy the adrenal cortex whilst sparing the mineralocorticoid-producing *zona glomerulosa*. Complete destruction of the cortex does not result, and the therapy is maintained using interval dosing. The second seeks to non-selectively destroy the three layers of the cortex, and life-long supplementation is required. In neither case is there any effect on the pituitary pathology, and in fact any pituitary adenoma may actually enlarge due to lack of negative feedback.
- **Selective destruction:**
  - Induction phase: 25 mg/kg SID for 7-10 days.
  - Maintenance dosage: 25 mg/kg SID once every 7-10 days.
  - The substitution with low doses of prednisolone (0.25 mg/kg) on the days of o,p´DDD application helps to prevent the risk of hypoadrenocorticism with addisonian crisis which is the most common side effect of this therapy.
  - **ACTH stimulation tests are recommended** at the end of the induction phase and periodically during the maintenance phase. Ideally, the resting level should be between 1.0 and 3 μg/dl (27-80 nmol/l), and the stimulated level should be between 3.0 and 5.0 μg/dl (80-135 nmol/l).
  - The *treatment regime needs to be adapted individually* due to variations in the response to therapy. For this purpose the owner needs to observe abnormalities in his dog’s normal behaviour, particularly during the first two or three weeks of therapy.
  - With any medical treatment, deficiency of glucocorticoids (lethargy, anorexia) or mineralocorticoids (Addisonian crisis) are possible complications.
  - If PU/PD is observed and persists after more than 10 days of therapy the daily dosage of o,p´DDD may still be too low to achieve a therapeutic effect. Normal consumption of water is about 20 ml/kg/d.
  - Signs of an Addisonian crisis include apathy, weakness, vomiting, trembling, diarrhoea, collapse, elevated numbers of lymphocytes, low glucose and elevated urea, creatinine and total protein serum levels, serum Na:K ratio less than 24:1 (normal 33:1), ECG: spiked T wave, flat P wave and prolonged Q-T interval.
  - **Non-selective destruction:**
    - The following *treatment schedule* is *more aggressive* and aims at the complete quick destruction of both adrenal cortices and lifetime substitution therapy for the iatrogenic adrenocortical insufficiency:
    - 50-75 mg/kg o,p´DDD daily for 25 days, divided in 3-4 doses, with food.
    - On the third day supplementation begins:
      - Cortisone, 2 mg/kg daily.
      - Fludrocortisone, 0,0125 mg/kg daily.
      - Sodium chloride, 0,1 mg/kg daily.
      - All doses are divided into 2-3 administrations.
      - After 25-30 days, the cortisone is reduced to 0,5-1 mg/kg daily and fludrocortisone and/or salt are adjusted to the needs depending on the results of measurements of Na and K plasma levels. This regimen has a lower incidence of relapses (30%) compared with the “classic” schedule. For all types of therapy, good owner compliance is mandatory. He/she must be well informed about the pathogenesis of the disease, possible side effects of the treatment, re-examinations and costs. Written instructions are highly recommended.
      - Since the primary lesion in the pituitary gland is not affected by these therapies, the relapse rate is high (more than 50%) and the risk of tumour growth is not reduced.
**Inhibition of cortisol production.**

- **Trilostane:**
  - Trilostane competitively inhibits the enzyme 3-beta-hydroxysteroid-dehydrogenase which processes the conversion of pregnenolone to progesterone, reducing steroid hormone levels.
  - This is the first choice for treatment of PDH and the only licensed drug in Europe and the USA.
  - Published data shows that response rates are as good or better than those with o,p’DDD, with mean survival rates of 930 days.
  - Unlike o,p’DDD, trilostane is usually given daily for life, although some 5-10% of cases appear to be “cured” after 1-2 years, and therapy can be withdrawn.
  - Various treatment protocols with trilostane exist. At the present time a twice daily administration is recommended.
  - The dose necessary to effect control varies widely, and on a per/kg basis is much higher in small dogs than large breeds.
  - The dosage of 6 mg/kg/day, divided twice daily is a usual starting dose which may result in a resolution of clinical signs and routine laboratory abnormalities.
  - Some dogs may need higher dosages to achieve good response.
  - Monitoring should consist of: clinical evaluation, assessment of water intake, and ACTH stimulation tests should be undertaken at 7, 28 and 56 days to assess the response. Similarly to treatment with o,p’DDD, well-controlled animals have a basal cortisol of 1-3 μg/dl (< 27-80 nmol/l) which stimulates to around 3-5 μg/dl (80-135 nmol/l). A resting value of < 1 μg/dl and/or lack of adequate stimulation indicates excessive suppression, and the dosage should be reduced. As the duration of effect of trilostane is considerably less than 24 hours, the test should be performed 2-4 hours after administration at which time it will show peak activity.
  - Monitoring by measuring urinary cortisol/creatinine ratios is not reliable.

- **Ketoconazole:**
  - Ketoconazole inhibits the production of cortisol by the adrenal glands. This effect is reversible. The recommended dose is 10 mg/kg daily *per os*, divided in 2-3 doses.
  - As ketoconazole has multiple side effects, the patients must be re-examined in short intervals, particularly at the beginning of the therapy. Liver parameters need to be monitored closely (every 2-4 weeks) due to the hepatotoxicity of this drug.
  - The first ACTH stimulation test for the control of cortisol production by the adrenal glands is recommended after 3-4 weeks of therapy and then depending on the course.
  - Ketoconazole is less effective than trilostane or o,p´DDD.

**ADH**

- The *therapy of choice* for ADH is the *adrenalectomy of the affected gland*. It is imperative to supplement with both glucocorticoids and mineralocorticoids during and immediately post surgery, and for up to 10 days afterwards on a gradually reducing dose.

- If this is not possible or the tumour is inoperable, o,p’DDD may be used at the same therapeutic schedule as described for PDH (50-75 mg/kg daily for 25 days initially and then for at least 3 months once weekly).

- After this time, ultrasonography and X rays are recommended to *re-evaluate the tumour size* and decide about the continuation of the therapy.

- So long as the tumor is not malignant, and there bare no secondaries, this approach leads to a complete and permanent cure.

**Iatrogenic hypercorticisim (ICS)**

- **Gradual corticosteroid withdrawal.**
  - If the withdrawal is too brisk, a deficiency of either glucocorticoids and/or mineralocorticoids may result where there has been long-term suppression due to exogenous steroid administration. Short-term supplementation may be necessary, and will certainly be required if the patient is subjected to any stressful situations, such as even minor surgery.

**Summary**

<table>
<thead>
<tr>
<th>PDH</th>
<th>ADH</th>
<th>Iatrogenic hypercorticisim (ICS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Hypophysectomy</td>
<td>• Unilateral adrenalectomy</td>
<td>• Corticosteroid withdrawal</td>
</tr>
<tr>
<td>• Radiation therapy of pituitary gland or conservative therapy (o,p’DDD, trilostane or ketoconazole)</td>
<td>• Conservative therapy with o,p’DDD</td>
<td></td>
</tr>
</tbody>
</table>

Clinical Handbook on Canine Dermatology
**Prognosis**

- **Iatrogenic hypercorticism (ICS).**
  - Good in most cases, so long as they are carefully managed.

- **Pituitary-dependent hyperadrenocorticism (PDH).**
  - Very variable response to therapy.

- **Hyperadrenocorticism caused by adrenocortical tumours (ADH).**
  - Excellent if unilateral tumour was removed and no metastasis have occurred.
  - Guarded to poor in other cases.

**Follow-up**

- **ICS.**
  - Side effects of complete withdrawal of corticosteroid therapy such as glucocorticoid insufficiency or an Addisonian crisis are uncommon, but the owner and clinician must be alert for the warning signs.
  - It is still helpful to re-examine the animal and to investigate primary diseases camouflaged by the cortisone-therapy:
    - usually allergic skin diseases,
    - ectoparasites infestations...

  - A thorough workup will be required after cessation of the corticosteroid therapy:
    - secondary cutaneous infections (bacteria and/or yeast), may be more evident upon withdrawal of the corticosteroids,
    - “steroid diabetes” is uncommon

- **SCS.**
  - Extensive monitoring of each case is mandatory.

**Conclusions**

- **Iatrogenic hypercorticism** is frequently observed due to abuse of corticosteroids. In most cases the diagnosis and treatment are uncomplicated. But the diagnosis of the disease for which the corticosteroid therapy was instituted in the first place, such as allergic skin disease, must be taken up.

- The spontaneous types of Cushing’s syndrome are less common, are frequently overlooked particularly in early stages and need extensive tests to confirm the diagnosis. Several types of treatment exist depending on the type of disease (pituitary dependent or caused by adrenocortical tumours) and depending on the individual animal.

- The outcome depends often also on an early enough diagnosis as well as on the patience of the owner of the animal and her/his ability to co-operate. Many dogs’ lives can be prolonged or saved.
Hypothyroidism

Introduction / Definition

- A disease syndrome caused by insufficient production of thyroid hormone and leading to a wide variety of clinical signs. Classically, there is lethargy, weight gain and alopecia. Secondary skin infection may occur.
- Hypothyroidism in dogs is rare. Amongst the endocrine disorders, Cushing’s syndrome is much more common.
- “False positive” laboratory test results are common and may reflect euthyroid sick syndrome.

Aetiology / Pathogenesis

- **Aetiology.**
  - Primary hypothyroidism. Nearly all cases (over 90%) are caused by naturally occurring disease. The principal causes are lymphocytic thyroiditis, an immune-mediated disease associated with infiltration into the thyroid gland of lymphocytes, plasma cells and macrophages, and idiopathic thyroid necrosis and atrophy.
  - Secondary hypothyroidism is a rare cause of hypothyroidism. It is chiefly associated with pituitary neoplasia and pituitary dwarfism, leading to a deficiency of thyroid stimulating hormone (TSH) and consequent lack of secretion of hormone by the thyroid gland. Suppression of TSH can also be caused by glucocorticoid administration or by thyroid supplementation of euthyroid dogs.

- **Pathogenesis.**
  - Overall effects. The clinical signs of hypothyroidism are a consequence of the lack of circulating thyroid hormones and the effects of this on metabolism in different organs.
  - Effects on the skin. Thyroid hormone is required for initiation of anagen hair growth. In hypothyroidism, growing hairs proceed to telogen and are shed but lack of replacement by anagen hairs leads to alopecia. In body areas subjected to friction, hair shedding is promoted and hair loss becomes obvious at an earlier stage. An uncommon consequence of this disruption of the hair growth cycle is hypertrichosis.
  - Effects on protein and lipid synthesis commonly lead to dullness of the coat, brittle hairs and severe keratoseborrhoeic disorder (scaling and a dry or greasy skin).
Clinical signs

- Signs of hypothyroidism are diverse and very variable and affect a variety of organ systems. Those associated with organs other than the skin are listed in the table.

Some clinical signs of hypothyroidism affecting systems other than the skin:

<table>
<thead>
<tr>
<th>Metabolic</th>
<th>Cardiovascular</th>
<th>Haematological</th>
<th>Ocular</th>
<th>Gastrointestinal</th>
<th>Neuromuscular</th>
<th>Reproductive</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lethargy</td>
<td>Bradycardia</td>
<td>Anaemia</td>
<td>Corneal lipid deposition</td>
<td>Constipation</td>
<td>Seizure</td>
<td>Anoestru</td>
</tr>
<tr>
<td>Mental depression</td>
<td>Cardiac arrhythmia</td>
<td>Coagulopathy</td>
<td>Corneal ulceration</td>
<td>Diarrhoea</td>
<td>Ataxia</td>
<td>Silent oestrus</td>
</tr>
<tr>
<td>Weight gain</td>
<td>Hyperlipidaemia</td>
<td>Hyperpigmentation</td>
<td>Uveitis</td>
<td></td>
<td>Circling</td>
<td>Prolonged oestral bleeding</td>
</tr>
<tr>
<td>Thermophilia</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Vestibular signs</td>
<td>Testicular atrophy</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Weakness</td>
<td>Decreased libido</td>
</tr>
</tbody>
</table>

- Cutaneous signs of hypothyroidism. Classically, there is a bilaterally symmetrical, non-pruritic alopecia focussed on the trunk and tending to first affect areas subjected to pressure or friction (flanks, perineum, tail, ventrum etc.) but which may become widespread. The coat is easily epilated and tends to be dry and brittle. Regrowth either does not occur or is slow after clipping. Sometimes lack of hair shedding leads to marked thickening of the coat (hypertrichosis) and the retained hairs may become bleached, particularly at the tips of the hairs.

- Commonly there is scaling and the skin surface may be dry or greasy (seborrhoea); hyperpigmentation may be present. The skin may feel cool and may show myxoedema. Predisposition to bruising, poor healing of skin wounds and excessive scarring may also occur.

- Cutaneous infection.
  - Dogs with hypothyroidism are more susceptible to skin infections: regional or generalised superficial or deep pyoderma, bacterial and/or yeast overgrowths (*Staphylococcus pseudintermedius* and/or *Malassezia pachydermatis*) and otitis externa.
  - In the presence of skin infection or seborrhoea, pruritus may be an important sign.
Diagnosis

History.
- Classically there is lethargy, dullness, unwillingness to exercise or poor exercise tolerance and a tendency to gain weight without an increase in calorific intake. Heat-seeking during cold weather may also have been observed. These signs are more clearly shown in adult dogs but can be insidious in onset and may only be noticed by observant owners.
- A dull coat, hair loss including a “rat tail”, seborrhea and recurrent skin infections which respond well to antimicrobial therapy may also be reported. Neurological, reproductive or gastrointestinal abnormalities as listed in the table may also be reported.
- Hair loss appears first and is most prominent in areas of skin subjected to traumatic stress, particularly friction. Such areas include the ventral trunk, the tail and perineal area, and pressure points including the flanks. Hair loss may be observed on the neck, in friction areas between the skin and the collar. In cases of extensive hair loss, the head and feet are sometimes spared.

Clinical elements.
- Signs can be very variable.
- Systemic signs such as lethargy, mental dullness, thermophilia, obesity and bradycardia are important indicators but any of the signs listed in the table may be observed.
- Cutaneous signs. Bilaterally symmetrical alopecia, possibly including “rat tail” points towards endocrine disease.
- Skin thickening (myxoedema), particularly of the face and forehead and associated with drooping of the eyelids and thickened facial skin folds may be apparent giving a “tragic facial expression”.
- Skin infection, including superficial and deep pyoderma, Malassezia dermatitis, and / or otitis externa may be present.

Major elements of history and clinical signs.
- Alopecia, dry and brittle coat.
- Recurrent skin infection.
- Dullness, lethargy, weight gain.

Differential diagnosis.
- This is very wide and should include: other endocrinopathies; follicular dystrophy/dysplasia; superficial and deep pyoderma; bacterial overgrowth; Malassezia dermatitis; and otitis externa.

Bilaterally symmetrical alopecia in a Labrit affected by hypothyroidism. Courtesy of: D. Pin
Tragic facial expression in a Labrit affected by hypothyroidism. Courtesy of: D. Pin
Complementary examinations.

- **Trichography.** Examination of hair pluck samples can be helpful. Hairs plucked from areas of hypotrichosis are mounted in liquid paraffin and examined microscopically. Ensure that the hairs are plucked firmly so that anagen hairs, which are less easily removed, are not left behind. The presence exclusively of telogen hairs suggests failure of anagen initiation and may point towards endocrinopathy. Broken hairs are indicative of pruritus, which may be present in hypothyroidism when there is secondary infection or as a consequence of dry or scaly skin.

- **Screening tests** are useful in animals with clinical signs consistent with hypothyroidism. All such tests must be related to normal values and in dynamic hormone tests to expected response values provided by the diagnostic laboratories carrying out the laboratory procedures.

  - **Routine blood tests** in hypothyroid dogs may show mild, non-regenerative anaemia (30 to 50% of cases) and hypercholesterolaemia (70% or more of cases). Raised levels of alanine aminotransferase, alkaline phosphatase and creatine kinase may be found. These are useful pointers but are not diagnostic.

  - **Serum total thyroxine** (total T\textsubscript{4}) measurement is a useful screening method. Levels are generally below normal in hypothyroidism. However, total T\textsubscript{4} levels normally fluctuate in healthy dogs, and are depressed by therapy with a variety of drugs, malnutrition and in non-thyroidal disease. Thus dogs with normal thyroid function may have low T\textsubscript{4} levels.

  - **Serum free thyroxine** (free T\textsubscript{4}) is less affected than total T\textsubscript{4} by factors such as non-thyroidal disease, malnutrition and drug therapy and is thus a better screening test of thyroid function but only when levels are measured by the equilibrium dialysis method. Less accurate methods have no advantage over measurement of total T\textsubscript{4}.

  - **Canine TSH assay.** Serum TSH levels are elevated in about 40% of dogs with primary hypothyroidism but may also be present in about 15% of euthyroid dogs. Thus TSH assay alone is not a reliable test. Assessment of both TSH and T\textsubscript{4} levels increases the diagnostic accuracy. Low total or free T\textsubscript{4} together with high TSH is strongly supportive of a diagnosis of hypothyroidism. As with all these assays, specific cut-off levels should be provided by the laboratory carrying out the assay.

  - **Dynamic thyroid function tests.** Both TSH and TRH (Thyrotropin Releasing Hormone) response tests are used.

    - **TSH response test.** This involves measurement of serum total T\textsubscript{4} prior to and six hours following intravenous administration of TSH or 50-100 μg recombinant human TSH). Post-TSH levels of T\textsubscript{4} below 20 nmol/L are diagnostic for hypothyroidism whilst levels above 30 nmol/L are normal, in assays from most laboratories. This test is generally accepted as the most reliable but as pharmaceutical grade TSH is not available, chemical grade product has to be used and there have been anecdotal reports of adverse reactions to above TSH. Thus authorisation, including owner’s consent, should be obtained before it is used. The TSH response test is especially useful in cases where there is non-thyroidal disease or where drugs affecting thyroid function tests are being used. In such cases administration of 150 μg recombinant human TSH is recommended.
• **TRH response test.** This involves measurement of serum total T4 prior to and four hours following intravenous administration of TRH (0.2 to 0.1 mg/kg). A normal response is assessed as elevation of T4 to 1.5 times the baseline level (+50%). A normal response is indicative of euthyroid status but because some normal dogs may not respond, lack of response cannot be used to confirm hypothyroidism. Side effects which may be observed following TRH administration include depression, rapid breathing, vomiting, defecation and salivation. May be less reliable than the TSH response test.

• **Sources of errors.** A variety of factors can affect serum thyroid hormone levels and thus complicate diagnosis. These factors should be considered and their effects eliminated, if possible, when performing thyroid function tests.

  • **Breed effects.** Breeds known to have thyroid hormone levels at the bottom end or below generally accepted normal ranges include Alaskan sled dogs, Scottish deerhounds and greyhounds.

  • **Drug effects.** Glucocorticoids and sulphonamides may depress thyroid hormone levels. A variety of other drugs is suspected of having such effects. T4 administration tends to cause thyroid atrophy and after prolonged administration may affect thyroid function tests for periods of weeks to months; at least 8 weeks should be allowed after withdrawing T4 before carrying out such tests.

  • **Non-thyroidal illness.** Serum T3 and T4 are reduced by illness and severe malnutrition but free T4 measured by equilibrium dialysis may remain unaffected. Serum TSH levels may also be elevated in up to about 30% of those dogs. Thus tests should be delayed until recovery from the illness is achieved. The TSH response test is less affected by non-thyroidal illness and should be employed to confirm hypothyroidism when the results of other tests are not conclusive.

  • **Autoimmune thyroiditis.** This condition may occur in 50% of cases of hypothyroidism. Autoantibodies to thyroid hormone can affect assays and may raise or lower the values given by different assay procedures. However, free T4 assays using equilibrium dialysis are not affected by autoantibodies to T4.

### Treatment

**Topical treatment.**

- In cases of pyoderma, bacterial overgrowth and *Malassezia* dermatitis, treatment with antimicrobial shampoos should be considered in conjunction with systemic therapy, where appropriate (refer to the corresponding monographs). Where there is seborrhoea, the use of antiseborrhoeic shampoos and with dry skin, moisturisers, will be beneficial.

**Systemic treatment.**

- **Treatment with T4** will generally be effective; both primary and secondary hypothyroidism are treated in the same way. Doses of 0.01-0.04 mg/kg (10-40 μg/kg) daily are given once daily or divided at 12-hour intervals. Improved demeanour should occur within 4 weeks but resolution of skin problems may take as long as six months. Treatment should be continued for at least three months to determine whether therapy is likely to be effective. Once a good response has been obtained, consideration should be given to reducing the frequency of administration, from twice to once daily, as well as the dose level. Treatment with T3 is seldom necessary and is less convenient; doses of 4-6 μg/kg every 8 hours are used.
• **Post-pill tests.** Testing of the level of $T_4$ achieved after administration of the therapeutic dose may be necessary in cases which fail to respond to therapy initially or those in which clinical signs recur. Following treatment, blood levels should be within the normal range in 4-6 hours. Higher peak levels will be expected with once daily therapy. In cases with poor responses in which 6-hour levels are adequate, levels should be reassessed at 24 hours. If levels at 24 hours are too low then consideration should be given to dosing twice daily and, where necessary, raising the dose. **Where the post-pill levels of $T_4$ are substantially above normal and there is a lack of clinical response, the diagnosis of hypothyroidism should be questioned.**

• **Dogs with other diseases** may need special consideration. Where there are other endocrinopathies, these will need to be treated as well. In **hypoadrenocorticism**, the adrenal insufficiency **should be effectively treated before administration of $T_4$**. In diabetes mellitus, dosage of insulin may need to be adjusted when $T_4$ therapy is instituted. In heart disease, dosage with $T_4$ should be started at a low level and gradually increased. A suggested regimen is to begin with $5 \mu g/kg$ every 12 hours and to increase the dose every two weeks by $5 \mu g/kg$ every 12 hours until a therapeutic dose is reached.

**Additional treatment.**

• **Systemic antibiotics.** Bactericidal antibiotics are advisable to treat any concurrent bacterial infection. Normal dose rates are used (see monographs on pyoderma) and the response is usually very good. Unless thyroid supplementation is instituted, recurrence of the pyoderma will occur. See monograph on complex approach pyoderma.

**Comments.**

• Concurrent topical therapy is an important aspect of the treatment whilst waiting for skin and coat problems to resolve.

**Prognosis**

⇒ In the great majority of cases, supplementation with $T_4$ is effective and leads to resolution of disease related to the hypothyroidism.
⇒ Therapy needs to be lifelong.

**Follow-up**

⇒ Post-pill testing and adjustment of the dose of thyroid supplementation may be required should the clinical signs recur.

**Conclusions**

⇒ Hypothyroidism is a disease that may be difficult to diagnose in some cases because of the very wide range of clinical signs and problems with thyroid hormone assessment.
⇒ Once a diagnosis has been made, recovery is normally complete and the dog can have a good quality of life provided that appropriate supplementation is maintained.
Otitis externa

This monograph is not an exhaustive list of clinical signs and causes of otitis externa, but a chronological and clinical method of investigation to allow management of otitis externa cases as specific dermatological problems. Even if it is generally not necessary, it is recommended that the patient be sedated to facilitate better examination and sampling quality.

**HISTORY AND CLINICAL EXAMINATION**

1/ Specific history of otitis

- Must be registered in the sections “otitis” and “response to therapy”.
- These elements help to categorize each case in terms of severity.
- Local pathology: degree of pruritus (relief when the ears are massaged...), pain (whining...), neurological involvement.
- Affected pets in contact (in case of ear mites).
- Duration of otitis externa.
- Number of relapses.
- Previous treatments: type of therapy, products used, duration, efficacy / response.
- Grooming and cleaning routine: method, products used.

2/ Preliminary clinical examination

- Head and ear position (head tilt, ear usually held low if painful, or neurological problem if there is an otitis media).
- Gentle palpation of the ear canal (EC), note: level of pruritus, pain, hyperplasia, amount of discharge.
- Pinnae and EC examination, note:
  - Primary and secondary lesions, hyperplasia...
  - Discharge or pus - amount and nature.
  - Odour.
- Look for an audito-podal reflex by stimulating the external orifice of the ear canal (suggestive sign of ear mites).
- Assess the hearing.

Clinical examination : ear canal palpation and area to stimulate to trigger an auditopodal reflex. Courtesy of: D. Pin
OTOSCOPY AND SAMPLING

3/Sampling for mycology and/or bacteriology
• Must be done with a sterile swab.

4/Exudate sampling for direct microscopic examination
• Must be done with a swab or a curette.
• Parasitic investigation (eggs, larva and adults):
  • *Otodectes cynotis* in ear mite infestation (1/10 of otitis cases),
  • *Demodex canis* or *cati* (rare).
• Refer to the slide preparation and examination in the “sarcoptic mange” monograph.

5/Exudate or pus sampling for cytological microscopic examination after fast-staining
• Look for perpetuating microbial factors: yeasts (mainly *Malassezia pachydermatis*), cocci (*Staphylococcus pseudintermedius* in particular), rods (*Pseudomonas aeruginosa* most frequently)...
• Presence of neutrophils indicating the presence of pus in suppurative otitis externa.
• Refer to the procedures in the “Pyoderma” and “Malassezia dermatitis” monographs.

6/Otoscopy / Video-otoscopy
• General aspect of the ear canal lining including assessment of amount and quality of discharge, presence of ulcers and/or nodular lesions.
• Appearance of the tympanic membrane.
• Look for foreign body
• Video-otoscopy has same use as otoscopy but with better accuracy (requires a good quality video-otoscope). It can be very demonstrative for the owner and allows to take pictures and assess the healing.
• Clinical method of investigation
7/Ear canal cleansing and further otoscopy / video-otoscopy

- If the ear canal is severely occluded.
- When the tympanic membrane has not been visualized, in particular in severe and/or chronic cases where an otitis media may be suspected.
- It allows a thorough assessment of the local disease; ear canal inside layer, presence of nodular and/or ulcerative lesions, presence of foreign body and observation of the tympanic membrane - on which any lesions would suggest otitis media (however, absence of tympanic membrane lesions does not exclude a suspicion of otitis media).

8/Exploration and management of an otitis media

- Major suspicion in cases of chronic otitis externa, particularly when suppurative, with or without rupture of the tympanic membrane.
- Diagnosis:
  - otitis media likely when the tympanic membrane is ruptured,
  - CT-Scan imaging: very useful diagnosis method,
  - myringotomy and tympanic bulla aspect • open mouth tympanic bulla radiographs (older method, much less accurate than CT-Scan).
- Management:
  - when the tympanic membrane is ruptured or after a myringotomy: sampling for culture and sensitivity (+ mycology and sensitivity to antifungal agents),
  - refer to “therapy”.

Suppurative otitis before cleansing. Courtesy of: D. Pin

Suppurative otitis after cleansing. Courtesy of: D. Pin

Cleansing: bulb flushing. Courtesy of: D. N. Carlotti
9/Use of samples from the ear canal

• Send to a laboratory for bacteriology / mycology in cases of:
  • otitis media,
  • chronic suppurative otitis,
  • severe recurrent otitis.
• Sensitivity must include antibiotics usually effective on the bacteria found in otitis:
  • important if a systemic treatment is prescribed,
  • provides useful information but is less determinant regarding choice of topical treatment (local antibiotic concentrations are far in excess of the systemic concentrations assayed in the in vitro sensitivity determination).

10/Therapy: in clinic and by the owner

Foreign Body Otitis
• Withdrawal of the foreign body.
• Secondary infection treatment.

Parasitic Erythematous-Ceruminous Otitis externa (ear mites)
• Topical acaracidal therapy.
• Systemic acaracidal therapy (refer to systemic therapy of sarcoptic mange).
• Concomitant antiseptic flushing solution can be used when Malassezia sp. is also present.

Infectious Erythematous-Ceruminous or Suppurative Otitis externa (yeast and/or bacteria)
• Topical therapy with a preparation containing antibiotic, antifungal and anti-inflammatory agents, known to be effective.
• Prerequisite initial cleansing in the practice in presence of copious discharge and exudates: ear flushing using adapted devices, most often under general anaesthesia, with an antiseptic and cleaning solution.
• Prerequisite regular cleansing by the owner in presence of copious discharge and exudate:
  • with an antiseptic cleaning solution applied generously,
  • gentle massage of the ear canal,
  • wipe off the secretions and excess product that appear at the external orifice of the ear canal with a paper towel,
  • repeat the procedure several times if necessary,
  • the therapeutic preparation should preferably be applied a few minutes later.
• Inflammation can persist once the infection is controlled:
  • preventive use of an antiseptic ear cleanser,
  • use of a therapeutic preparation containing corticosteroids in case of severe inflammation,
  • the basic therapy is the treatment or control of the associated disease.
Otitis media

- Topical cleansing with an antiseptic solution, under sedation, when the tympanic membrane is ruptured or after myringotomy.
- Topical therapy with a preparation containing antibiotic, antifungal and anti-inflammatory (corticosteroid) agents.
- Antibiotic systemic therapy at high dosage.

Antimicrobial topical products

- An antiseptic topical cleanser can be used in the first intention and can be excellent for maintenance and long-term prevention. The addition of sugars (glycotechnology) helps in reducing microbial adherence.
- A therapeutic product specific for acute otitis externa must include a broad spectrum antibiotic, an antifungal agent (effective against Malassezia) and a steroidal anti-inflammatory agent:
  - treat by courses of 1 to 2 weeks, and at least one week more after healing and/or microbial clearance (no microbial agents revealed on cytology),
  - avoid prolonged administration, treat when infection is established until recovery and elimination of the infection (repeat regularly cytological controls).

Systemic therapy

- Antibiotic and/or antifungal agents: useful in case of otitis media (at a high dosage) in order to increase the local level of the active ingredients delivered by the topical therapy (which remains vital).
- Corticosteroids: sometimes useful (and in some severe cases of otitis externa) to reduce thickening of the integument and narrowing of the ear canal in chronic otitis externa and therefore to avoid surgery (which would not solve the problem on the long term). Exploration and control of the associated dermatosis is the next critical step.
**Conclusions**

- Prevalence of otitis externa

<table>
<thead>
<tr>
<th>Type</th>
<th>Average prevalence</th>
<th>Unilateral (U) or Bilateral (B)</th>
<th>Primary causes</th>
<th>Perpetuating factors to control</th>
</tr>
</thead>
<tbody>
<tr>
<td>FBO (Otitis due to a Foreign Body)</td>
<td>ε</td>
<td>U</td>
<td>Foreign Body (FB)</td>
<td>Possible secondary infection</td>
</tr>
<tr>
<td>PECO (Parasitic Erythematoceruminous Otitis)</td>
<td>1/10</td>
<td>B at 95%</td>
<td>Otodectes ++++, others ±</td>
<td>Possible secondary infection</td>
</tr>
<tr>
<td>NPECO / IECO (Non Parasitic / Infectious Erythematoceruminous Otitis)</td>
<td>2/3</td>
<td>B at 75%</td>
<td>Allergic dermatitis in 85% of cases: mainly atopic dermatitis</td>
<td>Mainly yeast (Malassezia pachydermatis ++++) and cocci (Staphylococcus pseudintemedius ++++, streptococci +, others ±)</td>
</tr>
<tr>
<td>SO (Suppurative Otitis)</td>
<td>1/4</td>
<td>U at 50%</td>
<td>ECO complication (refer to its causes) FB otitis complication</td>
<td>Bacteria and particularly rods (Pseudomonas aeruginosa +++, Proteus spp +, others ±)</td>
</tr>
</tbody>
</table>

**Adult dog**

Infectious Erythematoceruminous Otitis externa are commonly found, and can degenerate into suppurative otitis externa in the most chronic (and/or poorly controlled) cases. Otitis externa, especially Erythematoceruminous Otitis externa, is generally found in adult dogs as they are quasi systematically associated with a general dermatosis (in particular an allergic dermatitis) which concern only adult dogs.

**Young dog**

The young dog is mainly affected by parasitic otitis externa, which may become secondarily infected, or less commonly, by foreign body otitis externa.

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**Clinical Handbook on Canine Dermatology**
Part.3

Applications and use of skin biopsies

Clinical Handbook on Canine Dermatology
Introduction

Skin biopsy followed by histopathological analysis is one of the most helpful procedures in the diagnosis of dermatological problems.

This chapter is entirely devoted to the description of skin biopsies, their indications and techniques. Its goal is to demonstrate the major indications for this procedure and to give some basic rules that will enable optimum information to be derived from the histological examination.

Skin biopsies are particularly helpful in the diagnosis of the rarer dermatoses, for which the clinical approach is more difficult. They are not necessarily indicated for the more common dermatoses for which the clinical approach is a prerequisite, but biopsies are never useless.

The role of the clinician and the role of the pathologist

The analysis of skin biopsies requires close teamwork between the clinician and the pathologist.

The role of the clinician is very important as he/she is responsible for the first step: the choice of the area to biopsy.

The role of the pathologist is of course equally important and critical, as he/she has to interpret, without seeing the animal, a histological picture at a fixed time point in what is an evolving pathological process. It is usually a combination of images, observed on histopathological sections from one or several specimens that enables the pathologist to distinguish one (or several) differential(s) with more or less certainty.

When skin biopsies are performed:
- The first objective may be to reach the diagnosis with the highest positive predictive value (PPV).
- Of equal importance, the second objective could be to rule out one (or several) differentials.
  In this case, the goal is to derive the highest negative predictive value (NPV).
- These predictive values depend as much on the pathologist as the clinician and in particular, the PPV of the former and the NPV of the latter.

The clinician is responsible for the type, the quality and the number of samples submitted to the pathologist who must have particular expertise in dermatopathology, and the submission must be accompanied by a complete history, a report of the dermatological examination, and a map of the lesions with the biopsy sites indicated.

The pathologist’s interpretation and conclusions may be useless to the clinician unless these few basic rules are followed.

When to biopsy?

General information / Client information

Performing a skin biopsy is never useless: at best it may give a definitive diagnosis and at worst it will help by pointing the clinician in the right direction.

When this procedure is advised, the owner must be made aware that the results will not necessarily provide a precise diagnosis, but the information obtained will nonetheless be extremely useful in the case management. Also important is the fact that the cost/benefit ratio is high, and that the results can be obtained relatively quickly.

The skin biopsy should be scheduled for a specific appointment to ensure that it is performed optimally, and to emphasize the importance of the procedure to the owner.

Also an indication of the likely total cost must be given, emphasising the potential benefits relative to that cost.
For the most frequent dermatoses (groups 1 and 2), the clinical approach is essential. Nevertheless, for those in group 1 (specific infectious and parasitic dermatoses), the histopathology results are sometimes diagnostic and if not they may help rule out some differentials. For those in group 2 (most frequent primary dermatoses - allergies and endocrine disorders), histopathological examination is seldom diagnostic. However, histopathology may help the clinician in the diagnostic approach to such dermatoses as well as excluding the presence of secondary dermatoses (pyoderma in particular).

For all the other, less common (group 3) dermatoses, histopathology is essential to establish the final diagnosis. This is because they are often severe requiring long and expensive treatment which may have side effects, and thus a definitive diagnosis, obtainable only by skin biopsies, is essential.

Skin biopsies are thus particularly indicated:
- When a specific condition is suspected:
  - for which the analysis of skin biopsies is recognised to be diagnostic,
  - when one wishes to support one’s diagnosis for a group 1 or 2 dermatosis.
- When the lesions appear atypical, or have a sudden and dramatic onset.
- For any erosive dermatosis (e.g.: generalised ulcerative lesions).
- If there is any suspicion of neoplasia.
- When any dermatosis belonging to group 3 is suspected. In this case, histopathology almost always gives a diagnosis.
- When the performance of other tests have not enabled a diagnosis.
- When a treatment, apparently correctly followed, has not resolved the problem.
- When one wishes to confirm the clinical approach taken towards such dermatoses, in particular for the most frequent primary dermatoses (group 2). It may first of all confirm the absence or regression of secondary dermatoses (group 1b) and give an indication for possible further diagnostic tests helpful in the differential diagnosis of, for example, allergies or endocrine disorders.

In all cases, and more particularly for the group 1 and 2 dermatoses, the establishment of the diagnosis is the responsibility of the clinician, taking into account the history, the clinical examination and possible other additional tests.
What to biopsy?

**How many samples?**
- When it has been decided to undertake skin biopsies, with the owner’s agreement, there should be no hesitation in taking a number of samples: the cost is identical, the time required almost the same and the value of the information obtained is often directly proportional to the number of samples.
- Different areas are often in different stages of the disease process. The performance of numerous biopsies therefore gives a better “sampling” of the various phases of the pathological mechanism in question. This can be enormously helpful to the pathologist.
- The clinician will thus have to endeavour to sample different areas of different clinical aspects (ranging from the most severely affected areas to the normal).
- Taking five biopsies is an acceptable average.

**Nature of the samples.**
- Primary lesions of all types must be sampled first (papular, pustular, nodules, or even erythema...). As they result from the principal pathogenic processes, they will be more reliably diagnostic.
- As a rule, biopsy *all* suspect lesions, particularly when the primary lesions are not easily identifiable. A crust examination, for example, will sometimes give extremely useful information.

How to biopsy?

**Material.**
- Ready availability of all the necessary instruments and materials will facilitate high standards and ensure that biopsies can be undertaken whenever there is any indication.
- The goal is not to work under aseptic but clean conditions. However, sterile instruments and materials must be used.
- A small surgery box should be reserved exclusively for skin biopsies and contain only the necessary instruments.
- The materials necessary are not highly specialised:
  - the appropriate amount of a local anaesthetic (2% lidocaïne), enough for 1 to 2 ml for each biopsy site, sterile syringes and needles; a general anaesthetic should also be available,
  - suture, absorbable or non-absorbable,
  - a pair of sterile gloves,
  - absorbent paper towel on which to dab the samples,
  - a small cold sterilisation tray containing the 6mm sterile punches, in good condition,
  - a small sterile surgical box containing: a new blade, a scalpel handle, two pairs of thin scissors, straight and curved, dissecting forceps, a small clamp, a very small pair of eye forceps, gauze pads, suture material (rat-toothed forceps and needle holder),
  - 10% formalin-filled vials,
  - and the necessary labels and documents for appropriate submission.
**Patient preparation.**
- The biopsy sites must remain perfectly intact, as the skin surface material is often of great value to the pathologist.
  - avoid clipping (or plucking!) the hair before sampling, as important material could be removed
  - avoid preliminary scrubbing.
- Before anaesthesia it is preferable to circle with an indelible marker the sites that will be sampled.
- Note the sites on the lesion map and indicate precisely the nature of the samples (type of lesion, normal area immediately adjacent to such lesion...).

**Local anaesthesia.**
- Most of the time, skin biopsies may be performed under local anaesthesia with appropriate physical restraint:
  - a sub-cutaneous injection of 1 ml (or 2 ml) of 2% lidocaïne per site is sufficient,
  - gently fold the skin, without damaging the lesions,
  - the anaesthetic is injected under the biopsy site making sure that the product penetrates under the skin,
  - introduce the needle into the immediate periphery of the biopsy site and reposition the needle carefully, several times, depositing the anaesthetic in several places,
  - wait approximately 5 to 10 minutes before performing the biopsy.

**General anaesthesia.**
- General anaesthesia may be useful for fractious animals.
- It is generally necessary when sampling the feet (footpads, nails), face (periorbital, muzzle, nasal planum), ano-genital areas, the medial aspect of the pinnae or the tail.

**Punch biopsies.**
- This is the simplest and quickest technique.
- In general, a 6 mm punch provides good sampling material with minimal trauma.
- Hold the skin flat without stretching it.
- Apply the punch perpendicular to the skin surface.
- Exert sufficient pressure to the punch with the hand while “rolling” it with a continuous rotational movement between the thumb and the index finger.
- The pressure must be maintained until one feels the skin loosen as the blade of the punch penetrates the subcutaneous tissue.
- Pick up the sample very carefully by the under surface, taking care not to crush or even squeeze it:
  - the fragment sampled generally emerges when the punch is withdrawn, then the subcutaneous fat by which it is still attached just has to be cut,
  - if not, gently lift the sample by introducing closed forceps into the wound.
- The specimen may be picked up by the subcutaneous fat.
- In order to remove the blood, gently blot the distal surface of the sample on an absorbent paper towel.
- Then put it immediately into the fixative.
- The biopsy punch technique has the enormous advantage of allowing numerous samplings, generally under local anaesthesia, and is well accepted by the owner.
- However, it does not allow the inclusion of diseased and normal tissue within the same sample (except if the lesion is very small).
- If a comparison has to be made between the diseased skin and normal skin of the same region, an adjacent normal area has to be biopsied separately (refer to the chapter concerning the sites to biopsy).
**Elliptical wedge biopsies.**

- This biopsy technique is useful when the lesions to be sampled are large (nodules, neoplasia...) and/or deep (panniculitis) and/or very fragile (big pustules, bullae, vesicles...).
- This method is also used when surgical resection of the lesion is indicated (nodule, tumour...).
- However, in order to enable adequate fixation by the formalin, the specimen submitted for histopathologic analysis must not be too large; a representative piece will suffice.
- Another interesting feature of this method is that the section defines a long axis if one requires the specimen to be cut prior to bring embedded (see chapter concerning the precise selection of sites to biopsy).
- The dermis and epidermis must be sectioned simultaneously, firmly and continuously.
- Pinch and slightly elevate the specimen by the subcutaneous fat using a pair of fine eye forceps.
- Trim away the deep tissue with fine iris scissors.
- Gently blot the distal surface of the specimen on an absorbent paper towel in order to eliminate excess blood.
- Immediately place the sample into the fixative.
- This technique is theoretically the best, but presents certain disadvantages: it takes more time, generates larger wounds, does not facilitate multiple sampling, and ultimately is less acceptable to the owner.
- It is on the other hand the only technique allowing the inclusion of lesional tissue as well as normal tissue in the same sample (see chapter concerning the precise selection of sites to biopsy).

**Specimen conservation.**

- Place each sample in formalin (10% phosphate-buffered) immediately after removal, thus preventing desiccation and autolysis.
- Do not use other fixatives (such as Bouin’s fluid).
- Some specialist laboratories provide vials containing the correct fixative.
- All specimens may be placed in the same vial, but the total volume of fixative must be at least ten times greater than the total volume of the specimens.
- Distribution of multiple samples in different vials may be useful for sample identification: a vial, for example, for each site biopsied.

**Haemostasis, antisepsis and suture.**

- Do not worry about the bleeding until the samples are completely removed and placed in the fixation vials!
- Haemostasis is generally readily effected by simple compression. It may sometimes be more difficult in certain areas (facial, pedal, auricular).
- Dab gently with an antiseptic solution and suture the wounds when all biopsies have been taken. If a wound keeps bleeding, compress firmly with clean gauze before suturing. The use of haemostats is rarely necessary.
- For 6mm punch sites, a single interrupted suture is generally sufficient.
- For 4mm punch sites, sutures may not be necessary.
- For elliptical wedge biopsies, suture (preferably simple interrupted) appropriate to sample size.
Sample dispatch.

- The mailing must include the case history and clinical data (including a map of the lesions).
- It is important to indicate clearly the sites and the type of lesion biopsied on the map.
- Distributing the samples in multiple vials is also useful.
- The pathologist will then be able to identify easily the origin of the different biopsies submitted.
- The simplest procedure is to send a photocopy of the history, clinical records and lesion map filled in during the consultation (including accurate indication of the sites and nature of the lesions biopsied).

- It is also necessary to indicate which other tests have been undertaken, with results if available, and details of all treatment given together with any response.
- Skin biopsies should be submitted to a reliable, reputable laboratory, making sure a dermatopathologist will interpret the samples.

**Lesion map**

- = site and aspect of each sample must be precisely indicated on the lesion map.

**Clinical record**

- Owner:
- Patient's name:

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Clinical Handbook on Canine Dermatology
Practical guidelines

**General method.**
- Search for several lesions of the same type; choose the best specimen - not just the first one found.
- Care must be taken not to damage the lesion while sampling, especially primary lesions such as pustules, which can be very fragile.
- Ensure that all parts of the biopsy sample are placed in the fixation vial, e.g. crust. All material provided are (normally!) analysed by the laboratory and observed by the pathologist (a crust may carry useful information).
- Keep in mind that the pathologist will bisect the specimen through its long axis, symmetrically and perpendicularly to the skin surface (Fig 1).

![Fig1 Diagram]

**Captions for the diagrams**
- = patient skin
- = punch biopsy
- = elliptical wedge biopsy
- = area apparently healthy
- = extensive lesional area
- = lesion considered as significant
- = histological section
- = correct biopsy procedure
- = biopsy procedure incorrect
**Sampling of well-demarcated lesions.**

- Most of the time, a punch can be used.
  - When the biopsy is performed with a punch, the specimen is cylindrical and the section will be generally done randomly according to one of the diameters.
  - It is thus very important to accurately focus the punch on the chosen lesion (Fig2).
  - If the lesion is not centred, the bisection will probably miss the lesion and normal tissue will be processed (Fig3).
  - Furthermore, focusing the punch will considerably limit the risk of damaging the lesion that will then be preserved intact (a critical factor for enabling a diagnosis with a skin biopsy).

In the case of a larger lesion, sample it entirely by performing an **elliptical wedge biopsy**, keeping in mind that the long axis must pass through the lesion (Fig4). This technique is a prerequisite with fragile and/or transient lesions (bullae, vesicles, or even furuncles...).
**Sampling extensive lesional areas.**

- The biopsy must be performed within the lesion (Fig5).
- **With a punch,** it is preferable not to biopsy surrounding normal skin, for there is a great risk that such a sectioning such a biopsy may miss the lesion (Fig6).
  
  ![Fig5](image1)
  ![Fig6](image2)

- **It is better** to perform an **elliptical wedge biopsy with a scalpel,** the long axis will be chosen perpendicular to a tangent of the lesion (Fig7). Otherwise two or three punch biopsies are necessary.
  - The section, done along the long axis of the specimen, will then allow the processing of the lesional area with the adjacent cutaneous area.
  - If well-demarcated lesions considered significant are present within the extensive lesion, at least one of them should be included in the biopsy specimen, along its long axis.

![Fig7](image3)
Key points to keep in mind

- Histopathology is an important part of the diagnostic work up of the dermatological case.
- Skilled teamwork between the pathologist and the clinician is essential to obtain the best information from the skin biopsies.
- For this same reason, the choice of the pathologist to whom the samples will be sent is crucial.
- It is strongly recommended to have the skin biopsies examined by a veterinary dermatologist proficient in dermatohistopathology or by a veterinary pathologist specialising in dermatohistopathology.
- It should be made clear on the report which pathologist has examined the biopsies in order to facilitate any later discussions.
- However, the clinician should be able to rely on the pathologist and the resultant report if the few basic rules which were described are followed, i.e. if the biopsy samples are:
  1°) in sufficient quantity,
  2°) of suitable quality,
  3°) of relevant nature (i.e. taken from areas truly representative of the skin condition),
  4°) and accompanied by:
    • a well described history of the case,
    • concise description of the results of the dermatological examination (a detailed lesional map is an excellent method),
    • the results of other laboratory examinations and any therapeutic trials carried out,
    • the results of all previous treatments,
    • and possible differential diagnosis indicated from the history and clinical examination.
- Particularly, the biopsies must:
  • be multiple,
  • include all primary lesions, as well as secondary lesions,
  • as far as possible, include the whole lesion and part of the immediately adjacent area,
  • otherwise, the centre of the lesion, a peripheral area, an area bordering normal skin and even normal skin itself.
- The number of sites to biopsy may seem numerous, but:
  • first of all, there is usually no additional fee for several specimens,
  • then, the greater the number of samples, the better are the chances of finding diagnostic sections,
  • finally and most importantly, it is quite obvious that not every biopsy submitted along these lines will have the same diagnostic value. Nevertheless, when the likely diagnosis is unclear, it is better to provide samples that will not be useful, rather than failing to provide samples that could have been crucial in establishing the diagnosis!
  • Of course, with experience, or in the case of a classical condition with limited differentials, the selection of the sites to biopsy may be refined and the number of samples reduced.
  • It is also important to specify if one wishes a precise diagnostic confirmation of a strong clinical suspicion and/or if one wishes on the contrary to rule out such or such differential for which the clinical suspicion is weak, or even if one wishes to rule out a rare dermatosis (group 3).
  • Not having a precise idea (or any idea at all!) of the cause of the skin condition is important information which will still be of value to the pathologist!

Conclusions

- Well conducted histopathology is:
  • Seldom useless, even for the most frequent dermatoses (for which the clinical approach is essential), particularly for those of group 1. The skin biopsy results will at least allow the clinician to rule out certain diagnostic differentials (more particularly from the group 3).
  • Suggestive in many cases (for example for the group 2 dermatoses, for which once again, the clinical approach is essential).
  • Diagnostic for a great number of dermatoses (belonging in particular to group 3).

- All in all, histopathological analysis is as revealing by what it shows as by what it does not show!
Part.4

Elements of physiology and structure of healthy canine skin

Clinical Handbook on Canine Dermatology
Physiology of the dog's skin

The skin should be considered as a vital organ, providing whole body protection (for other organs and tissues): it has connections with other organs and with the interior of the body (usually via the mucosae) at the eyes, tympanic membrane, nose, mouth, and ano-genital area. The skin is also responsible for many physiological functions, with continuous renewal of the epidermis and of the haircoat via the follicular cycle.

General functions

**Enclosing barrier:**
The most important function of skin is to make possible an internal environment for all other organs by maintaining an effective barrier to the loss of water, electrolytes, and macromolecules.

**Environmental protection:**
A related function is the exclusion of external injurious agents, chemical, physical, and microbiologic, from entrance into the internal environment.

**Motion and shape:**
The flexibility, elasticity, and toughness of the skin allow motion and provide shape and form.

**Adnexa production:**
Skin produces important keratinised structures such as hair, claws, and the horny layer of the epidermis.

**Temperature regulation:**
Skin plays a role in the regulation of body temperature through its support of the hair coat and regulation of cutaneous blood supply.

**Storage:**
The skin is a reservoir of electrolytes, water, vitamins, fat, carbohydrates, proteins, and other materials.

**Indicator:**
The skin may be an important indicator of general health, internal disease, and the effects of substances applied topically or taken internally.

**Immunoregulation:**
Keratinocytes, Langerhans' cells, and lymphocytes together provide the skin with an immunosurveillance capability that effectively protects against the development of cutaneous neoplasms and persistent infections.

**Pigmentation:**
Processes in the skin (melanin formation, vascularity, and keratinisation) help determine the colour of the coat and skin. Pigmentation of the skin helps prevent damage from solar radiation.

**Antimicrobial action:**
The skin surface (superficial lipid film) and epidermis (antimicrobial peptides) have antibacterial and antifungal properties.

**Sensory perception:**
Skin is a primary sense organ for touch, pressure, pain, itch, heat, and cold.

**Secretion:**
Skin is a secretory organ by virtue of its apocrine (epitrichial), eccrine (atrichial), and sebaceous glands.

**Excretion:**
The skin functions in a limited way as an excretory organ.

**Vitamin D production:**
Vitamin D is produced in the skin through stimulation by solar radiation. Vitamin D3 is formed in the epidermis and hydroxylated in the liver and again in the kidney. It is important in the regulation of epidermal proliferation and differentiation.

Normal dog’s skin structure

Cutting of the dog’s skin showing the structure. Courtesy of: P. Jasmin
Epidemis renewal

Complete renewal of the epidermis in 3 weeks

- Natural elimination of invisible cornified dead cells
- Differentiation & migration
- Multiplication

Follicular cycle of the dog’s skin

Growth of a new hair under and beside the old one; the “dead” hair staying in the hair follicle. The “dead” hair is pushed out by the new one and falls to the ground. A complete mosaic shedding occurs 3 to 4 times a year, with hormonal & nycthemeral rhythm influences.

“dead” hair

“old” hair

“new” hair
Fundamental differences between canine and human skin

1. Cutaneous pH difference: up to 7.5 in dogs / 5.5 in human
2. Epidermal thickness: 3-5 layers in dogs / 10-15 in human
3. Quicker epidermal turnover in dogs: 20 days / 28 days in human
4. Follicular cyclical: in dogs / continuous in human
5. No exocrine sudoral glands in dogs (except in footpads and eyelids)