Equine sarcoids - part 1: clinical presentation and epidemiology

Equine sarcoïden – deel 1: klinisch voorkomen en epidemiologie

L. Bogaert, A. Martens, P. Depoorter, F. Gasthuys

Department of Surgery and Anesthesiology of Domestic Animals, Faculty of Veterinary Medicine, Ghent University, Salisburylaan 133, B-9820 Merelbeke, Belgium

lies.bogaert@UGent.be

ABSTRACT

Equine sarcoids are the most common skin tumors in horses and other equids. In their pathogenesis, the bovine papillomavirus (BPV) plays a major role. Many clinical manifestations have been described, ranging from small single lesions to multiple aggressively growing masses. Histopathologically, it is considered as a biphasic tumor with epidermal hyperplasia and subepidermal proliferation of transformed fibroblasts. The diagnosis can be made clinically, histopathologically and/or by detection of BPV DNA. Sarcoïd can appear on any part of the body, but they are mostly localized on the ventral abdomen, the paragenital region, head and limbs. Sarcoïds occur independent of breed, coat color, sex or age, but they develop more commonly in young adults and certain families and breeds are more vulnerable than others. Transmission of BPV is supposed to happen from cattle to horse or from horse to horse, possibly via insects.

SAMENVATTING


INTRODUCTION

Equine sarcoids are the most common skin tumors in horses and other equids like donkeys, mules and zebras (Jackson, 1936; Ragland, 1970; Lazary et al., 1985; Nel et al., 2006). They were described for the first time by Jackson (1936), who considered the tumors as sarcoma-like mixed (fibro-epithelial) tumors. Equine sarcoids are observed worldwide, independent of breed, coat color, sex or age (Ragland, 1970; Tarwid et al., 1985). Nevertheless, certain breeds and families are more vulnerable than others (Ragland et al., 1966; Brostrom et al., 1988; Lazary, 1988; Gerber, 1989; Marti et al., 1993) and sarcoïds occur most commonly in young adults, between 3 and 6 years (Marti et al., 1993; Torrontegui and Reid, 1994).

Clinically and pathologically, sarcoïds present most of the features of a true neoplasm. The predominant cell type is a malignant/transformed fibroblast. Sarcoïds generally have a high capacity for local tissue invasion into the dermis and subcutis but true metastatic dissemination does not occur (Pascoe and Knottenbelt, 1999). The clinical presentation of an equine sarcoïd can vary a lot between different individuals and even within the same individual. The number of tumors per horse varies from one single sarcoïd over a few to more than 100 lesions (Pascoe and Knottenbelt, 1999). A sarcoïd can remain stable for years or it can show a rapid and aggressive growth with infiltration of surrounding skin (Ragland, 1970; Marti et al., 1993). The reason for these large differences in clinical presentation is presently not known. The infiltrative behavior together with the infectious etiology is responsible for the frequent recurrences after treatment. Sarcoïds themselves are never lethal, but failure to treat them and the functional impairment they cause may result in euthanasia or slaughtering. The economic repercussion can be great: besides the cosmetic aspect, the normal use of the horse can be hindered, especially when tumors are located at the level of the girth and bridle, distal limbs, corner of the mouth and eyelids. Moreover a genetic component
has a role in the development of the disease, diminishing the breeding value of affected animals (LaZary et al., 1985; Meredith et al., 1986; Gerber et al., 1988). Diseased horses are therefore difficult to sell (Gerber, 1989).

In the pathogenesis, the bovine papillomavirus (BPV) plays a major role. Papillomaviruses are normally strictly species-specific, but equine sarcoids result from a natural cross-species infection. Although no intact viral particles have been demonstrated in sarcoids so far, DNA, RNA and proteins of the virus can be found (Amtmann et al., 1980; Trenfield et al., 1985; Otten et al., 1993; Teilke et al., 1994; Nasir and Reid, 1999; Carr et al., 2001b). Moreover, many similarities can be observed between equine sarcoids and papillomatosis in other species: fast growth, tendency for multiplicity and spreading by contact over different parts of the body (Jackson, 1936). Epizootics have been described in closed herds of horses and zebras, pointing to an infectious agent (Ragland et al., 1966; Nel et al., 2006).

CLINICAL PRESENTATION OF EQUINE SARCOIDS

Clinical types

Many clinical manifestations of equine sarcoids are known. They can be single or multiple, small to very large, stable or aggressively growing. At cross-section sarcoids appear as dermal thickening with a pale yellow color and a firm texture due to fibroblastic proliferation and a small number of capillaries within the tumor (Foy et al., 2002). The epidermis varies from thick, rough and hyperkeratotic to thinned or ulcerated. A classification based on morphological features has been proposed by Pascoe and Knottenbelt (1999) and will be followed throughout this paper.

Verrucous sarcoids (Figure 1) have a typical wart-like appearance with a rough, thickened, scabby surface above the fibroblastic part of the tumor (Knottenbelt et al., 1995). They can appear as small exophytic growing masses or as flat, often extended, scaly areas of skin with multiple smaller wart-like lesions on the surface. Common differential diagnoses for this sarcoid type are warts (induced by equine papillomavirus), aural plaque, chronic sweet itch and chronic blistering (Pascoe and Knottenbelt, 1999).

Nodular sarcoids (Figure 2) are subcutaneous, easily moveable nodules, often but not always spherical, covered by intact, apparently normal skin (Foy et al., 2002). In some cases, however, the overlying skin can be thinner, shiny and adherent to the tumor (Knottenbelt et al., 1995). Differential diagnoses include eosinophilic granuloma, melanoma and other nodular diseases (Knottenbelt et al., 1995).

Occult sarcoids (Figure 3) are flat circular to oval areas characterized by alopecia and a roughened or scaly appearance (Knottenbelt et al., 1995). Sometimes small cutaneous nodules can be observed (Pascoe and Knottenbelt, 1999). In some cases, occult sarcoids can be very subtle, displaying no more than a slightly thickened skin with a thin hair coat and slight changes in pigmentation. This relatively benign type can evolve rapidly towards a more aggressive type, either spontaneously or following injury such as biopsy or an erroneous treatment. Common differential diagnoses are dermatophytosis, chronic skin rubbing, alopecia areata and vitiligo (Knottenbelt et al., 1995).

Fibroblastic sarcoids (Figure 4) are large fibrous masses with an ulcerated surface. This is the most aggressive sarcoid type and can evolve from any other type after accidental or iatrogenic manipulation, including biopsy (Knottenbelt et al., 1995; Foy et al., 2002). In these cases it is observed that the original tumor starts to grow very quickly and begins to ulcerate. Skin wounds, especially at the distal limbs, are also at risk for development of fibroblastic sarcoids. Both dehiscence of the suture line after surgery without an apparent reason as well as the presence of a slow healing hypergranulating wound can be suggestive of sarcoid development (Pascoe and Knottenbelt, 1999). Fibroblastic sarcoids are liable to trauma, haemorrhage and local infection with bacteria or maggots. Fibroblastic sarcoids can be sessile with an invasive character or pedunculated with a small base. Differential diagnoses are exuberant granulation tissue, botryomycosis, squamous cell carcinoma and habronemiasis in warmer climates (Knottenbelt et al., 1995).

Mixed sarcoids (Figure 5) are a combination of two or more of the above mentioned types. They can represent a progressive or transient state between the verrucous/occult types and the fibroblastic/nodular types (Pascoe and Knottenbelt, 1999).

Malevolent sarcoids are very rare and have been described only by one group (Knottenbelt et al., 1995). These tumors infiltrate in lymphatic vessels resulting in multiple nodular or fibroblastic masses along these vessels. Local lymph nodes might also be involved. This type of sarcoid usually evolves from any of the other types following repeated injury, although spontaneous transformation is also possible.

In our clinic, sarcoids infiltrating the underlying muscles have occasionally been observed. These sarcoids can arise spontaneously or following failure of treatment resulting in deterioration of the tumor.

This classification into different types is also reflecting the clinical behavior of the tumors: occult types are the most stable tumors, nodular and verrucous sarcoids display a moderate growth, fibroblastic sarcoids are often fast growing and malevolent and infiltrating sarcoids, although rarely seen, are the most aggressive sarcoid types.

HISTOPATHOLOGICAL PROPERTIES

Jackson (1936), who described the equine sarcoid for the first time, considered it as a biphasic tumor with an epidermal and dermal component. Typical changes are subepidermal proliferation of spindle-shaped to fusiform fibroblasts showing hyperchromasia with a moderate to high cell density (Tarwid et al., 1985; Goodrich et al., 1998). These immature fibroblasts show a higher density in the superficial part of the tumor compared to the deeper layers (Martens et al., 2000). The fibroblasts are fusiform to spindle-shaped, forming whorls, interlacing bundles and
haphazard arrays with one another (Goodrich et al., 1998). At the dermo-epidermal junction fibroblasts are oriented perpendicular to the basement membrane, which is known as a picket-fence pattern (Ragland, 1970; Tarwid et al., 1985; Scott and Miller, 2003; Pulley and Stannard, 1990; Marti et al., 1993). The mitotic rate is invariably low (Goodrich et al., 1998; Martens et al., 2000). The amount of collagen varies considerably between tumors (Goodrich et al., 1998). In between the fibroblasts of the deeper layers of the dermis, large polygonal cells with large basophilic nuclei can be found. These are infiltrating dendritic cells that have an antigen presenting function suggesting an immune mediated defence against virus infected cells (von Tscharner et al., 1996; Lepage et al., 1998). Sometimes small epithelial inclusion cysts are found (Ragland, 1970; Tarwid et al., 1985; Pulley and Stannard, 1990). In the most typical cases, pseudoepitheliomatous hyperplasia and hyperkeratosis are observed. There is marked formation of rete pegs, which are broad invaginations (up to more than 20 cells) of epidermal cells into the dermis (Goodrich et al., 1998). However, in less typical sarcoids the epidermal component can be normal, atrophic or even absent (Marti et al., 1993; Lepage et al., 1998; Martens et al., 2000). In situ hybridization for BPV DNA of tissue sections from sarcoids reveals viral DNA in the dermal layer within the fibroblasts but not within the epithelial cells (Lory et al., 1993; Teifke et al., 1994; von Tscharner et al., 1996). Epithelial changes seen in sarcoids are likely the result of growth-promoting factors expressed by the neoplastic fibroblasts that stimulate proliferation of surrounding epithelial cells (Carr et al., 2001a).

The histological properties seem to be mainly dependent of the clinical type (Martens et al., 2000). Most of the verrucous and mixed sarcoids display the histological features as described above (Figure 6). In verrucous sarcoids, the epithelial component is much more important than the dermal, sometimes only lying as a small band of active fibroblasts

Figure 1. Verrucous sarcoid on the right mandible.

Figure 2. Nodular sarcoid on the upper eyelid.

Figure 3. Occult sarcoids in the paragenital region of a mare (ventral abdomen and medial side of the thigh).

Figure 4. Fibroblastic sarcoid on the dorsolateral aspect of the metacarpus.

Figure 5. Mixed sarcoid (verrucous – fibroblastic) on the upper eyelid.
against the epidermis (Pulley and Stannard, 1990). In the fibroblastic type there is always partial or total ulceration of the epidermis with infiltration of polymorphonuclear cells (Martens et al., 2000). The epidermis just next to the ulcerative lesions is more or less hyperplastic. In nodular sarcoids the epidermis is often thinned (Martens et al., 2000). If rete pegs are present, they are short. If the dermal proliferation is not making contact with the epidermis, the latter is normal. In occult sarcoids (Figure 7) the epidermis is usually normal or only displaying slight changes. The only typical aspect of this sarcoid type is the increased density of subepidermal fibroblasts infiltrating between a reduced number of hair follicles and sweat glands (Martens et al., 2000). They do not show a typical morphology or specific whorling distribution pattern. The density of dermal fibroblasts is also lower compared to the other types of sarcoids. The only common property for all types of sarcoids is the increased density of dermal fibroblasts compared to normal skin (Martens et al., 2000). Nodular sarcoids can mimic schwannomas although their clinical appearance and the presence of BPV demonstrate the sarcoid origin of the tumor (Vanheerden, in preparation).

DIAGNOSIS

Diagnosis can be made in three ways: clinical examination, histopathology and detection of BPV DNA. A thorough clinical examination combined with a focused anamnesis (duration of problems, localization of lesions, age, breed, evolution, multiplicity of lesions…) should be sufficient in the majority of cases. Lack of clinical experience or atypical tumor characteristics may cause confusion and necessitate lab-assisted diagnosis. Histopathological examination is often diagnostic, but it should be remarked that taking a biopsy, in particular of small stable lesions, may induce rapid growth and ulceration. If a non-excisional biopsy must be performed, sites within the mass must be carefully chosen to minimize the confounding factors of surrounding inflammation and granulation and to include intact epidermis (Goodrich et al., 1998). Possible deterioration of the sarcoid following biopsy is the reason why taking a biopsy of verrucous, occult and small nodular lesions is contra-indicated, even if making a definitive diagnosis is not possible in such cases (Pascoe and Knottenbelt, 1999). Another possibility is to perform a full surgical excision as if the lesion was a sarcoid, including excision of wide margins of normal skin and non-touch approach, followed by histological confirmation afterwards. This allows the pathologist to observe the range of morphological characteristics of the tumor allowing a correct diagnosis (Goodrich et al., 1998).

A more recent approach in diagnosing equine sarcoids is the detection of BPV DNA in lesions by means of polymerase chain reaction (PCR). This can be performed on histopathological samples of tissue suspected of equine sarcoid but not displaying the typical histological features (Angelos et al., 1991). A new approach is the detection of BPV DNA by means of PCR in the limited amount of material obtained by swabbing or scraping the lesions. This technique is especially valuable for the diagnosis of equine sarcoid tissue in non-healing wounds and in case of recurrences after former surgery (Martens et al., 2001). PCR detection of BPV DNA has many advantages: it is not invasive, sampling is easy and the trauma to the tumor is minimal. Disadvantages are the unsuitability for diagnosing occult sarcoids, the lower sensitivity compared to clinical diagnosis and the low specificity due to high prevalence of BPV DNA in the normal skin of horses (Martens et al., 2001; Bogaert et al., 2007a).

EPIDEMIOLOGICAL FACTORS

Prevalence

The equine sarcoid is the most common tumor in horses, donkeys, mules and zebras. Depending on the study, they represent 12 to 67 % of all equine tumors and 70 % of all skin tumors in horses (Jackson, 1936;
Ragland, 1970; Miller and Campbell, 1982; Teifke et al., 1994; Lepage et al., 1998). A prevalence of 0.6 to 2% in clinical populations has been reported (Ragland, 1970; Mohammed et al., 1992; Goodrich et al., 1998). In our clinic, 2.2% of the patients are admitted for treatment of equine sarcoids. This percentage is biased, since it can be higher than the true prevalence because veterinary referral hospitals can be specialized in sarcoid treatment and thus have a high case load. On the other hand, not all horses with sarcoids are referred to the clinic: small tumors are often treated at home by local practitioners or are left untreated. Recent studies conducted on three-year-old horses presented for field tests in Switzerland showed a prevalence of 12% in the Swiss Warmblood Horse and the Freiberger (Mele et al., 2007; Studer et al., 2007), which might be a more accurate estimation of the true population prevalence. In populations of Cape mountain zebras in South Africa the prevalence because veterinary referral hospitals can be specialized in sarcoid treatment and thus have a high case load. On the other hand, not all horses with sarcoids are referred to the clinic: small tumors are often treated at home by local practitioners or are left untreated. Recent studies conducted on three-year-old horses presented for field tests in Switzerland showed a prevalence of 12% in the Swiss Warmblood Horse and the Freiberger (Mele et al., 2007; Studer et al., 2007), which might be a more accurate estimation of the true population prevalence. In populations of Cape mountain zebras in South Africa the prevalence of equine sarcoids mounted up to 53%, but this high prevalence could be influenced by a high degree of inbreeding (Zimmerman, 2004; Nel et al., 2006).

LOCALIZATION OF TUMORS

Sarcoids can appear on any part of the body, but they are mostly localized on the ventral abdomen, the paragenital region, head and limbs (Jackson, 1936; Pulley and Stannard, 1990; Torrontegui and Reid, 1994; Goodrich et al., 1998; Piscopo, 1999; Bogaert et al., 2007b). On the head, most tumors develop at the eyelids, ears and commissure of the lips (Foy et al., 2002). Sarcoids are least common on the dorsum of the trunk (Knottenbelt et al., 1995). In northern regions, lesions occur predominantly on the head and abdomen while in warmer climates the limbs are more often involved (Marti et al., 1993). Apart from these predilection sites, sarcoids can develop at any site in injured skin (Torrontegui and Reid, 1994; Foy et al., 2002). The location of sarcoids has a significant relationship to their size: sarcoids on the head are mostly smaller whereas sarcoids on the limbs are larger when compared to all other sites (Brostrom, 1995). Sarcoids can appear singly or in clusters (Brostrom, 1995; Goodrich et al., 1998). Horses with multiple lesions tend to have larger-sized tumors compared to horses with a single sarcoid (Brostrom, 1995).

RISK FACTORS

Age

In contrast to most tumors observed in man and animals, the equine sarcoid predominantly develops in young adults. The majority of affected horses are younger than 6 years, with a peak incidence between 3 and 6 years (Miller and Campbell, 1982; Scott and Miller, 2003; Marti et al., 1993; Torrontegui and Reid, 1994; Brostrom, 1995; Piscopo, 1999; Foy et al., 2002). The mean age of sarcoid development is 3.5 to 4 years (Brostrom, 1995; Studer et al., 2007). Nevertheless, also younger and older horses can develop sarcoids. According to Mohammed et al. (1992) a gradual increase in incidence is observed up to the age of 15 years, followed by a decreasing incidence. A study on a large group of donkeys revealed that mainly males between 0.5 and 3 years were at high risk (Reid and Gettinby, 1996). An obvious reason for this age distribution has not been demonstrated yet, but possibly older horses acquire a certain level of immunity which results in spontaneous regression of existing tumors and prevention of new tumor development. Another hypothesis is that genetically susceptible animals develop tumors early in life (Torrontegui and Reid, 1994).

Sex

Most authors do not assume that there is a gender predisposition for sarcoid development (Ragland, 1970; Miller and Campbell, 1982; Pulley and Stannard, 1990; Torrontegui and Reid, 1994). However, others claim that geldings are more susceptible than stallions and mares. Mohammed et al. (1992) observed that the risk for geldings to develop sarcoids was twice as high compared to stallions and mares. In donkeys, a predisposition in males was observed (Reid et al., 1994; Reid and Gettinby, 1996). A possible explanation is that in castrated males a wound is created which is a possible entrance for BPV (Reid et al., 1994). On the other hand, in spite of the frequent localization of sarcoids in the paragenital region, the castration wound itself is only seldom affected. Another hypothesis is that female animals have protective factors, rather than a predisposition in males (Reid and Mohammed, 1997).

Breed

There is clear breed predisposition for the development of equine sarcoids (Marti et al., 1993). In North-America, Quarter Horses are twice as likely to develop sarcoids compared to Thoroughbreds (Angelos et al., 1988). Also Appaloosas and Arabian Horses are more frequently affected than Thoroughbreds (Angelos et al., 1988; Mohammed et al., 1992). Standardbreds on the other hand are rather resistant (Meredith et al., 1986; Mohammed et al., 1992; Brostrom, 1995). A possible explanation is that Quarter Horses and Appaloosas are often used on large cattle farms which results in a higher risk to get in contact with BPV (Mohammed et al., 1992). Moreover they often work on rough surfaces resulting in frequent injury to the limbs and the possibility for sarcoid development in these wounds (Brostrom, 1995). Another more plausible explanation for breed predisposition can be found in the different genetic background of these breeds (Lazary et al., 1985; Meredith et al., 1986).

Genetic background

The most important genes known to play a role in sarcoid development are those of the major histocompatibility complex (MHC), but also other genes are involved. The genes of the MHC code for proteins involved in the immune response and for protein components of the complement system (Piscopo, 1999). In horses these proteins are called equine leucocyte antigens (ELA). Three major classes of MHC genes
exist (Lazary et al., 1994): class I genes code for glycoproteins on the cell wall of most nucleated cells. They have a role in recognition and killing of virus infected cells. Seventeen internationally accepted alleles (A1-10, A14-15, A19, W16-18, W20) and 5 regional variants (Be22, Be24-26, Be108) are distinguished serologically in the horse. Class II proteins are expressed on the cell surface of antigen presenting cells. Within this class the DQB gene is highly polymorphic. In the horse 5 internationally accepted alleles (W13, W22-23) and 2 local variants (BeVI, Be200) can be determined serologically (Lazary, 1988; Hesford et al., 1989). To date, the complete sequence coding the ELA-DQB gene is known (Szalai et al., 1994) and more than 23 DQB sequences have been reported in domestic horses (Gyllensten et al., 1990; Szalai et al., 1993; Horin and Mattiasovic, 2002), but not all reported sequences can be assigned to a serological specificity (Villegas-Castagnasso et al., 2003). Recent evidence suggests the presence of at least two copies of the horse DQB gene (Horin and Mattiasovic, 2002; Villegas-Castagnasso et al., 2003), one of them probably being a pseudogene. This genome organisation hampers an easy classification of DQB variants with modern molecular techniques such as PCR-RFLP. Class III genes encode several proteins involved in the complement system, but these are not associated with predisposition for sarcoids.

In most breeds such as the Swiss Warmblood, the Irish Warmblood, the Swedish Halfbred and the Selle Français, the presence of equine sarcoids is strongly correlated with ELA W13 (Meredith et al., 1986; Brostrom et al., 1988; Gerber et al., 1988; Lazary et al., 1994; Brostrom, 1995). A higher percentage of recurrences is also observed in the presence of this allele (Brostrom, 1995). Standardbreds lack W13, which might be a reason for the low prevalence of equine sarcoids in this breed (Meredith et al., 1986). In the Freiberger, a Swiss draft horse, W13 is also lacking, but a higher sensitivity for sarcoid development is correlated with Be108, a local variant (Lazary et al., 1994). In the Selle Français, A3 is correlated with a higher prevalence of sarcoids, but this could be due to linkage disequilibrium with W13 (Lazary et al., 1994). Indeed, ELA A3W13 is frequently seen as one of the paternal haplotypes in families where equine sarcoids are more commonly observed than in others (Brostrom et al., 1988; Gerber et al., 1988). A5 would be correlated with early onset of sarcoids (Brostrom, 1995). The association of equine sarcoids with certain antigens is however not absolute. A large proportion of horses are carriers of W13, but only a small part of them will develop sarcoids (Goodrich et al., 1998). On the other hand, also horses lacking W13 may be affected by sarcoids. Maybe these alleles are only markers of other susceptibility genes in linkage disequilibrium with the MHC, with a more direct influence on the pathogenesis (Brostrom et al., 1988; Gerber et al., 1988; Lazary et al., 1994). Still it seems plausible that genes coding for proteins regulating the immune system can also influence the susceptibility for certain diseases (Piscopo, 1999). Indeed, in virus-infected tumors the virus-infected cells express new cell wall antigens, both virus specific and virus induced non-viral products. These foreign antigens should normally be recognized by the immune system resulting in destruction of tumor cells. If a failure occurs in the immune system, recognition and destruction of tumoral cells may also fail. In man and rabbits, a linkage between papillomavirus induced cancer and MHC alleles has also been observed (Wank and Thomssen, 1991; Han et al., 1992).

In Arabian Horses a significant correlation exists between presence of equine sarcoids and heterozygosity for the defective DNA protein kinase catalytic subunit (DNA-PKcs) allele, which is in homozygous condition responsible for severe combined immunodeficiency. In a population of Arabian Horses, 8.7 % of the animals were carriers of the defective DNA-PKcs allele, compared to 18.6 % of the equine sarcoid affected animals (Ding et al., 2002).

TRANSMISSION

The way horses get infected with BPV is not yet clarified. Possible transmission routes are direct contact with cattle (Jackson, 1936), indirect transmission from cattle to horses (e.g. housing of horses in cattle stables, transmission by caretakers of the animals) and transmission from horses with sarcoids to other horses (direct or indirect via caretaker, tack, grooming equipment, common rubbing posts) (Bogaert et al., 2005; Bogaert et al., 2007a). In donkeys it is known that animals having close contact with affected animals are at higher risk for development of sarcoids (Reid et al., 1994). Epidemiological data, multiplicity of tumors on one horse, spontaneous development of sarcoids on intact skin and absence of contact with cattle or affected horses suggest that flies or other insects may play an important role as mechanical vector in BPV infection of the horse (Voss, 1969; Reid et al., 1994; Torrontegui and Reid, 1994; Knottenbelt et al., 1995; Pascoe and Knottenbelt, 1999). Recently, Kemp-Symonds and Kirk (2007) have demonstrated the presence of BPV-1 and -2 in Musca autumnalis face flies infesting sarcoid affected horses. Moreover, it is observed that in one specific region identical variants of BPV are seen both in horses and in cattle, also pointing into the direction of a flyborne transmission (Otten et al., 1993). In case of a skin wound, BPV coincidentally present in the environment of the horse can immediately come into contact with the subepidermal fibroblasts via insects (Reid et al., 1994; Torrontegui and Reid, 1994; Knottenbelt et al., 1995; Pascoe and Knottenbelt, 1999). Furthermore, cell growth is already stimulated as part of wound regeneration. This can eventually lead to malignant transformation of these cells (Phelps et al., 1999; Foy et al., 2002).

ACKNOWLEDGMENTS

Prof. Dr. Hilde De Cock (University of Antwerp) is greatly acknowledged to provide us with Figures 6 and 7.
LITERATURE


Op de Vakgroep Heelkunde en Anesthesie van de Huisdieren van de Faculteit Diergeneeskunde te Merelbeke loopt momenteel een onderzoek naar de transmissie van het bovienephalomavirus. Hiervoor zijn we op zoek naar proefpaarden met equine sarcoïden, evenals naar paarden die vroeger reeds behandeld werden voor sarcoïden maar nu sarcoïd-vrij zijn. Deze paarden zullen gebruikt worden om de exacte manier van transmissie te ontrafelen, uitgaande van viraal materiaal afkomstig van zowel runderen als van paarden. Opgelet: deze paarden zullen worden overgekocht door de vakgroep en kunnen dus niet in het bezit blijven van de eigenaar! Indien u in uw cliënteel dergelijke dieren hebt die anders toch moeten geëuthanaseerd of geslacht worden, kunt u ons steeds contacteren voor overname van deze dieren.

Contact: Pieter Depoorter (pieter.depoorter@UGent.be), Lies Bogaert (lies.bogaert@UGent.be)