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Rosa Fernández1*, Sebastian Kvist1, Jennifer Lenihan1, Gonzalo Giribet1, Alexander Ziegler2

1 Museum of Comparative Zoology, Department of Organismic and Evolutionary Biology, Harvard University, Cambridge, Massachusetts, United States of America, 2 Ziegler Biosolutions, Waldshut-Tiengen, Germany

Abstract

In spite of the high relevance of lumbricid earthworms (‘Oligochaeta’: Lumbricidae) for soil structure and functioning, the taxonomy of this group of terrestrial invertebrates remains in a quasi-chaotic state. Earthworm taxonomy traditionally relies on the interpretation of external and internal morphological characters, but the acquisition of these data is often hampered by tedious dissections or restricted access to valuable and rare museum specimens. The present state of affairs, in conjunction with the difficulty of establishing primary homologies for multiple morphological features, has led to an almost unrivaled instability in the taxonomy and systematics of certain earthworm groups, including Lumbricidae. As a potential remedy, we apply for the first time a non-destructive imaging technique to lumbricids and explore the future application of this approach to earthworm taxonomy. High-resolution micro-computed tomography (μCT) scanning of freshly fixed and museum specimens was carried out using two cosmopolitan species, Aporrectodea caliginosa and A. trapezoides. By combining two-dimensional and three-dimensional dataset visualization techniques, we demonstrate that the morphological features commonly used in earthworm taxonomy can now be analyzed without the need for dissection, whether freshly fixed or museum specimens collected more than 60 years ago are studied. Our analyses show that μCT in combination with soft tissue staining can be successfully applied to lumbricid earthworms. An extension of the approach to other families is poised to strengthen earthworm taxonomy by providing a versatile tool to resolve the taxonomic chaos currently present in this ecologically important, but taxonomically neglected group of terrestrial invertebrates.


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* E-mail: rfernandezgarcia@g.harvard.edu

Introduction

In one of the first comprehensive investigations into earthworm (Clitellata: ‘Oligochaeta’) taxonomy, written more than a century ago, Michaelsen [1] featured the prophetic words sine systemate chaos in the title of his treatise. This seminal work, together with Stephenson’s contribution to earthworm systematics published thirty years later [2], has constituted the backbone of the classical phylogenetic system of earthworms. However, in many higher earthworm taxa, the systems proposed by Michaelsen and Stephenson were based on a small number of key characters, which unfortunately failed to create a sound classification of the group. Since then, earthworm systematics has been gradually eroded by an ongoing controversy over how to classify various groups of these organisms. References to the instability of earthworm taxonomy are frequent in the literature (e.g., [3]), and studies have emphasized the lack of robust classifications on various taxonomic levels (e.g., ‘the family-level classification of the megascolecid earthworms is in chaos’ [4]).

In particular, this unfortunate state of affairs holds true for Lumbricidae Claus, 1876. The lumbricid system of classification has been referred to as an ‘unequalled chaos’ [5,6]. Despite their biological, ecological and economic importance, the taxonomic status and evolutionary relationships of most lumbricid genera are still under debate. The lack of agreement concerning the ranking of diagnostic morphological characters in lumbricid earthworms has led to the creation of numerous synonyms at the species or genus level. For instance, the overlap of certain morphological features (e.g., shape of prostomium; position of chaetae; position and shape of sexual organs including clitellum, tubercula pubertatis, testes, ovaries, and spermathecae) has complicated
species identification. Unfortunately, overlap in these key characters in closely- and distantly-related species is pervasive [6–13]. The instability of lumbricid earthworm taxonomy can be linked to three major issues: a high rate of synonymy, the establishment of ‘catch-all’ polyphyletic genera, and the existence of broad ‘species complexes’ [6–9,14].

During the last years, newly developed techniques such as DNA-based systematics, DNA barcoding, or integrative taxonomy [15–17] have been used to complement the classical, strictly morphology-based taxonomy in earthworms. Earthworm identification requires detailed anatomical studies, which have traditionally been performed through specimen dissection [7]. Also, earthworm dissections can be a time-consuming affair, require a high degree of expertise and, importantly, often cannot be applied to valuable and rare museum specimens. In order to increase the speed of advancement, taxonomists are in dire need of quick, reproducible, and non-destructive techniques that would permit the investigation of internal earthworm anatomy more reliably and on a larger scale.

A number of imaging techniques can be employed to successfully visualize internal structures of zoological specimens non-invasively, including minute samples. The list of methodologies comprises optical projection tomography [18], magnetic resonance imaging [19], synchrotron-radiation micro-computed tomography [20,21], autofluorescence imaging [22], and microcomputed tomography [23]. Recent reviews provide a comprehensive overview of non-invasive imaging techniques and their application to terrestrial and marine organisms [24,25]. Because of its high speed, low cost, ease of use, and the achievable high dataset resolution, micro-computed tomography (μCT) has recently evolved to become an important imaging tool in zoological studies. Based on the physical principle of X-ray imaging, μCT is particularly suitable for the visualization of mineralized tissues, and has thus been broadly applied in paleontology (e.g., [26,27]). However, electron-dense contrast agents such as iodine, lead, tungsten, or osmium can be employed to visualize soft parts as well [28–34].

Using this approach, samples pertaining to a wide range of extant fauna have been successfully analyzed in the past, with the animals under study either being scanned in the form of isolated body parts or as whole, intact specimens. These taxa include Poriëra [35], Chelicerrata [36–39], Hexapoda [40–46], Bryozoa [47], Polychaeta [48], Mollusca [49,50], and Vertebrata [29–34,51–57]. Preliminary analyses of a clitellate (Porifera) and several other phyla are ongoing. These include 3% dimethyl sulfoxide (DMSO) to increase cell membrane permeability, in ethanol (at 95% concentration). In order to provide a further tool for earthworm taxonomy, the present study aims to assess the usefulness of μCT scanning in combination with soft tissue staining to identify morphological characters of importance for earthworm taxonomy, and to compare the application of this non-destructive imaging approach to museum and freshly fixed specimens. Furthermore, the quality of μCT-derived data in comparison with results derived from traditional dissection techniques are discussed.

Materials and Methods

Specimens

Six specimens were used for μCT scanning: three freshly fixed specimens of Aporrectodea caliginosa collected in 2013 in Cambridge, Massachusetts, USA; a museum specimen collected in 1967 in Mérida, Venezuela; a freshly fixed specimen of A. trapezoides collected in 2013 in Menorca, Balearic Islands, Spain; and a museum specimen collected in 1945, in Oregon, USA. Freshly fixed specimens were collected either by digging 20–40 cm into soil or by surface collection following rainfall. The two species are not listed as endangered or protected, and no specific collecting permits were required for the four localities. All specimens were identified following the key provided in [3]. Table 1 gives an overview of the specimens employed for imaging, as well as GPS coordinates for some of the localities. All the freshly fixed and museum specimens that were scanned in this study are deposited in the collection of the Department of Invertebrate Zoology (IZ), Museum of Comparative Zoology (MCZ), Cambridge, Massachusetts, USA. Specimens information and associated images can be found on MCZbase (http://mczbase.mcz.harvard.edu).

Specimen staining

The six specimens intended for scanning were stained in 50 mL Falcon tubes (BD Biosciences, San Jose, California, USA) on a rocking table, using either a 10% iodine solution (Logol’s solution, iodine potassium iodide, I2KI) in ethanol (at 95% concentration) or a 0.3% phosphotungstic acid (PTA) solution, which also included 3% dimethyl sulfoxide (DMSO) to increase cell membrane permeability, in ethanol (at 95% concentration). Following an initial trial phase with different solutions as well as various lengths of staining and scanning times, specimens were finally stained for a total of three to four weeks using the PTA solution. Iodine staining was rejected due to the potentially significant specimen shrinkage that may occur at higher concentrations of I2KI [62]. Once stained with the PTA solution, specimens were placed in plastic drinking straws with 6 mm diameter, which were heat-sealed at the bottom, filled with the solution in which the worms had been kept during staining, and closed at the top end using Parafilm (Pechiney Plastic Packaging Co., Chicago, Illinois, USA). It is important to note that, typically, stained specimens are submerged in clean ethanol for scanning (e.g., [28,47]) However, we noticed that performing the scan with the organisms still submerged in the PTA solution used for staining did not lead to negative results. We assume that most of the staining molecules must have been absorbed by the tissue. In order to increase the isotropic voxel resolution of the three-dimensional (3D) datasets, only the first ca. 35 segments of each specimen were...
Table 1. List of specimens used for μCT scanning.

<table>
<thead>
<tr>
<th>Species</th>
<th>Specimen data</th>
<th>Specimen size</th>
<th>Staining parameters</th>
<th>Scanning parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aporrectodea caliginosa</td>
<td>MCZ IZ 25152; collected March 10, 2013, Cambridge, Massachusetts, USA; GPS: 42.378658, 71.11616; fixed in 95% EtOH, stored in 95% EtOH</td>
<td>4.1 mm diameter, 6.4 cm length</td>
<td>10% I2KI for 4 days</td>
<td>65 kV, 100 μA, 50 min 4 s, 8.88 μm</td>
</tr>
<tr>
<td></td>
<td>See above</td>
<td></td>
<td>10% I2KI for 10 days</td>
<td>65 kV, 100 μA, 32 min 58 s, 8.88 μm</td>
</tr>
<tr>
<td>Aporrectodea caliginosa</td>
<td>MCZ IZ 25150; collected March 10, 2013, Cambridge, Massachusetts, USA; fixed in 95% EtOH, stored in 95% EtOH</td>
<td>3.6 mm diameter, 8.1 cm length</td>
<td>0.3% PTA for 4 days</td>
<td>70 kV, 100 μA, 48 min 40 s, 7.11 μm</td>
</tr>
<tr>
<td>Aporrectodea trapezoides</td>
<td>MCZ IZ 24805; collected March 10, 2013, Cambridge, Massachusetts, USA; fixed in 95% EtOH, stored in 95% EtOH</td>
<td>4.5 mm diameter, 6.1 cm length</td>
<td>0.3% PTA for 21 days</td>
<td>70 kV, 100 μA, 1 h 20 min 54 s, 9.95 μm</td>
</tr>
<tr>
<td>Aporrectodea trapezoides</td>
<td>MCZ IZ 24804; collected January 3, 2013, Menorca, Spain; GPS: 40.01056, 3.87817; fixed in 95% EtOH, stored in 95% EtOH</td>
<td>5.2 mm diameter, 8.0 cm length</td>
<td>0.3% PTA for 28 days</td>
<td>70 kV, 100 μA, 1 h 37 min 1 s, 13.15 μm</td>
</tr>
<tr>
<td>Aporrectodea trapezoides</td>
<td>MCZ IZ 95901; collected December 14, 1945, Oregon, USA; fixed in formalin, stored in 70% EtOH</td>
<td>4.5 mm diameter, 10.8 cm length</td>
<td>0.3% PTA for 28 days</td>
<td>70 kV, 100 μA, 1 h 37 min 5 s, 8.17 μm</td>
</tr>
</tbody>
</table>

Scanning parameters were: source voltage (kV), source current (μA), scanning time (h, min, s), and isotropic voxel resolution of the reconstructed 3D dataset (μm). The embedded hyperlinks provide direct access to the specimen entries in MCZbase (e.g., MCZ IZ 25152).
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scanned. These segments harbor all internal and external structures commonly used as diagnostic characters in earthworm taxonomy.

Micro-computed tomography

Imaging was performed using a SkyScan 1173 μCT scanner (Bruker MicroCT, Kontich, Belgium) equipped with a Hamat-matsu 130/300 tungsten X-ray source and a FlatPanel Sensor camera detector with 2240×2240 pixels. Scanning parameters were as follows: source voltage = 65–70 kV, source current = 100 μA, exposure time = 1,000 ms, frames averaged = 2–6, frames acquired over 180° = 960, filter = no, binning = no, flat field correction = activated, and scanning time = about 45–100 min. Reconstruction of the raw data was accomplished using the software provided with the scanner (NRecon 1.6.6.0, Bruker MicroCT, Kontich, Belgium). Various settings were employed to enhance image contrast and to compensate for ring and streak artifacts. These dataset reconstruction parameters were: smoothing = no, ring artifact correction = 5–11, and beam hardening correction = activated.

The four most representative μCT scans have been deposited in the voxel repository GigaDB [63]. In addition, an accompanying publication provides further information about data availability, quality, and requirements [64].

Dissection and photography

In order to compare our μCT scanning results with traditional dissections commonly carried out by earthworm taxonomists, freshly fixed specimens identified as A. caliginosa, collected at the same time and location as the conspecific specimens used for μCT scanning, were dissected. These dissections were performed by making a dorsal incision to the body wall and exposing the internal organs. This destructive approach involves removal of a piece of integument to expose the internal organs, and partial or complete removal of some internal structures, such as the seminal vesicles or the septa. Auto-montage images of the dissected animals were acquired using a Leica MZ12.5 stereomicroscope (Leica Microsystems, Wetzlar, Germany) with an attached JVC KY-F75U digital imaging camera (JVC, Wayne, New Jersey, USA). Image stacks consisting of 17 separate photographs (exposure = 250 ms) were merged using the software Auto-Montage Pro 5.02.0096 (Syncroscopy, Frederick, Maryland, USA) with the following settings: method = fixed, optimize = precision, and patch size = 95.

3D visualization and modeling

The acquired 3D μCT datasets were analyzed using computer systems equipped with a 64-bit Windows operating system (Windows 7, Microsoft Co., Redmond, Washington, USA), a multi-core CPU, as well as a minimum of 6 GB of main and 1 GB of video RAM. Two-dimensional (2D) image slicing was accomplished with the free software DataViewer [http://www.skyscan.be/products/downloads.htm] and ImageJ 1.44p [http://imagej.nih.gov/ij/] using the plug-in Volume Viewer 2.01 [http://rsb.info.nih.gov/ij/plugins/volume-viewer.html]. 3D rendering was performed by employing the commercial software Amira 5.2 (FEI Co., Hillsboro, Oregon, USA) as well as the free software Drishti (http://sf.anu.edu.au/Vizlab/drishti/index.shtml). The interactive 3D PDF model was created by manually segmenting, then
surface-rendering and smoothing selected structures in Amira 5.2, assembling them in the commercial Adobe 3D Reviewer software (Adobe Systems, San Jose, California, USA), before finally exporting them as an interactive PDF file [65,66].

Results

The first section of the Results provides a description of structures identifiable in the µCT dataset of the freshly fixed specimen MCZ IZ 24805 (*Aporrectodea caliginosa*), which had been PTA-stained for four weeks. File S1 shows virtual 2D sections through the entire dataset. Following this first section, we present results concerning the morphological differences between the two *Aporrectodea* species, and a comparison of the performance of the approach on freshly fixed and museum material. In addition, we provide information on scanning artifacts encountered during the experiments. All acquired µCT datasets were analyzed qualitatively using 3D volume rendering techniques in combination with 2D virtual dataset slicing tools. In addition, a 3D PDF model, which permits interactive access to selected internal and external structures is available as File S2. The accompanying descriptive notes are provided as File S3.

Identification of characters conventionally used in earthworm taxonomy

**External features.** A classical 2D X-ray projection image of the anterior part of the PTA-stained, freshly fixed specimen of *A. caliginosa* reveals full penetration of the staining agent throughout all tissues (Fig. 1A). The general shape of the anterior body, its segments, as well as the intersegmental furrows can be readily identified in the µCT-based 3D volume renderings of the external surface (Fig. 1B-D). The prostomium, which surrounds the mouth, is epilobous and extends into a third of the first segment, the peristomium (Fig. 1B, C). The clitellum lies dorsally and begins to form at segment XXVII. The precise position of pores, often relevant for taxonomic identification, tends to be difficult to identify using traditional techniques. The µCT dataset allows unambiguous identification of most of these structures, in particular when 3D renderings (Fig. 1B-D) and virtual 2D sections (Fig. 2) are used in combination. On the dorsal side, the dorsal pores cannot be identified in the 3D renderings (Fig. 1B), but are visible in the 2D virtual sections (Fig. 2A). A number of features can be discerned on the ventral side (Fig. 1C), in particular the female (XIV) and male pores (XV). The latter constitute a large slit between chaetae bc and are surrounded by a glandular crescent protruding into the neighboring segments. Furthermore, the papillae of chaetae ab in segments IX, X, and XI can be identified. The nephridiopores are too small to be seen in the 2D virtual sections (Fig. 2). The course of the sediment as well as its composition throughout the digestive tract can be readily observed (Fig. 2). The course of the sediment as well as its composition throughout the digestive tract can be traced without difficulty – see File S4 for animated 3D volume renderings of the entire dataset. The anterior foregut is composed of the mouth (surrounded by the prostomium), the buccal cavity (I–III), a muscular pharynx (III–V) with dozens of pharyngeal muscles that can be traced all the way to their attachment sites on the dorsal and lateral body wall, and a thickened, muscular esophagus (VI–XI). The calciferous gland is characterized by the presence of two well-developed pouches with multiple layers in segment X (Fig. 3C) and the absence of lateral enlargements in segment XIII. The posterior foregut is composed of a wide crop (XV–XVI), followed by a muscular gizzard (XVII–XVIII). This latter structure particularly well in the 2D virtual sections. The chaetigerous sacs and muscles are in general difficult to discern, but those of the sexual chaetae can be identified in the virtual 2D sections.

**Body wall.** A transverse section through a given segment permits identification of parts of the body wall. Beginning from the outside, the cuticle is followed by the epidermis, which surrounds the circular and longitudinal muscle layers, and finally, the peritoneum, which borders the coelomic cavity. While the cuticle, the epidermis, and the two muscle layers can be differentiated in most sections (Fig. 2), the peritoneum is too thin to be discerned with certainty at the given dataset resolution (9.95 µm isotropic).

**Digestive system.** Most compartments of the digestive tract can be readily identified (Fig. 2). The course of the sediment as well as its composition throughout the digestive tract can be traced without difficulty – see File S4 for animated 3D volume renderings of the entire dataset. The anterior foregut is composed of the mouth (surrounded by the prostomium), the buccal cavity (I–III), a muscular pharynx (III–V) with dozens of pharyngeal muscles that can be traced all the way to their attachment sites on the dorsal and lateral body wall, and a thickened, muscular esophagus (VI–XI). The calciferous gland is characterized by the presence of two well-developed pouches with multiple layers in segment X (Fig. 3C) and the absence of lateral enlargements in segment XIII. The posterior foregut is composed of a wide crop (XV–XVI), followed by a muscular gizzard (XVII–XVIII). This latter structure

![Figure 1. Visualization of the µCT dataset of a PTA-stained specimen of *Aporrectodea caliginosa* (MCZ IZ 24805, freshly fixed specimen). Anterior facing upwards (A–C) or to the right (D). (A) 2D X-ray projection image of the dataset. (B) Dorsal view of a false-color volume rendering of the dataset. (C) Ventral view. (D) Lateral view. Abbreviations: Cab, chaetae ab; Ccd, chaetae cd; Cli, clitellum; Fpo, female pore; Isf, intersegmental furrow; Mpo, male pore; Per, peristomium; Pro, prostomium; Seg, segment; Spg, sperm groove; Tpu, tubercula pupertatis. Roman numerals denote segment numbers. doi:10.1371/journal.pone.0096617.g001](image)
terminates in a pre-intestinal valve. Following the foregut, which is entirely lined by cuticle, the midgut is composed of the intestine (starting at segment XIX). The intestinal walls are thinner than those of most parts of the foregut. A particularly saccular part of the intestine with numerous transverse folds can be found in the segments directly following the gizzard (XIX–XXVI). A prominent spade-shaped typhlosole starts to form in segment XXII. The elements of the hindgut (i.e., rectum and anus) are outside the field of view of the present dataset.

Vascular system. The intense staining of hemoglobin inside the vascular system facilitates the detection of blood vessels. They appear as clearly contoured dilations with a bright filling (Figs. 2, 3). The dorsal blood vessel can be traced through almost the entire length of the dataset. The ventral blood vessel can equally be traced all the way to the end of the field of view. In segments VI–XI, paired lateral hearts of varying sizes connect the dorsal blood vessel to its ventral counterpart. In the posterior part of the dataset, dorsoparietal vessels join the dorsal blood vessel with the small subneural vessel. The lateral neural and the ventroparietal vessels, as well as the dorsointestinal and ventrointestinal vessels can be identified in some of the virtual 2D sections, but the given resolution of the dataset (or the lack of hemoglobin in these parts of the vessels) somewhat limit their visibility.

Excretory system. Throughout most segments, metanephridia can be observed. These structures can be identified as structures with J-shaped bladders (Fig. 2B) that are hooking caudally. The nephrostome is too small to be clearly visualized at the given dataset resolution, while the nephridiopores are visible.

Reproductive system. Two pairs of sperm-filled spermathecae can be observed in intersegments IX–X and X–XI (Fig. 2). Their pores open in-between segments IX and X as well as segments X and XI. The testes are barely visible, but their sperm funnels are clearly shown. From these, a pair of vasa deferentia leading to the male pores is seen in segment XV. The massive, paired seminal vesicles can be seen in segments IX to XII, occupying the majority of the coelomic cavity. Paired metagynous ovaries in segment XIII and paired ovisacs in segment XIV (both containing several eggs) can be identified and are closely associated with the oviducts, which in turn lead to the female pores of segment XIV.

Nervous system. The brain is composed of paired supraesophageal (or cerebral) ganglia that lie dorsal to the buccal cavity.
in segment III (Fig. 2). The paired cerebral commissure connects the supraesophageal ganglia through segment IV with the paired subesophageal ganglia lying in segment V. The large ventral nerve cord starts to form in segment VI. The segmental nerves emanating from the ventral nerve cord can be seen in some sections, but never extending further than chaetae cd. The finer nerves are too small to be seen at the given dataset resolution.

**Aporrectodea caliginosa or Aporrectodea trapezoides? Differences between the two analyzed species**

Two pairs of sperm-filled spermathecae can be observed in intersegments IX–X and X–XI of the freshly fixed specimen of *A. caliginosa* (Fig. 2), while only empty spermathecae can be observed in the two specimens of *A. trapezoides* as well as the museum specimen of *A. caliginosa* (Fig. 3). In addition, not only the spermathecae, but also the seminal vesicles are much smaller in these three latter specimens compared to the freshly fixed individual of *A. caliginosa*. In the freshly fixed specimen of *A. caliginosa*, spermathecae and seminal vesicles occupy a large part of the coelomic cavity in segments IX–XII. While the presence of empty spermathecae may not be an indicator for parthenogenetic reproduction, several anatomical differences seem to indicate that the museum specimen labeled as *A. caliginosa* (MCZ IZ 95557) could be a parthenogenetic individual. However, because *A. caliginosa* is a sexual species (no parthenogenetic specimens have ever been described), this specimen could be a case of misidentification, and the sample may actually be *A. trapezoides* (or any senior synonym in use at the time of identification).

Ultimately, the solution for the problematic taxonomy of these two extremely similar species will rest on the inspection of the topotype of *A. caliginosa*, which is presumably located at the Muséum National d’Histoire Naturelle in Paris, France, and on the designation of a neotype of *A. trapezoides* from the possible type locality of this species (i.e., Montpellier, France) [3].

**Comparison of the performance of μCT scanning between freshly fixed and museum specimens**

In terms of the efficacy of the staining procedure and the resulting image quality, we found no significant differences between freshly fixed specimens and those stored in formaldehyde or ethanol for extended periods of time. Similarly, we did not find any differences in the performance of the approach between the two species used in the present study. The only factor that seems to affect the method chosen here is body size. Due to the relatively slow rate of diffusion of the staining agent, any increase in specimen width will result in a longer staining time. Our initial trial phase with different durations of staining led to insufficient penetration of specimens stained for about a week. This problem was successfully solved by staining all specimens for longer periods of time (i.e., three to four weeks).

The only notable difference between freshly fixed and museum specimens was that, depending on the fixation protocol originally used, some museum specimens were more contracted than the freshly fixed ones, resulting in occasional differences in organ size and shape. Furthermore, we found that it is of importance to properly arrange the animals in a standardized orientation prior to scanning. Therefore, all specimens were scanned following dorso-ventral and rostro-caudal straightening inside plastic straws. This position closely resembles the orientation of earthworms during dissection, facilitating a comparison of results derived from both techniques.

**Artifacts**

Artifacts related to the specimen’s biology, the staining procedure, or the scanning methodology may affect dataset quality. Pockets of air (Fig. 4A) were found inside the digestive tract of most specimens (Figs. 2A, B; 3A, B). These air pockets appear as black voids, because the staining solution does not diffuse into them. However, these artifacts are easily identified, do not affect volume rendering, and do not compromise the integrity of the surrounding anatomical features. Another problem encountered was the presence of pronounced streak artifacts caused by electron-dense sediment particles with higher X-ray attenuation than that of the surrounding tissues (Fig. 4B).

Furthermore, imperfect staining was a prominent artifact seen in all specimens treated with short staining times (Fig. 4C). This artifact is caused by the relatively slow molecular diffusion rate of the staining solution into the specimen. The almost circular void at the core of the animal shows that the size of the staining molecule and animal width are the main properties governing the transfer rate of the staining solution into the specimen. The thickest specimen analyzed (3.2 mm diameter, center specimen in Fig. 5) showed imperfect staining, despite four weeks of consecutive immersion in the staining solution. Based on the six specimens scanned, a diffusion rate of approximately 1 mm per week was estimated for the PTA solution used.

Another type of artifact frequently observed was the presence of concentric rings (Fig. 4D), which are caused by faulty detector elements on the pixel detector array. Algorithms can be employed to remove most ring artifacts during dataset reconstruction. However, an excessive presence of staining molecules within the liquid surrounding the specimen during scanning cannot be compensated for (Fig. 4E). Finally, movement of specimens during scanning resulted in datasets with blurred imagery and consequently a reduced isotropic voxel resolution (Fig. 4F).

**Discussion**

The present study is the first to explore the systematic application of a non-destructive imaging technique to earthworm specimens. One of the main advantages of the use of modern imaging techniques is that the acquired data are digital in nature and can thus be employed for numerous advanced visualization methods that would have been impossible using non-digital approaches. An example is provided in Figure 5, which shows a direct comparison of results derived from traditional earthworm dissection techniques (Fig. 5A, B) with a virtual dissection of a µCT dataset (Fig. 5C). One of the most conspicuous benefits of the virtual dissection is the possibility of observing internal organs in their natural anatomical context, which is often not the case when traditional dissection techniques are employed. Moreover, the
specimens can be visualized and rotated in 3D, allowing for a better understanding of their complex morphology.

Furthermore, the present study demonstrates that the time span between initial specimen fixation and scanning does not represent a limiting factor for the performance of the technique. However, the fixation event itself is of importance, because variations in organ size and shape due to shrinkage or inflation may have occurred. Nonetheless, this effect would certainly also influence morphological results derived from traditional dissections of such specimens. Differences in the performance of the approach when applied to separate earthworm species can also be discarded, because staining properties of the same earthworm organs are likely to be similar, regardless of the species. However, one of the potential limitations of the application of μCT scanning in combination with soft tissue staining is the presence of artifacts (Fig. 4).

With regard to artifacts, the size of the specimen is the most important criterion, because the relatively slow diffusion rate of the various chemical agents used to stain biological tissues [28,30,31,47,70–72] can reduce the applicability of this approach to larger specimens. However, given the relatively small diameters of most earthworm species (in the millimeters to few centimeters...
on zoological studies now offer access to μCT systems, thus demonstrating the wide availability of this methodology. In this context, μCT scanning in combination with soft tissue staining constitutes a versatile tool for multidisciplinary approaches including morphological, ecological, and molecular methods that aim to assess the validity of characters of taxonomic relevance in earthworms.

Conclusions

Earthworm taxonomy is in dire need of tools that can be used to provide further insights into the details of animal anatomy and morphology and that allow for the non-destructive analysis of type material or other valuable and rare museum specimens. Our study shows that μCT in combination with soft tissue staining can be successfully applied to earthworm taxa—whether freshly fixed or museum specimens are used—in order to investigate internal and external features that would otherwise be impossible to determine without dissection. At the isotropic voxel resolutions that were achieved, we were able to identify numerous internal structures such as spermathecae, seminal vesicles, metanephridia, ganglia, and sperm funnels, as well as features of the vascular and digestive systems. In addition, external features that may be difficult to identify through traditional dissection techniques (e.g., dorsal pores, nephridiopores, or pores of the spermathecae) are clearly discernible in the datasets. The protocols used in the present study can be employed to produce high-resolution imagery of hundreds of specimens in a much shorter timeframe than traditional protocols would permit. Moreover, the approach can likely be applied to various other taxa with similar anatomical characteristics, such as Hirudinida, Onychophora, Pentastomida, Sipuncula, Echiura, Priapulida, or Phoronida. The future advancement of earthworm taxonomy will likely come to rely upon non-invasive imaging techniques that allow rapid generation of massive amounts of digital data—an approach that is already commonplace in several other biological disciplines.

Supporting Information

File S1 Video file showing the full image stack of the μCT-scanned, PTA-stained specimen of Aporrectodea caliginosa (MCZ IZ 24805, freshly fixed specimen). (PDF)

File S2 Interactive 3D PDF model of selected earthworm structures. The 3D model is based on a μCT-scanned, PTA-stained specimen of Aporrectodea caliginosa (MCZ IZ 24805, freshly fixed specimen). Left-click to activate the embedded multimedia content (requires the use of Adobe Reader 9 or higher on Windows, Mac, and Linux systems). Use the ‘+/-’ or ‘toggle full-screen’ options in order to maximize window size. Various pre-saved views can be accessed through the menu inside the viewer window or by opening the model hierarchy using the model tree icon. Grey, body wall; pink, digestive tract; red, circulatory system; brown, pharyngeal musculature; yellow, nervous system; orange, metanephridium; green, muscular septum; cyan, seminal vesicle; blue, spermatheca. (PDF)

File S3 Text file providing a step-by-step walkthrough of the pre-saved views shown in the interactive 3D PDF model (File S2). The views mentioned in this description can be accessed through the menu inside the viewer window or by opening the model hierarchy using the model tree icon. (PDF)
File S4 Video file showing a virtual dissection of the anterior part of a µCT-scanned, PTA-stained specimen of Aporrectodea caliginosa (MCZ IZ 24805, freshly fixed specimen). The video presents volume-rendered, false-colored coronal, sagittal, and transverse 3D views through the body of the specimen. Left-click to activate the embedded multimedia content (requires the use of Adobe Reader 9 or higher on Windows, Mac, and Linux systems). (PDF)

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Author Contributions
Conceived and designed the experiments: RF SK JL AZ. Performed the experiments: RF SK JL. Analyzed the data: RF SK JL AZ. Contributed reagents/materials/analysis tools: RF SK JL GG AZ. Wrote the paper: RF SK JL GG AZ.

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