

Review Article

Equine proliferative enteropathy – a review of recent developments

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Summary

Equine proliferative enteropathy (EPE) is a disease of foals caused by the obligate intracellular organism *Lawsonia intracellularis*. This emerging disease affects mainly weaning foals and causes fever, lethargy, peripheral oedema, diarrhoea, colic and weight loss. The diagnosis of EPE may be challenging and relies on the presence of hypoproteinaemia, thickening of segments of the small intestinal wall observed upon abdominal ultrasonography, positive serology and molecular detection of *L. intracellularis* in faeces. Although the clinical entity, diagnostic approach and treatment of EPE are well established and described, the epidemiology for this disease has remained largely unaddressed. This article focuses on new developments in the field of EPE, including epidemiology, pathophysiology, clinical signs, diagnosis, treatment and prevention.

The Summary is available in Chinese – see Supporting information.

Keywords: horse; *Lawsonia intracellularis*; equine proliferative enteropathy; pathophysiology; clinical management

Introduction

Lawsonia intracellularis is the aetiological agent of the recently recognised and emerging intestinal disease in horses called equine proliferative enteropathy (EPE). *Lawsonia intracellularis* is an obligate intracellular, motile, curved, Gram-negative bacterium that resides freely within the apical cytoplasm of infected intestinal enterocytes [1]. The genus *Lawsonia* was placed within the *Desulfobionaceae* family based on phylogenetic analysis of 16S rDNA sequences and, to date, the *Lawsonia* genus contains only one species [2]. Similar to the *Desulfobionaceae*, *L. intracellularis* is anaerobic but differs from other members of the family in its inability to reduce sulphate. *In vivo*, it causes proliferation of the affected enterocytes, resulting in a thickened small and sometimes large intestine. *Lawsonia intracellularis* can only be grown *in vitro* in cell culture and requires dividing cells and a specific atmosphere for infection and growth [1]. Peak infection occurs between 6 and 7 days of incubation. A single polar flagellum and darting motility are observed with extracellular *L. intracellularis* organisms in cell culture. Besides horses, *L. intracellularis* infects many species of domestic and wild animals, including pigs, hamsters, rabbits, foxes, deer, ferrets, ostriches and nonhuman primates. The first published report of proliferative enteropathy in horses was in 1982 by Duhamel and Wheeldon [3]. Since 1996, several reports of sporadic cases and outbreaks on breeding farms have been described [4–21]. In the last few years, reported cases of EPE have been increasing, occurring primarily in post weaning foals and occasionally in young adults [20]. The disease has almost reached a worldwide occurrence and has been reported in the USA, Canada, Europe, South Africa, Australia, Brazil and Japan.

Pathogen characterisation

Molecular investigations of *L. intracellularis* isolates from proliferative enteropathy lesions of a variety of animal species including horses and hamsters, showed 98% homology of the 16S-rDNA gene to pig isolates [22]. Moreover, phenotypic characterisation of outer membrane proteins and immunoblots of different *L. intracellularis* isolates using several antibodies and more sensitive molecular characterisations of the *L. intracellularis* genome demonstrated only minor differences among isolates [23]. None of these differences appear to be antigenically relevant. Recently, the whole genome of a porcine *L. intracellularis* isolate was sequenced and analysed for the presence of variable number tandem repeat (VNTR) sequences. Variable number tandem repeat sequences in the genomes of prokaryotes are often associated with a high level of polymorphism and enable bacterial strain differentiation with substantial discriminatory power. Use of these

L. intracellularis VNTR sequences provides a sensitive method for analysis of the genetic relatedness of *L. intracellularis* bacteria or DNA obtained from various temporal and geographical locations and from various animal species. This provided insight into the phylogenetic relatedness of these isolates. Molecular VNTR sequence profiles of *L. intracellularis* isolates from various documented outbreaks of proliferative enteropathy occurring in pigs, horses, ostriches, spider monkeys, ferrets and rabbits were analysed. Patterns emerged that provide some insight into the sources and phylogenetic relatedness of *L. intracellularis* isolates from different species. The VNTR sequence types obtained from pigs were very different from those obtained from horses or other nonpig species. The VNTR patterns distributed the isolates into 2 distinct groups, those obtained from pigs and those obtained from horses and other nonpig species. Little or no genetic variation was found between isolates from within outbreaks for any animal species or in multiple temporal samples taken from the same outbreak site. Slight variations between isolates obtained from outbreaks at different geographic locations were found and though these differences were minor, they were consistent and stable enough that outbreak isolates could be identified and traced.

Epidemiology

In pig populations, the chronic form of the disease is maintained by subclinically affected pigs or is established environmentally in some sites, allowing transmission of *L. intracellularis* to serial flows of pigs [24]. Mice and rats have been shown to be important reservoirs of *L. intracellularis* on pig farms with prevalences of PCR positive animals varying substantially between farms (4–83%) [25,26]. Rodents appear to be suitable reservoir hosts due to their susceptibility to *L. intracellularis*, their close contact to domestic animals and high reproductive rate, which maintains *L. intracellularis* across generations. The source of infection has not been determined for horses. Exposure to pig faeces has been speculated as a potential source of infection for horses since the first reported cases of EPE. However, in most cases of EPE, no history or evidence of direct or indirect exposure to pigs or pig faeces has been reported. Further, multi-locus VNTR profiles of pig and equine isolates differ greatly. A recent experimental study has shown host specificity for *L. intracellularis* isolates cultured from pig or horse intestines [27]. The study showed that clinical signs, longer periods of shedding and stronger serological immune responses were observed in animals infected with species-specific isolates. Previous studies have shown that a variety of wild and domestic animals including dogs, cats, rabbits, opossums, skunks, mice and coyotes, can shed *L. intracellularis* on farms with diagnosed EPE cases [28,29]. On a



Fig 1: Heavy contamination of hay with faeces from cottontail rabbits at a farm in northern California with endemic occurrence of equine proliferative enteropathy.

recently identified farm in northern California endemic for EPE, 7.5% of faecal samples and 27% of serum samples from cottontail rabbits tested PCR positive and seropositive for *L. intracellularis*, respectively [29]. Of interest was that on this farm, a large population of cottontail rabbits lived in the hay barn and had direct access to the hay fed to the horses. An epidemiological investigation on this farm showed rabbit faeces on top of hay bales but also in the feeders of the weanling foals, suggesting that the foals developing EPE were likely exposed to *L. intracellularis* via the oral ingestion of infected rabbit faeces (Figs 1, 2). Similar to rodents, lagomorphs may represent an effective reservoir/amplifier host due to their large population, their close contact to horses and short reproductive cycle. It still remains to be determined how *L. intracellularis* became endemic in the rabbit population on this farm.

Faeco-oral transmission of *L. intracellularis* has been documented in naïve foals housed with clinically infected foals experimentally challenged with an equine isolate of *L. intracellularis* [30]. A recent study has shown that faeces from experimentally infected rabbits with an equine isolate of *L. intracellularis* served as infectious material for weanling foals [31]. Although infected rabbits and foals remained asymptomatic, infection was supported by faecal shedding of *L. intracellularis* and detection of specific antibodies to *L. intracellularis*. Although the natural infectious dose for foals has not been determined, pigs receiving as low as 10^5 *L. intracellularis* have been shown to develop infection [32]. Recent



Fig 2: Faeces from cottontail rabbits found at the bottom of a hay feeder in a weanling pasture from a farm in northern California with endemic occurrence of EPE.

work suggested that 1 g of infectious faeces would suffice to deliver the above challenge dose [26]. It is the authors' opinion that on most farms, the initial transmission of *L. intracellularis* occurs via the accidental ingestion of infectious faeces from one of the described or yet undetermined amplifiers/hosts. Amplification of *L. intracellularis* and environmental contamination leading to exposure rates of up to 100% of resident foals are likely to occur secondary to the shedding of large quantities of *L. intracellularis* from either clinically or subclinically infected foals.

In piglets, large group size, weaning, transportation, diet change and mixing have been associated with clinical disease [1]. Predisposing factors such as the stress of weaning, overcrowding, decline in *L. intracellularis*-specific colostral antibodies, endoparasitism and introduction of new animals have been suggested in the development of EPE in foals [7]. In pigs, infection and faecal shedding of *L. intracellularis* may persist for as long as 12 weeks [32]. This is in sharp contrast to experimentally infected foals showing onset and duration of faecal shedding ranging from 10 to 14 days and 17 to 27 days, respectively [30,33]. Previous work has shown that *L. intracellularis* is likely to survive in environmental conditions for 1–2 weeks at 5–15°C [34].

Pathogenesis and immune responses

The pathogenesis and pathophysiology of EPE has remained poorly investigated and most of the information available has been extrapolated from experimentally infected hamsters, pigs and rabbits. Comprehensive studies of lesion development and evolution have been conducted in pigs [35] and hamsters [36]. Morphological studies of early lesions in experimentally infected animals indicate that enterocyte hyperplasia is directly preceded by infection and replication of the intracellular organism [35,36]. *In vivo*, the onset of hyperplasia associated with proliferative enteropathy follows an increased number of intracellular *L. intracellularis* in enterocytes. Likewise, resolution of the lesions is closely related to disappearance of the intracellular organisms, indicating a correlation between the 2 events [1]. The means by which *L. intracellularis* produces hyperplasia or how the host cells respond to this infection are unknown. No other cytopathological effects on infected enterocytes are seen *in vivo* or *in vitro*. Inflammation is a factor only in later stage lesions and is not characteristic of the primary lesion. The peak of lesion development in both pigs and horses is about 21 days after infection but may vary according to the dose of *L. intracellularis*.

While the dynamics of *L. intracellularis* infection have been well characterised, little is known about the genetic basis for its virulence, pathogenesis or physiology. Bacterial isolates at low passage induce clinical and pathological changes typical of proliferative enteropathy but isolates at high passage in cell culture are attenuated and do not induce typical lesions. Recently, the differential transcriptional profiles of a pathogenic (low passage) and a nonpathogenic (high passage) isolate were determined in order to define potential virulence factors [37]. The pathogenic isolate showed a wider transcriptional landscape made up of plasmid-encoded genes, regulatory factors and ATP-binding cassette transporter genes. Loss of pathogenicity *in vivo* of the high passage isolate appears to be due to gene silencing mechanisms. Further, this study identified a number of genes expressed in pathogenic *L. intracellularis* that will make ideal targets for development of further diagnostic and control measures.

In horses, lesions are most commonly seen in the ileum, near the ileal-caecal junction and appear as a thickening of the mucosa. Gross lesions are not evident in all cases of EPE and may often be overlooked. Intestines show an irregular, patchy subserosal oedema. The ileal mucosa is thickened with deep folds and chronically affected animals may have patches of pseudomembrane covering the mucosa (Fig 3). Hypertrophy and thickening of the muscularis mucosa may occur in chronically affected or recovering animals. Histologically, adenomatous proliferation occurs among the epithelial cells in the crypts of the small intestine, in association with the presence of curved, intracellular bacteria in the apical cytoplasm of these enterocytes [3,4,7]. Severe proliferative enteropathy is diagnosed by the demonstration of hyperplasia of the crypt glands with an increased number of mitotic figures and marked reduction or absence of goblet cells in routine haematoxylin and eosin preparations; however, for visualisation of the bacteria in the cytoplasm of enterocytes, special stains are

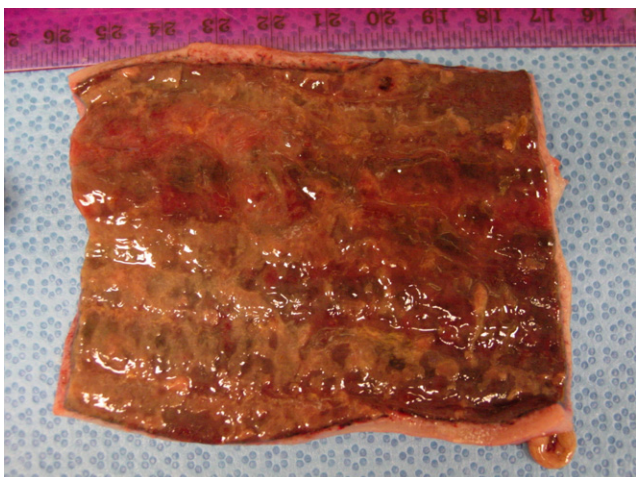


Fig 3: Gross lesions of equine proliferative enteropathy of an 11-month-old Thoroughbred colt with proliferative enteropathy showing a diffusely thickened intestinal wall and patches of pseudomembrane covering the mucosa.

necessary. The histological lesions of proliferative enteropathy are unique and inflammation is not normally a hallmark of disease. Warthin-Starry silver stain allows the detection of the bacteria in histological sections (Fig 4), improving the diagnostic sensitivity, but the technique has limitations when applied to autolysed and necrotic samples [1]. Immunohistochemistry procedures, using *post mortem* tissue or biopsy material with an antibody specific for *L. intracellularis*, are considered the gold standard for histological confirmation of EPE (Fig 5). Recently, Page *et al.* [38] reported on fatal necrotising enteritis associated with *L. intracellularis* in 4 foals. In these 6- to 8-month-old foals, the clinical entity was acute and followed by rapid clinical deterioration and death due to endotoxaemia, bacteraemia or disseminated intravascular coagulation.

Cell-mediated immune response appears to be an important feature in protecting animals from reinfection with *L. intracellularis* [39,40]. Descriptive immunocytological studies of intestinal tissue sections of pigs affected by proliferative enteropathy, reveal a mild infiltration of cytotoxic T cells, macrophages and B lymphocytes carrying major histocompatibility complex (MHC) class II structure at the beginning of the cell-mediated immune response [41]. Immunohistochemical studies of intestinal sections of naturally infected pigs also demonstrated a large accumulation of immunoglobulin A (IgA) in the apical cytoplasm of proliferating enterocytes

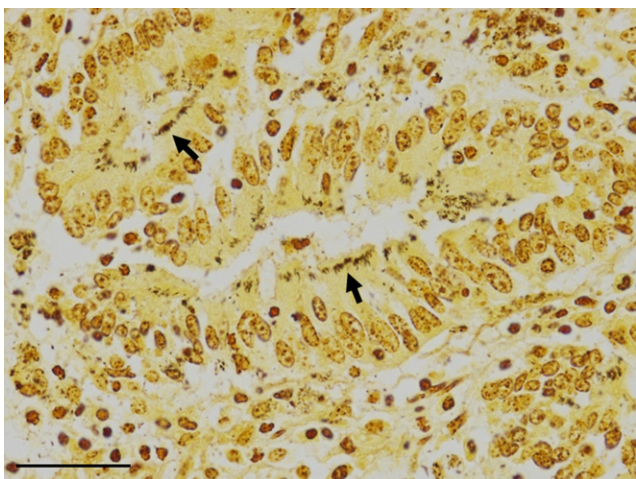


Fig 4: Ileal mucosa from a 6-month-old Quarter Horse filly with proliferative enteropathy. Aggregates of bacteria (arrows) are present in the cytoplasm of epithelial cells of hyperplastic crypts (Warthin-Starry silver stain; bar = 30 µm).

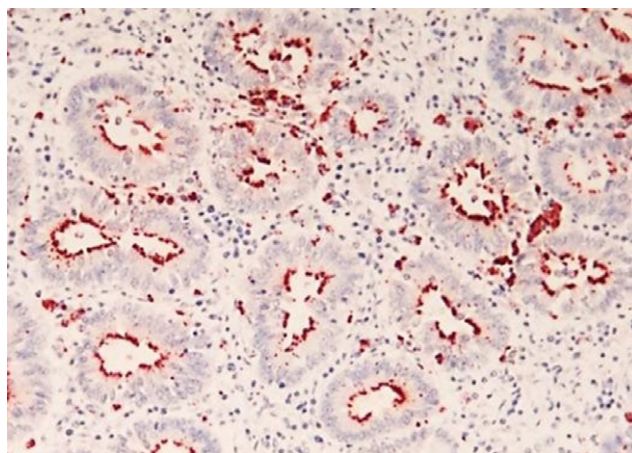


Fig 5: Immunohistochemical stained section of small intestine from an 8-month-old foal with proliferative enteropathy. *Lawsonia intracellularis* specific antibody stains the bacteria lining the apical cytoplasm of the affected crypts (red areas).

[41]. Further, interferon gamma is produced by peripheral blood mononuclear cells (PBMCs) of both pigs and horses following specific stimulation [39,42,43] and *L. intracellularis*-specific IgA is detected in intestinal lavages of challenged pigs [40]. Similarly, interferon gamma played a role in limiting intracellular infection and increased cellular proliferation in experimentally infected mice [44].

Clinical and laboratory findings

There are characteristic case details, seasonality, clinical signs and blood work abnormalities associated with EPE. The disease is generally manifested in foals less than one year of age and in North America is often seen between August and February [16]. Although the disease is commonly seen in weanling foals 4–7 months of age, cases of EPE have been seen in young adults [20]. Lethargy, anorexia, fever (>38.5°C), peripheral oedema (ventrum, sheath, throatlatch and distal limbs; Fig 6), weight loss (Fig 7), colic and diarrhoea are among the most common clinical findings in affected foals. Early clinical signs are generally unspecific and include mild depression, partial anorexia and fever. Although diarrhoea is commonly seen in affected foals and can vary from cow pie to watery, some affected foals may have normal faecal character. Foals with EPE may also have concurrent disorders such as respiratory tract infections, gastric

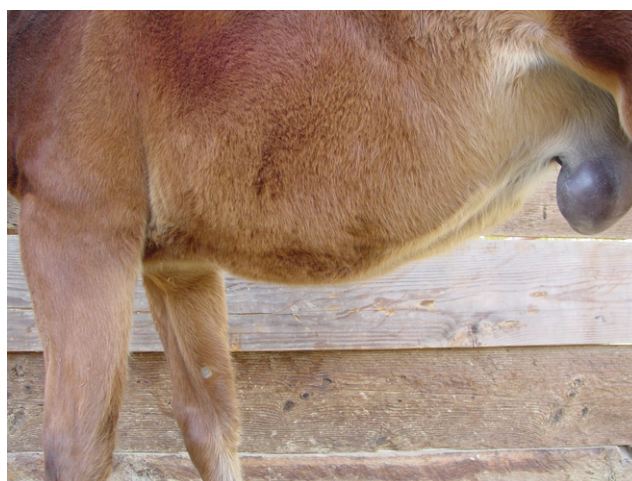


Fig 6: Six-month-old Quarter Horse colt with proliferative enteropathy displaying ventral and scrotal oedema.



Fig 7: Severe weight loss in a 5-month-old Quarter Horse colt with proliferative enteropathy.

ulceration and intestinal parasitism. One must keep in mind that signs of EPE may resemble those of more common gastrointestinal disorders such as parasitism, bacterial infections (*Clostridium* spp., *Salmonella* spp., *Rhodococcus equi*, *Neorickettsia risticii*), rotavirus, coronavirus, ulcerations, sand accumulation, intestinal obstruction and intoxication with plants, chemicals and pharmacologic agents such as nonsteroidal anti-inflammatory drugs or antimicrobials. Similar to pigs, the disease can be subclinical in foals and be manifested by a self-limiting and transient decrease of total serum protein concentration coupled with decreased daily weight gain when compared with unaffected foals [30,33]. It will remain to be determined if growth retardation or unthriftiness are associated with subclinical infection.

The most consistent laboratory finding of clinical EPE is hypoproteinaemia due to hypoalbuminaemia. Total protein is generally less than 50 g/l and albumin is usually less than 20 g/l. In a recent case report [16], hypoalbuminaemia was the only consistent clinicopathological abnormality of 57 affected foals with albumin concentrations ranging from 9 to 33 g/l (normal reference range 27–42 g/l). The exact mechanisms by which hypoalbuminaemia develops in affected foals has not been investigated. It appears that a combination of decreased feed intake, coupled with malabsorption and protein-losing enteropathy due to the proliferative nature of the disease may represent likely mechanisms by which low albumin occurs [45]. Experimentally infected hamsters had significantly lower intestinal absorption of glucose, potassium and chloride, suggesting that malabsorption is the main mechanism involved in the physiopathology of the diarrhoea in *L. intracellularis*-infected animals [46].

Affected foals may also demonstrate nonspecific blood abnormalities such as anaemia or haemoconcentration, leucocytosis or neutropenia, hyperfibrinogenaemia, increased activity of muscle enzymes and electrolyte abnormalities (hypocalcaemia, hypochloraemia and hyponatraemia). Urine analysis to rule out protein-losing nephropathy and cytological evaluation of abdominal fluid to rule out protein lost to a third space are generally unremarkable.

Diagnosis

A presumptive diagnosis of EPE is generally made based on age of the affected animal, clinical signs, hypoproteinaemia/hypoalbuminaemia, presence of thickened small intestinal loops on ultrasonographic evaluation and ruling out other causes of enteropathy and protein losses. Abdominal ultrasonography, although not very sensitive, may show segments of thickened small intestine (Fig 8) and excessive abdominal fluid. In these cases, abdominocentesis will yield a noninflammatory

transudate. An *ante mortem* diagnosis is generally confirmed via PCR detection of *L. intracellularis* in faeces or rectal swab and/or serology [47].

It is essential to combine both molecular and serological diagnostic testing since these modalities have high analytical specificity but variable sensitivity depending on the situation. Negative PCR results can be expected if the faecal samples are collected from foals with prior antimicrobial treatment or during advanced disease stage when *L. intracellularis* organisms are no longer expected in the faeces. Negative serological results can be expected in the early stage of the disease, when humoral immune responses are not yet strong enough to be detectable by serology. Further, differences in sensitivity among different PCR and serological assays can lead to divergent results. Among PCR assays, the use of real-time platform has been shown to yield the best sensitivity and to reduce the likelihood of cross- or carry-over contamination (i.e. false positive results) [48–50]. Several serological assays, including indirect fluorescent antibody test (IFAT), enzyme-linked immunosorbent assay (ELISA) and immunoperoxidase monolayer assay (IPMA), have all been validated and established for pigs [51–54]. A preliminary comparative study using equine serum samples has shown that various serology assays are accurate; however, the IPMA is the most specific when determining the presence of anti-*L. intracellularis* antibodies in adult horses with EPE (C.J. Gebhart, personal communication).

Supportive and medical treatment

It is important to treat affected animals early before lesions become advanced and result in marked weight loss and critically low serum protein concentrations. Treatment of EPE in horses involves the use of antimicrobials such as macrolides, alone or in combination with rifampin, chloramphenicol, oxytetracycline or doxycycline administered for 2–3 weeks. The choice of antimicrobial in the treatment of EPE should take into account the risk of inducing disturbance of the gastrointestinal flora and renal toxicity. This is especially a concern when treating older foals with severe hypoalbuminaemia. National and international policies on antimicrobial use in horses should also be considered [55]. Supportive care such as i.v. fluids, plasma transfusion, parenteral nutrients and anti-ulcer drugs are commonly used to treat affected foals. Concurrent medical conditions should also be addressed. Rapid clinical improvement following treatment is to be expected; however, it may take weeks for the hypoproteinaemia to resolve. Spontaneous recovery of clinically infected foals has not been documented and up to 93% of treated foals usually survive the disease [16]. However, in more recent years the authors have seen more severe cases presented to veterinary hospitals with a poorer outcome. Generally nonsurvivors develop complications associated with



Fig 8: An ultrasound image showing a thickened section of small intestinal wall in a 7-month-old Thoroughbred colt with proliferative enteropathy. The wall thickness measured between 5.4 and 6.0 mm (normal wall thickness \leq 3 mm).

gastrointestinal translocation, gastrointestinal perforation or kidney failure [38]. At 2 different veterinary hospitals, the mortality rate in the past 2 years ranged from 13% (2/15) to 31% (4/13; N.M. Slovis and N. Pusterla, personal communication). Recent work by Frazer [16] showed that clinically affected and successfully treated foals sold for an average of 68% of the average price of unaffected foals by the same stallion [16]. However, their monetary earnings from racing were not significantly different from other horses [56].

Monitoring and disease prevention

Based on clinical observations, it appears that the exposure rate to *L. intracellularis* is higher than the clinical attack rate. That said, and assuming that index cases are only the tip of the iceberg, it is always advisable to test herd mates to determine their exposure and clinical status. This is best achieved by collecting blood to determine the level of anti-*Lawsonia intracellularis* antibodies by serology and to measure total protein concentration by refractometry. Another more expensive alternative is to measure total protein and/or albumin concentrations by chemical analysis. The use of PCR testing of faeces from healthy herd mates is not advised in this situation due to the expense of testing and low rate of positivity. Also, the results from previous epidemiological studies show that healthy herd mates rarely shed detectable *L. intracellularis* [47,57]. Daily clinical monitoring of all herd mates is also recommended in order to recognise early stages of disease. This is best achieved via daily physical examination, including rectal temperature, and the regular assessment of weight, allowing the calculation of daily weight gain. A positive titre by IPMA (≥ 60) in a healthy herd mate with no hypoproteinaemia should be viewed as past exposure with no apparent disease or possibly early, not yet clinically apparent, EPE. Seropositive or seronegative clinically healthy herd mates with hypoproteinaemia (<50 g/l) or hypoalbuminaemia (<30 g/l) should undergo further diagnostic testing (cell blood count, abdominal ultrasound examination, faecal PCR) to determine if *L. intracellularis* infection is the cause of the hypoproteinaemia. Treating foals with suspected EPE based only on clinical findings and hypoproteinaemia/hypoalbuminaemia is not recommended, knowing the risk associated with the use of antimicrobials. Healthy seronegative herd mates with no hypoproteinaemia should continue to be monitored daily for clinical signs and bimonthly for hypoproteinaemia and/or hypoalbuminaemia and detectable antibodies to *L. intracellularis*. Any foal developing clinical signs of EPE should undergo a thorough diagnostic assessment. Further, clinically affected foals or foals with suspected clinical EPE should be separated from the rest of the healthy herd mates in order to decrease environmental contamination until their shedding status has been determined by PCR. It has been previously shown that experimentally infected foals start shedding *L. intracellularis* 5–17 days prior to developing hypoproteinaemia and clinical signs [30]. It is this prodromal stage of subclinically infected foals that is likely responsible for the environmental contamination (including potential carriers, vectors and multipliers) and exposure of susceptible foals.

The monitoring of a herd with endemic status follows guidelines similar to those for herds with diagnosed index cases. This includes the regular physical evaluation of resident foals and bimonthly assessment of total protein concentration and serological status. Monitoring for exposure to *L. intracellularis* and hypoproteinaemia/hypoalbuminaemia should begin at least 4 weeks prior to the historical first detection of clinical cases. Bimonthly data, including concentration of total serum solids or albumin and weight gains, should be evaluated for each foal and compared with the previous data in order to determine decreasing trends potentially associated with early disease. Recent work performed in central Kentucky has shown a seasonality to EPE cases with peak cases recorded in November and December [16]. Year to year variations depending on climatic conditions can be expected; however, most of the EPE cases are seen between August and February in the northern hemisphere, as recently weaned foals are primarily affected. Considering the cost of treating a foal with clinical EPE, this monitoring programme is cost effective, especially if measurement of total serum solids concentrations can be performed by farm personnel. The lack of epidemiological data regarding potential natural reservoir hosts as well as the paucity of information pertaining to the biology of *L. intracellularis* precludes the

institution of any management changes on endemic farms. Early recognition of clinical cases and separating them from the rest of the susceptible foals until full recovery or cessation of faecal shedding appears to be a logical biosecurity measure to prevent spread and environmental contamination. Further, maintaining good pest control and preventing nonequine domestic and wild animals from gaining access to feed and feeding areas may potentially minimise the risk of disease spread.

Prevention strategies have been best described in pigs using in-feed antimicrobials and a commercially available *L. intracellularis* vaccine [39,58,59]. Recent work has shown that detectable humoral and cellular responses can be measured in foals administered an avirulent live *L. intracellularis* vaccine [33,60–62]. The recently established vaccine protocol has shown that the intrarectal administration of 30 ml of either the lyophilised or the frozen-thawed formulation of the avirulent *L. intracellularis* vaccine given twice, 30 days apart, yielded the strongest immunological responses [60]. The *L. intracellularis* vaccine has been shown to be safe and administration well tolerated by the foals. Further, the avirulent *L. intracellularis* vaccine has not been associated with the induction of clinical disease in pigs or foals. Faecal shedding for up to 12 days has been documented following intrarectal vaccine administration in foals [60]. Using the above mentioned protocols, vaccine efficacy has been evaluated in the field and, more recently, under experimental conditions. A field efficacy trial performed on EPE endemic farms in central Kentucky in 2008 showed that vaccinated foals maintained higher daily weight gains and higher total protein concentrations when compared with a nonvaccinated, naturally seroconverted group [63]. Due to the low incidence of disease reported on the study farms, no difference in attack rate between vaccinated and nonvaccinated foals could be determined. The overall decreased disease prevalence in the study population may have been associated with the ongoing vaccine trial on these farms, as disease prevalence in central Kentucky did not change in 2009 compared with 2008. Potential explanations for the decreased number of clinical cases were the elimination of so called 'super shedders' and possible exposure of nonvaccinated foals to *L. intracellularis* vaccine organism shed in the faeces of recently vaccinated foals. Under experimental conditions, weanling foals vaccinated intrarectally with an avirulent live vaccine against *L. intracellularis* were protected against clinical and subclinical EPE following challenge exposure with a virulent *L. intracellularis* isolate of equine origin [33]. This was determined by lack of clinical disease, absence of hypoproteinaemia and ultrasonographic abnormalities compatible with EPE and a significant reduction in *L. intracellularis* faecal shedding in vaccinated foals compared with nonvaccinated foals. Further, average daily weight gains from the vaccinated foals over the entire study period were similar to the control foals and significantly higher when compared with the nonvaccinated foals, highlighting the benefit of the vaccine in the prevention of subclinical disease. The extra-label use of the *L. intracellularis* vaccine should be considered on naive and endemic farms in an attempt to reduce or prevent EPE. Timing of vaccine administration should again be synchronised with historical disease occurrence. Further, routine monitoring for clinical signs and hypoproteinaemia/hypoalbuminaemia is still recommended even when vaccine prophylaxis is used.

Conclusions

Equine proliferative enteropathy is an emerging disease with an increasing number of cases reported each year. It also appears that the disease is becoming endemic on an increasing number of horse farms. Equine proliferative enteropathy commonly affects foals less than 12 months of age and is clinically characterised by lethargy, anorexia, fever, peripheral oedema, weight loss, colic and diarrhoea. A presumptive diagnosis of EPE is generally made based on age of the affected animal, clinical signs, hypoproteinaemia/hypoalbuminaemia and presence of thickened small intestinal loops on ultrasonographic evaluation. An *ante mortem* diagnosis is generally confirmed via PCR detection of *L. intracellularis* in faeces or rectal swab and/or serology. Treatment of affected animals consists of supportive and medical treatment including antimicrobials, i.v. fluids, plasma transfusion, parenteral nutrients and

anti-ulcer drugs. Concurrent medical conditions should also be addressed. Treatment of affected foals generally results in clinical improvement, especially if treatment is initiated early in the course of the disease.

The progressive nature of EPE makes this disease a prime candidate to be monitored for in endemic herds. Monitoring strategies combine daily physical evaluation combined with regular measurement of serum protein/albumin concentration and antibodies against *L. intracellularis*. Until new information regarding the epidemiology of this disease is available, preventive strategies are restricted to the use of an avirulent *L. intracellularis* vaccine. The use of the *L. intracellularis* vaccine should be considered on naïve and endemic farms in an attempt to reduce or prevent EPE.

Authors' declaration of interests

No conflicts of interest have been declared.

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Supporting information

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Summary in Chinese.