

A web-based histology atlas for the freshwater sentinel species

Daphnia magna

Short title: Web-based *Daphnia* histology atlas

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Abstract

Daphnia are keystone species of freshwater habitats used as model organisms in ecology and evolutionary biology. Their small size, wide geographic distribution, and sensitivity to chemicals make them useful as environmental sentinels in regulatory toxicology and chemical risk assessment. Biomolecular (-omic) assessments of responses to chemical toxicity, which reveal detailed molecular signatures, become more powerful when correlated with other phenotypic outcomes (such as behavioral, physiological, or histopathological) for comparative validation and regulatory relevance. However, the lack of histopathology or tissue phenotype characterization of this species presently limits our ability to access cellular mechanisms of toxicity. Here, we address the central concept that interpreting aberrant tissue phenotypes requires a basic understanding of species normal microanatomy. We introduce the female and male *Daphnia* Histology Reference Atlas (DaHRA) for the baseline knowledge of *Daphnia magna* microanatomy. Additionally, we also included developmental stages of female *Daphnia* in this current atlas. This interactive web-based resource of adult *Daphnia* features overlaid vectorized demarcation of anatomical structures whose labels comply with an anatomical ontology created for this atlas. We demonstrate the potential utility of DaHRA for toxicological investigations by presenting aberrant phenotypes of acetaminophen-exposed *D. magna*. We envision DaHRA to facilitate the effort of integrating molecular and phenotypic data from the scientific community as we seek to understand how genes, chemicals, and environment interactions determine organismal phenotype.

Keywords: *Daphnia magna*, sentinel, microanatomy, atlas, phenotypes, sexual dimorphism, histopathology, toxicology

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25 **1. Introduction**

26 Environmental pollution is the leading cause of premature morbidity and mortality globally
 27 (Fuller et al., 2022; Landrigan et al., 2018; Naidu et al., 2021). Habitat loss, climate change, and
 28 pollution also impact biodiversity, with more than 60% of ecosystems services being diminished
 29 in the last two decades (Cardinale et al., 2012). New approach methodologies (Stucki et al.,
 30 2022) for assessing chemical toxicity are developed to improve regulatory outcomes by replacing
 31 outdated, data-poor methods dependent upon apical endpoints (such as death or reproductive
 32 failure) with data-rich molecular data (e.g. transcriptomic and metabolomic) (Harrill et al., 2021;
 33 Hines et al., 2010; Palmer et al., 2020). These data are robust at measuring biomolecular activity
 34 that are potentially indicative of chemical modes of action. However, these data lack spatial
 35 resolution, context within the whole organism, or correlation with associated abnormal tissue
 36 phenotypes that can be highly informative with regard to potential human toxicity (European
 37 Chemicals Agency, 2020). Since understanding abnormal tissue phenotypes requires knowledge
 38 of normal microanatomy (microscopic anatomy or histology), the primary objective of this paper
 39 is to provide a resource for visualizing and interpreting tissue phenotypes for the model species
 40 *D. magna* – an organism used globally to set regulatory limits on potentially hazardous chemical
 41 substances in the environment (United States Environmental Protection Agency, 1996, 2002;
 42 Organisation for Economic Co-operation and Development, 2004, 2018), and one of five models
 43 being used to uncover evolutionarily conserved toxicity pathways (“The Precision Toxicology
 44 initiative,” 2023).

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The water flea *Daphnia* is a keystone branchiopod crustacean (order *Cladocera*) in freshwater lotic ecosystems worldwide and an established model in ecology, evolutionary biology, and ecotoxicology (Ebert, 2022; Miner et al., 2012; Stollewerk, 2010). They are responsive to environmental change and adapt via evolutionary mechanisms and plasticity (Cuenca-Cambronero et al., 2021; Stoks et al., 2016; Walsh et al., 2018). Relevant to ecotoxicity testing is their short generation time that enables the experimental manipulation of large populations and a parthenogenetic life cycle that allows the rearing of populations of identical clones (Hebert and Ward, 1972). The latter property has the unique advantage of facilitating the concurrent study of molecular and phenotypic responses to multiple environmental insults, including chemical pollutants (Abdullahi et al., 2022; Cuenca Cambronero et al., 2018). *Daphnia magna* is a model species for ecotoxicogenomics (Kim et al., 2015; Shaw et al., 2008). Recently, its hologenome (Chaturvedi et al., 2023), reference genome (Byeon et al., 2022; Lee et al., 2019) and transcriptome (Campos et al., 2018; Jankowski et al., 2022; Orsini et al., 2016) have been published, elevating this species to the ranks of other biomedical model species for ecological genomics. The full potential of this species is best realized when correlations can be established between molecular, and tissue- and cell- specific phenotypes.

Histopathology, the microscopic examination of diseased tissues, enables the identification of targets of toxicity and diseases, bridging phenotypes and biomolecular perturbations induced by environmental insults (Majno and Joris, 2004; Wester and Canton, 1991). Histopathology-based toxicological studies in fish (Huang et al., 2021; Manjunatha et al., 2022; Ramírez-Duarte et al., 2008) and bivalves (Fraga et al., 2022; Joshy et al., 2022) have been useful for water quality monitoring and assessment. The application of histopathology to millimeter-size sentinel species

used in ecotoxicology would enable the analysis of tissue-specific toxicity phenotypes in the context of the whole animal. However, identification of affected cell and tissue types requires prior knowledge of normal microanatomy; the use of web-based atlases maximizes accessibility across research and educational communities (Copper et al., 2018; Graham et al., 2015; van der Ven et al., 2003).

Here we present the first curated web-based female and male Daphnia Histology Reference Atlas (DaHRA; RRID:SCR_024913), further broadening the discovery capacity of this sentinel species. We have optimized methods for *D. magna* histology and created a collection of digitized histological images for adult female and male *D. magna* in three anatomical planes to illustrate sexual dimorphism associated with environmentally induced phenotypic plasticity. We also present a subset of developmental stages showcasing some representative developmental events. As proof-of-concept, we also present histological alterations in *D. magna* caused by exposure to toxic levels of a common pharmaceutical painkiller, acetaminophen, to demonstrate how our platform can facilitate whole-organism visualization and comparison for an experiment. This resource is made open-access and interactive, allowing smooth magnification with a dynamic scale bar. Anatomical structures are highlighted and labeled in compliance with an anatomical ontology we generated for the atlas, providing researchers and chemical risk managers with an unprecedented tool to navigate the microanatomy of *D. magna*. This atlas has the potential to support both tissue-specific and whole-organism phenotyping, informing (eco)toxicology, genetic and phenomic studies.

2. Material and methods

2.1 *Daphnia magna* culturing

A commercial clone of *D. magna* was purchased from Carolina Biological (NC, USA) and raised in "Aachener Daphnien-Medium" or ADaM at room temperature ($20^{\circ}\text{C} \pm 1^{\circ}\text{C}$) under a 16-hours light/8-hours (hrs) dark photoperiod. *D. magna* cultures were fed three times weekly with 3.0×10^7 cells/ml of green microalgae (*Raphidocelis subcapitata*) and once a week with 0.1 mg/mL of dissolved bakers' yeast. The animal density was maintained at about 20 neonates, 10 juveniles and 6 to 8 reproducing adults per liter to prevent overcrowding. Under these conditions, animals reached maturity at 6 to 8 days post-birth and reproduced parthenogenetically every 3 days after sexual maturation with an average of 15 neonates per brood from the second brood onwards. Production of males was induced by overcrowding (>10 reproducing adults per liter) and shorter photoperiod (8 hrs) (Zhang and Baer, 2000).

2.2 Chemical Exposure

In order to have pronounced abnormal tissue phenotypes as a proof-of-concept to demonstrate the utilization of this atlas, we used a wide range of Lethal Concentration (LC) 50 values documented in the literature for acetaminophen (de Oliveira et al., 2016; Du et al., 2016). Reproducing female *D. magna* (approximately 10 days old and carrying the second parthenogenetic brood 2 hrs post-ovulation) were exposed to 5 concentrations of acetaminophen (5, 15, 25, 35, 50 ug/mL). Gravid *D. magna* were used for this exposure to evaluate the toxic effects of acetaminophen on both adults and developing embryos. The exposures lasted for 72 hrs and were conducted with two adult females in 200 ml medium. The medium was replenished, and the animals fed daily. After 72 hrs exposure, each surviving animal was prepared for histological observations as described in the following.

2.3 Histological Processing

2.3.1 Fixation and decalcification

Exposed and control clones of *D. magna* were fixed with 20X Bouin's solution (Newcomer Supply, WI) and incubated for 48 hrs at room temperature (about 21°C) on a low-speed orbital shaker (Corning LSE) set to 55 revolutions per minute (RPM). The fixation is done to preserve tissues from decay due to autolysis or putrefaction. After the fixation step, samples were washed twice with 1X phosphate-buffered saline (PBS) for 10 min. This washing step was followed by decalcification in 20X sample volume of pre-chilled 6% formic acid (Sigma-Aldrich, MO) for 24 hrs on the orbital shaker set to 55 RPM. Samples were then rinsed in 70% ethanol for one minute and immersed in fresh 70% ethanol for 30 min before agarose embedding. We tested fixation using 4% Paraformaldehyde (PFA) in 0.1M phosphate buffer (pH 7.4) (Bioenno LifeSciences, CA) and 10% Buffered Formalin Phosphate (NBF; Fisher Scientific, ON) with different fixation times and temperatures (See Table S1, Supplementary Material). The *D. magna* samples fixed using PFA and NBF (n=23) showed "ballooning", a severe fixation artifact causing the carapace to 'puff-up' (See Figure S1, Supplementary Material). It was concluded after comparison of histological sections generated using these fixatives that Bouin's solution is the best fixative for *D. magna* and was used to fix all samples used in this atlas (See Figure S2, Supplementary Material).

2.3.2 Agarose embedding

Visualization of histological sections in each of the three standard anatomical planes (coronal, sagittal, and transverse) is critical for understanding organismal anatomy. Therefore, the ability to generate consistent sections in each of these planes is essential. Agarose embedding using a mold or an array facilitates consistent positioning and orientation of millimeter-size samples for sectioning (Copper et al., 2018; Santana et al., 2023). A mold was designed (See Figure S3, Supplementary Material) and 3D-printed for casting an agarose block with triangular wells that could hold up to 18 adult *D. magna* for concurrent tissue processing and sectioning (See Figure S3, Supplementary Material). To create an agarose block, laboratory labeling tape (VWR) was wrapped tightly around the mold. Then, 2.5 mL of 1 % agarose (Sigma-Aldrich, MO) at 55 °C was pipetted onto the mold and allowed to solidify at room temperature. The agarose block was removed gently from the mold. Each fixed *D. magna* sample was pipetted with a small volume of ethanol and transferred into the well of the agarose block using a single-use plastic transfer pipette. Samples designated for the sagittal plane sectioning were laid on their sides with a swimming antenna in the wells and all rostra facing the same direction (see Figure S4 for *Daphnia* anatomy and File S1 for *Daphnia* anatomy glossary, Supplementary Material). Samples designated for coronal and transverse orientation were laid on their back in the wells. Once all samples were positioned in individual wells, excess ethanol was carefully dried off using lint-free Kimwipes without touching the samples. Each sample was first topped off with one drop of molten 1 % agarose (about 50 °C) without disturbing the sample, followed by a thin layer of 1% agarose to completely cover the sample. After the agarose layer solidified (about 5 min at room temperature), the block was trimmed as needed, placed into a tissue cassette, and stored in 70 % ethanol for tissue processing.

2.3.3 Tissue processing, sectioning, and staining

All samples were dehydrated in graded ethanol and infiltrated with Formula R paraffin (Leica Biosystems #3801450) in RMC Model 1530 automated closed reagent type tissue processor (See Table S2, Supplementary Material). Following this step, they were serially sectioned at 5 μ m on a Leica RM2255 automated rotary microtome. Sections were then stained with Harris' hematoxylin and eosin in an auto-stainer (Sakura Tissue Tek DRS 2000, IMEB, CA) following a protocol adapted from Copper et al. (2018) where the duration of hematoxylin staining was extended from 3 to 7 min to achieve better contrast for *Daphnia* samples (See Table S3, Supplementary Material). Cover glasses No. 1 (Platinum Line) were used for cover-slipping.

2.4 Histology slide digitization

All slides were screened using an Olympus BX41 microscope and 10X and 20X objective lenses. Those selected for the atlas were scanned at 40X using an Aperio AT2 slide scanner (Leica Biosystems, IL) and images were saved in TIFF format. 40X scanning was performed using 20X objective lens (0.075 n.a. Plan Apo) with 2X optical magnification changer, yielding a digital resolution of 0.25-micron per pixel. The images of *D. magna* samples included in the atlas were cropped using Aperio ImageScope (version 12.4.3.5008). Three channels (Red, Green, Blue) of these digital slides were stacked using Fiji (Schindelin et al., 2012) or ImageJ (Schneider et al., 2012). Then, image processing was performed in Adobe Photoshop (version 22.1.1) where images were rotated and set to have the same canvas size; the image background was removed using "Remove Background"; the "Exposure" was adjusted to fall between 0.1 to 0.25 and the same value was used for each set of images; and "Levels" were adjusted using preset "Midtone Darker". Each set of digital slides was then pyramidally tiled for the web-based viewer.

2.5 Digital labeling of anatomical structures

The anatomical ontology, consisting of a list of anatomical terms organized by groups (organ systems) and subgroups (tissues and cell types), was created for the atlas (See Table S4, Supplementary Material). We cross-referenced the extensive work of Fryer (Fryer, 1991) with other published literature (Agar, 1950; Auld et al., 2010; Bednarska, 2006; Benzie, 2005; Binder, 1931; Christensen et al., 2018; Consi et al., 1987; Ebert, 2005; Edwards, 1980; Goldmann et al., 1999; Halcrow, 1976; Hiruta and Tochinnai, 2014; Kikuchi, 1983; Kress et al., 2016; McCool et al., 2011; Metschnikoff, 1884; Quaglia et al., 1976; Rossi, 1980; Schultz and Kennedy, 1976; Smirnov, 2013; Stein et al., 1966; Steinsland, 1982; Weiss et al., 2012; Wuerz et al., 2017; Zaffagnini and Zeni, 1987, 1986; Zeni and Franchini, 1990) and decided on the commonly used *Daphnia* anatomical terms. Annotation and labels for each anatomical structure presented on the atlas were created using Adobe Illustrator (version 25.1). One image at a time, each anatomical structure was annotated by outlining the structure using the “Curvature” and assigned a color corresponding with the anatomical ontology. Annotation and labels of each structure were saved under “Layers”. After completion of the labeling of all anatomical structures on a given image, the annotations were exported in single scalable vector graphic (SVG) file format to be used as input for the web-based viewer.

2.6 Building the web-based digital slide visualization platform

To improve accessibility and usability, we developed an open-access, web-based digital slide viewing platform based on the open-access project OpenSeadragon (<https://openseadragon.github.io/>). This interface removes the need to download full-

resolution images. The viewer combines SVG files and digital scans into a seamless experience to provide user-friendly access to high-resolution data. The atlas' code was written in client-side JavaScript, HTML, and CSS. Pyramidally tiled images were parsed and visualized with OpenSeadragon. When the user opens an image, the viewer opens the corresponding SVG file containing all the anatomical labels and their corresponding annotations. The viewer parses all labels from the SVG file, plots the corresponding regions, and updates the ontology to note which regions are available on a particular image.

3. Results and discussion

3.1 *Daphnia* Histology Reference Atlas (DaHRA) presenting *D. magna* microanatomy

3.1.1 Interactive viewer

We developed the web-based atlas, DaHRA (<http://daphnia.io/anatomy/>), to be a user-friendly interface to access a collection of digitized histological sections of wildtype female and male *D. magna* in each of three standard anatomical planes (Figure 1A). DaHRA's interface allows users to visualize digital scans of whole-organism sections up to 40X objective magnification (0.25-micron per pixel resolution), providing sufficient resolution to recognize virtually all cell types with broader organismal context. Compared to histology atlases of other model organisms (for example, zebrafish (Copper et al., 2018) and mouse embryos (Armit et al., 2017)), DaHRA offers interactive visualization of normal microanatomy using overlaid vectorized demarcation of anatomical structures whose labels comply with an anatomical ontology created for this atlas. The anatomical ontology consisting anatomical structures can be found on the left side of the

viewer, with the anatomical terms arranged alphabetically within groups (Figure 1B). Annotations of the anatomical structures are presented as color overlays and indicated by check marks next to the anatomical terms. Unchecking the box hides the color overlays. Anatomical terms with underlined labels indicate nested substructures (for example, “microvilli” under “epithelial cell”, both under “midgut”). Hovering over an anatomical term in the ontology dynamically highlights the corresponding structure or structure groups in the viewer, temporarily hiding other checked structures.

In order to make the atlas as a central resource for *Daphnia* community, we also created “Reference” tab which lists published literature related to *Daphnia*’s specific organs or cell-types that were used to annotated this atlas, and “Resources” tab which contains a file annotating *Daphnia* gross normal anatomy, *Daphnia*-specific anatomical definitions, anatomy ontology curated for the atlas, and histology protocols optimized for *Daphnia* samples, including a casting mold stereolithography file . To facilitate collaboration, updates, and validation, a “Feedback” tab is provided for users to leave comments and suggestions. A video demonstrating the features of the atlas is also available on the atlas histology landing page.

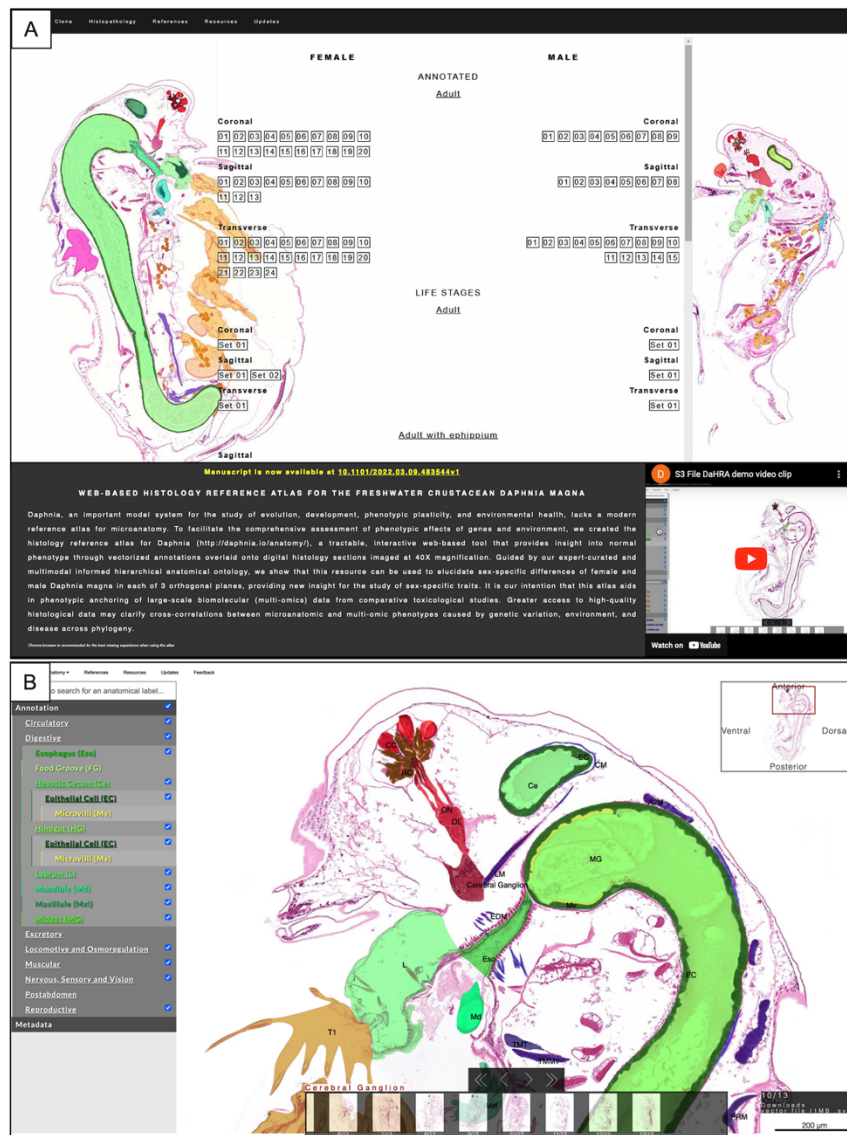


Figure 1. Interactive web-based viewer for DaHRA. (A) The landing page hosts annotated images of female and male commercial clone of *D. magna* with an instruction video describing the features of the atlas. Unannotated images of embryos at different stages, juvenile, and adults are also categorized under “Life stages”. (B) Interactive viewer displaying the expandable list of anatomical structures on the left; the checked boxes indicate the structures labeled in the image. The anatomical terms on the image are shown as acronyms to minimize obscuring of structures; hovering the mouse cursor over an acronym or its corresponding region will show the full term. Unchecking a box will hide the color overlay and annotation corresponding to the box.

3.1.2 *D. magna* male and female microanatomy

DaHRA presents the first microanatomical representation of female and male *D. magna* from the same clone. A hundred samples were sectioned for protocol optimization, and one sample per orientation of each sex was selected and annotated for the atlas. Scanned images of 10 *D. magna* are presented in the histology atlas. Organs and cell types included in the anatomical ontology are briefly described here with representative images from the three anatomical planes of the female (Figures 2A-C) and male *D. magna* (Figures 2D-F). The terminology used for the DaHRA anatomical ontology (a list of terms organized by groups and subgroups) was cross-referenced with published literature (Agar, 1950; Auld et al., 2010; Bednarska, 2006; Benzie, 2005; Binder, 1931; Christensen et al., 2018; Consi et al., 1987; Ebert, 2005; Edwards, 1980; Fryer, 1991; Goldmann et al., 1999; Halcrow, 1976; Hiruta and Tochinai, 2014; Kikuchi, 1983; Kress et al., 2016; McCooles et al., 2011; Metschnikoff, 1884; Quaglia et al., 1976; Rossi, 1980; Schultz and Kennedy, 1976; Smirnov, 2013; Stein et al., 1966; Steinsland, 1982; Weiss et al., 2012; Wuerz et al., 2017; Zaffagnini and Zeni, 1987, 1986; Zeni and Franchini, 1990) for uniformity. We identified 50 anatomical structures and categorized them into 8 groups (circulatory, digestive, excretory, locomotive and respiration, muscular, nervous, sensory and vision, postabdomen, and reproductive), and can be expanded if/ when more structures are identified.

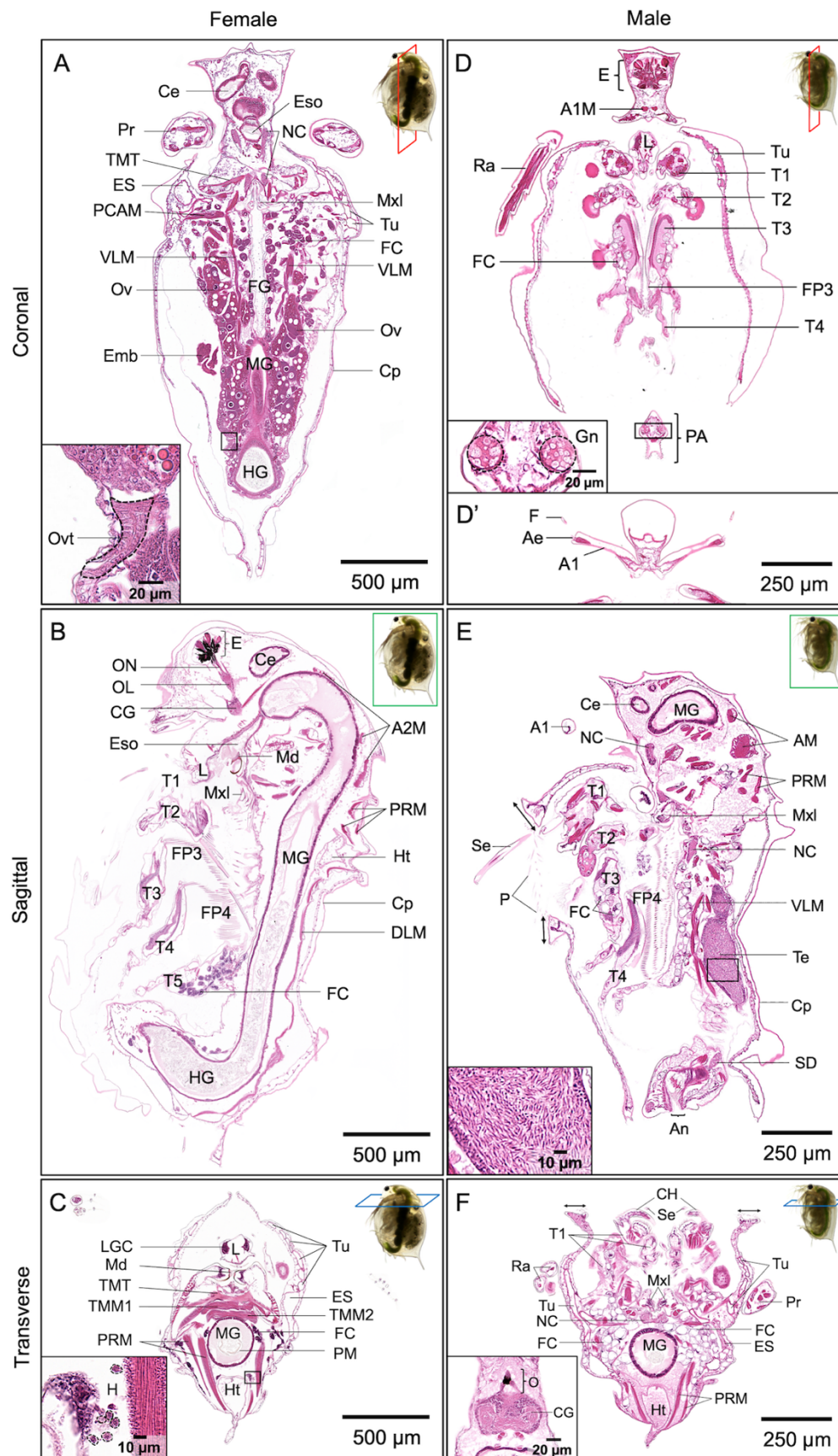


Figure 2. Representative microanatomical structures of female (A-C) and male (D-F') *D.*

***magna* in three orthogonal planes.** The coronal plane (panels A and D) displays most of the structures in pairs. Inset of panel A shows the oviduct (Ovt; dotted circle) and inset of panel D shows gonopores (Gn) in the male. The histology section for Panel D' is slightly ventral to that of panel D and displays prominent and elongated antennules (A1) with flagella (F) at the tips. The mid-sagittal plane of the female (panel B) includes connections between the compound eye (E) and the optic lobe (OL) and cerebral ganglia (CG) by optic nerves (ON). The labrum (L), maxillules (Mxl), and mandibles (Md) are anterior to the esophagus (Eso) that opens into the midgut (MG) and is followed by the hindgut (HG). This section also cuts through the five thoracic limbs (T1-5) and filter plates (FP3, FP4). The sagittal plane of the male (panel E) shows the elongated seta (Se) on the first thoracic limb, pubescence (P) at the wider ventral opening of the carapace, thickening of carapace at the ventral opening (arrows), one of the testes (Te), and a small portion of sperm duct (SD). Inset of panel E showing the spermatozoa in the testis. The transverse plane of the female (panel C) shows the asymmetrical paired mandibles (Md) with the transverse mandibular tendons (TMT), transverse mandibular muscles (TMM1), transverse muscles of mandibles (TMM2), and the posterior rotator muscles of mandibles (PRM). Inset of panel C displays several hemocytes (H) outlined by dotted circles. The transverse plane of male (panel F) displays the paired copulatory hooks (CH) on the first thoracic limbs (T1) and the thickening of the carapace (arrows) at the ventral opening. This also shows the abundance of fat cells (FC) which are quite different from those in the female. The histology section for Panel F', is slightly anterior to that of panel F and shows the pigmented ocellus (O) is connected to the cerebral ganglia (CG). A1M, antennule muscle; A2M, antennal muscle; Ae, aethetasc; An, anus; Ce, hepatic cecum; Cp, carapace; DLM, dorsal longitudinal muscle; ES, end sac of the maxillary

gland; FG, food groove; Ht, heart; LGC, labral gland cell; NC, nerve chord; PA, postabdomen; PCAM, posterior carapace adductor muscle; PM, peritrophic membrane; Ra, ramus of swimming antenna; Tu, tubule of the maxillary gland; VLM, ventral longitudinal muscle. Atlas links to each panel can be found in File S2, Supplementary Material.

We first summarize the sexually dimorphic traits of *D. magna* and follow with brief descriptions of the normal anatomy and microanatomy. Adult males have a smaller body size and much longer antennules than females, that bear a single long flagellum on the tip (Benzie, 2005)(Figure 2D'). Male antennules contain muscle tissue (Figure 2D) that is absent in females. The first thoracic limbs of the males are equipped with elongated setae (Figure 2E) and chitinized copulatory hooks (Figure 2F) that are used for clasping females during copulation. The male postabdomen has gonopores (Figure 2D inset) that are involved in transferring mature spermatozoa from the testes to the female in the region of the oviduct during copulation. Besides having a wider frontal opening, pubescence and thickened angular margins are also observed at the ventral margin of the carapace in males (indicated by arrows in Figures 2E and 2F). Fat cells in males are observed to be different from those of females. Male fat cells contain much larger lipid droplets, reduced and less granular cytoplasm, and smaller nucleoli than female fat cells (Figure 3).

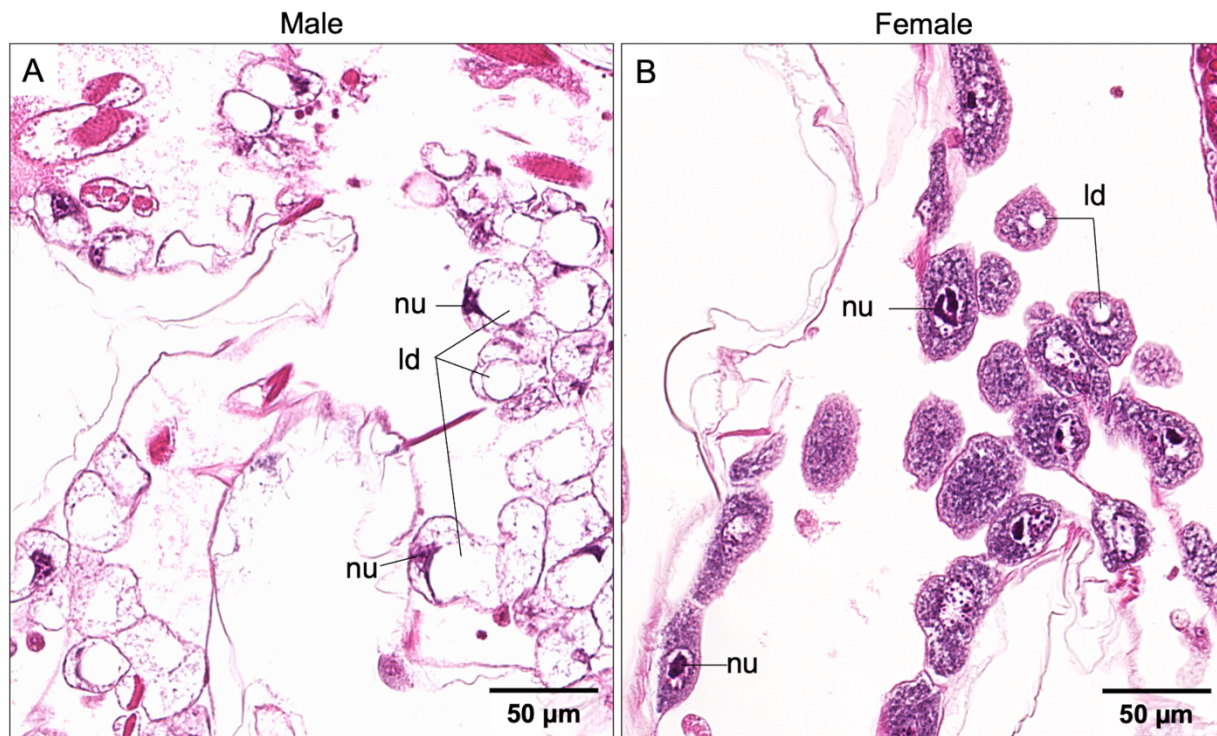


Figure 3. Comparison of male and female fat cells. (A) Fat cells in males contain larger lipid droplets (ld), reduced and less granular cytoplasm with smaller nucleoli (nu) situated at the cell periphery compared with those of females. (B) Fat cells in females have more granular cytoplasm with smaller lipid droplets (ld) and bigger nucleoli (nu) that often appeared subdivided.

Circulatory system. *Daphnia* have an open circulatory system and a myogenic heart (Stein et al., 1966; Steinsland, 1982). Since *Daphnia* are semi-transparent the beating of their hearts is easily visualized in live animals. Hemolymph (blood-like fluid) containing hemocytes (Auld et al., 2010; Metschnikoff, 1884) (Figure 2C inset) is pumped through the body cavity. In line with the literature, we also observe that the *Daphnia* heart has a pair of ostia anterior to the brood chamber, between the midgut and the dorsum (Figures 2B, C, and F). Hemoglobin is synthesized in fat cells and epithelial cells on epipodites of the thoracic limbs (Goldmann et al., 1999).

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327 *Digestive system.* *Daphnia* are filter feeders. Food particles are filtered through filter plates
328 consisting of setae on thoracic limbs 3 and 4, passed through maxillules and mandibles into the
329 esophagus, which is the first part of the digestive system (Figures 2A and B). The digestive
330 system also consists of paired hepatic ceca, midgut, and hindgut (Figures 2A-C, 2E-F) that are
331 lined with epithelial cells and microvilli, with the columnar epithelial cells in the midgut, and the
332 cuboidal cells in hepatic ceca and hindgut (Quaglia et al., 1976; Schultz and Kennedy, 1976).
333 The labrum houses labral glands that have been suggested to be involved in food ingestion and
334 endocrine function (Zaffagnini and Zeni, 1987; Zeni and Franchini, 1990) (Figures 2B-D).

335

336 *Excretory system.* The maxillary gland, also known as the shell gland, is the organ of excretion,
337 housed between the inner and outer walls of the carapace (Smirnov, 2013). It consists of an end
338 sac, a series of tubules, and an opening in the anterior brood chamber (Figures 2A, C, D, and F).

339

340 *Locomotive and osmoregulation system.* The swimming antennae are *Daphnia's* primary organ
341 of locomotion (Fryer, 1991). Each of the paired swimming antennae has a protopodite, two rami
342 bearing setae (Agar, 1950) (Figures 2C, D, and F), and antennal muscles. *Daphnia* have five
343 thoracic limbs (Benzie, 2005) (Figures 2B, D, and E) internal to the carapace. Movements of
344 thoracic limbs produce a constant current that brings food particles into the digestive tract and
345 facilitates osmotic regulation mediated by the epipodite on each thoracic limb (Kikuchi, 1983).
346 First thoracic limbs in males are different from those of female *Daphnia*; only the male has
347 chitinized copulatory hooks (Figure 2F) and longer setae (Figure 2E).

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Muscular system. The muscular system occupies a significant portion of the body (Binder, 1931; Fryer, 1991). The largest muscles are the ventral and dorsal longitudinal muscles that extend along the gut, three paired antennal muscles, transverse mandibular muscles, transverse muscles of mandibles, posterior rotator of the mandibles, carapace adductor muscles, followed by groups of muscles that allow the motion of thoracic limbs and postabdomen (Figure 2). Other small muscles include those around the compound eye, labrum, and esophagus (Consi et al., 1987). All muscles are striated and surrounded by sarcoplasm, that contains many nuclei and is mostly vacuolated. Sarcoplasm is particularly abundant and more vacuolated in the antennal muscles. Male antennules also have internal muscle fibers (Figure 2D) that appear to be absent in females.

Nervous, sensory, and vision systems. *Daphnia* have a pigmented compound eye consisting of 22 ommatidia (Figure 2B) and a light-sensing, pigmented nauplius eye or ocellus with four lens-like bodies (Weiss et al., 2012) (Figure 2F'). Each ommatidium contains eight reticular cells sending a parallel bundle of axons, collectively as the optic nerve into the optic lobe, which is then connected to the cerebral ganglia (Figure 2B). From the cerebral ganglia, two chains of nerve cords run along the thorax, underneath the gut, and to other anatomical structures (Kress et al., 2016; McCoole et al., 2011) (Figures 2A, E, and F). Both sexes have a pair of antennules bearing a group of 9 olfactory setae or aesthetascs (Klann and Stollewerk, 2017; Rieder, 1987) but the male antennules are more prominent and elongated, uniquely fitted with a flagellum at each tip (Figure 2D').

Reproductive system. The ovaries in females are paired tubular structures ending in oviducts (Figure 2A). *Daphnia* are generally cyclical parthenogens, which means that sexual (meiotic)

and clonal (ameiotic) reproduction alternate (Decaestecker et al., 2009). Under favorable environmental conditions, females produce parthenogenetic eggs that are genetically identical to themselves. During clonal reproduction, oogenesis occurs in clusters of four oocytes each. Only one definitive oocyte in each set accumulates yolk granules and lipid droplets during maturation while the others become nurse cells (Rossi, 1980) (Figure 4A). After maturation, parthenogenetic eggs are released into the brood chamber through the oviducts. Fully developed, free-swimming juveniles are extruded after 3 to 4 days. Sexual reproduction is cued by environmental changes such as shorter light photoperiod, lower temperature, and over-crowding, which triggers the parthenogenetic production of genetically identical males for mating with receptive females, the endpoint being two embryos that enter a state of diapause. Unlike parthenogenetic embryos, the development of these resting embryos arrests at the 3000-cell count and enters dormancy (Chen et al., 2018). The resting embryos are encased in a chitinous shell called an ephippium that protects them from harsh environmental conditions (Figure 4C), including freezing and desiccation. Dormancy in *Daphnia* can be exceptionally long, lasting decades and even centuries (Cáceres, 1998; Mergeay et al., 2004). The resting embryos hatch when cued by favorable environmental conditions.

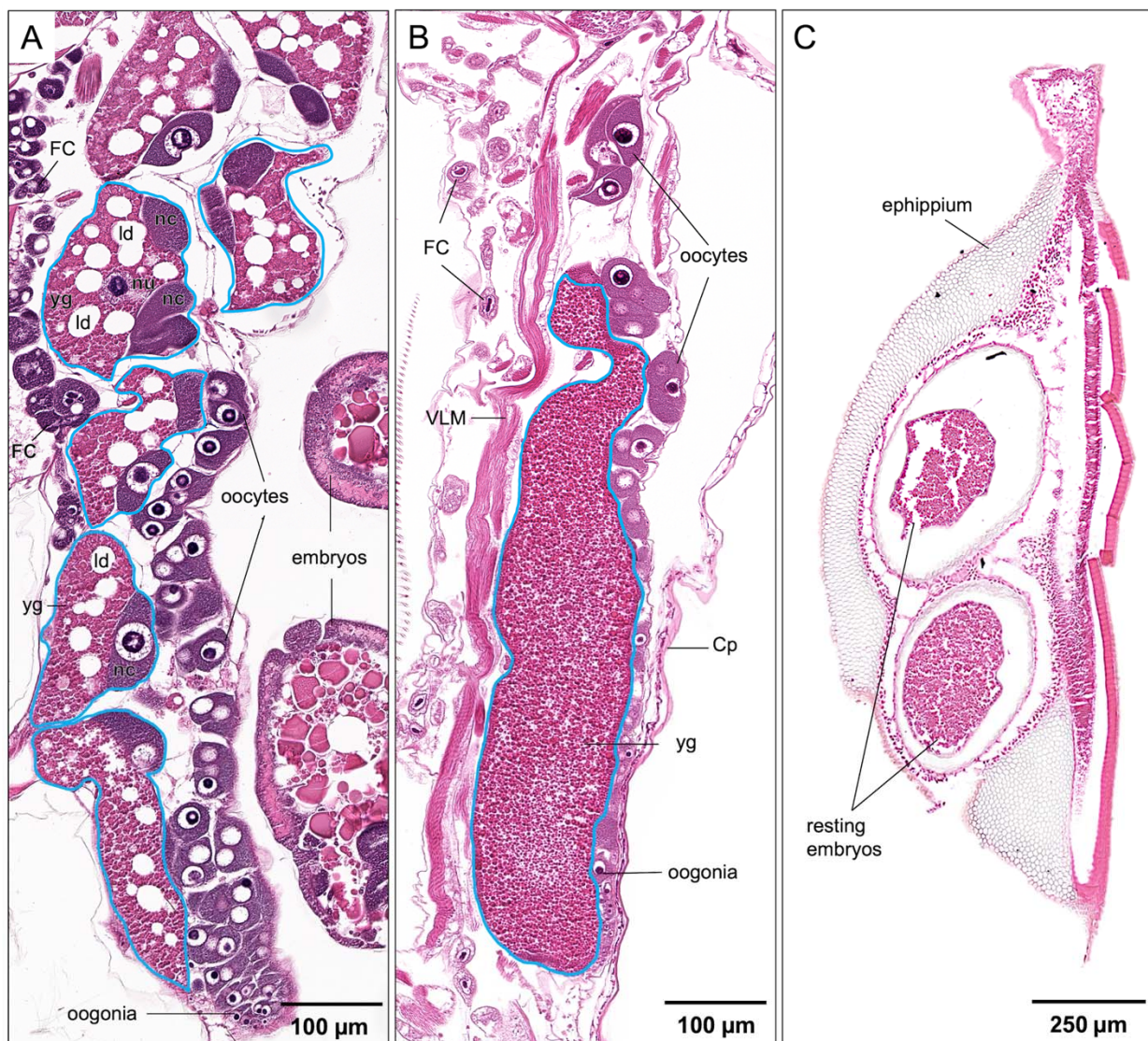


Figure 4. Comparison of parthenogenetic and sexual eggs. (A) The parthenogenetic eggs contain a large amount of lipid droplets (ld) and yolk granules (yg). (B) The sexual eggs contain a large proportion of fine yolk granules without lipid droplets. (C) Resting embryos encased in the ephippium. The top embryo shows artifact. Cp, carapace; FC, fat cell; nc, nurse cell; nu, the nucleus of oocyte; VLM, ventral longitudinal muscle. A solid blue circle indicates an individual egg.

Daphnia testes consist of two tubular structures connected by sperm ducts to gonopores or ejaculatory openings (Figures 2D and E). Spermatogenesis begins at the testes' walls, and mature spermatozoa are displaced inward toward the central region of the testes (Wuerz et al., 2017).

Fat cells are polyploid (Beaton and Hebert, 1989) and consist of a massive portion of lipid and glycogen (Zaffagnini and Zeni, 1986). They are typically found along the trunk, around ovaries or testes, and on the epipodites of the thoracic limbs (Figure 2). In females, these cells have a cytoplasm rich in RNA, one or several lipid droplets of various size, and one large nucleus within which a nucleolus of irregular shape resides (Zaffagnini and Zeni, 1986). The nucleolus often appears subdivided into two or more parts. They are most likely sites of vitellogenin synthesis (Zaffagnini and Zeni, 1986). Compared to female fat cells, male fat cells contain a much larger lipid droplet, reduced and less granular cytoplasm, and a smaller nucleus that is usually situated at the cell periphery (Figure 3).

3.1.3 *D. magna* developmental stages

Daphnia embryos develop in the brood chamber before being extruded. Embryos can also develop outside of the brood chamber, in the culture medium or distilled water. *Daphnia* embryogenesis is usually staged based on the time of development after oviposition (Green, 1956; Gulbrandsen and Johnsen, 1990; Threlkeld, 1979; Toyota et al., 2016), and morphological landmarks (Mittmann et al., 2014). Due to the difficulties in orienting and sectioning the minute individual *Daphnia* embryos, DaHRA presents a selection of images of embryos found in gravid adults. The stages of the embryos were determined under the dissecting microscope according to the developmental events used by Toyota et al (2016) before

the adult female were processed for histology sectioning. Developmental events corresponding with developmental stages are Stage 1: egg membrane is intact; Stage 2: egg chorion breaks down and eye spots are not yet visible; Stage 3: embryo bears two small pink or red eyes; Stage 4: embryo bears two brown or black eyes, and Stage 5: embryo bears a single median black eye. To-date, DaHRA showcases histology images of developmental stages 1, 3, 4, and 5.

At around 20°C, embryos develop within 3 days in the brood chamber after the oviposition. In our histology images, Stage 1 embryo contains yolk granules, lipid droplets, and peripheral cytoplasm (Figure 5A). During Stage 3, eye spots start to develop, and embryo will bear 2 pink eyes towards the end of Stage 3. Labrum, gut, thoracic limbs, and swimming antennae becoming distinguishable at this stage (Figure 5B). Eye pigment increases at Stage 4, and the embryo will bear two brown or black pigment cells (Figure 5C). Ocellus, cerebral ganglia and optic lobe are now distinguishable. Structures of the digestive system, such as mandibles, hepatic ceca, esophagus, midgut and hindgut can also be identified. Segments of thoracic limbs such as epipodites and exopodites are distinctly visible. Stage 5 embryo bears a single median eye, and all major anatomical structures are distinguishable as the body continues to elongate (Figure 5D). Currently, we have not been able to provide images for developmental Stage 2 which is a very short stage (about 4 hours).

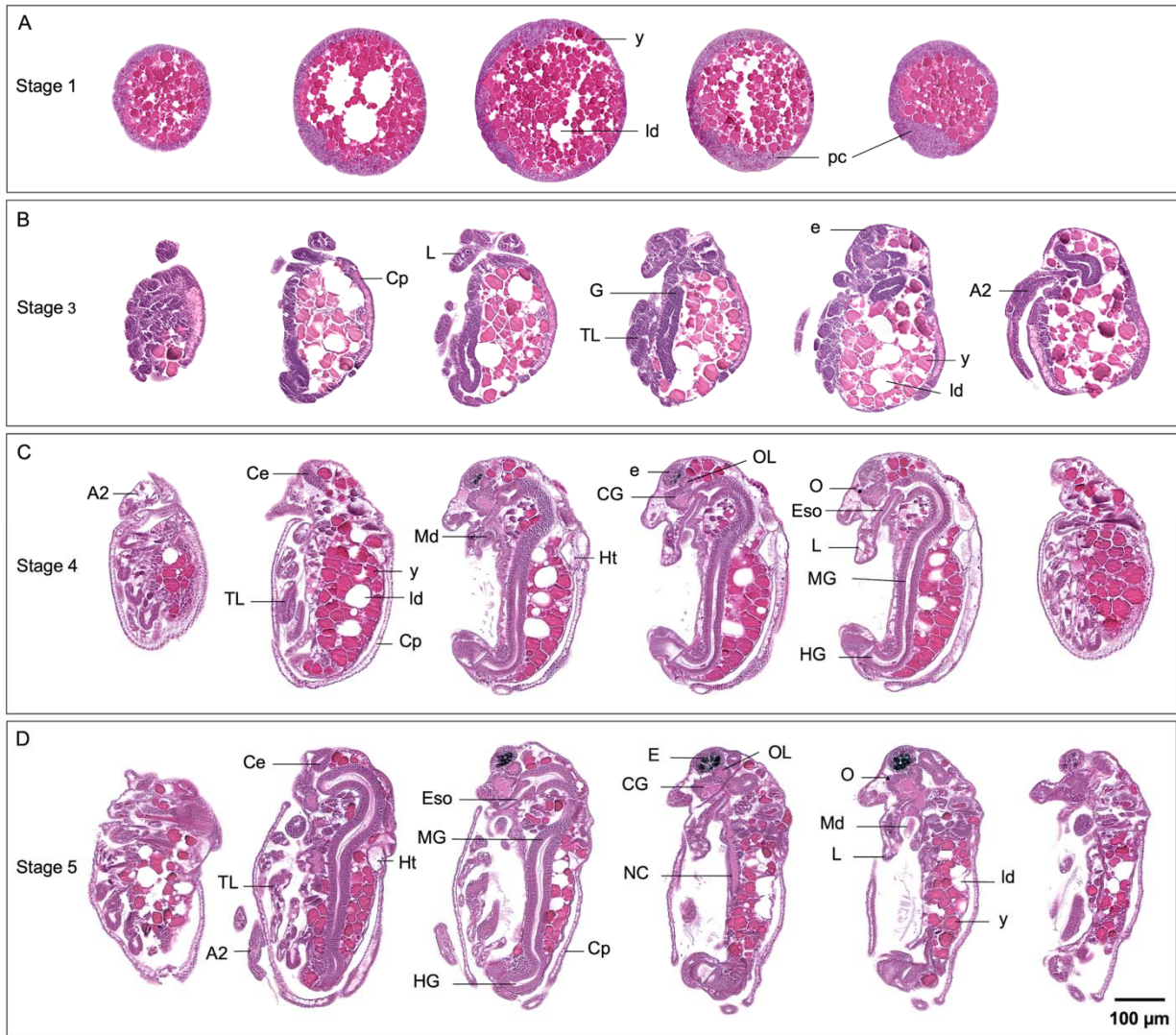


Figure 5. Representative developmental stages of parthenogenetic female *D. magna* in sagittal plane. (A) Stage 1 embryo in spherical form containing yolks (y), lipid droplets (ld), and peripheral cytoplasm (pc). Intact chorion or egg membrane is not visible here due to sectioning artifact. (B) Stage 3 embryo bears two pink eyes (e) with differentiation of thoracic limbs (TL), labrum (L), gut (G) and swimming antennae (A2). (C) Stage 4 embryo bears two brown or black eyes (e) with ocellus (O), optic lobe (OL) and cerebral ganglia (CG) being distinguishable. The labrum (L) is elongated, and segments of digestive tract are distinct. (D) Stage 5 embryo bears a single black median eye (E), and the body continues to elongate. A2,

swimming antennae; Ce, hepatic cecum; Cp, carapace; Eso, esophagus; HG, hindgut; Md, mandible, MG, midgut; NC, nerve chord.

3.1.4 *D. magna* clones

According to the diversity panel curated by Ebert research group, there are more than 200 clones of *D. magna* being cultured in the laboratories around the world (<https://evolution.unibas.ch/ebert/research/referencepanel/>). Clonal variation refers to the different genotypes of the same species, and they may differ strongly in their toxicological responses (Barata et al., 2002; Barber et al., 1990; Kim et al., 2023). While DaHRA showcases annotated histology from a commercial clone, we have also included histology images from a clone provided by the University of Birmingham, UK (UOB_LRV0_1) to demonstrate how this atlas can be expanded for the addition of new image datasets (<http://daphnia.io/anatomy/clone/>). This clone was revived from a biological archive of Lake Ring, a shallow lake in Denmark (Cuenca Cambronero et al., 2018). The clone had been sequenced to generate the first chromosomal-level genome assembly which 33,950 genes and 31,336 proteins had been annotated (Chaturvedi et al., 2023).

3.1.5 An example of *D. magna* histopathology

To demonstrate how our atlas facilitates visualization of abnormal tissue architecture in the context of the whole organism, we included acetaminophen-exposed *D. magna* that show strong histopathological tissue phenotypes. “Comparison” tool shows web-based comparisons at any magnification between the exposed and to the control animals in an experiment (See Figure S5, Supplementary Material; <http://daphnia.io/anatomy/treatments/>).

469
470 Besides its therapeutic effects, acetaminophen is known to induce toxicological outcomes if
471 overdose is taken over a period of time (Peranathan et al., 2024; Yoon et al., 2016). It is also a
472 contaminant found in surface waters and wastewater throughout the world (Phong Vo et al.,
473 2019). In our study, gravid *D. magna* were exposed to 5, 15, 25, 35, 50 µg/mL of
474 acetaminophen for 72 hrs to allow the evaluation on both adults and developing embryos. *D.*
475 *magna* exposed to 5 µg/mL acetaminophen showed no distinct changes compared to unexposed
476 controls. Exposure to 25 µg/mL and 35 µg/mL acetaminophen caused similar phenotypes, but
477 exposure to 50 µg/mL acetaminophen caused death by the end of the exposure. This exposure is
478 not intended for reporting toxicological effects of acetaminophen; therefore, no replicate and
479 error had been generated. *D. magna* exposed to 15 and 25 µg/mL acetaminophen were used to
480 illustrate histopathological phenotypes in a whole-organism context.

481
482 Histology of the exposed *D. magna* revealed morphological alterations in various organs and
483 tissue types. Excessive vacuolation was observed in the labral glands of *D. magna* exposed to
484 both 15 and 25 µg/mL acetaminophen (Figures 6B and C). Besides the absence of a peritrophic
485 membrane and partially digested food particles, the midgut and hindgut of the exposed *D. magna*
486 showed degeneration in the epithelium lining (Figures 6E and F), with the degree being
487 particularly conspicuous in the 25 µg/mL-exposed *D. magna* (Figure 6F). Extrusion and
488 sloughing of epithelial cells were also observed (Figures 6E and F). Cytoplasmic swelling was
489 noted in the midgut and hindgut epithelial cells and fat cells (Figures 6H and I).

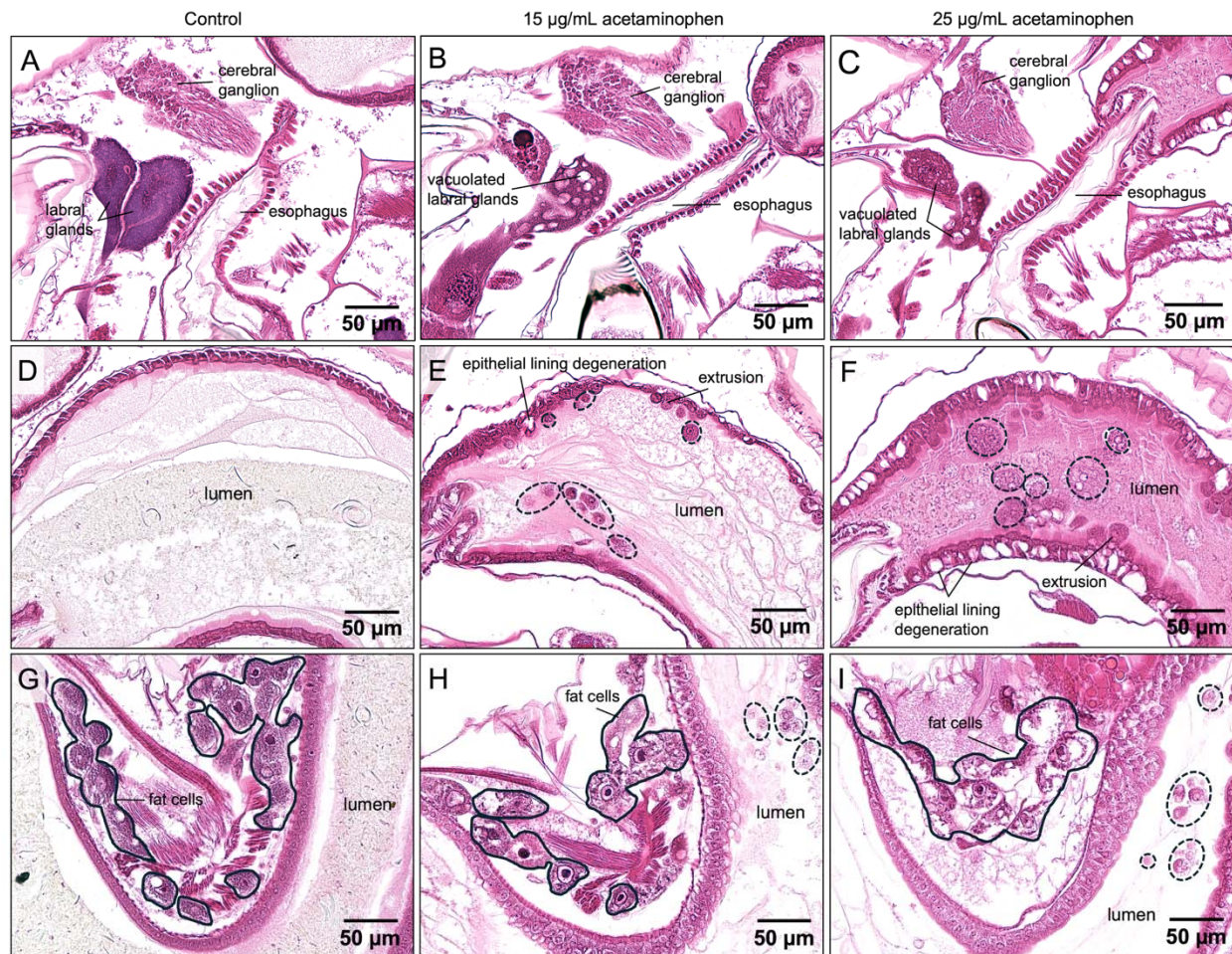


Figure 6. Histopathologic features of the digestive system and fat cells in *D. magna* exposed to 15 and 25 µg/mL of acetaminophen for 72 hrs. Compared to the control (A), vacuolization of labral glands were present in exposed *D. magna* (B, C). Extrusion, and sloughing of degenerated epithelial cells (dotted circles) was observed in exposed animals (E, F), with the degree of degeneration being particularly conspicuous in the epithelium lining of 25 µg/mL-exposed *D. magna* (F). Cytoplasmic swelling was observed in the fat cells and the hindgut epithelial lining of exposed *D. magna* (H, I). Sloughed epithelial cells (dotted circles) were prominent in the hindgut of the exposed *D. magna* (H, I).

A dead, and deformed embryos were found in the brood chamber of 25 µg/mL acetaminophen-exposed *D. magna* after 72 hrs of exposure. The deformed embryos remained in the chorion, showed development of the compound eye and gut precursors, but no visible elongation of body length, or development of the swimming antennae and thoracic limbs were evident after 72 hrs of exposure (Figure 7).

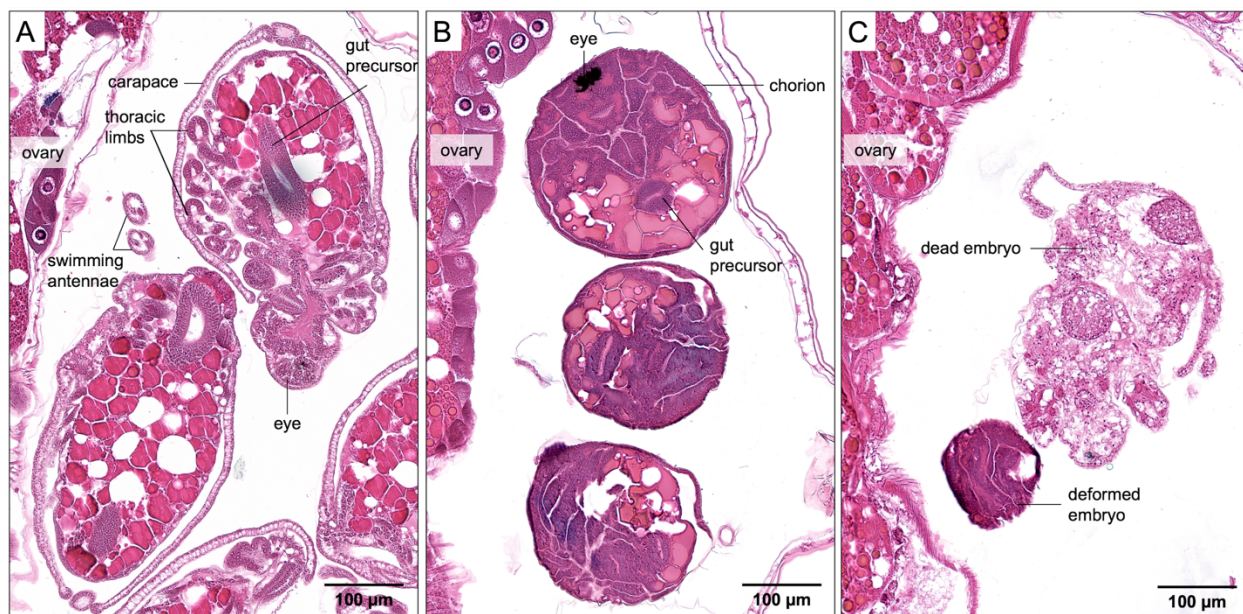


Figure 7. Embryotoxicity in the *D. magna* exposed to 25 µg/mL acetaminophen. (A) Normal embryos in the brood chamber of control compared to (B) deformed embryos remained in the chorion, showing some development of the compound eye and gut precursors but no visible development of the swimming antennae and thoracic limbs. (C) A dead embryo was also observed in the brood chamber of an exposed *D. magna*.

Toxicological effects are often quantified through apical endpoints (e.g. immobilization) in acute exposures (Organisation for Economic Co-operation and Development, 2004; United States Environmental Protection Agency, 2002) and fitness-linked life history traits in chronic

exposures (Organisation for Economic Co-operation and Development, 2018; United States Environmental Protection Agency, 1996). Few studies employ ultrastructural analysis of a target organ, usually the midgut, for the pathological assessment of chemical toxicity (Bacchetta et al., 2017; Bodar et al., 1990; Heinlaan et al., 2011; Yang et al., 2010). Acetaminophen exposure performed for this work was not intended for reporting the toxicological responses. However, the observation of embryotoxicity and histopathological change in multiple tissue types (fat cells and labral glands), including the whole digestive tract (ceca, midgut, and hindgut) in the exposed *D. magna* suggested the important role of non-targeted toxicity assessment for the complete detection of phenotypes across organ systems and during embryonic development. A combination of complete histopathological phenotyping in whole *Daphnia* with toxicological omics data (e.g., transcriptomics and metabolomics) and tissue-specific biomarkers (e.g., single cell and spatial transcriptomics) will enable a comprehensive evaluation of toxicological effects and reveal tissue-specific toxicity (Tian et al., 2024). Establishing these links is essential for biomolecular data to be regulatory relevant because hazards are classified by phenotypic outcomes (behavioral, physiological, and/or histopathological). Such integration will play an important role in discovering and applying adverse outcome pathways for next-generation risk assessment (United States Environmental Protection Agency, 2014) where links between biomarkers of adversity, causative agents, and organ-specific effects remain to be well-established. Using the principle of evolutionary and functional conservation of genes and pathways in organisms across the tree of life, hypotheses on targets of toxicity can be extrapolated across non-target species, including humans (“The Precision Toxicology initiative,” 2023).

3.2 Future directions

We have created an annotated, interactive atlas for web-based access to *D. magna* microanatomy using a single commercially available clone of *D. magna* to present reasonable examples of what “normal” microanatomy looks like. Clonal and intraspecific variation are important components in *Daphnia* biology (Ho et al., 2019; Miyakawa et al., 2015; Tolardo et al., 2016) where differential responses to toxicants and other stressors have been reported for physiological and reproduction parameters (Barata et al., 2002; Barber et al., 1990; Kim et al., 2023), but remain to be explored with regard to histopathological change and tissue-specific phenotypes.

A key aspect of this work is our commitment to open science, enabling broader participation of *Daphnia* community to utilize and further develop this atlas. A clone UOB_LRV0_1 has been included in the current phase of DaHRA, and through collaboration and additional resources, we anticipate this atlas to include additional datasets possibly covering clonal and/or intraspecific genetic variation, examples of how intraspecific variation influences tissue-specific phenotypes, and most importantly, observation of other histopathological change. It is our hope for this atlas to be an informational tool to the *Daphnia* community, but it is not intended and should not be utilized as “control” for chemical-exposure and toxicity-testing experiments.

Web-based atlases have served as a platform for the systematic integration of spatial and molecular data (Asp et al., 2019; Snyder et al., 2019; Thul and Lindskog, 2018; Yao et al., 2023). As integrative tissue-based or spatial atlases for human and mammalian models are constantly

being improved , DaHRA is envisioned as a potential integrative platform for anchoring *Daphnia* multi-omics data with tissue phenotypes. This will increase its potential value as a tool for exploring the spatial and chemical complexities of biological systems.

In summary, our first, open-source visualization platform for *D. magna* microanatomy provides a baseline knowledge of *Daphnia*'s cells, extracellular tissue, and their arrangements in health as a foundation for the detection of aberrant tissue phenotypes. Its user-friendly interface and global web-based access will facilitate broader contributions to (eco)toxicology, biology and beyond.

Supplementary Material (in order of citation)

Table S1. Fixatives and fixation parameters tested for best preservation of whole *D. magna* samples

Figure S1. “Ballooning” artifact observed in *D. magna* sample fixed with PFA and NBF

Figure S2. Comparison of histological sections generated with Bouin's, PFA and NBF

Figure S3. Agarose embedding of *D. magna* samples using agarose block with triangular wells

Figure S4. Anatomy of adult female and male *D. magna*

File S1. *Daphnia* anatomy glossary

Table S2. Tissue processing steps for serial dehydration and infiltration of *D. magna* samples with Formula R paraffin in tissue processor

Table S3. Automated steps for staining *D. magna* 5- μ m sections with Harris' hematoxylin and eosin in an auto-stainer

Table S4. Anatomical ontology for DaHRA

Figure S5. Overview of DaHRA displaying annotated histopathological data

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Competing Interests

The authors declare no competing interest.

Author Contributions

Mee Ngu: Data curation, Investigation, Methodology, Project administration, Visualization, Writing-original draft; Daniel Vanselow: Data curation, Visualization, Software; Carolyn Zaino: Data curation, Visualization; Alex Lin: Methodology; Jean Copper: Methodology; Margaret Beaton: Validation; Luisa Orsini: Conceptualization; John Colbourne: Conceptualization; Keith Cheng: Conceptualization, Funding Acquisition, Resources; Khai Ang: Conceptualization, Funding Acquisition, Investigation, Methodology, Resources, Supervision. All authors contributed to Writing- Review and Editing.

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