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ABSTRACT

Parasitic nematodes (roundworms) of livestock have major economic impact globally. In spite of the diseases caused by these nematodes and some advances in the design of new therapeutic agents (anthelmintics) and attempts to develop vaccines against some of them, there has been limited progress in the establishment of practical diagnostic techniques. The specific and sensitive diagnosis of gastrointestinal nematode infections of livestock underpins effective disease control, which is highly relevant now that anthelmintic resistance (AR) is a major problem. Traditional diagnostic techniques have major constraints, in terms of sensitivity and specificity. The purpose of this article is to provide a brief background on gastrointestinal nematodes (Strongylida) of livestock and their control; to summarize conventional methods used for the diagnosis and discuss their constraints; to review key molecular-diagnostic methods and recent progress in the development of advanced amplification-based and sequencing technologies and their implications for epidemiological investigations and the control of parasitic diseases.

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1. Introduction

The phylum Nematoda (roundworms) includes many parasites that are of major socio-economic importance. For instance, grazing ruminants are usually parasitized by one or more nematodes (order Strongylida) which can cause parasitic gastroenteritis (PGE) (Taylor, 2007). Various species of strongylid nematodes can vary considerably in their pathogenicity, geographical distribution and susceptibility to anthelmintic drugs (Dobson et al., 1996). Mixed infections involving multiple genera and species are common, and usually have a greater impact on the host than monospecific infections. In addition, the species composition of the parasites present in a host animal can have an important relationship with the severity of infection (Wimmer et al., 2004). Depending on the number, species and burden of parasitic nematodes, common signs of PGE include reduced weight gain or weight loss, anorexia, diarrhoea, reduced production and, in the case of blood-feeding species, anaemia and oedema, due to the loss of blood and/or plasma proteins (Kassai, 1999; Taylor et al., 2007). Therefore, the knowledge of the nematode species present in a particular geographical area, and their biology and epidemiology, have important implications for the control of PGE, particularly given the increasing problems of anthelmintic resistance (AR) in strongylid nematodes of livestock (Kaplan, 2004, Wolstenholme et al., 2004).

The accurate diagnosis of parasitic diseases and AR is central to these areas and the control of parasites. Traditional methods of diagnosis can be time consuming to perform and have limitations, in terms of their specificity and sensitivity (Gasser, 2006). In particular, in the case of mixed infections, the diagnosis of infection can be laborious and time consuming using techniques such as faecal egg counts (FEC) and larval culture and differentiation (MAFF, 1986). DNA techniques that rely on the amplification of nucleic acids, particularly those coupled to the polymerase chain reaction (PCR) (Saiki et al., 1988), are effective for the specific identification of parasites, and aid the diagnosis of infections from minute amounts of target template, if suitable genetic markers are employed. Such methods are likely to provide powerful alternative tools to traditional approaches, to underpin fundamental research into parasite epidemiology and to improve the control of parasitic disease (Gasser, 2006). The purpose of this article was to: (i) concisely review the biology and significance of gastrointestinal strongylid nematodes of small ruminants; (ii) discuss salient aspects of parasite control and AR; (iii) review traditional methods for the diagnosis of strongylid infections and discuss their limitations; (iv) summarize nucleic acid-based diagnostic techniques, emphasizing recent advances in the establishment of robotic PCR-based technology and its implications.

2. Gastrointestinal strongylid nematodes of small ruminants

2.1. Strongylids and their biology

The order Strongylida includes five superfamilies; the Diaphanocephaloidea, Ancylostomoidea, Strongyloidea, Trichostrongyloidea and Metastrongyloidea. The Strongylida are characterized by the presence of a copulatory bursa and are thus called bursate nematodes (Anderson, 2000). The first four of these superfamilies are monoxenous and predominantly live in the gastrointestinal tract of their vertebrate hosts (Fig. 1). Adult strongylid nematodes exist as females and males; the females produce relatively large numbers (depending on the species) of typically ovoid, strongylid eggs (70-150 µm), which are excreted in the faeces into the external environment. The first-stage larva (L1) develops inside the egg to then hatch (within 1-2 days, depending on environmental conditions) and develops through to the second-stage larva (L2). Both the L1s and L2s feed on bacteria and other microorganisms in the external environment (faeces). After the moults, the ensheathed third-stage larva (L3) develops (usually within 1-2 weeks,
depending on species, temperature, humidity, pH and/or other factors). The cuticular sheath around the L3 prevents it from feeding but protects it from relatively harsh environmental conditions. After the L3 is ingested by the animal and passes through the stomach(s), it exsheaths (xL3) and (after a tissue phase) develops through to the fourth-stage larva (L4) and subsequently the adult at the predilection site in the alimentary tract. The time from the xL3 to the production of eggs by the adult female is usually 3-4 weeks.

2.1. Key nematodes and aspects of disease

Important gastrointestinal strongylid nematodes that infect small ruminants are listed in Table 1. Key nematodes responsible for disease in grazing sheep include *Haemonchus contortus*, *Teladorsagia circumcincta* and intestinal species of *Trichostrongylus* (Besier and Love, 2003). Sheep are usually infected with one or more nematodes, but the severity of disease can vary considerably (e.g., Donald et al., 1978). Disease is predominately linked to factors, such as the species and number of worms infecting the host, the immunological and health state of the host, environmental factors, such as climate and pasture type, stress, stocking rate, management and/or diet (Kassai, 1999; Taylor et al., 2007). Three main groups of animals are susceptible to high intensity infections: (i) young, non-immune animals, (ii) adult, immuno-compromised animals, and (iii) animals exposed to large numbers of L3s from the environment (Zajac, 2006). Nematode populations in sheep are usually over-dispersed, with the majority of sheep having low and only few sheep high intensities of infection, respectively (Barger, 1985).

*Haemonchus contortus* is one of the most pathogenic and fecund strongylid nematodes of small ruminants. Adult females are capable of producing thousands of eggs per day, which can lead to rapid larval pasture contamination and associated outbreaks of haemonchosis (Levine, 1968). In sheep, the pre-patent period of *H. contortus* is 18-21 days. Worms are short-lived, surviving in their hosts for only a few months. The main pathogenic effects are caused by their blood feeding activity, resulting in anaemia which usually becomes apparent after ~ 2 weeks of infection (Baker et al., 1959). Acute disease is usually intensity-dependent and is associated with dark-stained faeces, oedema, weakness, reduced production of wool and muscle mass, or sometimes sudden death. In chronic disease, decreased food intake, weight loss and anaemia are most commonly observed (Kassai, 1999). Unlike many other gastrointestinal nematodes, *H. contortus* is not a primary cause of diarrhoea, and its effects on a flock are not always detected by routine observation (Zajac, 2006).

*Teladorsagia circumcincta* does not feed on blood, and the main pathogenic effect is caused by its larval stages. Larval development takes place in the gastric glands, leading to nodule formation in abomasal mucosa and extensive damage to parietal cells, in turn causing a decrease in hypochloric acid production (Levine, 1968). Subsequently, the increase in abomasal pH causes a failure of pepsinogen to convert to the active form pepsin, which results in elevated plasma pepsinogen levels and reduced protein digestion. The severity of the infection depends on concurrent infections, nutritional state of the host and also the ability to develop an immunogenic response (Stear et al., 2003). Commonly, moderate or subclinical infections occur, causing diarrhoea, poor weight gain, weight loss and reduced wool production (Zajac, 2006).

Species of *Trichostrongylus* represent an important genus in grazing small ruminants. Most species occur in the small intestine and mainly exert their pathogenic effects in lambs and weaners, but have also been reported to cause significant depression of wool growth in old animals (Donald et al., 1978). In Australasia, the three most common species are *T. colubriformis*, *T. vitrinus*, and *T. rugatus* (Beveridge et al., 1989a). Highest pathology is caused by the exsheathed L3s of *T. vitrinus*, which burrow between the intestinal villi and live in sub-epithelial tunnels (Beveridge et al., 1989b). Immature nematodes developing in these tunnels are released 10-12 days following infection. The liberation of young adults is associated with extensive damage to the epithelium, with signs of generalised enteritis, including haemorrhage, oedema and protein loss into the intestinal lumen, and subsequent hypoalbuminaemia and hypoproteinaemia (Taylor, 2007). Low
intensity infections with *Trichostrongylus* can be difficult to distinguish from malnutrition (Taylor, 2007) but, if present at high intensity infections usually cause “black scours” (watery diarrhoea which stains the fleece on the hindquarters) (Levine, 1968). *Trichostrongylus axei*, which inhabits the abomasum, appears to be less common and occurs usually in low intensity (Donald et al., 1978).

*Cooperia curticei, Nematodirus spathiger, N. filicollis, Oesophagostomum venulosum* are common nematodes of the small and/or large intestine, whilst *Chabertia ovina* and *Bunostomum trigonocephalum* (hookworm), are less common (Zajac, 2006). Each of these species have relatively low pathogenicity alone, but can contribute to PGE in grazing small ruminants. *Nematodirus battus* is of particular disease significance in some areas, such as the British Isles, where the mass-hatching of infective L3s occurs during spring, causing disease of young lambs (Taylor and Thomas, 1986); however, this latter species has not yet been reported in Australasia.

**3. Anthelmintic resistance (AR)**

The control of gastrointestinal nematodes relies largely on the use anthelmintics representing three main chemical groups: the benzimidazoles (BZ), the macrocyclic lactones (ML) and the imidazothiazoles/tetrahydropyrimidines (LV) (Besier and Love, 2003, Hoste and Torres-Acosta, 2011). While there has been a recent advance with the development of a new compound, monepantel, from an alternative drug class (amino-acetonitrile derivatives, AADs) (Kaminsky et al., 2008), success in the discovery of new anthelmintics has been extremely limited over the last decades (Kaplan, 2004).

The often excessive and frequent use of these drugs has led to a widespread problem with AR in livestock parasites (Taylor et al., 2009). AR has emerged as a major bionomic and economic problem globally, being currently most pronounced in nematodes of small ruminants (von Samson-Himmelstjerna, 2006, Waller, 1994, 1997). For instance, in Australia, it has been proposed that the prevalence and extent of resistance to all major classes of broad-spectrum anthelmintics is so widespread that it threatens the profitability of the whole sheep industry (Besier and Love, 2003). Therefore, monitoring the AR status of strongylid nematode populations in livestock must be a high priority, and should be an integral part of sustainable parasite control. Various methods, such as faecal egg count reduction test (FECRT), and egg hatch- and larval development assays, have been used for estimating levels of AR in strongylid nematodes of small ruminants, cattle and horses (Coles et al., 1992).

Advances in the diagnosis of AR have focused on the implementation of a standardized protocol for the egg hatch test (von Samson-Himmelstjerna et al., 2009) and a larval migration inhibition test (Demeler et al., 2010a). However, many of these assays are quite time consuming to conduct and suffer from a lack of reliability, sensitivity and reproducibility of test results (Taylor et al., 2002). Therefore, novel approaches of AR diagnosis are required.

Molecular methods have been proposed to provide new alternatives to commonly applied *in vivo* and *in vitro* techniques for the diagnosis of AR, and might be able to overcome some of their limitations (Beech et al., 2011, Demeler et al., 2010a,b, von Samson-Himmelstjerna, 2006). Crucial to the development of molecular diagnostic assays for AR is an in-depth knowledge of the mode of action of these chemicals, their target sites and mechanisms linked to reduced susceptibility to drugs in parasites (Beech et al., 2011, von Samson-Himmelstjerna, 2006, Wolstenholme et al., 2004).

At this stage, the BZ resistance in nematodes seems to be best understood at the molecular level, whilst much less is known about resistances against other classes of anthelmintics (Taylor et al., 2002). A single nucleotide polymorphism (SNP) at codon 200 of the beta-tubulin isotype 1 was believed to be linked to BZ resistance (Kwa et al., 1994, Wolstenholme et al., 2004) and has been demonstrated in resistant strains of *H. contortus* (see Geary et al., 1992), *T. colubriformis* (see Silvestre and Humbert, 2002) and *Te. circumcincta* (see Elard and Humbert, 1999) in sheep. At
least two more SNPs at position 167 and 198 have been detected, but appear to be less common in different species of trichostrongyloid nematodes (Beech et al., 2011, Wolstenholme et al., 2004). Besides the sequence changes in beta-tubulin, which are believed to be the major cause of BZ resistance, recent investigations have suggested a link to the drug transporter P-glycoprotein, hypothesized to play a role in the transport of the anthelmintic away from its site of action and may also select for resistance to MLs (Beech et al., 2011).

Based on current knowledge of the genetic basis of BZ resistance, allele-specific PCRs were developed to determine the genotype of adult worms of *H. contortus* (see Kwa et al., 1994) and *Te. circumcincta* (see Elard and Humbert, 1999). This work was extended by Silvestre and Humbert (2000) by combining the previously described PCR assays with a RFLP procedure, which allowed the phenetic characterisation and identification of L3s of *H. contortus, T. colubriformis* and *Te. circumcincta*. Alvarez-Sanchez et al. (2005) designed a real-time PCR (RT-PCR) assay to assess the frequency of the beta-tubulin isotype 1 allele (linked to codon 200) in nematode samples. As stated by the authors, the diagnosis of BZ resistance using this assay showed an agreement with phenotypic tests, including the egg hatch test and the faecal egg count reduction test (von Samson-Himmelstjerna, 2006).

In spite of these developments, there has been no detailed evaluation of the suitability of these assays using field samples containing mixed species of gastrointestinal parasites, which limits their practical utility at this stage. In addition, all currently employed molecular assays used adult nematodes (only available through necropsy of the host) or infective L3s (requires the culturing of eggs for 1-2 weeks), but none of them has yet been assessed for the detection of AR directly from (mixed populations of) eggs, which would significantly reduce the time required for diagnosis. In contrast to the BZs, the molecular mechanisms associated with resistance to LEV and ML anthelmintics are not yet deeply understood, and recent research has suggested that, in both cases, multiple genes (Beech et al., 2011) are involved in resistance and that resistance is often the result of changes in the parasite other than the immediate drug target, such as transporters and metabolism (Cvilink et al., 2009). Consequently, the multigenic nature of AR and the lack of reliable and universal markers represent a major obstacle to the development of molecular diagnostic tools for AR. No molecular test is yet available for these two groups of broad-spectrum anthelmintics.

### 4. Conventional diagnostic techniques and their limitations

Disease caused by gastrointestinal nematodes manifests itself in a range of clinical signs, including scouring, anaemia, loss of body-condition, and in severe cases death (Hungerford, 1990). The nature and extent of clinical manifestation is also influenced by factors, such as the species and number of worms present, the plane of nutrition and immunological/reproductive status of the host (Hungerford, 1990, Levine, 1968). A number of approaches have been developed for the interpretation of clinical signs linked to PGE; these include body condition- (Russel et al., 1969), “dag”- (Larsen et al., 1994) or anaemia-scoring (van Wyk and Bath, 2002). Although useful as indicators, these clinical approaches are subjective and lack specificity (van Wyk and Bath, 2002).

#### 4.1. Faecal egg counts (FEC)

The counting of eggs from faeces is the most common method for the diagnosis of gastrointestinal nematode infections. This method is inexpensive, easy to perform and does not require specialized instrumentation, making it suitable for use in most diagnostic settings. Important applications of this technique include estimating infection intensity (McKenna, 1987, McKenna and Simpson, 1987), estimating levels of contamination with helminth eggs (Gordon, 1967), evaluation the effectiveness of anthelmintics (Waller et al., 1989), determining the breeding
value of an animal when selecting for worm resistance (Woolaston, 1992), and guiding decisions regarding treatment and control (Brightling, 1988).

This method involves mixing faeces with a saturated salt or sugar solution (e.g., sodium nitrate or sucrose; specific gravity: 1.1-1.3) to float parasite eggs (with the exception of trematode eggs) on the surface of the suspension. An aliquot of this suspension is aspirated and eggs counted, and the number transformed into eggs per gram (EPG). Various methods have been developed, including the direct centrifugal flotation method (Lane, 1922), the Stoll dilution technique (Stoll, 1923), the McMaster method (Gordon and Whitlock, 1939) and the Wisconsin flotation method (Cox and Todd, 1962), of which the McMaster method appears to be most widely used (Nicholls and Obendorf, 1994). In the last decades, numerous modifications of these methods have been described (Levine et al., 1960, Raynaud, 1970, Roberts and O'Sullivan, 1950, Whitlock, 1948), and most teaching and research institutions use their own modifications of original protocols (Kassai, 1999). Modifications include the use of different flotation solutions (and specific gravities), sample dilutions and counting procedures, which achieve varying sensitivities and may complicate the comparisons of FEC results among different laboratories. In addition to these issues, factors, such as variation in biotic potential of different nematode species (e.g., Gordon, 1981, Le Jambre et al., 1971, Martin et al., 1985, McKenna, 1981, Roberts and Swan, 1981, Rowe et al., 2008, Stear and Bishop, 1999), water content (Gordon, 1953, 1981, Le Jambre et al., 2007) and storage/preservation conditions (Nielsen et al., 2010, Rinaldi et al., 2011, Whitlock, 1943) of faeces can each affect the interpretation of test results. Other considerations are that FECs (i) relate to patent but not pre-patent infections (Thienpont et al., 1986), (ii) do not provide any information on male or immature worms that may be present (McKenna, 1981), and (iii) can be influenced by variation in the excretion of eggs by adult worms (Villanua et al., 2006), age of the worm population, and/or the immunity, sex and age of the host (Thienpont et al., 1986). While there are some differences in morphology of eggs between some socioeconomically important nematodes (Georgi and McCulloch, 1989), specific identification is not reliable by routine microscopy (with the exception of, for example, Nematodirus spp.) (Lichtenfels et al., 1997).

FECs alone should not be used to guide treatment decisions, but should be interpreted in conjunction with information about the nutritional status, age and management of sheep in a flock (McKenna, 2002). However, according to common practice, a FEC of ≥ 200 EPG is regarded to indicate a “significant” intensity of infection (www.wormboss.com.au). The value of FEC results also depends on the parasite and host species involved. For example, FEC results for adult cattle are of limited diagnostic value, as they do not usually relate to worm burden (McKenna, 1981); FECs in cattle are usually low and require more sensitive flotation methods than for small ruminants (Mes et al., 2001); for species of Nematodirus, FECs are also regarded to be of limited value, as most pathological damage during infection is caused by the immature stages prior to egg-laying commences (McKenna, 1981). In addition, the detection limit of some flotation techniques is in the order 10-50 EPG (depending on protocol), which can represent a constraint for the diagnosis of AR by FECRT (Levecke et al., 2012).

Nonetheless, attempts have been made by the World Association for the Advancement of Veterinary Parasitology (WAAVP) to improve and implement FEC protocols for the assessment of AR in different species of animals (Coles et al., 2006). In addition, lectin staining for the identification of H. contortus eggs (Palmer and McCombe, 1996), computerized image recognition of strongylid eggs (Sommer, 1996) and automated egg enumeration (Mes et al., 2007) are interesting developments toward improved species identification and differentiation. However, the suitability of the latter two approaches requires rigorous evaluation for routine applications. With the development of FECPAK, a diagnostic test-kit for coproscopic examination (www.techiongroup.co.nz), efforts have been made to provide sheep farmers with a field-based FEC method. However, the implementation of such a method requires the cooperation by farmers, adequate training and quality assurance to ensure that diagnoses are accurate (McCoy, 2005). Also FLOTAC (Cringoli et al., 2010) seems to be a promising FEC method. Once validated for different
host and parasite species, this method might deliver FECs at increased sensitivity (i.e., 1 EPG) and could represent an alternative to current flotation techniques.

4.2. Larval culture (LC)

Larval culture involves incubating faecal samples containing eggs of strongyloid nematodes to allow L1s to hatch and then develop through to L3s; the latter are examined microscopically and differentiated morphologically/morphometrically. A number of protocols have been published which differ in the temperatures, times and media used for culture, and the approach of larval recovery (Dinaburg, 1942, Hubert and Kerboeuf, 1984, MAFF, 1986, Roberts and O’Sullivan, 1950, Whitlock, 1956). The most widely employed protocol includes an incubation at 27 °C for 7 days (MAFF, 1986).

Investigations of the ecology and developmental requirements of various species of gastrointestinal nematodes of livestock (Beveridge et al., 1989a, O’Connor et al., 2006) have shown that different species of strongyloid nematodes require distinct conditions, such as environmental temperature and relative humidity, to develop adequately. This aspect is particularly important to consider when larval culture results are used to estimate the contribution of different species to mixed infections. It has been demonstrated that one culture protocol can favour the development of one species over others (Dobson et al., 1992). For instance, Whitlock (1956) observed that the usual culture condition (27 °C for 7 days) is suitable for most species, but that the free-living stages of Teladorsagia species develop better at somewhat lower temperatures. This statement was supported by the findings of Dobson et al. (1992), who demonstrated that the developmental success of the infective larvae in faecal cultures was lower for Te. circumcincta than for T. colubriformis when cultured alone or concurrently, indicating that LC was unreliable for estimating the contribution of individual species in mixed infections. Berrie et al. (1988) reported similar findings for the bovine parasites H. placei, O. radiatum and Cooperia pectinata. In this study, the recovery of larvae of H. placei was significantly lower compared with the other two species under the same LC conditions. Based on their results, the authors stated that LC and subsequent larval differentiation are unsuitable for an accurate estimation of the proportions of individual species in animals with mixed infections and only provide an indication of the species present (Berrie et al., 1988).

Further variability in LC results have been attributed to differences in the composition of the culture medium, pH, humidity and oxygen (Hubert and Kerboeuf, 1984, Roberts and O’Sullivan, 1950). Therefore, it had been proposed that a defined medium might help to obtain more consistent results (Hubert and Kerboeuf, 1984). To test this proposal, Hubert and Kerboeuf (1984) established a modified method of LC using an “on-agar” approach to provide standardised conditions. LC on agar medium led to higher recoveries of larvae compared with traditional faecal cultures, but lengthy preparation times and increased laboratory requirements appeared to limit the routine application of this method.

In addition to variation in results relating to LC conditions, the differentiation of cultured L3s provides challenges. Differentiation relies on morphological and morphometric parameters, such as the length of the tail sheath extension and total body length of L3s (Dikmans and Andrews, 1933, Gordon, 1933, MAFF, 1986, McMurtry et al., 2000, van Wyk et al., 2004). Various keys for the identification of L3s have been published (Dikmans and Andrews, 1933, Gordon, 1933, MAFF, 1986), but there is an overlap in the body length measurements between some species, and substantial variation in the length of L3s has been reported (McMurtry et al., 2000).

Van Wyk et al. (2004) used the mean length of the tail sheath extension to differentiate L3s of Teladorsagia and/or Trichostrongylus from the larvae of Haemonchus and Chabertia and/or Oesophagostomum. However, although useful to differentiate genera, without the requirement to measure every single larva (thus being more time efficient), this approach has the disadvantage that it does not allow the unequivocal differentiation of all genera. For instance, Teladorsagia and
Trichostrongylus (being the most common genera in winter rainfall areas) cannot be differentiated based on sheath extension length. To further refine the differentiation of these two genera, other morphological features are required. Lancaster and Hong (1987) suggested that the presence of an inflexion (“shoulder”) at the cranial extremity of Teladorsagia larvae was an informative morphological feature. However, this feature is subtle and its detection is subjective. Gordon (1933) proposed the differentiation of Teladorsagia and Trichostrongylus L3s based on the body length. Based on the measurements of 1,000 larvae of each genus, a body length of > 720 μm was used to infer Teladorsagia and ≤ 720 μm for Trichostrongylus. Although practical, this approach requires individual larvae to be measured and does not take into account variability in the length of developing larvae (as a consequence of culture conditions, climate/season, food source for developing larvae and/or immune status of the host) (McMurtry et al., 2000).

McMurtry et al. (2000) described an approach for the differentiation of Teladorsagia from Trichostrongylus L3s, which involves the exsheathment of cultured larvae with sodium hypochlorite and counting of tubercles at the posterior end of the exsheathed L3 using a microscope. As claimed by the authors, this approach allows the differentiation among populations of T. axei, T. colubriformis, T. vitrinus and Te. circumcincta. However, the authors acknowledged that there is a degree of variability in the number of tubercles and that the tails of Te. circumcincta and T. axei lack these structures (McMurtry et al., 2000).

Interestingly, L3s of the large intestinal nematodes Oesophagostomum and Chabertia cannot be differentiated morphologically/morphometrically under a light microscope, which has prevented epidemiological studies of the distribution and prevalence of these genera (and species). A less commonly used method for larval differentiation involves the culture and morphological identification of L1s (Whitlock, 1959). This technique has the advantage of being rapid, since the time required for the development of the L1 stage is shorter; however, the same issues in relation to culture conditions and identification apply to L1s and L2s as to L3s (Lichtenfels et al., 1997). Although routinely used in most parasitology diagnostic laboratories, the technique of LC coupled to larval differentiation by microscopy is time-consuming, laborious to perform, suffers from inaccuracy (see Johnson et al., 1996, Lichtenfels et al., 1997) and cannot readily be automated.

### 4.3. Immunological and biochemical methods

In addition to conventional copro-diagnostic methods, various immunological and biochemical methods have been assessed or established, aimed at the specific diagnosis of infection. These methods rely mainly on the detection and measurement of parameters, such as pepsinogen, gastrin or specific antibody in serum, which might be indicative of parasite infections.

#### 4.3.1. Immunological detection

A number of immunological methods, including those that are based on the detection of an immune response in an infected animal, and those for the detection of parasite antigens, have been developed for the specific diagnosis of parasitic infections (e.g., Engvall and Ruitenberg, 1974, Fletcher, 1965, Ogunremi et al., 2008). Based on the target molecule (antigen or antibody), such methods can be classified as either “direct” or “indirect”.

Direct immunological methods provide direct evidence of an infection and can be based on the detection of parasite antigens present in the circulation and/or excreta from infected hosts. Parasitic extracts have a complex composition and contain molecules that are sometimes shared by other parasites (i.e., are cross-reactive) (Cohen and Sadun, 1976). Shared antigenic composition of closely related parasite species represents a challenge, particularly for nematodes, and often leads to cross-reactivity in immunological tests (Eysker and Ploeger, 2000, Noordin et al., 2005). Also the presence of host materials associated with the parasite can complicate antigen purification and can sometimes interfere with the specificity of a diagnostic assay. Furthermore, the stage of a
parasite, used as an antigen source, can influence immuno-diagnostic results, as parasites undergo significant structural and biochemical changes during their development (Cohen and Sadun, 1976). As an example, the antigenic composition of larval stages differs from that of adults (Williams and Soulsby, 1970) and can give rise to variation in diagnostic specificity and sensitivity (McLaren et al., 1978).

Johnson et al. (1996) described an immunodiagnostic assay for the quantitative detection of excretory/secretory parasite antigens in host faeces (coproantigens). These authors evaluated the usefulness of this approach in a murine model system using *Heligmosomoides polygyrus*, a trichostrongyloid gastrointestinal nematode related to the common nematode species infecting ruminants. The authors also suggested that the enzyme-linked immunosorbent assay (ELISA) was useful for the detection of parasite antigens in the host faeces and might have potential for the detection of pre-patent infections. The diagnostic performance of this assay was promising under experimental conditions, but cross-reactivity, faecal components interfering with the reactivity and the loss of antigens in faeces were reported (Johnson et al., 1996).

Indirect immunological methods are usually based on the detection of anti-parasite antibodies or cell-mediated immune responses in infected hosts. A variety of methods has been developed and applied to the diagnosis of nematode infections, such as the complement fixation test, indirect immunofluorescence, indirect haemagglutination and ELISA, of which the latter has been most commonly used (Doenhoff et al., 2004). However, parasitic helminths possess a huge variety of antigens, and there is limited information on which stages and antigens are actually responsible for eliciting immune responses (Berghen et al., 1993). Antibody detection from serum has several disadvantages, including that it cannot distinguish between a current and past infection, which is a major challenge when evaluating the effects of chemotherapy, does often not reflect infection intensity and sometimes achieves poor specificity, particularly in disease-endemic areas (Doenhoff et al., 2004).

The detection of anti-*Ostertagia* antibodies in the serum of cattle has been found to be useful for epidemiological and cross-sectional studies, but only of limited utility for diagnosis on an individual animal basis (Berghen et al., 1993). Although anti-*Ostertagia* antibodies are detectable in milk samples by ELISA, there are also some limitations to this approach (Charlier et al., 2010). The response to parasitic infections is variable among host individuals, and it has been shown that serum antibody levels can be influenced by factors, such as milk yield, season, mastitis, the number of pregnancies of a cow, stage of lactation and genetic constitution (Gasbarre et al., 1993, Kloosterman et al., 1993, Sanchez et al., 2004). Also the use of bulk milk samples has been investigated, which has the advantage of being an inexpensive and user-friendly approach (Charlier et al., 2010). However, bulk milk samples taken only a few weeks apart can show significant variation in test results, depending on calving patterns, number of cows contributing to the milk in a tank (i.e., dilution effect) and their relative milk yields (Pritchard, 2001).

### 4.3.2. Gastrin or pepsinogen detection

Gastrin is a hormone produced by G-cells in the stomach. Gastrin stimulates parietal cells to secrete acid, and also stimulates pepsinogen secretion, stomach motility and blood circulation in gastric vessels. It was proposed that strongylid nematodes can directly stimulate G-cells, causing an increased gastrin production (Berghen et al., 1993). However, as shown for pepsinogen, the specificity of this approach was questioned (Berghen et al., 1993), because other parasites or factors, such as diet, lactation and/or abomasal lesions, can also effect gastrin levels. Furthermore, in an experimental context, it has been shown that high infective doses need to be administered to parasite-naïve calves to induce a significant rise in blood gastrin (Berghen et al., 1993).

Pepsinogen is a pro-enzyme produced by chief cells of the gastric fundus. It is converted to its active form by acid produced by parietal cells. When parasitized glands of the gastric mucosa are destroyed, the hydrochloric acid production of parietal cells decreases, causing a rise in
abomasal pH and resulting less conversion of pepsinogen to active pepsin (Levine, 1968). Accumulating pepsinogen can escape between disrupted cell junctions into the blood. Therefore, an increase in serum pepsinogen concentration has been regarded to relate to mucosal damage by developing larval stages of Ostertagia (Levine, 1968). Berghen et al. (1993) reviewed the value and application of pepsinogen, gastrin and antibody responses as diagnostic indicators for ostertagiasis and identified a number of potentially limiting factors. The authors suggested that other parasitic or non-parasitic diseases can be responsible for a moderate rise in pepsinogen concentrations in blood, thus limiting the specificity of this approach.

4.5. Post mortem diagnosis

The post mortem diagnosis of infection is usually employed in parasitology to determine the number of nematodes present in the gastrointestinal tract (= intensity of infection), for epidemiological studies or to assess anthelmintic efficacy. These techniques involve the opening and washing of respective parts of the gastrointestinal tract and the examination of subsamples to estimate the total numbers of nematodes present. Various techniques have been described (Eysker and Kooyman, 1993, MAFF, 1986, Robertson and Elliott, 1966); the main differences among them are in the counting of nematodes, the soaking or not of the organ in water or saline (mainly used to recover immature stages), and the proportion of the total volume and the number of aliquots examined (Gaba et al., 2006). Other differences are in the length of the intestinal section examined (proximal 10 meters of small intestine versus the entire length) and the mesh size of the sieve employed to remove plant debris from the washes (McKenna, 2008).

The common practice of examining the proximal 10 m of the small intestine is based on the observation that most intestinal Trichostrongylus spp. occur within the first 6 m of the small intestine (Beveridge and Barker, 1983). McKenna (2008) stated that processing only the first 10 m of the small intestine led to a recovery of < 50% of the worms located in the entire length, resulting in serious underestimates of the total number of worms present. However, the results reported were based on the necropsy of only 15 sheep, and a recovery of less than half of the total number was observed only in a few individual sheep, whereas in most infected sheep trichostrongylid nematodes were located in the proximal 10 m of the small intestine (cf. McKenna, 2008). Therefore, it can be concluded that the improvement of accuracy achieved by processing the entire small intestine is marginal and involves a significant increase in the amount of labour and time required for processing.

Eysker and Kooyman (1993) described a method that involves three parts (contents, immediate water wash of the organ and the saline wash after 5 h of soaking the organ). The disadvantage of this method is that it involves more labour at necropsy, but it has the distinct advantage that worms are separated from the bulk of the gut contents, allowing a rapid worm count. Gaba et al. (2006) assessed their approach for H. contortus and Te. circumcincta and suggested that the estimation of infection intensity, based on gut washes alone, is reliable. However, a prerequisite is that gut sections (e.g., abomasum) are processed rapidly (within 15 min) after the death of the sheep, as some worms progressively start migrating into the contents (Gaba et al., 2006). Gaba et al. (2006) also stated that immediate washing of the gut is insufficient for extracting T. axei or larvae from the mucosa.

Similarly, the selection of mesh size of the sieves used is dictated by the purpose of the counting procedure. The use of a smaller mesh size enables a higher recovery of early L4s, but has the disadvantage that more debris remains in the subsample examined, resulting in prolonged time for counting (McKenna, 2008). Therefore, a small sieve size (e.g., 38 µm aperture) is only required if L4s are counted. If studies are conducted to confirm AR (reflected by a reduced efficacy against adult worms), larger mesh sizes (e.g., 250 µm aperture) can be used (McKenna, 2008).
4.6. The need for standardization

Surprisingly, the performance of most diagnostic tests used routinely for the diagnosis or parasitic infections or disease have not been validated against standards of the Office International des Epizooties (Conraths and Schares, 2006, OIE, 2004). The validation of the performance of any diagnostic test (cf. Tables 2 and 3) is critical and involves the characterisation of basic parameters and can be achieved in number of steps (OIE, 2004). As a first step, a test suitable for a particular use has to be selected, developed and optimised. Subsequently, validation parameters have to be determined, such as analytical sensitivity and analytical specificity (Conraths and Schares, 2006). Following this initial assessment, the diagnostic sensitivity and specificity are determined by examining a larger number of samples for which the true disease or infection status of the animals being tested is known (determined by a “gold standard”). After a test has been evaluated, it may be considered validated (Conraths and Schares, 2006), but a continuous monitoring of test performance during routine application is also advisable in both commercial and research settings.

5. Nucleic acid-based methods for diagnosis

Clearly, conventional methods of diagnosis (reviewed in section 4) have some limitations, in terms of sensitivity and/or specificity. In addition, they can be time consuming and costly to perform. DNA technologies have enabled the development of new, sensitive and specific diagnostic methods that have found applications in parasitology. The ability to specifically identify and study parasites (irrespective of life-cycle stage) using DNA methods has provided new insights into parasite biology, epidemiology and ecology, and has important implications for the specific diagnosis, treatment and control of parasitic diseases (Gasser, 2006). In particular, methods that rely on the enzymatic amplification of nucleic acids can overcome some of the limitations of traditional approaches (Gasser, 2006). Methods that employ the polymerase chain reaction (PCR) (Mullis et al., 1986, Saiki et al., 1988) can selectively amplify in vitro target DNA sequences from complex genomes or matrices, and have led to advances in many areas of the biological sciences.

PCR involves the heat denaturation of double-stranded DNA, followed by a decrease in temperature to allow oligonucleotide primers to bind (= anneal) to their complementary sequence on sense and antisense strands of the target template. Then, the temperature is increased again to enhance the enzymatic activity of a thermostable DNA polymerase, which extends the complementary strands from the primer sites. These synthesis steps are usually repeated 20-40 times in an automated thermal cycler, resulting in an exponential increase in target DNA copies. The major advantage of this methodology is that it enables the study of parasite DNA from minute amounts of template, which would otherwise be insufficient for conventional analysis. The value of this technology in the field of diagnostic veterinary parasitology lies in its ability to specifically identify parasites, detect infection and analyse genetic variation, which are particularly important, given the increasing problems of AR in parasitic nematodes (Gasser, 2006, Gasser et al., 2008).
5.1. Sample processing and PCR inhibition

The selection of the most suitable sample preparation method depends on the type of sample and the purpose of the PCR analysis, as there is no universal method that suits all sample matrices and/or applications (Hoorfar et al., 2004). The main goals of sample preparation are to (i) concentrate the target organisms and the template for subsequent PCR, (ii) eliminate possible PCR-inhibitors, and (iii) produce a homogenous sample for specific and sensitive enzymatic amplification (Rådström et al., 2004). Complex biological (e.g., faecal) samples can contain a wide range of inhibitory substances (e.g., bile salts, collagen, haeme, humic acids and polysaccharides), which are capable of reducing or preventing PCR amplification (Rådström et al., 2004, Wilson, 1997). Different samples can have very different compositions, and the presence of substances potentially inhibitory to the PCR often varies depending on the sample type and composition (Hoorfar et al., 2004, Wilson, 1997). ‘Spike-controls’ (natural or synthetic nucleotide sequences/fragments introduced into samples) can be used to assess the presence of inhibitory substances in the amplification mixture and the efficiency of the DNA isolation and/or PCR reaction (Ninove et al., 2011). Therefore, the selection and evaluation of the sample preparation approach and a suitable reaction mixture, including polymerases and primers, are critical to obtain PCR-compatible samples of comparable composition, irrespective of the variation in the original matrix (e.g., batch-to-batch variation) (Hoorfar et al., 2004).

5.2. Genetic markers for specific identification or detection

The key to developing a reliable PCR method for the specific diagnosis of infection is the definition of one or more suitable DNA targets (genetic marker or locus) based on DNA sequencing. Since different genes evolve at different rates, the DNA region selected should be sufficiently variable in sequence to allow the identification of parasites to the taxonomic level required. For specific identification, the target DNA should display no or minor sequence variation within a species and differ sufficiently in sequence to consistently allow the delineation among species. In contrast, for the purpose of identifying population variants (subspecies, genotypes or “strains”), a considerable degree of variation in the sequence should exist within a species. A range of target regions in the nuclear and mitochondrial genomes have been employed to achieve the identification of parasites to species or sub-specific genotypes (Anderson et al., 1998, Blouin, 2002, Chilton, 2004, Gasser, 2006). In nuclear ribosomal genes and spacers, there is often less sequence variation among individuals within a population and between populations, which makes them suitable as species-specific markers. Hence, in the case of genetic markers for the specific identification of strongyloid nematodes of livestock, most of the focus has been on employing nuclear ribosomal DNA (rDNA).

Although some success was achieved with other DNA targets (e.g., Callaghan and Beh, 1994, 1996, Christensen et al., 1994, Roos and Grant, 1993, Zarlenga et al., 1994), most studies have consistently shown that the first (ITS-1) and second (ITS-2) internal transcribed spacers (ITS) of nuclear rDNA provide reliable genetic markers for the specific identification of a range of strongyloid nematodes of livestock, including species of Haemonchus, Teladorsagia and Ostertagia (abomasum); Trichostrongylus (abomasum or small intestine), Cooperia, Nematodirus, Bunostomum (small intestine); Oesophagostomum and Chabertia (large intestine); Dictyocaulus, Protostrongylus and Metastrongylus (lung) (reviewed by Gasser, 2006, Gasser et al., 2008).

A comparison of the ITS sequences from a range of strongyloid nematodes has shown that the ITS-1 (364-522 bp) is usually larger in size than the ITS-2 (215-484 bp) (see Chilton, 2004). For instance, the ITS-1 region of Ostertagia ostertagi and O. lyrata (801 bp) (Zarlenga et al., 1998b) is longer than that of other trichostrongylids, including congeners, due to the presence of an internal 204 bp fragment, which is repeated twice (Zarlenga et al., 1998a,b, Zarlenga and Higgins, 2001). No major differences have been detected among species of Teladorsagia/Ostertagia in the
lengths of their ITS-2 sequences (Chilton et al., 2001, Stevenson et al., 1996). The G+C content (39-50%) of the ITS-1 sequence of species studied is usually greater than of their ITS-2 (29-45%). The ITS-2 sequences of some species can be relatively A+T-rich (60-70%), which may relate to structural aspects of the precursor rRNA molecule. In addition, studies to date, show that the magnitude of sequence variation in both the ITS-1 and ITS-2 within a species is less (usually <1.5%) than the levels of sequence differences among species (Gasser, 2006), providing the basis for the specific identification of strongylids and diagnosis of infections.

6. Conventional PCR tools

ITS-1 and/or ITS-2 provide useful genetic markers for the development of diagnostic PCR-based tools for strongylid nematodes (Gasser et al., 2006, 2008); in addition to being species-specific in sequence, they are short (usually ≤ 800 bp), repetitive and undergo homogenisation (Elder and Turner, 1995, Gasser, 2006), the latter factors ultimately determining the efficiency, “sensitivity” and specificity of any PCR amplification procedure.

PCR-based SSCP analysis has provided a useful approach for the specific identification of strongylid nematodes using markers ITS-1 and/or ITS-2 and for detecting cryptic (=morphologically similar but genetically distinct) species at any stage of development (Gasser et al., 2006, Gasser and Chilton, 2001). Although there has been a considerable focus on nematodes of humans, there have been some applications to strongylids of livestock (reviewed by Gasser et al., 2006, 2008).

Oligonucleotide primers have been designed to specific regions flanking and/or within the ITS-1 or ITS-2 for diagnostic applications (reviewed by Chilton, 2004, Gasser, 2006, Gasser et al., 2008). Using rDNA targets, this strategy has also been employed for the development of PCR assays for the genus- or species-specific identification of different developmental stages of strongylid nematodes. For instance, Zarlenga et al. (1998a) described the development of a semi-quantitative PCR assay for the diagnosis of patent Ostertagia ostertagi infection in cattle. Conserved oligonucleotide primers were used in PCR to amplify a ~1 kb rDNA region, spanning the ITS-1 and part of the 5.8S rRNA gene, from *O. ostertagi*, whereas amplicons of ~600 bp were amplified from *H. contortus*, *C. oncophora* and *Oe. radiatum*. When DNA samples derived from adult nematodes of the different genera were mixed and amplified simultaneously, there was no evidence of inhibition in the PCR, and *O. ostertagi*-specific amplicons were readily detected electrophoretically. There was a correlation between the intensity of the ~1 kb and 600 bp amplicons on gels and the percentage of *O. ostertagi* DNA within the mix of heterologous species. There was also a strong correlation between the percentage of *O. ostertagi* DNA and percentage of *O. ostertagi* eggs in the faeces. Effective amplification was achieved from 5% of the genomic DNA isolated from a single *O. ostertagi* egg. Hence, the establishment of this PCR assay had major implications for diagnosis of patent *O. ostertagi* infection in cattle as well as for studying the epidemiology of this parasite. Other studies (reviewed by Gasser, 2006, Gasser et al., 2008) have demonstrated the diagnostic utility of PCR assays using species-specific ITS oligonucleotide primers or probes, even when the sequences of related species differ by a single nucleotide (Hung et al., 1999). For instance, Zarlenga et al. (2001) extended previous work to develop a multiplex PCR assay for the specific detection and differentiation of economically important gastrointestinal strongylid nematodes (including *H. placei*, *O. ostertagi*, *Trichostrongylus* spp., *C. oncophora/Co. surinabada* and *Oe. radiatum*) of cattle.
7. Real-time PCR (RT-PCR)

7.1. Principle

RT-PCR was developed in the early 1990s (Higuchi et al., 1992) and allows enzymatic amplification to be monitored in real time in vitro. All current RT-PCR systems detect the amplification using fluorescent dyes or probes. The predominant advantages of real-time PCR over conventional PCR are that it allows high throughput analysis in a “closed-tube” format, not requiring handling or electrophoresis following amplification, that it can be employed for quantitation over a wide “dynamic range” and that it can be used to differentiate amplicons of varying sequence(s) by melting-curve analysis.

The principle of the original method was to incorporate a specific, intercalating dye (e.g., ethidium bromide) into the PCR to measure the change in fluorescence after each cycle using a digital camera and a fluorometer coupled to the reaction tube (Higuchi et al., 1993). The technique has been modified to include other (non-carcinogenic) dyes, such as SYBR Green I (Becker et al., 1996), LCGreen (Wittwer et al., 2003), SYTO9 (Monis et al., 2005a) and EvaGreen (Wang et al., 2006). Real-time PCR assays using such dyes enable the relative or absolute quantitation of amplicons by allowing the identification of the cycle (Ct) at which the amplification commences. One or more DNA standards (of differing concentrations) and test samples are subjected to cycling at the same time and their Ct values established and compared. Standard curves can be constructed based on the use of reference samples, and the relative amounts of template in test samples are calculated in relation to these curves.

Intercalating dyes, such as SYBR Green I, detect any double-stranded DNA, which is advantageous because they can be incorporated into any assay. However, a disadvantage is that the dye binds to all double-stranded DNA in a reaction, which includes primer dimers and non-specific products. This limitation can be overcome by acquiring fluorescence data at a temperature that denatures the non-specific products and leaves the specific products intact. The melting point of an amplicon is linked to the composition and length of the nucleotide sequence(s) within it, which means that a melting-curve analysis can be used to detect and/or characterize sequence variation within and among samples. Other recent advances include the complementary use of high resolution melting-curve (HRM) analysis following RT-PCR (e.g., Jeffery et al., 2007). Melting analysis using the dye LCGreen or SYTO9 has been reported to achieve acceptable levels of reproducibility, attaining better mutation detection capacity than SYBR Green I (Monis et al., 2005a, Wittwer et al., 2003). Alternative, more expensive detection systems (other than intercalating dyes) include Taqman probes (Heid et al., 1996), minor groove binder (MGB) Eclipse probes (Afonina et al., 2002), molecular beacons (Piatek et al., 1998) and fluorescence resonance energy transfer (FRET) (Chen and Kwok, 1999), ensuring specificity in the PCR through exclusive binding to the target sequence (Monis et al., 2005b).

7.2. RT-PCR assays for the diagnosis of strongylid nematode infections

In spite of promising results of RT-PCR for the diagnosis and quantification of selected protozoan and metazoan parasites (Bell and Ranford-Cartwright, 2002, Monis et al., 2005b, van Lieshout and Verweij, 2010, Zarlenaga and Higgins, 2001), to date, relatively little research has focused on its use for the diagnosis of strongylid infections of livestock (cf. Gasser, 2006, Gasser et al., 2008). There have been attempts to use RT-PCR for the specific diagnosis and/or quantification of helminth eggs or larvae from the faeces from infected hosts. First efforts were made by von Samson-Himmelstjerna et al. (2002, 2003), who developed RT-PCR assays for the diagnosis and quantification of ovine gastro-intestinal nematodes, including *H. contortus*, *O. leptospicularis*, *T. colubriformis*, *Co. curticei* and for small strongyles (cyathostomins) of horses.
However, these assays were used for the identification of larval or adult nematodes only, which limited their utility for routine diagnostic applications.

Harmon et al. (2007) evaluated the use of RT-PCR to quantify eggs of *H. contortus* from sheep faeces and examined various aspects, such as the influence of faecal inhibitors on PCR, the effects of competing and non-competing DNA in multiplex reactions and the impact of embryonic development within the egg on the PCR result. The assay developed showed linear quantifiable amplification of DNA obtained from egg quantities ranging from five to 75 eggs, whereas DNA from higher egg numbers of 75-1000 eggs did not show significant differences in Ct, limiting the quantitative capacity of the assay to a narrow detection range (Harmon et al., 2007). During this study an impact of egg embryonic development on Ct values has only been observed between 0 and 6 h of development at 21 °C, whilst later time periods at 6, 12, and 30 h did not show statistical differences in Ct when compared to each other (Harmon et al., 2007). Non-competitive DNA, derived from environmental sources, did not appear to have a negative impact on amplification, but in multiplex reactions, the presence of high amounts of competing *Trichostrongylus* DNA hindered the amplification of a different target species whose DNA is present at much smaller amounts (Harmon et al., 2007).

The storage of faecal samples is often necessary in a practical context, but the possible impact of egg embryonation during prolonged storage is known to be a critical factor relating to the accuracy of quantifying egg numbers by RT-PCR (Harmon et al., 2007; Bott et al., 2009). Therefore, approaches to circumvent this issue should be developed, which could possibly involve allowing maximum development to occur prior to DNA isolation (Harmon et al., 2007). It has been proposed that the method used for DNA extraction and the presence of PCR inhibitors might be responsible for discrepancies in the linear correlation between DNA amount and number of nematode eggs (Harmon et al., 2007). Harmon et al. (2007) suggested to account for the variability arising from DNA extraction and the presence of faecal inhibitors through the use of multiplex PCR systems that quantify, in relative terms, egg numbers using Ct values, and include an exogenous DNA template to standardize Ct values and assess every sample individually for faecal inhibitors (Harmon et al., 2007). Additional work in this area had been undertaken by two other research teams, who developed RT-PCR assays for the diagnosis of infections with the human hookworms *Ancylostoma duodenale*, *Necator americanus*, and the nodule worm *Oe. bifurcum* (Verweij et al., 2007) and the equine parasite *Strongylus vulgaris* (see Nielsen et al., 2008). These assays employed specific primers and *TaqMan* probes to target the ITS-2 region of nuclear ribosomal DNA. Verweij et al. (2007) suggested that false-negative RT-PCR results (*n* = 2) in relation to LC might be explained by the amount of faeces used for DNA isolation being 30-times less than that used to set up LC. However, both assays were reported by the authors to be of high analytical specificity and of a sensitivity superior to that of LC. Inhibition by faecal components was not evident. A potential limitation of these studies was that the specificity of these assays was based exclusively on the design and use of *TaqMan* probes. However, HRM or sequencing was not used to verify the identity or specificity of the amplicons produced.

Bott et al. (2009) established a combined microscopicRT-PCR method that allows the semi-quantification of strongylid infections in sheep. During this study specific oligonucleotide primers were designed to ITS-2 and 28S rDNA regions of seven key genera or species of strongylids of sheep, including *H. contortus*, *Te. circumcincta*, *Trichostrongylus* spp., *Co. oncophora*, *C. ovina*, *Oe. columbianum* and *Oe. venulosum*, and used in separate PCR reactions. To determine relative proportions of species/genera contributing to a FEC, standard curves were prepared for the RT-PCRs for individual species and demonstrated a log-linear relationship over four orders of magnitude. The Ct values obtained from species-specific PCR reactions showed a linear correlation to the numbers of eggs present per gram of faeces and demonstrated the applicability of this PCR approach for semi-quantification of target species. For some of the primer pairs used in this study, as little as 0.1-2 pg of DNA was sufficient to achieve specific amplification from the respective species, which equates to a proportion of genomic DNA which can be isolated from a single egg.
(Bott et al., 2009). In the evaluation of this PCR assay, all amplicons generated from specific primer pairs were examined by HRM and sequence analysis to confirm their identity. Designed primers were critically assessed for their analytical specificity (i.e., against a broad range of parasites that are known to be detectable from the faeces of infected sheep, including lungworms), and there was no evidence of non-specific amplification. However, Bott et al. (2009) stated that, due to possible sequence polymorphism or heterogeneity of ITS-2 among or within individuals from different geographical locations, the performance of the PCR platform might need additional assessment, if applied in different countries or regions.

Bott et al. (2009) also discussed points that needed consideration for future applications, such as the effect of faecal consistency on FEC and PCR results. The testing of loose/diarrhoeic and desiccated faecal samples can lead to an under- and over-estimation, respectively, of FECs (Le Jambre et al., 2007) and likely variability in semi-quantitative PCR results. Furthermore, Bott et al. (2009) discussed the need for rapid DNA isolation from faecal samples following their collection. The results of previous studies (Harmon et al., 2007) indicated that mitosis during the larval development leads to an increase of ITS-2 copy number and results in enhanced amplification during the RT-PCR. Because the storage and transport of samples at ambient temperatures is often necessary for practical reasons, approaches for the preservation of faecal material, for example, ethanol fixation might be applied. In addition, the direct extraction of DNA from faeces, using commercially available kits, has been proposed as an alternative to methods that involve the concentration of eggs by faecal flotation prior to DNA isolation (cf. Bott et al., 2009, Nielsen et al., 2008). However, such direct extraction methods need to be critically assessed for their ability to remove potential inhibitors (e.g., humic acids, phenolic compounds or polysaccharides) from faeces. A faecal flotation and egg isolation approach has been shown to remove such inhibitors (Bott et al., 2009), and has the advantage that it provides a FEC, which can be compared with a PCR result but which is not possible employing a direct DNA isolation-coupled PCR method. Furthermore, flotation allows for a concentration of eggs from multiple grams of faeces prior to DNA isolation and PCR, thus increasing the likelihood of amplifying DNA from very small numbers of nematode eggs (Bott et al., 2009, Nielsen et al., 2008). By contrast, only small amounts of faeces (e.g., ~0.25 grams) are used in most commercially-available, direct faecal DNA isolation methods, limiting the ‘sensitivity’ of subsequent PCR. Noting these limitations, the combined microscopic-molecular method (Bott et al., 2009) was established for the specific diagnosis of patent strongylid infections in sheep, and future work is required to evaluate the performance of this method for the specific diagnosis of infections with immature (pre-patent) or hypobiotic stages (e.g., Te. circumcincta or H. contortus) of nematodes and to compare it with direct amplification from DNA isolated directly from faecal samples.

7.3. Critical evaluation and application of RT-PCR to assess the composition of strongylid nematode populations in sheep

Roebert et al. (2011) critically evaluated the performance of the RT-PCR method (Bott et al., 2009) for the diagnosis of naturally acquired strongylid nematode infections in sheep (n = 470; in a temperate climatic zone of south-eastern Australia), using a panel of 100 ‘negative control’ samples from sheep known not to harbour parasitic helminths. The authors compared the diagnostic sensitivity and specificity of this RT-PCR assay with a conventional faecal flotation method. They also established a system to rank the contribution of particular strongylid nematodes to the faecal egg counts (FECs) from ‘mixed infections’ in individual sheep. The testing of faecal samples revealed that Te. circumcincta (80%) and Trichostrongylus spp. (66%) were most prevalent, followed by C. ovina (33%), Oe. venulosum (28%) and H. contortus (1%). For most sheep tested in this study, Te. circumcincta and Trichostrongylus spp. represented the largest proportion of strongylid eggs in faecal samples from individual sheep. This was the first large-scale prevalence survey of gastrointestinal nematodes in live sheep utilizing a molecular tool. The ability to rapidly rank strongylid nematodes according to their contribution to mixed infections
represents a major advantage over routine flotation methods. The conclusion from this study was that this RT-PCR tool might be able to replace the conventional technique of larval culture.

This assessment showed that the RT-PCR assay achieved high diagnostic sensitivity (98%) and specificity (100%), and the test results were in ‘good’ agreement (i.e., Kappa: 0.95) with those achieved using conventional faecal flotation. In addition, of 53 field samples which were test-negative based on coproscopy, 23 tested positive by PCR for one or more target nematodes (a result which was confirmed by sequencing), demonstrating better sensitivity of the molecular approach compared with coproscopic examination, and reinforcing that microscopy is not an adequate reference technique [i.e., is an imperfect gold standard (cf. Conraths and Schares, 2006)] for the detailed assessment of the sensitivity and specificity of a diagnostic assay. The results achieved demonstrated that the prevalences of key genera/species, such as *Te. circumcincta* and *Trichostrongylus* spp., known to be the dominant species infecting sheep in the winter rainfall environment of Victoria, Australia, were largely in accordance with information available in the published literature (e.g., Anderson, 1972, 1973). The application of an ordinal ranking system to estimate the contribution of individual parasites to observed FEC results showed that these genera/species were also responsible for the largest proportion of strongylid eggs in the faecal samples tested. Although known to be abundant in winter rainfall environments, *C. ovina* and *Oe. venulosum* were found at high prevalence (33.6% and 28.7%, respectively). An unexpected finding was that *Oe. venulosum* was the main contributor to the observed FEC results for one of nine farms tested, which has important implications for the interpretation of FECs and anthelmintic control.

According to common practice (Brightling, 1988), FEC results of ≥ 200 eggs per gram (EPG) are considered to relate to a ‘significant’ worm burden, and without further considerations of the species present and their reproductive capacity, give an indication for anthelmintic treatment. This common practice, which involves frequent and, in many cases, unnecessary or excessive administration of anthelmintic drugs can promote AR development in gastrointestinal nematodes of sheep and other hosts, as recent evidence has shown (Kaplan, 2004, Wolstenholme et al., 2004). Given that *Oe. venulosum* is recognized to be less pathogenic than most strongylids of the upper alimentary tract (Donald et al., 1978) but has high fecundity (Gordon, 1981), FEC results (e.g., > 200 EPG) in which *Oe. venulosum* is the sole or main contributor would be misinterpreted, and sheep harbouring this relatively non-pathogenic would be treated unnecessarily. Therefore, the specific/generic identification of infecting nematodes assists the interpretation of FEC results and, subsequently, treatment decisions, thus, directly contributing to efforts of preserving the efficacy of currently available anthelmintics.

The RT-PCR assay (Roeber et al., 2011), coupled to conventional coproscopy, and the microscopic detection of *Nematodirus*, *Trichuris* and *Moniezia* eggs in faecal samples revealed that the majority of sheep investigated were parasitized by two to four helminth taxa per animal. Data from this study provided new and important insights into the composition and distribution of nematodes, which would not have been achievable in such detail using any of the currently used coprological methods. The results indicated that this tool should be applicable in other climatic regions and/or major sheep producing countries in the world. Depending on the nematode species infecting sheep in a particular climatic zone, minor modifications could be made to the molecular assay to adapt it for the diagnosis of infections with other important parasites (e.g., hookworm or lungworm) and to provide opportunities to study their biology, prevalence and distributions. For instance, as the life-cycles of some lungworms, such as *Muellerius capillaris* and *Protostrongylus rufescens*, involve invertebrate intermediate hosts, such as snails and slugs, an adapted RT-PCR assay could be used to examine the prevalence and relative intensity of these parasites in their intermediate hosts and to study their ecology.

The ability to identify helminth species and to rank them according to their contribution to FEC results (Roeber et al., 2011) represents a novel approach that is time- and cost-efficient compared with classical diagnostic techniques, and enables a better interpretation of FEC results, particularly in relation to the anthelmintic treatment of infected sheep (cf. McKenna, 1996, 1997).
This advance in the diagnosis of gastrointestinal nematode infections could directly and significantly contribute to enhanced parasite control.

7.4. Evaluation of RT-PCR to replace larval culture (LC) and support faecal egg count reduction testing (FECRT)

Roebber et al. (2012a) assessed the RT-PCR assay to support the diagnosis of AR in nematodes, in conjunction with conventional FECRT; in addition, a direct comparison of PCR results with those of larval culture (LC) and selective total worm counts (TWC; considered a “gold standard”) was undertaken. In this assessment, the molecular assay achieved a diagnostic sensitivity of 100% and specificity of 87.5% in relation to TWC. These percentages were similar to those achieved previously (Roebber et al., 2011) (diagnostic sensitivity 98% and specificity 100% in relation to FEC), demonstrating that the RT-PCR assay consistently achieved a high diagnostic performance. DNA sequencing results also demonstrated that this molecular assay had a better sensitivity than the routinely used TWC method and, together with FECRT, was of practical value for the detection of albendazole resistance in *Te. circumcincta* and *T. colubriformis* populations. However, although the PCR test results were in accordance with TWC, the direct comparison of molecular and LC results showed markedly different findings, depending on the recommended measurements used for larval differentiation (Dikmans and Andrews, 1933, Gordon, 1933, McMurtry et al., 2000). Using the morphometric criteria given by Dikmans and Andrews (1933), the majority of L3s from the LC of the albendazole treated group of sheep were identified as *Trichostrongylus*, whereas, using the measurements recommended by Gordon (1933) and McMurtry (2000), the same larvae were identified as *Teladorsagia*. This discrepancy emphasizes the complications and errors associated with the use of LC, which can readily lead to misinterpretations as to which nematodes are resistant to a particular drug. Overall, this study (Roebber et al., 2012a) demonstrated clearly that the molecular assay coupled to FECRT provides a rapid, efficient and universally applicable tool for the diagnosis of AR and the early detection of residual populations of worms in sheep following treatment. Future studies should be conducted to test sheep on different farms and the response of gastrointestinal nematodes to the treatment with other main groups of anthelmintics, such as imidazothiazoles/tetrahydropyrimidines or macrocyclic lactones, and also the newly developed monepantel (Zolvix, Novartis) (Kaminsky et al., 2008).

The movement of sheep and their gastrointestinal parasites between or among farms favours the spread of drug resistance (Blouin et al., 1995). Therefore, the routine use of the RT-PCR assay, in conjunction with FECRT, could provide a universally applicable method to test sheep before transport to and/or introduction on to a new farm, in order to reduce the spread of drug resistant populations of nematodes. In addition, the assay allows the identification of sheep shedding large numbers of parasite eggs in faeces and, in conjunction with information about the infecting helminth species, can be used to support “targeted, selective treatment” approaches (Kenyon et al., 2009). Such a strategy focuses on treating only sheep that will benefit most from an anthelmintic treatment, whereas sheep with low-intensity infections remain untreated to provide refugia for the dilution of resistance genes within a parasite population. Furthermore, the present RT-PCR assay had a similar or improved sensitivity compared with post mortem diagnosis, thus being able to replace the latter. In practice, this means that the presence of particular species/genera and their prevalence can be assessed reliably without the need to kill sheep.

8. Multiplexed-tandem PCR (MT-PCR) for specific diagnosis

8.1. Rationale and establishment of MT-PCR
A limitation of RT-PCR developed (Roeber et al., 2011, 2012a) was that it employs individual primer pairs for individual specific or generic amplifications and requires numerous, manual handling steps throughout the entire procedure. Therefore, the goal was to develop a user-friendly and practical platform that would allow the rapid testing of large numbers of samples at relatively low cost with limited manual intervention, and that could be introduced into a routine testing laboratory. Although conventional multiplex RT-PCR (e.g., Chamberlain et al., 1988) seemed to be a promising prospect at first, preliminary work conducted (Bott and Gasser, unpublished) showed that primer sets used in individual PCR runs (cf. Bott et al., 2009) could not be incorporated into a single reaction to achieve specific amplifications. Other restrictions of a conventional multiplex PCR approach are that each target sequence requires a probe for fluorescence-based detection at a particular wave-length and that such probes are costly and require the use of multi-channel RT-PCR thermocyclers - which usually have only four to six distinct wave-length channels, thus limiting the number of species/genera that can be detected.

To circumvent these issues, Roeber et al. (2012b) explored the use of multiplexed-tandem PCR (MT-PCR) (Stanley and Szewczuk, 2005). MT-PCR consists of two amplification phases: (i) a primary ‘target enrichment’ phase (through a small number of PCR cycles) conducted using multiplexed primer sets, and (ii) a subsequent analytical amplification phase (utilizing a diluted product from the primary amplification as a template), consisting of the targeted amplification, in tandem rather than by multiplex, of each genetic locus using specific, nested primers. Because the initial amplification phase is limited to 10-15 cycles, interactions between or among multiplexed primer sets is minimized, reducing competition or the generation of artefactual products and limiting amplification bias, which would otherwise prevent downstream quantification (Stanley and Szewczuk, 2005). Because the primary amplicons are diluted (e.g., 100-fold) prior to use as templates in the secondary phase, primer carry-over and PCR inhibition are substantially reduced. By conducting the secondary (analytical) amplification phase in tandem, the method can be coupled to a single-channel, RT-PCR thermocycler, allowing rapid screening and quantification employing one fluorogenic dye (e.g., SYTO-9), thus reducing the cost associated with detection.

Roeber et al. (2012b) established a high throughput MT-PCR assay for the diagnosis of nematode infections in sheep, and critically assessed its diagnostic sensitivity and specificity relative to RT-PCR as well as conventional LC using faecal samples from different flocks of sheep from a broad geographical range in Australia. The MT-PCR achieved high diagnostic specificity (87.5%) and sensitivity (100%) based on the testing of a panel of 100 faecal DNA samples from helminth-free sheep and 30 samples from sheep with infections confirmed by necropsy. This MT-PCR assay was then used to test 219 faecal samples from sheep with naturally acquired infections from various geographical localities within Australia, and results were compared with those of LC, using 139 of the 219 samples. The MT-PCR and LC results correlated significantly for most nematodes examined, but parasites of the large intestine were significantly under-represented in the LC results. The findings showed that Trichostrongylus spp. (87%), Te. circumcincta (80%) and H. contortus (67%) had the highest prevalences, followed by Oe. venulosum (51%) and C. ovina (12%). Importantly, this MT-PCR allowed a species- or genus-specific diagnosis of patent nematode infections to be made within 24 h (compared with 7-10 days for LC).

The evaluation of two different pre-PCR genomic DNA isolation methods showed that a combined egg flotation and column-purification approach achieved better sensitivity, overall, compared with direct faecal DNA isolation (Roeber et al., 2012b). The testing of samples from sheep from different geographical locations in Australia showed that the prevalence of the key nematodes investigated were consistent with their presumed distribution, based on historical data and many years of studies conducted using routine diagnostic procedures (Donald et al., 1978). Epidemiological information had not been provided in such detail prior to the use of a molecular diagnostic tool, and the results achieved by routine LC and the molecular diagnostic platform showed significant correlations ($r_s = 0.69-0.83$) for most nematodes. However, C. ovina and Oe. venulosum were both much less frequently detected in LC than by the molecular assays. Thus, the
evidence showed that LC was unreliable for the diagnosis of infections with these two nematodes, and did not allow their prevalence and distribution to be studied with any confidence. Importantly, the inability of LC to detect the latter two species subsequently led to an overestimation of other nematodes, namely *H. contortus*, *Trichostrongylus* and *Teladorsagia*, present in the LCs examined, thus providing a biased result. This finding also highlights the utility of the present molecular-diagnostic platform for epidemiological studies and suggests that results obtained previously using traditional diagnostic techniques might have led to inaccurate information on the prevalence and distribution of particular species of gastrointestinal nematodes. The findings of Roeber et al. (2012a,b) showed that *C. ovina* and *Oe. venulosum* may contribute much more significantly to infections in sheep than currently acknowledged. Given that the pathogenicity of these parasites is considered low (Donald et al., 1978), it is likely that the inability of classical techniques to reliably identify and quantify these parasites has led to unnecessary anthelmintic treatments at increased cost to the farmer and an increased risk of AR in nematode populations.

8.2. Applications and implications

An important future application of the MT-PCR platform could be the testing of pasture samples for infective larvae of strongylid nematodes. This approach has significant implications for studying the epidemiology and ecology (i.e., the seasonal occurrence of species and length of larval survival) of gastrointestinal nematodes in livestock, and enables assessments of the extent of contamination and risk of infection. For this purpose, pasture foliage could be collected, washed and concentrated; DNA could then be isolated from the larval suspension and molecular testing conducted. However, as the number of cells increases during the mitotic division in developing eggs and larvae, the number of ITS-2 copies also increases, which means that the MT-PCR platform will need to be calibrated to accommodate this change in copy number. This PCR platform permits different sensitivity settings (based on the number of amplification cycles in the primary PCR) and, whilst a medium sensitivity (15 cycles of primary amplification) was shown to produce the best results for the detection of eggs from faeces, this setting could be adjusted for L3s from pasture samples.

There are numerous other possible applications of a high throughput MT-PCR assay. For instance, such an assay could also be used to assist in evaluating the efficacy of vaccines by monitoring FEC, either at the stage of initial vaccine development or, later, in field trials, to predict levels of protection against particular parasite species in individual host animals of different genetic backgrounds (i.e., different breeds of sheep). Immunological methods could also be used in concert with FECRT to measure protective effects of anti-nematode vaccines for sheep following challenge infection/s (with reference to well-defined controls). Moreover, to further refine the diagnosis of AR, genomic DNA regions linked to resistance (e.g., beta-tubulin gene) could be integrated as targets into the assay. However, as AR is frequently polygenic (Beech et al., 2011) and/or can also be associated with drug transport mechanisms (Cvilink et al., 2009), most mechanisms of resistance are only partially understood for many drug classes (cf. Taylor et al., 2002), and more research needs to focus on identifying reliable DNA markers for the specific detection of AR.

There is now major potential to extend the use of MT-PCR assay to other socioeconomically important pathogens, including helminths (e.g., flukes and lungworms), protists (e.g., *Cryptosporidium*, *Eimeria* and *Giardia*), bacteria (e.g., *Campylobacter*, *Escherichia*, *Salmonella* and *Yersinia*) and viruses (e.g., Rota- and Corona-viruses). Furthermore, such a platform could also be adapted, with modifications being made to the DNA isolation procedure and primers used, for the diagnosis of infections with blood pathogens (e.g., *Babesia* and *Theileria*) or the detection of pathogens in environmental samples. In addition, although we have focused on the development of this platform for the detection and identification of pathogens of socioeconomic importance in sheep (Roeber et al., 2012b), a similar approach would be of major benefit assessing
such pathogens in other hosts, including other small ruminants, cattle, horses and even humans. For example, estimating the number of nematode eggs in the faeces from adult cattle is challenging, as eggs are generally present in low numbers and, thus, require more sensitive techniques of diagnosis (Agneessens et al., 2000, Mes et al., 2001). Therefore, the MT-PCR-coupled diagnostic platform could be further enhanced by automating the counting of eggs in faecal samples prior to DNA isolation. Approaches that involve the recovery of nematode eggs from faeces and the automated counting of all eggs present in a sample (as opposed to the counting of eggs from sub- aliquots of a sample and their extrapolation to the total sample volume) by image recognition have been developed (Mes et al., 2001, 2007). Consequently, with only minor modifications, the MT-PCR-based platform could provide a major advance in the diagnosis of nematode infections in cattle.

The MT-PCR platform (Roeber et al., 2012b) could also be further enhanced through the use a robotic DNA isolation procedure. Automated DNA isolation platforms are readily available from different manufacturers and include easyMAG (BioMerieux, Mary l’Etoile, France), m2000sp (Abbott, Abbott Park, IL, USA), MagNA Pure LC 2.0 (Roche) and QiaSymphony (Qiagen, Hilden, Germany), but their suitability for the isolation of DNA from nematode eggs from faecal samples would need to be critically assessed before they could be incorporated into the present MT-PCR system. Further advantages of this approach include the possibility for large-scale, targeted collections and long-term storage of DNA samples. Unlike traditional copro-diagnostic testing, in which samples are usually examined only once to enumerate parasite eggs or larvae, and are then discarded. The use of DNA samples has the advantage that the sample material collected can be stored (at – 80 °C) in a clean and space efficient manner (i.e., either frozen or in dehydrated form), allowing the testing of the samples many years or even decades later. This provides the opportunity of carrying out (even retrospectively) studies of “new” or “emerging” pathogens. These are important applications in times of changing environmental conditions (e.g., climate change, urbanization and associated changes in pathogen transmission, distribution and epidemiology), increased problems with AR as well as the increased mobility of humans and/or animals between/among different regions or countries.

9. Prospects for field-based assays

PCR-based technologies have become central to the diagnosis of parasitic and other infectious diseases. However, most PCR methods are laboratory based and often require relatively expensive and specialized instrumentation and reagents, a clean laboratory to avoid contamination during processing and testing and skilled operators with some technical knowledge in molecular technologies. Therefore, most PCR assays are not applicable to use in the field, which represent a limitation for application in resource-poor countries or in situations where a rapid ‘on-the-spot’ diagnosis is required. Recent developments provide possibilities for the miniaturization and automation of devices for diagnostic testing.

The time required to perform PCR (typically 90-120 min) depends, to large extent, on the ability of the instrument to cycle rapidly through the denaturation, annealing and extension steps (deMello, 2003). By reducing the thermal mass of the instrument, the total reaction times can be significantly shortened. Two different parameters can be altered to achieve this: (i) the physical dimensions of a system can be reduced, in order to reduce the thermal mass of the instrument, and (ii) a sample can be moved through multiple reaction zones which are held at specific temperatures required for the in vitro amplification of nucleic acids (deMello, 2003). The latter approach allows the heating or cooling of small fluid elements which move through different temperature zones within 100 msec, thus allowing ultra-rapid reaction times and making the functional integration of PCR into micro-chips possible (deMello, 2003). Obeid et al. (2003) developed such a microchip that demonstrated efficient amplification of DNA within 5 min and, additionally, provides a separate channel for reverse transcription, in which RNA samples can be transcribed into DNA.
before entering the PCR zone of the chip for amplification. Furthermore, the continuous injection of small sample volumes (1-2 µl), separated by water and air plugs, allows the simultaneous amplification of multiple samples without cross-contamination (deMello, 2003).

Isothermal amplification methods provide prospects for the design of simple, portable and low-energy consuming operating systems (Asiello and Baeumner, 2011). Some of these methods include nucleic acid sequence-based amplification (NASBA) (Compton, 1991), loop-mediated isothermal amplification (LAMP) (Notomi et al., 2000), helicase-dependent amplification (HDA) (Vincent et al., 2004), rolling circle amplification (RCA) (Liu et al., 1996) and strand displacement amplification (SDA) (Walker et al., 1992a,b). Advantages of such methods seem to be that they reduce costs, simplify the use of amplification reactions by eliminating laboratory equipment, prevent contamination and provide the potential to run several reactions in parallel. Furthermore, miniaturized test systems can be combined with integrated steps of sample preparation and detection of amplification products, thus providing scope for the design of monolithic diagnostic systems, also referred to as micro-total analysis systems (µTAS) or labs-on-a-chip (LOC) (Asiello and Baeumner, 2011). Although, these technologies appear to be in their early phase of development and require further research to translate into practical diagnostics, there are some reports demonstrating their utility. For example, Liu et al., 2011 developed a point-of-care diagnostic system, which is equipped with an integrated membrane for isolation, concentration and purification of nucleic acids. In this system, the amplification process is carried out using a LAMP procedure coupled to real time detection of amplicons using a fluorescence reader. The authors assessed their system for the detection of human immunodeficiency virus (HIV-1) from oral fluids and demonstrated a detection limit of less than 10 HIV particles (Liu et al., 2011).

A number of LAMP assays have already been reported for a range of metazoan parasites, including taenid cestodes (Nkouawa et al., 2010), Fasciola spp., Opisthorchis viverrini, Paragonimus westermani (see Ai et al., 2010, Arimatsu et al., 2012, Chen et al., 2011a) and Angiostrongylus cantonensis and Wucheria bancrofti (see Chen et al., 2011b, Takagi et al., 2011). There are reports of LAMP assays for, for example, Toxoplasma (see Hu et al., 2012), Plasmodium (see Sirichaisinthop et al., 2011), Theileria annulata (Liu et al., 2012) or Eimeria (Barkway et al., 2011). However, these assays are still laboratory-based and, to date, there is no point-of-care diagnostic system for routine use. Although a portable system for the detection of animal diseases has been developed and was assessed for foot-and-mouth disease (FMD) (Seise et al., 2011), the steps of RNA isolation from the virus responsible for this disease still require laboratory work. These recent achievements indicate some potential for novel diagnostic systems with prospects for rapid, field-based diagnosis.

10. Future diagnostic applications of advanced sequencing technologies

Further advances in molecular diagnostics are expected from the rapidly developing sequencing technologies. In the past, studies that investigated the diversity of microorganisms in a natural sample involved the cloning and subsequent Sanger sequencing of selected genes (commonly 16S rDNA) to produce profiles of diversity in environmental samples (Tyson et al., 2004). These so-called metagenomic studies (i.e., the study of genetic material recovered directly from environmental samples) showed that cultivation-based methods are unable to detect the vast majority of microorganisms and have allowed novel insights into a previously hidden diversity of microbial life (Hugenholtz et al., 1998). Following the rapid reduction in cost of DNA sequencing (Pushkarev et al., 2009) and the development of high-throughput sequencing technologies (e.g., 454 pyrosequencing, Illumina- and SOLID sequencing platforms) (Bentley et al., 2008, Harris et al., 2008, Mardis, 2008), these technologies have become accessible to many research groups and have enabled the direct sequencing of microbe and parasite genomes from environmental samples. However, these novel approaches of DNA sequencing create read lengths which are significantly shorter than those produced by Sanger sequencing and a much larger number of sequence reads,
ultimately leading to the generation of large amounts of data for analyses (Desai et al., 2012). Analyses of such data sets to obtain biologically meaningful information increased complexity and the need for enhanced computing power and presently represents a bottleneck to this approach (Desai et al., 2012). Nonetheless, a number of studies show the value of metagenomic approaches for assessing the diversity of microbial communities in the gut of humans and other mammals (e.g., Hess et al., 2011, Lamendella et al., 2011, Lepage et al., 2012, Qin et al., 2010) or in the marine environment (e.g., Breitbart et al., 2002, Venter et al., 2004, Yooseph et al., 2010) and have led to the discovery of large numbers of previously unknown microorganisms.

Another technological advance is the DNA sequencing using nano-pores. This type of analysis is emerging and involves the use of a voltage to drive molecules through a nano-scale pore in a membrane between two electrolytes. This allows the analysis of charged polymers (single-stranded DNA, double-stranded DNA or RNA) by monitoring the change of the ionic current as single molecules pass through it (Schneider and Dekker, 2012). Nano-pore sequencing has the advantages that it does not require the labeling of nucleotides, amplification prior to sequencing, can be applied to single molecules and is capable for high throughput DNA analysis (Venkatesan and Bashir, 2011). Further benefits are that it is low cost, requires low reagent volumes and generates long reads, which appears to make it suitable for de novo sequencing (Venkatesan and Bashir, 2011). It has been proposed that nano-pore-based diagnostic tools could detect target molecules at extremely low concentrations and from minute sample volumes, detect simultaneously multiple biomarkers or genes, eliminate the need for time-consuming amplification and conversion steps, thus providing a rapid analysis at a low cost (Venkatesan and Bashir, 2011). In addition to the sequencing through nano-pores, with the aim of rapid and affordable DNA sequencing, a number of other approaches have been developed, including the single-molecule evanescent field detection of sequencing-by-synthesis in arrays of nano-chambers (Pacific Biosciences) (Eid et al., 2009), sequencing by ligation on self-assembled DNA nano-arrays (Complete Genomics) (Drmanac et al., 2010), and the detection of H+ ions released during sequencing-by-synthesis on silicon field-effect transistors from multiple polymerase template reactions (Ion Torrent) (Rothberg et al., 2011). Clearly, the rapid emergence of a range of exciting sequencing technologies provides new prospects for diagnostic applications. Current evidence suggests that some of these advanced technologies will change the face of molecular diagnostics in the near future.

11. Conclusions

The accurate diagnosis of gastrointestinal nematode infections of livestock underpins investigations of the biology, ecology and epidemiology of parasites and supports the monitoring of emerging problems with anthelmintic resistance (AR). Current, routinely used methods of diagnosis rely on the detection or morphological identification of the infective stages (eggs and/or larvae) of these nematodes in host faeces. Until recently, these traditional techniques, which can be time-consuming and laborious, have not undergone any substantial technological advancement. As eggs and larvae of numerous genera and species of nematodes infecting livestock lack distinctive morphological features, traditional approaches are not able to achieve a species- or even genus-specific diagnosis in the live animal, making it challenging to conduct reliable studies of the biology, epidemiology and ecology of parasites, unless expensive and laborious post mortem investigations are carried out. This situation has also hampered investigations of the occurrence and distribution of AR in strongylid nematodes of livestock, which represents a global problem.

Advances in PCR-based methods and the availability of specific genetic markers in the internal transcribed spacers of nuclear rDNA have provided the opportunity of developing enhanced PCR-based tools for diagnosis (reviewed by Gasser, 2006; Gasser et al., 2008). Recent studies (Roeber et al., 2011, 2012a,b) showed that RT-PCR and MT-PCR assays can replace the
inaccurate and time-consuming method of LC. This high throughput MT-PCR (Roeber et al., 2012b) takes < 1 day to perform, compared with at least a week for LC, thus reducing the time that the farmer has to wait for a diagnosis. From a service provision perspective, this platform does not require detailed technical expertise of the operator, has high sensitivity and specificity, and has broad applicability, in that it can be used to carry out large-scale epidemiological studies, to support the diagnosis of drug resistance, and will be applicable or adaptable to other parasites and/or hosts. In addition, the MT-PCR platform delivers, rapidly, objective and detailed results to a genus or species level, which is of major value for enhanced control. Overall, the MT-PCR established (Roeber et al., 2012b) essentially meets the international standards (OIE, 2004; Conraths and Schares, 2006) required for use in a laboratory setting for research or routine diagnostic purposes and has significant advantages over classical methods, particularly in relation to the interpretation of FEC results and recommendations about anthelmintic treatment. This test improves the diagnosis of infections with nematode species, which are problematic to detect or identify by traditional coprological techniques, either because of their morphological/morphometric similarity with other species/genera (i.e., Teladorsagia and Trichostrongylus, Chabertia ovina and Oesophagostomum venulosum) or their unfavourable development under standard culture conditions. Current evidence indicates that this MT-PCR assay is highly adaptable, allowing the development of a wide range of next-generation diagnostic tools to underpin the control of socioeconomically important infectious diseases of animals and the detection and monitoring of drug resistance.

Clearly, a number of nucleic acid-based methods of diagnosis have significant advantages, particularly in terms of sensitivity and specificity, reproducibility and repeatability. These features make them suitable to be incorporated into surveillance systems, which are based on a “from pasture/stable-to-table” approach, such as the Hazard Analysis Critical Control Point (HACCP). The routine application of advanced diagnostic platforms, including practical and high performance field-based assays and, particularly, high throughput sequencing technologies as well as, now provide major scope for better disease surveillance and for detailed epidemiological investigations, shorter response times to tackle and control disease outbreaks (as the diagnosis using molecular tools usually takes less time compared with culture- or microscopy-based approaches) and, ultimately, to provide greater protection for the consumer of animal products. Epidemiological data obtained using such methods would help government authorities, such as the OIE, Food and Agricultural Organization (FAO) and/or the World Health Organization (WHO) in the detailed tracking and mapping of disease outbreaks or spreads, to make forecasts about their occurrence and to implement appropriate contingency plans and guidelines for the effective and sustainable control of parasitic and other infectious diseases. This focus will be accompanied by an expansion in revolutionary, new diagnostic technologies and important biotechnological outcomes.

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References


Barkway C, Pocock R, Vrba V, Blake D. Loop-mediated isothermal amplification (LAMP) assays for the species-specific detection of Eimeria that infect chickens. BMC Vet Res 2011;7:67-


Hoorfar J, Wolffs P, Radström P. Diagnostic PCR: validation and sample preparation are two sides of the same coin. APMIS 2004;112:808-14.


Hung GC, Gasser RB, Beveridge I, Chilton NB. Species-specific amplification by PCR of ribosomal DNA from some equine strongyles. Parasitology 1999;119:69-80.


Lamendella R, Domingo JWS, Ghosh S, Martinson J, Oerther D. Comparative fecal metagenomics unveils unique functional capacity of the swine gut. BMC Microbiol 2011;11:103-.

Lancaster MB, Hong C. Differentiation of third stage larvae of 'ovine Ostertagia' type and Trichostrongylus species. Vet Rec 1987;120:503.


McKenna PB. Further potential limitations of the undifferentiated faecal egg count reduction test for the detection of anthelmintic resistance in sheep. NZ Vet J 1997;45:244-6.
Monis PT, Giglio S, Saint CP. Comparison of SYTO9 and SYBR Green I for real-time polymerase chain reaction and investigation of the effect of dye concentration on amplification and DNA melting curve analysis. Anal Biochem 2005a;340:24-34.
Obeid P, Christopoulos T, Crabtree HJ, Backhouse C. Microfabricated device for DNA and RNA amplification by continuous-flow polymerase chain reaction and reverse


Stevenson LA, Gasser RB, Chilton NB. The ITS-2 rDNA of *Teladorsagia circumcincta*, *T. trifurcata* and *T. davitiani* (Nematoda: Trichostrongylidae) indicates that these taxa are one species. Int J Parasitol 1996;26:1123-6.


Fig. 1. Generalized life cycle representing key gastrointestinal strongylid nematodes of small ruminants (adapted from Demeler, 2005). First-, second- and third-stage larvae (L1, L2 and L3, respectively) are ‘free-living’ in the environment. The fourth larval (L4) and adult stages (dioecious) are ‘parasitic’ in the gastrointestinal tract of the host. Disease in the host animal is caused by the adult and/or L4 stages, and depends on the species of nematode; intensity of infection; species, age and immunological/health status of the host; host response against the parasite; stress and other environmental and management factors (Kassai, 1999; Taylor et al., 2007).
Table 1

<table>
<thead>
<tr>
<th>Family</th>
<th>Species</th>
<th>Morphometrics/morphology</th>
<th>Pre-patent period (days)</th>
<th>Location in the host</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trichostrongyldae</td>
<td>Haemonchus contortus</td>
<td>♂ 10-20 White ovaries coiled around red intestines. Presence of vulvar flap depends on strain.</td>
<td>18-21</td>
<td>Abomasum</td>
</tr>
<tr>
<td></td>
<td></td>
<td>♀ 18-30</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Teladorsagia circumcincta</td>
<td>♂ 7-8 Small head and buccal cavity. Presence of vulvar flap can be present.</td>
<td>15-21</td>
<td>Abomasum</td>
</tr>
<tr>
<td></td>
<td></td>
<td>♀ 10-12</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Trichostrongylus axei</td>
<td>♂ 2-6 Dissimilar spicules of unequal length.</td>
<td>15-23</td>
<td>Abomasum</td>
</tr>
<tr>
<td></td>
<td></td>
<td>♀ 3-8</td>
<td></td>
<td>or stomach</td>
</tr>
<tr>
<td></td>
<td>T. colubriformis</td>
<td>♂ 4-8 Equal length spicules with triangular tip.</td>
<td>15-23</td>
<td>Anterior small intestine</td>
</tr>
<tr>
<td></td>
<td></td>
<td>♀ 5-9</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>T. vitrinus</td>
<td>♂ 4-7 Thick spicules of equal length.</td>
<td>15-23</td>
<td>Anterior small intestine</td>
</tr>
<tr>
<td></td>
<td></td>
<td>♀ 5-8</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>T. rugatus</td>
<td>♂ 4-7 Dissimilar spicules of unequal length.</td>
<td>15-23</td>
<td>Small intestine</td>
</tr>
<tr>
<td></td>
<td></td>
<td>♀ 6-7</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cooperia curticei</td>
<td>♂ 4-5 Transverse striaion of cuticle, watch-spring-like body posture, and presence of a small cephalic vesicle are characteristic.</td>
<td>14-15</td>
<td>Small intestine</td>
</tr>
<tr>
<td></td>
<td></td>
<td>♀ 5-6</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Nematodirus spathiger</td>
<td>♂ 10-19 Small but distinct cephalic vesicle. Vary long spicules ending in a spoon-shaped terminal piece.</td>
<td>18</td>
<td>Small intestine</td>
</tr>
<tr>
<td></td>
<td></td>
<td>♀ 15-29</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>N. ilicolls</td>
<td>♂ 10-15 Small but distinct cephalic vesicle.</td>
<td>18</td>
<td>Small intestine</td>
</tr>
<tr>
<td></td>
<td></td>
<td>♀ 15-20</td>
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<tr>
<td>Ancylostomatidae</td>
<td>Bunostomum trigonocephalum</td>
<td>♂ 12-17 Anterior end is bend dorsally. Buccal capsule is subglobular without teeth.</td>
<td>40-70</td>
<td>Small intestine</td>
</tr>
<tr>
<td></td>
<td></td>
<td>♀ 19-28</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chabertidae</td>
<td>Oesophagostomum columbianum</td>
<td>♂ 12-16 Have two leaf crowns and a shallow buccal capsule. Cervical papillae are well developed.</td>
<td>40-45</td>
<td>Large intestine</td>
</tr>
<tr>
<td></td>
<td></td>
<td>♀ 14-18</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>O. venulosum</td>
<td>♂ 11-16 Cervical papillae are situated posterior to the oesophagus.</td>
<td>40-45</td>
<td>Large intestine</td>
</tr>
<tr>
<td></td>
<td></td>
<td>♀ 13-24</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Chabertia ovina</td>
<td>♂ 13-14 Mouth is directed antero-ventrally. Buccal capsule is subglobular without teeth.</td>
<td>42-50</td>
<td>Large intestine</td>
</tr>
<tr>
<td></td>
<td></td>
<td>♀ 17-20</td>
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</tr>
</tbody>
</table>
Table 2
Key validation parameters employed for the assessment of a diagnostic test (based on Conraths and Schares, 2006, Pfeiffer, 2010, Thrusfield, 2005).

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
<th>Method of assessment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensitivity</td>
<td>The proportion of animals with the disease and which test positive.</td>
<td>Assessment of these two parameters requires an independent, valid criterion termed a</td>
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<td></td>
<td></td>
<td>&quot;gold standard&quot; used to define the true disease status of an animal.</td>
</tr>
<tr>
<td>Specificity</td>
<td>The proportion of animals without the disease and which test negative.</td>
<td></td>
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<tr>
<td>Agreement</td>
<td>The agreement in results between two diagnostic test, with one of the tests</td>
<td>Frequently assessed by Kappa test, which measures the proportion of agreement beyond</td>
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<tr>
<td></td>
<td>being a generally accepted diagnostic method.</td>
<td>that to be expected by chance.</td>
</tr>
<tr>
<td>Accuracy</td>
<td>Refers to the concordance between test results and the 'true' clinical state.</td>
<td>Depends on the number of 'false positives' and 'false negatives', in comparison with</td>
</tr>
<tr>
<td></td>
<td></td>
<td>the true infection state as determined by the &quot;gold standard&quot;.</td>
</tr>
<tr>
<td>Reliability</td>
<td>The extent to which test results are consistent in repeat experiments.</td>
<td>This includes the assessment of repeatability, reproducibility, inter- and intra-assay</td>
</tr>
<tr>
<td></td>
<td></td>
<td>variability.</td>
</tr>
</tbody>
</table>

Table 3
Stages of validation of a diagnostic test (adapted from Conraths and Schares, 2006).

Stages of test validation

1. Feasibility studies
2. Assay development and standardisation
   - optimisation of reagents, protocols and equipment
   - preliminary estimate of repeatability
   - determination of analytical sensitivity and specificity
3. Determination of assay performance characteristics
   - diagnostic sensitivity and specificity
   - repeatability and reproducability
4. Monitoring the validity of assay performance
5. Maintenance and enhancement of validation criteria
Advances in the diagnosis of key gastrointestinal nematode infections of livestock, with an emphasis on small ruminants


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