Epidemiology, Diagnosis and Public Health Importance of Trichinellosis

Muluken Yayeh, Gemechis Yadesa, Mareyes Erara, Shumye Fantahun, Alemseged Gebru and Mastewal Birhan

University of Gondar, College of Veterinary Medicine and Animal Sciences, Ethiopia

Email: yayehmulu@gmail.com

Supporting Information

ABSTRACT: Trichinellosis is a parasitic zoonosis caused by Trichinella following ingestion of raw or under cooked meat containing Trichinella larvae. Nematode worms of the genus Trichinella are one of the most prevalent zoonotic pathogens in the world. The parasite infects domestic and wild animals and has a worldwide distribution. The life cycle of the parasite consists of a domestic cycle in mainly pigs and a sylvatic cycle in a wider range of animals such as bears and wild boar. Humans become infected after eating raw or undercooked meat from domestic pigs, horses or game containing Trichinella larvae. There are twelve genotypes within the genus Trichinella, eight of which have been designated as species from which T. spiralis is the most pathogenic one. Host animals ingesting even high numbers of Trichinella larvae from infectious meat will not develop clinical symptoms. In humans, the clinical picture is usually illustrated by an intestinal stage within the first or second week after infection and later muscular stage with periorbital oedema, myalgia or muscle weakness as the major symptoms. The severity of the clinical course depends firstly on parasitic factors, such as the species implicated and the number of infective larvae ingested, and secondly on host factors, such as sex, age, and immune status. In practice, treatment with anthelmintics and immunosuppressive drugs is used only with human patients, not with animals. Trichinella infection in humans is strongly associated with the consumption of raw or undercooked meat; thus, cultural factors such as traditional dishes based on raw or undercooked meat or meat-derived products.

Keyword: Human, Parasite, Pig, raw meat, Trichinellosis; Zoonosis

INTRODUCTION

Parasitic zoonosis includes both helminthic and protozoan infections. Amongst one thousand five hundred known infectious agents for human being, 287 are helminths (Chomel et al., 2008). Helminths are complex eukaryotic organisms with large genomes and complex multistage life cycles that involve numerous hosts (Hewitson et al., 2009). Nematode worms of the genus Trichinella are one of the most widespread zoonotic pathogens in the world. Infection by Trichinella species has been identified in domestic and/or wild animals of all continents, with the exception of Antarctica, where there is no record of the parasite (Murrell, 2006).

Clinical signs of trichinellosis are not generally recognised in animals, and its main importance is as a zoonosis. Trichinosis is a food-borne zoonotic disease caused by Trichinella species. Trichinosis in humans is caused by eating raw or undercooked meat from Trichinella-infected food animals or game (Gajadhar et al., 2006). Until recently, all Trichinella infections occurring in animals and humans were attributed to Trichinella spiralis. Today, eight species (T. spiralis, T. nativa, T. britovi, T. pseudospiralis, T. murrelli, T. nelsoni, T. papuae, and T. zimbabwensis and three genotypes (Trichinella T6, T8, T9) within two classes (encapsulated and non-encapsulated) are documented in this genus (Zarlenga et al., 2006).

It is a tissue-dwelling nematode acquired by the ingestion of raw or insufficiently cooked meat-products containing encapsulated larvae (La Rosa et al., 2000). The most important source of human infection worldwide is the domestic pig. In Europe, meats of horses and wild boars have played a significant role during outbreaks within the past three decades. Infection of humans occurs with the ingestion of Trichinella larvae that are encysted in muscle tissue of meat from domestic or wild animals (Bruschi et al., 2007).

Trichinella infection in the human host can be divided into two stages: an intestinal (or enteral) phase and a muscular (or parenteral or systemic) phase. Low-intensity infection can remain asymptomatic, but parasite burdens greater than a few hundred larvae can initially cause gastroenteritis associated with diarrhoea and abdominal pain approximately 2 days post infection (intestinal acute phase of disease). Subsequently, migrating larvae and their metabolites provoke an immediate reaction, with an inflammatory and allergic response, pyrexia, eyelid or facial oedema, and eosinophilia are the most prominent manifestations, occasionally complicated by myocarditis, thromboembolic disease, and encephalitis. Months or even years at the acute stage, chronic trichinellosis may yield persistent formation,
numbness, and excessive sweating as well as impaired muscle strength and conjunctivitis, which may continue up to 10 years post infection (Zarlenga et al., 2013).

The diagnosis of trichinellosis is based on history of consumption of potentially contaminated meat, the presence of compatible signs and symptoms, and identification of *Trichinella* larvae in biopsy muscle tissue or specific antibody in serum. These diagnostic methods in human host can be categorized two as direct and indirect. Under direct there are direct muscle biopsy while under indirect such as serology and molecular technique (Oivanen, 2005). Muscle biopsies are rarely performed, but they allow for the molecular identification of the *Trichinella* species or genotype, which is not possible with antibody testing (Oivanen, 2005).

Prompt treatment with anti-parasitic drugs can help prevent the development of trichinellosis by killing the adult worms and so preventing further release of larvae. Once the larvae have become established in skeletal muscle cells, treatment may not completely eliminate the infection and associated symptoms (Sun, 2015). Therefore, the administration of effective anthelmintic drugs at the stage of intestinal invasion or in the acute phase is critical for successful therapy. In addition, because of the predominantly zoonotic importance of infection, the main efforts in many countries have focused on the control or elimination of Trichinella from the food chain (Gottstein, 2009).

The increase in the report of Trichinellosis has been observed many eastern European countries, in Africa and Asia (Blaga et al., 2007; Azim et al., 2008). Human population growth and socioeconomic changes might have played a fundamental role in the disease emergence and spread in recent years (Macpherson, 2005). The increase in human population density, ecological change, and subsequent increased contact between humans and wild animals necessitates the importance having an update on potentially emerging diseases like trichinellosis. Therefore, the objective of this paper is to review the epidemiology and public health importance of trichinellosis.

**Taxonomy and Morphology of the Parasites**

**Taxonomy and Aetiology**

The taxonomy of the genus *Trichinella* has been presented with slightly varying details According to the traditional classification, the genus belongs to the phylum Nematode, roundworms, class Adenophorea, order Trichinellicida, and superfamily Trichinelloidea (Oivanen, 2005). The taxonomy has recently been challenged. On the basis of results from ribosomal deoxyribonucleic acid (DNA) sequences, the present higher-level classification of Nematode will need change in to two classes, Secernentea and Adenophorea (Oivanen, 2005). Within the genus *Trichinella* there are twelve genotypes have been identified, eight of which have been designated as species (Gajadhar et al., 2006; Murrell et al., 2000; Pozio and Zarlenga, 2005). *Trichinella spiralis* was recognized in London in 1835s. The parasite being detected in an autopsy of an Italian male corpse (Oivanen, 2005).

*Trichinella spiralis* (T1) is distributed in temperate regions world-wide and is commonly associated with domestic pigs. It is highly infective for domestic and sylvatic swine, mice and rats, but it can also be detected in other mammalian, carnivores and horses (Pozio and Zarlenga, 2005). This species is also the most important etiological agent to cause disease in humans (Pozio, 2006). Trichinella native is the species that are very widespread in arctic and subarctic areas of the northern hemisphere (Pozio, 2000). *Trichinella britovi* species differs from *T. spiralis* with weak infectivity for rats, moderate resistance to freezing, moderate infectivity for swine, slow nurse cell development and low in vitro production of NBL (Malakauskas and Kapel, 2003). *Trichinella nelsoni* has occasionally been detected in pigs (Suidae) and humans, although it has very low infectivity for pigs and rats. The infectivity for humans has not been long-established (Pozio, 2001).

*Trichinella murrelli* this species has very low reproductive capacity in pigs and rats, low NBL production in vitro, slow nurse cell development, and low resistance to freezing (Malakauskas and Kapel, 2003). *Trichinella pseudospiralis* strains three genotypic isolates were identified by multiplex polymerase chain reaction from different parts of the world (PCR) test (Zarlenga et al., 1999: La Rosa et al., 2001). *Trichinella papuae are where Muscle* larvae are non-encapsulated and lack freezing tolerance but can survive in +5°C storage for four weeks (Webster et al., 2002). *Trichinella zimbabwensis* is the first *Trichinella* strain isolated in reptiles in nature. In the laboratory, it can also infect rats, mice, pigs, baboons (Papio sp.), turtles, pythons, varans, and caimans. Its muscle larvae are non-encapsulated. It is not infective for birds, nor can it resist freezing (Pozio et al., 2004).

**Morphology of the Parasite**

*Trichinella* worms are the smallest nematode parasite of humans, they are the largest intracellular parasite and have been described as “the worm that would be virus” (Foreyt, 2013). The morphology of the parasite’s oesophagus is characteristic of the *Trichinellidae* family, and it occupies approximately one-third of the body length and is surrounded by large cells. Adult males are 1.4 to 1.6 mm in length and do not have spicules, but a pair of lateral flaps is found on each side of the cloacal opening and two pairs of papillae are between them. Females’ are 3 to 4 mm in length, and the vulva opens in the middle of the oesophageal region (Foreyt, 2013). Adult the length of *T. spiralis* NBL is 80-120 μm and the diameter 5-6 μm. The larvae do not increase in size until they enter the muscle cells. The larvae begin to grow in their nurse cells, reaching a length of 900-1280 μm and a diameter of 35-40 μm by 30days p.i. *Trichinella* adult females are a little longer and thicker than the males. Their length and diameter are 2460-3390 μm and 35-70 μm, respectively, while the resulting figures for males are 1040-1300 μm and 29-32 μm (Oivanen, 2005).
The life cycle of the parasites

The basic life cycle of *Trichinella* has been recognized since the middle of the 19th century. This genus is unique among parasitic nematodes in that all stages of the life cycle occur within a single host. In nature, the cycle is repeated when another host animal ingests the flesh of another host containing viable muscle-stage larvae (Oivanen, 2005). The generalized life cycle of *Trichinella* is described in figure 2. Enteral phase; 1: muscle tissues are digested in the stomach and larvae are released; 2: larvae penetrate the intestinal mucosa of the small intestine, reach the adult stage within 48 h post infection, male and female mate; 3: female worm releases new born larvae in the lymphatic vessels (from the fifth day post infection onwards; the length of New born production, from one to several weeks, is under the influence of the host immunity). Parenteral phase; 4: the new born larva reach the striated muscle and actively penetrate in the muscle cell; 5: the larva grows to the infective stage in the nurse cell (the former muscle cell); 6: after a period of time (weeks, months or years) a calcification process occurs (Pozio and Murrell, 2006).

Figure 1: Morphology of parasites (Foreyt, 2013)

![Morphology of adult male and female Trichinella worms. (Modified from Villala, 1970)](image)

Figure 2: *Trichinella* sp. life cycle (Pozio and Murrell, 2006). A: main sources of *Trichinella* spp. infections for humans; B: *Trichinella* spp. cycle in the host body.
Epidemiology

Geographic distribution

*Trichinella* species are present throughout most of the world in over 150 different hosts (Dick et al., 2001). In addition to *T. spiralis*, 7 other species in 4 genotypes, all of which are more commonly found in wild animals than in domestic pigs. *T. spiralis* is cosmopolitan, this species is also the most important etiological agent to cause disease in humans (Pozio et al., 2006). In the domestic cycle, pork scraps from *T. spiralis*-infected pigs are the main source of infection for synanthropic animals (e.g., rats, horses, stray cats, and dogs). Conversely to the domestic cycle, the sylvatic cycle of *T. spiralis* includes a broad range of wild carnivores, which may, however, become the origin of a life cycle beginning into a domestic host population (Dick, 2001). *Trichinella nativa* is found in Arctic and subarctic areas of America, Asia, Europe. *Trichinella* genotype T6 is also found in Canada, Alaska, Rocky Mountains, and Appalachian Mountains in the United States (Pozio, 2001). *Trichinella britovi* is found in the temperate areas of Europe and Asia, Northern and Western Africa, *Trichinella* T8 is found in South Africa and Namibia. *Trichinella murrelli* is found in the United States and Southern Canada, *Trichinella* genotype T9 Japan, *T. nelsoni* Eastern-Southern Africa, and *Trichinella* genotype T12 is found in Argentina, and all the above are encapsulated while the below are non-capsulated. Those are *T. pseudospiralis* which is cosmopolitan in Palearctic, Nearctic, Ocean land (Pozio, 2001) Cosmopolitan, *T. papuae* is found in the Papua New Guinea, Thailand and *T. zimbabwensis* which is found in Zimbabwe, Mozambique, Ethiopia, South Africa (Gottstein, 2009).

Host range

The epidemiology and systematics (i.e., the study of the diversification) of this zoonosis are now recognized to involve in addition to *T. spiralis*, 7 other species in 4 genotypes, all of which are more commonly found in wild animals than in domestic pigs (Pozio, 2009). *T. spiralis* is found in the Domestic and sylvatic mammal, while *Trichinella* T8, *T. murrelli*, *Trichinella* genotype T9. *Trichinella nativa and Trichinella* genotype T6 is found in the sylvatic carnivores. Another species, *T. britovi* is found in the Sylactic mammals and seldom domestic pigs, while *T. nelson* in the Sylactic mammals. *Trichinella* genotype T12, *T. Pseudo spiralis* Sylactic mammals and birds, domestic pigs, *T. papuae* Wild pigs, salt water crocodiles and *T. zimbabwensis* which is found in the Nile crocodiles, monitor lizards (Gottstein, 2009).

Main source of Infection in human

Domestic pigs and wild boars were the major sources of *Trichinella* spp. Infection for humans, but in recent years new infection sources, particularly from exotic hosts, have emerged (Boireau, 2000). The main source of infection in human, *T. spiralis* is found in the Domestic and sylvatic swine horses, while *T. nativa* is found in Bears, walruses. Others like *Trichinella* genotype T6 is found in Carnivores *T. britovi* is found in the Wild boars, domestic pigs horses, foxes, jackals *T. zimbabwensis*, *Trichinella* genotype T12, *Trichinella* genotyp T8 and *Trichinella* genotype T9. *Trichinella murrelli in Bears*, especially horses while *T. nelson* is found in the Warthogs, bush pigs Warthogs, bush pigs, *T. papuae* is found in the Wild pigs and *T. pseudospiralis* wild and domestic pigs (Gottstein, 2009).

Resistance of larvae in frozen muscle

Most of them are not resistance to the frozen muscle. *Trichinella* T8, *T. murrelli*, *T.pseud ospiralis* *T. papuae and T.zimbabwensis*, while others are *T. spiralis* resistance in horse muscles. *T. nativa* are resistance in carnivore muscles. *Trichinella* genotype T6 is resistance in carnivore muscles, *T. britovi* are resistance in carnivore and horse muscles and *Trichinella* genotype T12 Unknown (Gottstein, 2009). The epidemiology of trichinellosis is summarized as below in table 1.

Disease ecology

The usual source of trichinosis in humans is from eating pork products or meat from horses, dogs, or a variety of wildlife species, including wild pig, bear, walrus, and seal. *Trichinella* spp. is transmitted by two specific cycles, the domestic cycle and the sylvatic cycle (Dick and Pozio, 2001).

Domestic cycle

The domestic cycle is prevalent on small farms where disease control is not a primary objective in food production. Areas where infection is endemic are found throughout the world (Dupouy-Camet, 2000). The domestic cycle of transmission is primarily involves *T. spiralis* in a cycle of pig-to-pig transmission, and humans enter the cycle through eating pork. The infection can be highly pathogenic in humans. Synanthropic rats, mice, cats, dogs, and horses, as well as many wildlife species, also contribute to the cycle in many areas. Pigs maintain the cycle by eating pieces of infected meat scraps, eating infected rats or mice, biting the tails of infected pigs, cannibalizing dead pigs, ingesting feces from pigs that have recently eaten infected meat, or eating other species of infected mammals (Ortega-Pierres and others et al., 2000).

Sylvatic cycle

The sylvatic cycle of transmission predominantly involves predation, cannibalism, or scavenging behaviours of species of carnivorous wildlife. *Trichinella* spp. are transmitted when fresh, frozen, or decomposing carcasses or meat scraps are eaten (Dupouy-Camet, 2000). The species of *Trichinella* associated with the sylvatic cycle are *T. nativa*, *T. britovi*, *T. murrelli*, *T. nelsoni*, *T. pseudospiralis*, *T. papuae* and *T. zimbabwensis*. *T. spiralis* can also affect wildlife in temperate and tropical regions, but it does not survive in arctic and subarctic regions because larvae do not survive in a frozen carcass (Ortega-Pierres et al., 2000).

Clinical signs

The severity of the clinical course depends firstly on parasitic factors, such as the species involved and the number of infective larvae ingested, and secondly on host factors, such as sex, age, and immune status (Bruschi and Murrell,
2002). The chief clinical of trichinellosis were compatible in type and frequency with the classical trichinellosis syndrome, i.e., myalgia, diarrhoea, fever, facial oedema and headaches that, after treatment, disappeared within 2–8 weeks (Dupouy-Camet and Bruschi, 2007). The clinical signs of acute trichinellosis are characterized by two phases: an enteral and a parenteral phase, corresponding to the presence of parasites in the intestine and in the circulation and/or musculature, respectively (Oivanen, 2005). The most common signs during the enteral phase of a mild infection are transient diarrhoea and nausea. However, in moderate to severe infections, the first signs are upper abdominal pain, diarrhoea or constipation, vomiting, malaise, and mild fever. The enteral phase lasts for six weeks (Kocięcka, 2000). From the second to the sixth week post infection, the enteral phase is still present, but the dominating signs arise from the parenteral phase due to the migrating larvae and their indiscriminate penetration of different tissues. During the third week post infection the symptoms intensify due to invasion of muscle cells. Characteristic signs include weakness, pain, paralysis, and photo phobia. Edema is prominent and patients may have shortness of breath. Endocarditis, myocarditis, and cardiac failure have been reported. The signs of acute illness usually diminish from the fifth or sixth week post infection onwards (Kocięcka, 2000; Oivanen, 2005).

### Table 1 - Epidemiology of Trichinellosis

<table>
<thead>
<tr>
<th>Species or genotype</th>
<th>Geographical distribution</th>
<th>Main source of infection in human</th>
<th>Resistance of larvae in frozen muscle</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Encapsulated</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>T. spiralis</em></td>
<td>Cosmopolitan</td>
<td>Domestic and sylvatic mammals</td>
<td>Domestic and sylvatic swine horse</td>
</tr>
<tr>
<td><em>T. nativa</em></td>
<td>Arctic and subarctic areas of America, Asia, Europe</td>
<td>Sylvic carnivores</td>
<td>Bears, walruses</td>
</tr>
<tr>
<td>Trichinella genotype T6</td>
<td>Canada, Alaska, rocky mountains, and Appalachian Mountains in the united states</td>
<td>Sylvic carnivores</td>
<td>Carnivores</td>
</tr>
<tr>
<td><em>T. britovi</em></td>
<td>Temperate areas of Europe and Asia, northern and western Africa</td>
<td>Sylvic mammals and rarely domestic pigs</td>
<td>Wild boar, domestic pig , horse, foxes, jackal</td>
</tr>
<tr>
<td>Trichinella T8</td>
<td>South Africa and Namibia</td>
<td>Sylvic carnivores</td>
<td>None documented</td>
</tr>
<tr>
<td><em>T. murrelli</em></td>
<td>United states and southern Canada</td>
<td>Sylvic carnivores</td>
<td>Bears, horses</td>
</tr>
<tr>
<td>Trichinella genotype T9</td>
<td>Japan</td>
<td>Sylvic carnivores</td>
<td>None documented</td>
</tr>
<tr>
<td><em>T. nelson</em></td>
<td>Eastern-southern Africa</td>
<td>Sylvic mammals</td>
<td>Warthogs, bush pigs</td>
</tr>
<tr>
<td>Trichinella genotype T12</td>
<td>Argentina</td>
<td>Cougars</td>
<td>None documented</td>
</tr>
<tr>
<td><strong>Non encapsulated</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>T. pseudospiralis</em></td>
<td>Cosmopolitan</td>
<td>Sylvic mammals and birds, domestic pig</td>
<td>Domestic and wild pigs</td>
</tr>
<tr>
<td><em>T. papuae</em></td>
<td>Papua new guinea ,Thailand</td>
<td>Wild pig, salt water crocodile</td>
<td>Wild pig</td>
</tr>
<tr>
<td><em>T. zimbabwensis</em></td>
<td>Zimbabwe, Mozambique, south Africa, Ethiopia</td>
<td>Nile crocodiles, monitor lizards</td>
<td>None documented</td>
</tr>
</tbody>
</table>

**Source:** Gottstein (2009).

### Diagnosis

**Direct method**

Meat inspection for the detection of *Trichinella* larvae is designed to prevent clinical trichinellosis in humans but not to prevent infection. The identification of *Trichinella* larvae in muscle samples from pigs and other animal species intended for human consumption (e.g., horses, wild boars, and bears) is limited to post-mortem inspection of carcasses. Muscle biopsy is a traditional method applied to diagnose trichinellosis. Samples are usually taken from the *M. deltoideus*. Other possible sites are the *Musculus biceps brachii*, *Musculus gastrocnemius*, *M. pectoralis*, *M. glutus maximus*, and *Musculi intercostali* (Gamble, 2000). Muscle biopsy is recommended only in cases where serological results are unclear. In autopsy, the sampling site is the diaphragm (Bruschi and Murrell, 2002). Direct detection is also applied in wildlife monitoring, where indicator animals (e.g., foxes or raccoon dogs) are examined to assess the prevalence of *Trichinella* infection among the wildlife reservoir and the risk of introduction into domestic animals. Methods to detect *Trichinella* larvae in muscle samples need to be highly sensitive, and performance is greatly influenced by the sample size, the muscle type selected for sampling, and the specific method used (Nockler, 2000). In order to identify predilection sites, in particular, animal species that optimal for diagnostic investigations, several experimental studies using doses that mimic natural infections have been performed. Thus, in domestic swine, the three main predilection sites for *T. spiralis* are the diaphragm crus, the tongue, and the masseter (Gamble, 2000), and analogous results were observed in experimental *T. britovi* and *T. pseudospiralis* infection in this host species (Nockler, 2005). Some of the sampling sites recommended by the International Commission on Trichinellosis for different domestic and wild animals subjected to meat examination or epidemiological studies are summarized in **Table 2**.
Serology

Serology is considered to be appropriate for the surveillance and epidemiological investigations of *Trichinella* in domestic animals and wildlife (Dworkin, 1996). The indirect serological diagnostic methods can be used at both anti-mortem and post-mortem examination for *Trichinella*-specific antibodies. Several conventional sero diagnostic methods have been practiced in detecting *Trichinella* larvae. These include ELISA, immunofluorescence antibody test (IFAT), complement fixation test, and hemagglutination test and molecular technique (Oivanen, 2005).

**Enzyme Linked Immunosorbant Assay (ELISA)**

The ELISA method is relatively simple to apply, and it can be automated in *Trichinella* diagnostics. It is sufficiently sensitive to detect low-level infect ion (Nöckler et al., 2000). Traditionally ELISA has been applied to analyse antibodies in serum samples. According to some reports, samples of muscle juice can substitute for serum samples. This may be a practical solution if serum is unavailable. Results with muscle juice were hopeful in pigs but inconsistent in wild red foxes (Vercammen et al., 2002). However, this method cannot replace the direct methods at meat inspection because it can fail to detect early or very late stages of infections (Gamble et al., 2004). Infection levels as low as one larva/100 g of tissue is detectable by ELISA in pigs (Gamble et al., 2004). This high level of sensitivity makes serological testing by ELISA a useful method for detecting ongoing transmission of *Trichinella* infection at the farm or for more broadly based surveillance programmes. A disadvantage of serology for the detection of *trichinellosis* is the low rate of false-negative results observed in infected animals (OIE, 2012). For this reason, serological methods are not recommended for individual carcass testing. Serological responses in pigs persist for a long time after infection with no decline in titre; however, antibody has been reported to reject in horses within a few months following infection. The use of ELISA to detect the presence of parasite-specific antibodies provides a quick method that can be performed on serum, blood or tissue fluid collected before or after slaughter. The dilution used is different for serum than for tissue fluid (Nöckler et al., 2000). Antigens that are specifically secreted from the stichocyte cells of living L1 larvae and bear the TSL-1 carbohydrate epitope are recognised by *Trichinella*-infected animals. The specificity and sensitivity of ELISA is largely dependent on the quality of the antigen used in the test (Forbes et al., 2004; OIE, 2012).

**Molecular technique.** Since there are no morphological features to specify larvae, molecular diagnosis is used to yield the species or genotype diagnostically recovered. For this purpose, a multiplex PCR has been developed for the simple and unequivocal differentiation of *Trichinella* species and genotypes (Zarlenga, 1999). Polymerase chain reaction limited studies have shown that PCR can be used to detect the nucleic acid of larvae in the musculature of infected animals (Zarlenga et al., 2003). However, this method lacks sensitivity and is not practical for routine testing of food animals. Identification of the species or genotype of *Trichinella* recovered from muscle tissue is useful in understanding the epidemiology of the parasite in animals, in assessing the relative risk of human exposure and to trace back the infection to the farm of origin (OIE Terrestrial Manual, 2012). Specific primers have been developed that allow the identification of single larva collected from muscle tissues at the species and genotype level by PCR. This multiplex PCR is a sensitive, inexpensive, and rapid molecular approach that can unequivocally identify a single larva at the species and genotype levels (Pozio et al., 2003).

**Status of Trichinellosis in Ethiopia**

At least two confirmed outbreaks of trichinellosis had been reported in Ethiopia. One of the outbreaks was reported in Gojam administrative region. The outbreak was associated with ingestion of meat from a wild boar. In this outbreak, from 30 soldiers, 20 who ate the raw meat became ill and 5 of them were admitted to Hospital with distinctive history and clinical features the disease. The diagnosis was confirmed by deltoid muscle biopsy in all the 5 cases. Similar outbreak had been reported from Central Arsi (Kefenie et al., 1988; Kefenie and Bero, 1992).

**Public health importance of trichinella**

In humans, trichinosis is an important food-borne disease that can cause acute and chronic illness. Humans are only infected with *Trichinella* larvae through the ingestion of meat that has not been appropriately cooked. All species of *Trichinella*, except for the none encapsulated species (*T. pseudospiralis*, *T. papuae*, and *T. zimbabwensis*), can be highly...
pathogenic in humans (Kociecka, 2000). *T. spiralis* is apparently more pathogenic in humans than other species because more larvae are produced by the female worms (Foreyt, 2013). Recently, *T. papuae* has been implicated in outbreaks of human trichinosis (Khumjui et al., 2008). Clinical manifestations are often complex, and they depend on the age of the human host, the state of resistance, and the numbers of larvae ingested. Most clinical symptoms are present between 1 and 6 weeks after infection and the psychological effects of affected humans advance complicate the physical symptoms of the disease. Three stages of disease in humans have been described: the enteral or intestinal phase, the migratory or mucosal invasion phase, and the parenteral or convalescence phase (Foreyt, 2013).

Recently, *T. papuae* has been implicated in outbreaks of human trichinosis. Twenty-eight people in Thailand became sick after eating wild boar and suffered symptoms of trichinosis, and *T. papuae* was identified in a muscle biopsy from one of the patients (Khumjui et al., 2008). *T. papuae* was also suspected as the cause of an outbreak of trichinosis in eight people who had eaten raw soft-shelled turtles in Taiwan (Lo et al., 2009).

CONCLUSION AND RECOMMENDATION

Trichinellosis (trichinosis) is caused by nematodes (roundworms) of the genus *Trichinella*. The disease has a significant public health importance. All mammals are susceptible to infection, but the number of larvae required for infection varies according to the genetic constitute of the parasite and the host species. Trichinellosis is acquired by eating raw or undercooked meat that contains *Trichinella* larvae. Domestic animals can be infected by the consumption of infected raw tissues. *Trichinella* has a direct life cycle, which means it completes all stages of maturity in one host. Transmission from one host to another host can only occur by ingestion of muscle tissue which is infected with the encysted larval stage of the parasite. When ingested, muscle larvae excyst and enter tissues of the small intestine, where they undergo development to the adult stage. Male and female adult parasites mate and produce newborn larvae which leave the intestine and migrate, through the circulatory system, to striated muscle tissue. The severity of human trichinellosis is dependent upon the number of infected larvae ingested, the species of *Trichinella*, and the immune status of the human host. Muscle biopsy, ELISA and PCR method is important tool for diagnosis of infection.

Based on above conclusion the following recommendations are forwarded: 1) Education of the consumer about the risk of consumption of raw or undercooked meat and meat products from both domestic and wild pigs should be emphasized; 2) Strict quarantine should be exercised to control the slaughter and meat distribution of potentially infected animals.

DECLARATIONS

**Corresponding Author**
E-mail: Yayeh.mulu@gmail.com

**Consent to publish**
Not applicable.

**Competing Interests**
The authors declare that they have no competing interests.

**Authors' contributions**
M. Yayeh and M. Birhan conceived the study, coordinated the overall activity, and carried out the statistical analysis, drafted the manuscript. M. Birhan participated in drafting and reviewing the manuscript. M. Yayeh and M. Birhan conceived the study, coordinated the overall activity, and reviewed the manuscript. M. Yayeh and M. Birhan participated in drafting and reviewing the manuscript. All authors read and approved the final manuscript.

**Availability of data and materials**
Data will be made available up on request of the primary author.

**Acknowledgment**
The authors’ heartfelt thanks University of Gondar, Research and Community Service V/ President Office for finance and resource supporting.

**REFERENCES**


