Some helminth species belonging to the order Strongylida (Phylum, Nematoda; Class, Chromadorea) are parasites of the digestive tract of vertebrates, including man and domestic animals [1]. The eggs released in hosts’ feces are morphologically similar, with some differences in size, so it is difficult to identify the genus/species by conventional parasitological examination [2].

Strongylida eggs are thin-shelled and designed to release into the environment; after about 24 hours, Rhabditoid larvae survive and grow by actively feeding on the organic matter in the soil. These first-stage larvae, approximately 250 μm in length, undergo a cuticle molt 48 hours after hatching and grow until reaching near 400 μm, giving rise to the second-stage larvae. These continue to grow until reaching 500–700 μm, develop a new sheath, and molt to the third stage (filariform larvae), when they no longer grow or feed, being ready to penetrate the host, percutaneously or orally [3]. Most parasitic Strongylida species are directly transmitted and filariform larvae in the environment represent the infective stage. Within the vertebrate host, and after a new molt, fourth-stage larvae emerge migrating through different organs; most Strongylida species will
inhabit the digestive tract of the vertebrate host, where the adult worms copulate and females lay eggs which are shed with the host’s feces [3].

The order Strongylida has four families encompassing parasites that infect humans: (i) Ancylostomatidae, which includes hookworms such as Necator americanus and Ancylostoma spp., (ii) Chabertidae, which includes Oesophagostomum spp. and Ternidens deminutus, (iii) Trichostrongyliidae, to which Trichostrongylus spp. belong (and the enzootic species Hystrostrongylus rubidus, the red stomach worm of pigs, Haemonchus contortus, parasite of small ruminants, and the parasites of bovines Cooperia punctata and Ostertagia ostertagi), and (iv) Angiostrongyliidae, which includes Angiostrongylus cantonensis and Angiostrongylus costaricensis. Besides these families, Metastrongyliidae includes Metastrongylus spp., the lungworm of pigs, only accidentally transmitted to humans, and the family Syngamidae includes Syngamus trachea, which infects birds, and Mammmomonogamus laryngeus, a parasite of cats, ungulates, and orangutans, which can also rarely infect humans.

Some Strongylida parasites are zoonotic. Oesophagostomum bifurcum infects nonhuman primates and humans, being endemic in populations living in some regions of Togo and Ghana, being efficiently mitigated by collective chemotherapy with albendazole [4, 5]. Human infections with Trichostrongylus spp. have been reported in populations living in contact with goats and sheep or their feces in many regions [6, 7]. In humans, infections with Oesophagostomum spp. and Trichostrongylus spp. may be misdiagnosed as hookworm disease on parasitological stool examination [8, 9], mainly in regions where these infections are coendemic.

Although the hookworms which infect humans (N. americanus and Ancylostoma duodenale) are markedly anthropotonic, cross-host transmission of N. americanus among humans and nonhuman primates has been documented in Africa [10, 11]. There is scientific evidence that a potential zoonotic cycle of hookworm disease in the human-swine interface can be considered feasible. In El Salvador, living in close proximity to pigs was a recognized risk factor for hookworm infection [12]. In Ghana, viable N. americanus eggs were recovered from pigs’ feces [13]. Necator suli is a hookworm species described in pigs for almost a century [14], and investigations into its validity, occurrence, epidemiology, and zoonotic potential have not been forwarded.

The present work is part of a broader project, in which we assessed, through molecular tools (mitochondrial DNA sequencing), the zoonotic circulation of Strongylida and Ascaris in the human-swine interface in low-resource rural communities in the state of Piauí [15–18]. In this report, we present the results of the morphological study of Strongylida larvae obtained through cultivation of human and swine feces.

2. Materials and Methods

2.1. Setting and Study Design. The fieldwork was carried out in rural communities in Nossa Senhora de Nazaré, in the state of Piauí, northeastern Brazil. In the communities studied, pigs are raised extensively and roam freely on the streets, providing soil contamination with fecal matter (Figures 1(a) and 1(b)). The human population practices open defecation and sanitation conditions are poor. Households were visited by the research team to obtain sociodemographic data and human fecal samples in plastic bottles (n = 103 residents) without preservatives. Swine fecal samples (n = 27 pigs) were obtained after spontaneous defecation (n = 16) or necropsy (n = 11).

2.2. Parasitological Procedures. All fecal samples were screened for Strongylida eggs by conventional parasitological examination, including the Ritchie, Kato-Katz, and flotation in hypertonic glucose solution techniques [19–22]. Strongylida-positive fecal samples were submitted to coprocultures using the Harada–Mori technique [23]. Briefly, 1-2 g of feces was spread on a 150 × 15 mm filter paper. This strip was placed in a 20 × 200 mm Falcon tube containing 2-3 mL of distilled water. The tube was closed and kept in an upright position for approximately 10 days. Finally, the water was examined with a magnifying glass to check for the presence of larvae. Larvae were collected with Pasteur pipettes. The larvae were examined by light microscopy, and the morphological characteristics proposed in the taxonomic key presented in the chart were verified. Larvae were measured with ImageJ (U. S. National Institutes of Health, Bethesda, Maryland, USA). The ANOVA of the length of the larvae, with and without sheath, was performed and plotted in a graph using the software SPSS Statistics for Windows, version 26 (SPSS Inc., Chicago, Ill., USA).

3. Results

From a total of 103 residents who provided fecal samples, 12 (11.7%) were positive for Strongylida eggs, and 36 larvae were recovered, of which 22 could be submitted to morphological analysis. Regarding the 27 pigs studied, 16 had their feces collected after spontaneous defecation, and 11 had their feces collected at postslaughter necropsy. Of the total, 23 (85.2%) were positive for Strongylida eggs, and 87 larvae were recovered, of which 37 could be morphologically analyzed. All 22 larvae obtained from human feces were classified as hookworms (Ancylostomatidae). From a total of 37 larvae obtained from swine feces by coproculture, 23 (62.3%) were classified as belonging to the genus Oesophagostomum, 6 (16.2%) were Hystrostrongylus, and 4 (10.8%) were Trichostrongylus. Four larvae (10.8%) from pigs were classified as Strongyloides. The lengths of the analyzed larvae (mean and standard deviation) are shown in Figure 2.

Oesophagostomum larvae recovered from swine were significantly longer than hookworm larvae obtained from human feces. Trichostrongylus larvae also tended to be longer than hookworm larvae. The morphological characteristics of the larvae are shown in Figure 3. An identification key was proposed, based on morphological characteristics for the filariform larvae of some geohelminths of the order Strongylida in humans and swine (Supplementary Table 1).
4. Discussion

The present work demonstrates the feasibility of evaluating, through an accessible and inexpensive technique, the genus of Strongylida parasites infecting humans and domestic animals in scenarios where there could be zoonotic transmission and potential misidentification of eggs in fecal microscopic examinations.

No larvae of parasites usually identified in animals were detected among humans, that is, all larvae recovered from humans were morphologically characterized as hookworms. According to Hayashi et al. [24], the length of filariform (third stage) larvae can distinguish *N. americanus* (590–660 μm) from *Ancylostoma duodenalis* (660–720 μm).

The length of all hookworm larvae analyzed was compatible with *N. americanus*.

In previous works, we demonstrated that human hookworm disease is endemic in the studied area and that the species involved in the vast majority of cases was *N. americanus*, through DNA sequencing of the parasite’s cytochrome oxidase 1 coding gene (cox1) fragment [15, 25]. Nevertheless, in some hookworm-positive fecal samples, it was not possible to characterize the *Necator* species present in the material [15]. Previous reports on swine hookworms also motivated the investigation of the presence of *Necator* in pigs in the region [14, 26–29]. The results of the studies on pigs’ hookworms carried out in the 1920s and 1930s are conflicting, and it has not been clearly defined whether the
Figure 3: (a1)–(a4) Strongyloides larva obtained from swine. (a1). Long esophagus (approximately half the length of the larva), absence of sheath; (a3) straight intestine; (a4) forked tail end. (b1)–(b4) Hookworm larva obtained from human. (b1) Short esophagus (reaching 1/3 of the length of the larva), presence of sheath; (b2) marked esophageal bulb; (b3) straight intestine; (b4) tip of the tail. (c1)–(c4) Oesophagostomum larva obtained from swine. (c1) Short esophagus and presence of sheath; (c2) rounded cephalic end; (c3) accentuated sheath folds, zigzag intestine; (c4) long sheath. (d1)–(d4) Hyostrongylus larva obtained from swine. (d1) Short esophagus, presence of sheath; (d2) cephalic end in square shape; (d3) zigzag intestine; (d4) tip of the tail in finger shape. (e1)–(e4) Trichostrongylus larva obtained from swine. (e1) Short esophagus and presence of sheath; (e2) tapered cephalic end; (e3) zigzag intestine; (e4) short and blunt end of the sheath. A vertical bar in a1, b1, c1, d1, and e1 = 100 μm. Bar in the other images = 20 μm.
hookworm species *N. suillus* is valid or whether those infections were caused by *N. americanus*.

In the present study, the Strongylida genera infecting pigs were *Oesophagostomum*, *Hyostrongylus*, and *Trichostrongylus*, parasites usually present in swine extensively raised. In a previous work, we demonstrated that the *Oesophagostomum* species characterized in pigs in the region by cox1 partial sequencing were *O. dentatum*, *O. quadrispinulatum*, and *O. columbianum*, although the latter is a parasite usually found in goats and sheep [17].

*Hyostrongylus rubidis*, the red stomach worm, causes gastritis in pigs and is endemic in several regions, and its occurrence in swine has been described in Brazil [30]. Usually, its presence in herds is verified by the observation of gastric adult worms obtained after slaughter or suspected by the finding of Strongylida eggs in parasitological examinations of feces. The present report illustrates the feasibility of identifying *Hyostrongylus* larvae in swine Harada–Mori coprocultures.

Species belonging to the genus *Trichostrongylus*, mainly *T. colubriformis* and *T. axei* are usually found in small ruminants (goats and sheep) and also parasites and the digestive tract of pigs in many regions [31, 32]. In the studied locality, the close proximity in which small ruminants and pigs are raised may favor cross-host transmission of *Trichostrongylus*, which was detected infecting pigs in the present study. About 10% of the specimens studied had characteristics of *Strongyloides* filariform larvae. Swine strongyloidiasis, caused by *Strongyloides ransomi*, is a worldwide distributed disease more frequent in herds with extensive breeding, determining economic losses for breeders [33].

**5. Conclusions**

The morphological study of larvae obtained by coprocultures is a useful tool in the screening of intestinal helminthiasis and can help the characterization of zoonotic transmission cycles in a One Health approach in socioenvironmental scenarios of poor sanitation and close coexistence of humans and farm animals.

**Data Availability**

The data used to support the findings of this study are available from the corresponding author upon request.

**Ethical Approval**

This study was approved by the Ethics Committee of Instituto Oswaldo Cruz (Fiocruz) (CAAE: 12125713.5.0000.5248).

**Conflicts of Interest**

The authors declare that they have no conflicts of interest.

**Authors’ Contributions**

Polyanna A. A. Bacelar was involved in the planning of the experimental design, fecal sample collection, data analysis, and interpretation. Kerla J. L. Monteiro was involved in the planning of the experimental design, fecal sample collection, and data analysis. Jéssica P. Santos, Denilson de A. e Silva, Daniella N. Leal, Mayron M. Almeida, Brenda B. C. Evangelista, and Francisco M. de Oliveira-Neto were involved in fecal sample collection and data analysis. Filipe A. Carvalho-Costa was involved in the data analysis and the drafting of the manuscript.

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**Supplementary Materials**

Supplemental Table 1: Identification key based on morphological characters for the filaroid larvae of some geo-helminths of the order Strongylida in humans and swine [3, 34]. (Supplementary Materials)

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