A web-based histology atlas for the freshwater Cladocera species Daphnia magna

Short title: Web-based Daphnia histology atlas

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1 Abstract

2 Daphnia are keystone species of freshwater habitats used as model organisms in ecology and 3 evolution. They are also routinely used as environmental sentinels in regulatory toxicology and 4 are increasingly contributing to new approach methodologies (NAM) for chemical risk assessments Yet, it is challenging to establish causal links between biomolecular (omics) 5 6 responses to chemical exposure and their toxicity phenotypes without a baseline knowledge of 7 tissue- and cell-morphology of healthy individuals. Here, we introduce the Daphnia Histology 8 Reference Atlas (DaHRA, http://daphnia.io/anatomy/), which provides a baseline of wildtype 9 anatomical and microanatomical structures of female and male *Daphnia magna*. This interactive 10 web-based resource features overlaid vectorized demarcation of anatomical structures that 11 compliant with an anatomical ontology created for this atlas. Since sex is environmentally 12 induced in *Daphnia*, DaHRA is a map of sexual dimorphism by phenotypic plasticity. We also 13 benchmark this tool for mechanistic toxicology by exposing *Daphnia* to acetaminophen and use 14 the atlas to document its effects in organs, tissues, and cell-types. DaHRA represents an essential 15 step towards correlating phenotypes with the discovery power of hypothesis-free, molecular 16 backdrop against which pathology can be interpreted, thereby offering a platform to elucidate 17 how genetic variation and external perturbations cascade through multiple biological scales to 18 influence phenotype.

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

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Synopsis: Whole-organism *Daphnia* atlas as foundation for unbiased phenotyping, and its utility
in characterizing sexual dimorphism and effects of chemical toxicity.

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26 **1. Introduction**

27

Despite regulatory restrictions on hazardous chemicals, chemical pollution is the leading cause of premature deaths and morbidities globally (1–3). Adding to this, habitat loss, climate change, and pollution are also leading environmental factors impacting biodiversity, with more than 60% of ecosystems services being reduced or wiped out in the last two decades (4). The introduction

32 of new approach methodologies (NAMs) for assessing chemicals for their toxicity is designed to 33 improve regulatory outcomes by replacing outdated, data-poor methods that rely on observing 34 apical endpoints (such as death or reproductive failure in selected animals) with modern methods 35 at revealing the pathways to toxicity (5) These methods include data-rich techniques such as 36 transcriptomics and metabolomics (omics) that measure changes in the abundance of all gene 37 transcripts and endogenous metabolites upon exposure that are indicative of the chemical modes 38 of action (6–8). Although these data are robust at measuring biomolecular activity linked to a 39 huge array of potential adverse health effects, these data must be anchored to their phenotypic 40 effects for regulatory relevance (9). Yet despite omics producing high content mechanistic 41 information that enables data-driven discoveries, there are few analogs to discover causal links to 42 phenotypes. A primary objective of this paper is to provide a necessary resource for obtaining 43 and interpreting high content phenotypic data for the model species D. magna, which is used 44 globally to set regulatory limits on potentially hazardous chemical substances in the environment 45 and one of five models used uncover the evolutionary origins of toxicity (10).

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47 The water flea *Daphnia* is a keystone branchiopod crustacean (order Cladocera) in freshwater 48 lotic ecosystems worldwide and is an established model in ecology, evolution, and ecotoxicology (11-13). They are responsive to environmental change and they adapt via evolutionary 49 50 mechanisms and plasticity (14–16). This plasticity includes the sex of daphniids, which is 51 determined by environmental conditions. Relevant to ecotoxicity testing is their short generation 52 time that enables the experimental manipulation of large populations; and a parthenogenetic life 53 cycle that allows the rearing of populations of identical clones from single genotypes (17). The 54 latter property has the unique advantage of allowing the concurrent study of molecular and 55 phenotypic responses to multiple environmental insults, including chemical pollutants. Daphnia magna is the species of choice in ecotoxicogenomics (18,19). Recently, its hologenome (20), 56 57 reference genome (21,22) and transcriptome (23-25) have been published, elevating this species 58 to the ranks of other biomedical model species for ecological genomics. Yet, the full potential of this species cannot be entirely realized without the correlation between molecular, and tissue-59 60 and cell- specific phenotypic responses.

62 Histopathology, the histological study of tissue-specific changes, enables the identification of 63 targets of toxicity and diseases, bridging phenotypes and biomolecular perturbations induced by 64 environmental insults (26,27). Histopathology-based toxicological studies in fish (28–30) and bivalves (31,32) have shown to be useful for water quality monitoring and assessment. The 65 66 application of histopathology to millimeter-size sentinel species used in ecotoxicology would enable the analysis of tissue-specific toxicity phenotypes in the whole animal. However, 67 68 identification of affected cell and tissue types requires prior knowledge of normal structure, 69 which is generally only possible through atlases.

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71 Here, we present the first curated web-based histology atlas for both female and male D. magna, 72 further broadening the discovery capacity of this sentinel species. We optimized methods for D. 73 magna histology and created a collection of digitized histological images for adult female and 74 male *D. magna* in three anatomical planes to elucidate sexual dimorphism by environmentally 75 induced phenotypic plasticity. We included the first proof-of-concept application of this atlas by 76 comparing tissue of clonal replicates of the same genotype of D. magna under control conditions 77 (no chemical exposure) with replicates exposed to an over-the-counter painkiller, 78 acetaminophen. This common pharmaceutical substance is found in surface waters and 79 wastewater throughout the world (33). Besides its therapeutic effects, it is known to induce 80 toxicological outcomes if overdose (34,35). This benchmarked resource for mechanistic 81 toxicology using *Daphnia* is made open-access and interactive, allowing smooth magnification 82 with a dynamic scale bar. Anatomical structures are highlighted and labeled corresponding with 83 an anatomical ontology, providing researchers and chemical risk managers with an 84 unprecedented tool to navigate both normal and abnormal microanatomical structures of organs, 85 tissues, and cell-types. This resource has the potential to support tissue-specific and whole-86 organism phenotyping, informing (eco)toxicology, genetics and phenomics studies.

87

88 2. Material and methods

89 2.1 *Daphnia magna* culturing

90 A commercial strain of *D. magna* was purchased from Carolina Biological (NC, USA) and raised

91 in "Aachener Daphnien-Medium" or ADaM at room temperature ($20^{\circ}C \pm 1^{\circ}C$) under a 16-hour

light/8-hour dark photoperiod. D. magna cultures were fed three times weekly with 3.0 x 10^7 92 93 cells/ml of green microalgae (Raphidocelis subcapitata) and once a week with 0.1 mg/mL of 94 dissolved bakers' yeast. The animal density was maintained at about 20 neonates, 10 juveniles 95 and 6 to 8 reproducing adults per liter to prevent overcrowding. Under these conditions, animals 96 reached maturity at 6 to 8 days post-birth and reproduced parthenogenetically every 3 days after 97 sexual maturation with an average of 15 neonates per brood from the second brood onwards. 98 Production of males was induced by overcrowding (>10 reproducing adults per liter) and shorter 99 photoperiod (8 hours) (36).

100

101 2.2 Chemical Exposure

102 Reproducing female *D. magna* (approximately 10 days old and carrying the second 103 parthenogenetic brood 2 hours post-ovulation) were exposed to 5 concentrations of 104 acetaminophen (5, 15, 25, 35, 50 mg/L). Gravid *D. magna* were used for this exposure to 105 evaluate the toxic effects of acetaminophen on both adults and developing embryos. The 106 exposures lasted for 72 h and were conducted with two adult females in 200 ml medium. The 107 medium was replenished, and the animals fed daily. After 72 h exposure, each surviving animal 108 was prepared for histological observations as described in the following.

109

110 2.3 Histological Processing

111 2.3.1 Fixation and decalcification

112 Exposed and control clones of *D. magna* were fixed with 20X Bouin's solution (Newcomer 113 Supply, WI) and incubated for 48 h at room temperature (about 21°C) on a low-speed orbital 114 shaker (Corning LSE) set to 55 revolutions per minute (RPM). The fixation is done to preserve 115 tissues from decay due to autolysis or putrefaction. After the fixation step, samples were washed 116 twice with 1X phosphate-buffered saline (PBS) for 10 min. This washing step was followed by 117 decalcification in 20X sample volume of pre-chilled 6% formic acid (Sigma-Aldrich, MO) for 24 118 h on the orbital shaker set to 55 RPM. Samples were then rinsed in 70% ethanol for one minute 119 and immersed in fresh 70% ethanol for 30 min before agarose embedding. Different fixation 120 methods were compared before opting for the Bouin's solution. We tested fixation using 4% 121 Paraformaldehyde in 0.1M phosphate buffer (pH 7.4) (Bioenno LifeSciences, CA) and 10%

Buffered Formalin Phosphate (Fisher Scientific, ON) with different fixation times andtemperatures (Supplementary Table 1).

124 2.3.2 Agarose embedding

125 Agarose embedding using a mold or an array facilitates consistent positioning and orientation of 126 millimeter-size samples for sectioning (37,38). Adapted from Sabaliauskas et al. (2006)(39), a 127 mold was designed and 3D -printed for casting an agarose block with wells that could hold up to 128 18 adult *D. magna* for concurrent tissue processing and sectioning. To create an agarose block, 129 laboratory labeling tape (VWR) was wrapped tightly around the mold. Then, 2.5 mL of 1 % 130 agarose (Sigma-Aldrich, MO) at 55 °C was pipetted onto the mold and allowed to solidify at 131 room temperature. The agarose block was removed gently from the mold. Each fixed D. magna 132 sample was pipetted with a small volume of ethanol and transferred into the well of the agarose 133 block using a single-use plastic transfer pipette. Samples designated for the sagittal plane 134 sectioning were laid on their sides with a swimming antenna in the wells and all rostra facing the 135 same direction (see Supplementary Figure 1 for Daphnia anatomy and Supplementary Text 1 for 136 Daphnia anatomy glossary). Samples designated for coronal and transverse orientation were laid 137 on their back in the wells. Once all samples were positioned in individual wells, excess ethanol 138 was carefully dried off using lint-free Kimwipes without touching the samples. Each sample was 139 first topped-off with one drop of molten 1 % agarose (about 50 °C) without disturbing the 140 sample, followed by a thin layer of 1% agarose to completely cover the sample. After the agarose 141 layer solidified (about 5 min at room temperature), the block was trimmed as needed, placed into 142 a tissue cassette, and stored in 70 % ethanol for tissue processing.

143

144 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 (Supplementary Table 2). Following this step, they were serially sectioned at 5 µm 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) (40) where the duration of hematoxylin staining was extended from 3

to 7 min to achieve better contrast for samples fixed with Bouin's solution (Supplementary Table3). Cover glasses No. 1 (Platinum Line) were used for cover-slipping.

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154 2.4 Histology slide digitization

155 All slides were screened using an Olympus BX41 microscope and 10X and 20X objective lenses. 156 Those selected for the atlas were scanned at 40X using an Aperio AT2 slide scanner (Leica 157 Biosystems, IL) and images were saved in TIFF format. 40X scanning was performed using 20X 158 objective lens (0.075 n.a. Plan Apo) with 2X optical magnification changer, yielding a digital 159 resolution of 0.25-micron per pixel. The images of *D. magna* samples included in the atlas were 160 cropped using Aperio ImageScope (version 12.4.3.5008). Three channels (Red, Green, Blue) of 161 these digital slides were stacked using Fiji (41) or ImageJ (42). Then, image processing was 162 performed in Adobe Photoshop (version 22.1.1) where images were rotated and set to have the 163 same canvas size; the image background was removed using "Remove Background"; the 164 "Exposure" was adjusted to fall between 0.1 to 0.25 and the same value was used for each set of 165 images; and "Levels" were adjusted using preset "Midtones Darker". Each set of digital slides 166 was then pyramidally tiled for the web-based viewer.

167

168 2.5 Digital labeling of anatomical structures

The anatomical ontology, consisting of a list of anatomical terms organized by groups (organ 169 170 systems) and subgroups (tissues and cell types), was created for the atlas (Supplementary File 1). 171 We cross-referenced the extensive work of Fryer (43) with other published literature (44–70) and 172 decided on the commonly used Daphnia anatomical terms. Annotation and labels for each 173 anatomical structure presented on the atlas were created using Adobe Illustrator (version 25.1). 174 One image at a time, each anatomical structure was annotated by outlining the structure using the 175 "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 176 177 anatomical structures on a given image, the annotation was exported in a single scalable vector 178 graphic (SVG) to be used as input for the web-based viewer.

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- 180

181 2.6 Building the web-based digital slide visualization platform

182 The file size of a set of digitized images is > 4 GB, making access challenging for users with 183 standard computational resources. To improve accessibility and usability we developed an open-184 access, web-based digital slide viewing platform based on the open-access project 185 OpenSeadragon (https://openseadragon.github.io/). This interface removes the need to download 186 full-resolution images. The viewer combined annotations and digital scans into a seamless 187 experience to provide user-friendly access to high-resolution data. The atlas' code was written in 188 client-side JavaScript, HTML, and CSS. Pyramidally tiled images were parsed and visualized 189 with OpenSeadragon. When the user opens an image, the viewer will open the corresponding 190 SVG file containing all the anatomical labels and their corresponding shape vector information. 191 The viewer parses all labels from the SVG file, plotting the corresponding regions, and updates 192 the ontology to note which regions are available to visualize on a particular image. 193

194 **3. Results and discussion**

3.1 *Daphnia magna* histology atlas presenting *Daphnia* microanatomy
and enabling the discovery of aberrant phenotypes

197 3.1.1 Interactive viewer for <u>Daphnia Histology Reference Atlas</u> (DaHRA)

We developed the <u>Daphnia Histology Reference Atlas</u> (DaHRA; http://daphnia.io/anatomy/), a user-friendly interface to access a collection of digitized histological sections, including wildtype female and male *D. magna* in three standard anatomical planes (Figure 1), and *D. magna* exposed to 25 mg/L acetaminophen (Figure 2; http://daphnia.io/anatomy/treatments/).

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The interface allows users to visualize digital scans (0.25-micron per pixel resolution) with and without annotations of anatomical structures. The anatomical ontology (Supplementary File 1) including of all the anatomical structures can be found on the left side of the viewer, with the anatomical terms arranged alphabetically within 8 groups: circulatory, digestive, excretory, locomotive and respiration, muscular, nervous, sensory and vision, post-abdomen, and reproductive (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. A collection of abnormal anatomical structures of *D. magna* exposed to 25 mg/L acetaminophen is shown under "Histopathology" (Figure 2B).

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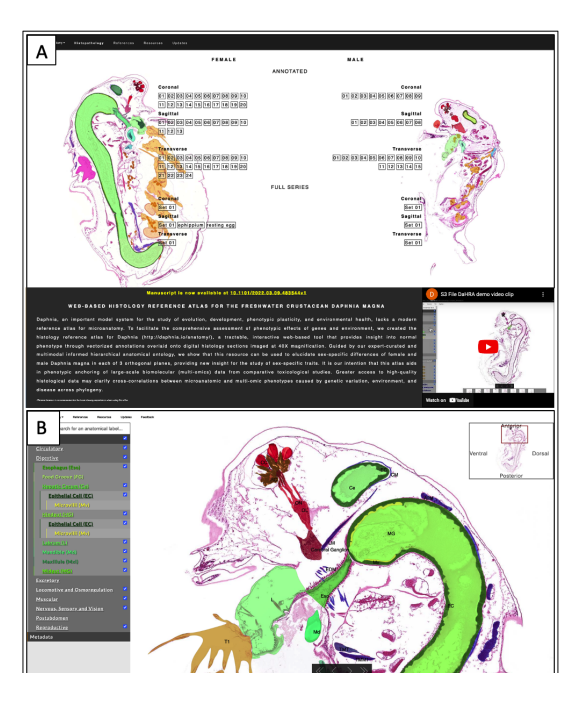


Figure 1. Overview of the interactive web-based viewer for DaHRA. (A) Landing page hosting all annotated and unannotated histology images of the same female and male strain of *D. magna* with an instruction video describes the features of the atlas. (B) Interactive viewer displaying the expandable list 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; hovering the mouse cursor over an acronym or its corresponding region will show the full term.

224 Unchecking a box will hide the color overlay and annotation corresponding to the box.

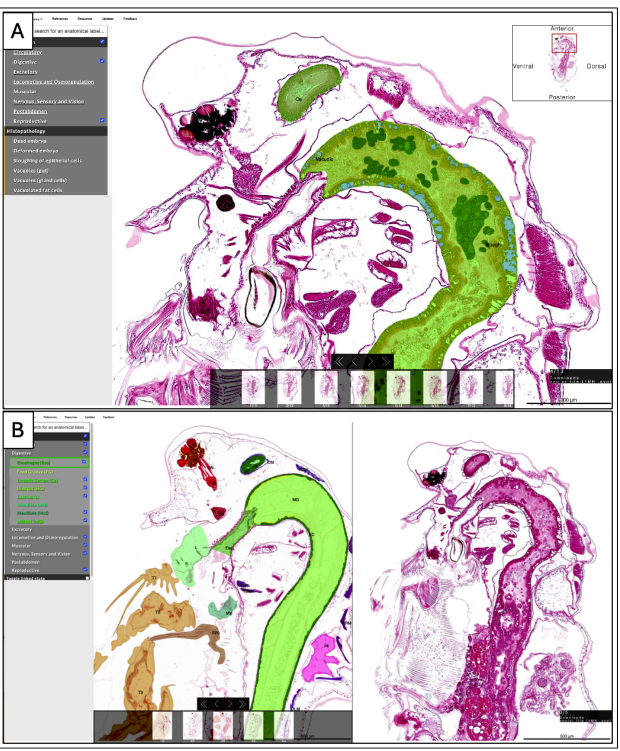


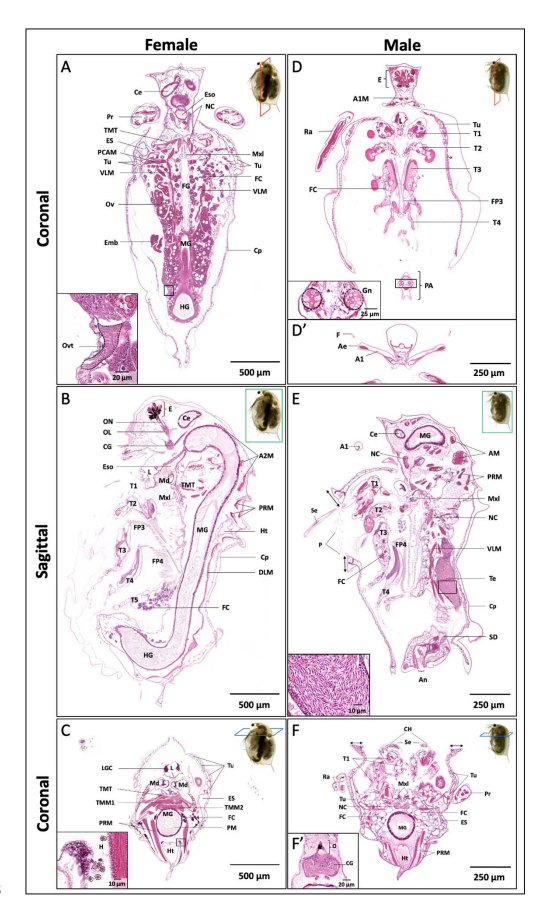
Figure 2. Overview of the atlas displaying annotated histopathological data. (A) The checked boxes in the anatomical ontology indicate the affected structures overlaid with 227 annotation and the observed abnormalities listed under "histopathology". (B) Viewer comparing 228 229 the non-exposed (left) to the acetaminophen-exposed (right) D. magna. 230

232 3.1.2 *Daphnia magna* male and female microanatomy

233 DaHRA presents the first microanatomical representation of female and male D. magna from the 234 same genotype. All organs and cell types included in the anatomical ontology are briefly 235 described here with representative images from the three anatomical planes of the female (Figure 236 3A-C) and male D. magna (Figure 3D-F). The terminology used for the DaHRA anatomical 237 ontology (a list of terms organized by groups and subgroups) was cross-referenced with 238 published literature (43-69) for uniformity. We identified 50 anatomical structures, and 239 categorized them in 8 groups (circulatory, digestive, excretory, locomotive and respiration, 240 muscular, nervous, sensory and vision, postabdomen, and reproductive), and can be expended if/ 241 when more structures are identified.

242

243 Here, we first summarize the sexually dimorphic traits in D. magna and follow with a brief 244 description of the normal anatomy and microanatomy. Apart from the obvious differences in 245 body size and reproduction organs, D. magna presents other sexual dimorphisms that can be 246 clearly visualized and compared on the atlas. Generally, the body size and first antennae or 247 antennules of *D. magna* differ between the sexes. Adult males have a smaller body size but much 248 longer antennules bearing a single long peg or flagellum on the tip (Figure 3D'). Male antennules 249 are also composed of muscle tissue (Figure 3D) that is absent in females. The first thoracic limbs 250 of the males are equipped with elongated setae (Figure 3E) and chitinized copulatory hooks 251 (Figure 3F) that are used for clasping females during copulation. The male postabdomen has 252 gonopores (Figure 3D inset) that are involved in transferring mature spermatozoa from the testes 253 to the female in the region of the oviduct during copulation. Besides having a wider frontal 254 opening, pubescence and thickened angular margins are also observed at the ventral margin of 255 the carapace in males (indicated by arrows in Figures 3E and 3F). Fat cells in males are different 256 from those found in females. Male fat cells contain much larger lipid droplets, reduced and less 257 granular cytoplasm, and smaller nucleoli than female fat cells (Figure 4).



259 Figure 3. Representative microanatomical structures of female (A-C) and male (D-F') D. 260 *magna* in the three orthogonal planes. The coronal plane (panel A and D) displays most of the 261 structures in pairs. Inset of panel A shows the oviduct (Ovt; dotted circle) and inset of panel D 262 shows gonopores (Gn) in the male. Panel D', slightly ventral to panel D, displays the prominent 263 and elongated antennules (A1) with flagella (F) at the tips. The sagittal plane of the female 264 (panel B) displays the connection of the compound eye (E) to the optic lobe (OL) and cerebral 265 ganglia (CG) by optic nerves (ON). The labrum (L), maxillules (Mxl), and mandibles (Md) are 266 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). 267 268 The sagittal plane of the male (panel E) shows the elongated seta (Se) on the first thoracic limb. 269 pubescence (P) at the wider ventral opening of the carapace, thickening of carapace at the ventral 270 opening (arrows), one of the testes (Te), and a small portion of sperm duct (SD). Inset of panel E 271 showing the spermatozoa in the testis. The transverse plane of the female (panel C) shows the 272 asymmetrical paired mandibles (Md) with the transverse mandibular tendons (TMT), transverse 273 mandibular muscles (TMM1), transverse muscles of mandibles (TMM2), and the posterior 274 rotator muscles of mandibles (PRM). Inset of panel C displays several hemocytes (H) outlined 275 by dotted circles. The transverse plane of male (panel F) displays the paired copulatory hooks 276 (CH) on the first thoracic limbs (T1) and the thickening of the carapace (arrows) at the ventral 277 opening. This also shows the abundance of fat cells (FC) which are quite different from those in 278 the female. Panel F', slightly above that of panel F, shows the pigmented ocellus (O) is 279 connected to the cerebral ganglion (CG). A1M, antennule muscle; A2M, antennal muscle; Ae, 280 aethetascs; An, anus; Ce, hepatic cecum; Cp, carapace; DLM, dorsal longitudinal muscle; ES, 281 end sac of the maxillary gland; FG, food groove; Ht, heart; LGC, labral gland cell; NC, nerve 282 chord; PA, postabdomen; PCAM, posterior carapace adductor muscle; PM, peritrophic 283 membrane; Ra, ramus of swimming antenna; Tu, tubule of the maxillary gland; VLM, ventral 284 longitudinal muscle.

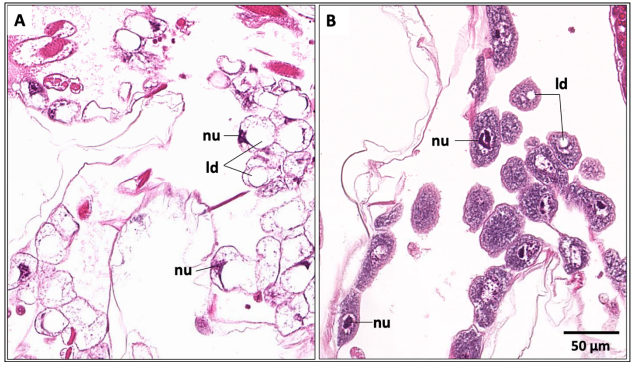


Figure 4. Comparison of male and female fat cells. (A) Fat cells in males consist of larger lipid droplets (ld), reduced and less granular cytoplasm with smaller nucleoli (nu) situated at the cell periphery. (B) Fat cells in females have more granular cytoplasm with smaller lipid droplets (ld) and bigger nucleoli (nu) often appeared subdivided and situated in the cell centers.

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Circulatory system. Daphnia have an open circulatory system and a myogenic heart (63,64). As *Daphnia* are semi-transparent, the beating heart can be easily observed. Hemolymph (blood-like fluid) containing hemocytes (Figure 3C inset) (45,59) is pumped through the body cavity. In line with literature records, we observe that the *Daphnia* heart has a pair of ostia situated in the immediate anterior of the brood chamber, between the midgut and the dorsal surface (Figure 3B, 6C, 6F). Synthesis of hemoglobin happens in fat cells and epipodite cells (on thoracic limbs)(53).

298 Digestive system. Daphnia are filter feeders. Food particles are filtered through filter plates (FP3 299 and FP4) consisting of setae on thoracic limbs 3 and 4, passed through maxillules and mandibles 300 into the esophagus, which is the first part of the digestive system (Figure 3A and B). The 301 digestive system also consists of paired hepatic ceca, midgut, and hindgut (Figures 6A-C, 6E-F) 302 that are lined with epithelial cells and microvilli, with the columnar epithelial cells in the midgut, 303 and the cuboidal cells in hepatic ceca and hindgut (61,65). The labrum houses labral glands that 304 have been suggested to be involved in food ingestion and endocrine function (69,70) (Figure 3B-305 D).

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Excretory system. The maxillary gland, also known as the shell gland, is the organ of excretion,
housed between the inner and outer walls of the carapace (62). It consists of an end sac, a series
of tubules, and an opening that is situated within the anterior part of the brood chamber (Figure
3A, C, D and F).

311

312 Locomotive and osmoregulation system. The second pair of antennae, usually referred to as 313 swimming antennae, is the primary organ of locomotion (43). Each swimming antenna has a 314 protopodite, two rami bearing setae (44) (Figure 3C, D, and F), and is supported by antennal 315 muscles. *Daphnia* have five thoracic limbs (47) (Figure 3B, D, and E). Movements of thoracic 316 limbs produce a constant current that brings food particles into the digestive tract (43,46) and 317 facilitates osmotic regulation, which occurs in the epipodite on each thoracic limb (56). First 318 thoracic limbs in male are different from the female Daphnia, with only the male having the 319 chitinized copulatory hooks (Figure 3F) and longer setae (Figure 3E).

320

321 *Muscular system.* The muscular system is very prominent and occupies a significant portion of 322 the body (43,48). The largest muscles are ventral and dorsal longitudinal muscles that extend 323 along the gut, three paired antennal muscles, transverse mandibular muscles, transverse muscles 324 of mandibles, posterior rotator of the mandibles, carapace adductor muscles, followed by groups 325 of muscles that allow the motion of thoracic limbs and postabdomen (Figure 3). Other small 326 muscles include those around the compound eye, labrum, and esophagus (50). All muscles are 327 striated and surrounded by sarcoplasm, which contains many nuclei and is mostly vacuolated. 328 Sarcoplasm is particularly abundant and more vacuolated in the antennal muscles. Male 329 antennules also carry muscles (Figure 3D) that are absent in the females.

330

Nervous, sensory, and vision systems. Daphnia have a pigmented compound eye consisting of 22 ommatidia (Figure 3B) and a light-sensing, pigmented nauplius eye or ocellus with four lens-like bodies (Figure 3F') (66). Each ommatidium contains eight retinular 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 3B). The cerebral ganglia are connected to two chains of nerve cords that run along the thorax, underneath the gut, and reach other anatomical structures (57,58)

(Figure 3A, E, and F). Both sexes have a pair of antennules bearing a group of 9 olfactory setae
or aesthetascs (71,72) but the male antennules are more prominent and elongated, uniquely fitted
with flagella at their tips (Figure 3D').

340

341 *Reproductive system.* The ovaries in females are paired tubular structures ended with oviducts 342 (Figure 3A). Daphnia are cyclical parthenogens, which means that sexual (meiotic) and clonal 343 (ameiotic) reproduction alternate (73). Under favorable environmental conditions, females 344 produce parthenogenetic eggs that are genetically identical to themselves. During clonal 345 reproduction, happens in clusters where each cluster of four oocytes are formed. Only one definitive oocyte will accumulate yolk granules and lipid droplets during maturation and the 346 347 others will transform into nurse cells (60) (Figure 5A). After maturation, parthenogenetic eggs 348 (Figure 5A) are released into the brood chamber through oviducts and fully developed, free-349 swimming juveniles are extruded after 3 to 4 days. Sexual reproduction is cued by environmental 350 change, such as photoperiod, temperature, and over-crowding, which triggers the 351 parthenogenetic production of genetically identical males for mating with receptive females for 352 sexual recombination; the end point are two embryos that ultimately enter a state of diapause. 353 Unlike parthenogenetic embryos, the development of these resting embryos is arrested at the 354 3000-cell count and enters dormancy (74). The resting embryos are encased in a chitin shell 355 called an ephippium that protects them from harsh environmental conditions (Figure 5C) 356 including freezing and desiccation. Dormancy in *Daphnia* can be exceptionally long, lasting 357 decades and even centuries (75,76). The resting embryos hatch when cued by favorable 358 environmental conditions. A proportion of these resting embryos is not exposed to environmental 359 cues and remains buried in lake sediment, from where it can be isolated, revived and maintained 360 in the laboratory through parthenogenetic reproduction (77,78).

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Testes of male *Daphnia* consist of two long tubular structures connected to gonopores or ejaculatory openings by sperm ducts (Figure 3D and E). Spermatogenesis begins at the testes' walls, and mature spermatozoa are displaced inward toward the central region of the testes (67).

Fat cells, which are polyploid (79) and consist of a massive portion of lipid and glycogen (68), are typically found along the trunk, around ovaries or testes, and on the epipodites of the thoracic

368 limbs (Figure 3). They are most likely sites of vitellogenin synthesis (68). In the females, these 369 cells contain one or several lipid droplets of various size and one large nucleolus of irregular 370 shape, which often appears sub divided into two or more pieces. Compared to the female fat 371 cells, male fat cells contain much larger lipid droplet, reduced and less granular cytoplasm, and a 372 smaller nucleolus that usually is situated at the cell periphery (Figure 4).

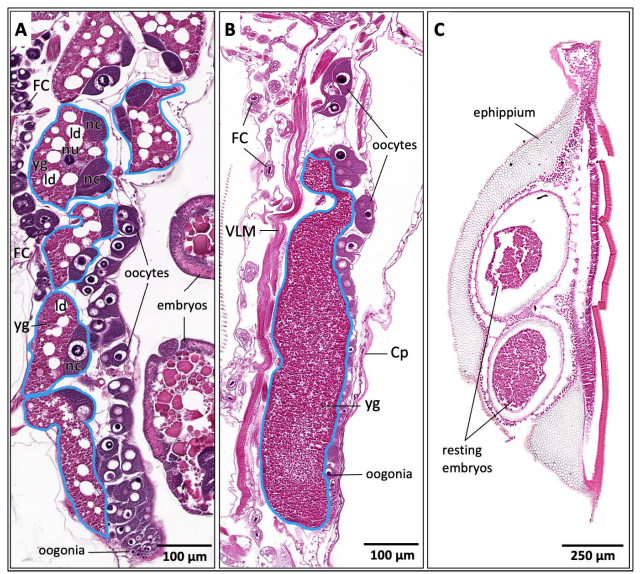




Figure 5. Comparison of parthenogenetic and sexual eggs. (A) The parthenogenetic eggs contain a large amount of lipid droplets (ld) and yolk granules (yg). (B) The sexual eggs contain a large proportion of fine yolk granules without lipid droplets. (C) Resting embryos encased in the ephippium. Top embryo shows artifact. Cp, carapace; FC, fat cells; nc, nurse cell; nu, the nucleus of oocyte; VLM, ventral longitudinal muscle. A solid blue circle indicates an individual egg.

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382 3.1.3 Documenting histopathological change using DaHRA

To illustrate a key application of the DaHRA, we included a female *D. magna* exposed to acetaminophen to demonstrate toxicity effects across multiple organs and tissue types. We compared tissue architecture in clonal replicates of an exposed and non-exposed *D. magna* to identify pathological tissue phenotypes linked to chemical exposure.

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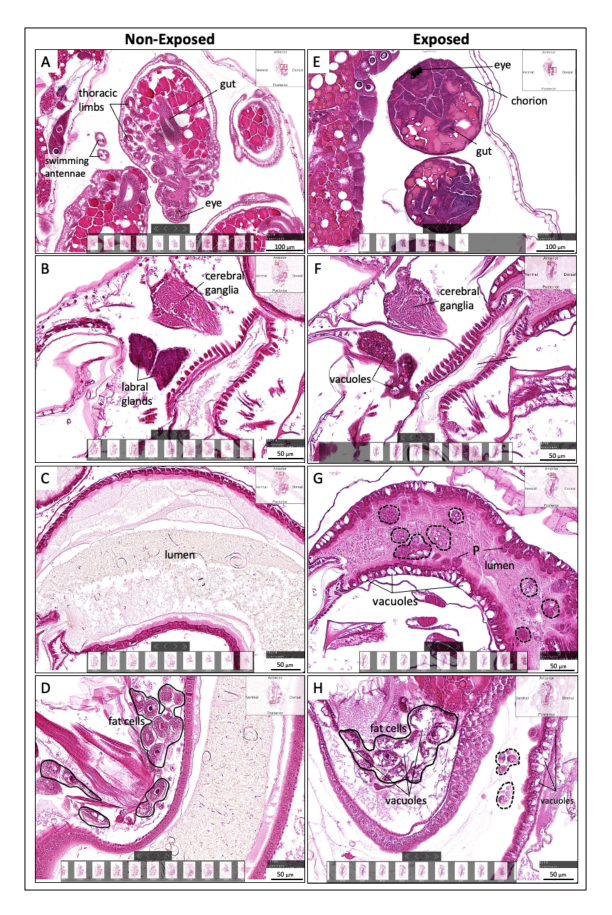
Adult gravid *D. magna* exposed to 25 mg/L of acetaminophen for 72 h produced dead or abnormal embryos. Specifically, the abnormal embryos remained in the chorion, showed development of the compound eye and the gut precursor, but no visible elongation of body length, or development of the swimming antennae and thoracic limbs (Figure 6E) after 72h in the brood chamber.

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Histology of the exposed *Daphnia* also revealed changes in various organs and cell types. Vacuoles were observed in most fat cells (Figure 6H) and across the digestive system, including the labral glands (Figure 6F), midgut (Figure 6G) and hindgut (Figure 6H). The midgut and hindgut of the exposed *D. magna* also showed excessive protruding and sloughing of degenerated epithelial cells (Figure 6G and H).

399

400 Typically, toxicological effects are quantified through apical endpoints (e.g. immobilization) in 401 acute exposures (80,81) and fitness-linked life history traits in chronic exposures (82,83). Few 402 studies employ ultrastructural analysis of a target organ, usually the midgut, for the 403 histopathological assessment of chemical toxicity (85-88). Sublethal concentration of non-404 steroidal drugs have previously been shown to reduce reproduction, growth and to induce 405 neurotoxicity in Daphnia (89–91). Furthermore, acetaminophen has been shown to induce age-406 dependent alterations of gene expression and dysregulation of metabolic pathways (92,93). The 407 tissue toxicity we observe in the adult D. magna and its embryos confirms these published 408 results, highlighting impact on multiple tissues, as well as embryonic failure.



410 Figure 6. Organ and tissue changes in *D. magna* exposed to acetaminophen (E-H) as 411 compared to a clonal replicate of the same non-exposed strain (A-D). (E) Malformed 412 embryos with some development of the compound eye and gut precursors but no visible 413 development of the swimming antennae and thoracic limbs. (F) Vacuoles in the labral glands. 414 (G) Vacuoles, excessive amount of protruding (P) and sloughing of degenerated epithelial cells 415 (dotted circles) in the midgut of exposed D. magna. (H) Alteration observed in the fat cells at the 416 postabdomen region; vacuoles, excessive protruding and sloughing of degenerated epithelial 417 cells (dotted circles) in the hindgut.

418

419 3.2 Method development for *Daphnia* histology

Invertebrates, especially chitinous aquatic invertebrates are challenging to handle for histology,
because of their high tissue water content and the exoskeleton, which has a different texture and
composition than soft tissue. We present here key adaptations of histology methods to *D. magna*,
paving the way to application in other chitinous invertebrates.

424

425 Sample fixation. The accurate representation of microanatomy depends upon the preservation of 426 tissue structure with minimal distortion and optimal staining allowing a clear distinction between 427 cell types and subcellular structures. To determine how to generate sections with minimal 428 distortions for the atlas, we tested three commonly used fixatives: Bouin's solution, 4% 429 Paraformaldehyde in 0.1M phosphate buffer (PFA), and 10% Neutral Buffered Formalin (NBF). 430 Bouin's and PFA are commonly used to fix D. magna. NBF was also tested because it is the 431 most used fixative for routine histology of mammalian tissues. The observation of tissue 432 obtained from these three fixatives showed the clearest results with minimal distortion when 433 samples were fixed with Bouin's solution at room temperature for 48 hours. This fixative 434 preserved soft tissues, such as the gut, which are more challenging to maintain intact and allowed 435 us to visualize microvilli and epithelial cell nuclei distinctively (Figure 7A). Nurse cells (nc), 436 oocytes (oo) and yolk granules (yg) were observed intact in the ovaries (Figure 7B) and cellular 437 features across the developing embryos (Figure 7C) were also more clearly visible in samples 438 fixed with Bouin's as compared to PFA and NBF.

439

The *D. magna* samples (N=23) fixed using PFA and NBF showed "ballooning", a severe fixation artifact causing the carapace to 'puff-up' (Supplementary Figure 2). Inconsistent tissue preservation and distortion in some microanatomical structures were also observed when using

443 these fixatives Particularly, the epithelial cells were not as distinct as when using the Bouin's

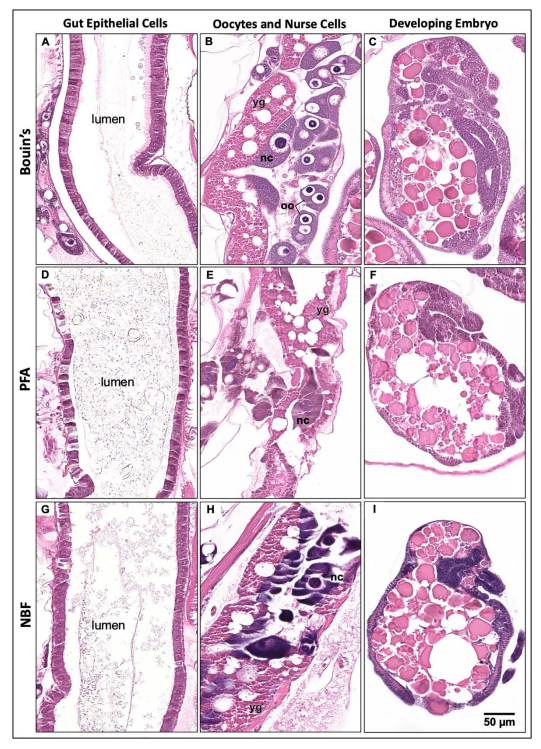
444 fixative, and the microvilli were generally not visible in the gut (Figure 7D and 7G). In the

445 ovaries, oocytes and nurse cells were compromised and challenging to be identified due to the

tissue damage (Figure 7E and 7H). The developing embryos also showed less cell clarity (Figure

447 7H and 7I). In sum, Bouin's solution was observed to be the best fixative for *D. magna* whole-

448 organism histology and was used to fix all samples used in this atlas.



449

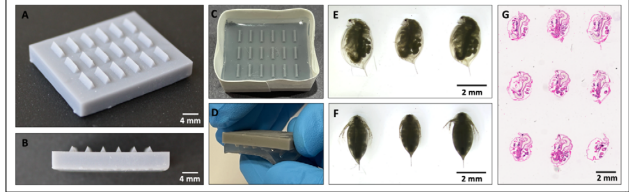
Figure 7. Comparison of histological sections generated with different fixatives: Bouin's, PFA and NBF. Bouin's fixed samples showing intact gut sections, with microvilli and epithelial cell nuclei (A), as well as nurse cells (nc), oocytes (oo) and yolk granules (yg) clearly visible in

the ovary (**B**). In comparison, PFA or NBF fixed samples show less cellular clarity due to the low preservation of histological sections. Cellular details across the developing embryos are

455 clearly visible in samples fixed with Bouin's (C) as compared to other two fixatives (F and I).

456 Sample embedding. Visualization of anatomical sections in each of the three standard anatomical 457 planes (coronal, sagittal, and transverse) is critical for understanding organismal anatomy. 458 Therefore, the ability to generate consistent sections in each of these planes is essential. Agarose 459 pre-embedding using mold or array facilitates consistent positioning and orientation of 460 millimeter-size samples for sectioning (37,38). We have applied this criterion to D. magna 461 sections, using custom-designed plastic mold to create agarose blocks with wells that hold 462 individual fixed D. magna samples prior to tissue processing and paraffin embedding. We tested 463 several designs of casting molds (Supplementary Figure 3) and found that agarose blocks cast 464 using triangular mold (Figure 8, STL file in Supplementary File 2) allowed consistent and 465 precise positioning of adult *D. magna* samples (see Materials and Methods for details). Using the 466 mold for agarose pre-embedding, we were able to embed up to 18 samples per paraffin block for 467 sectioning.

468



469 470 Figure 8. Agarose embedding using casting mold. (A) Top view and (B) side view of 471 triangular mold printed by stereo-lithography (3D-SLA) at 25 µm resolution for a smooth surface 472 allowing easy removal of agarose blocks. (C) Casting 1% agarose block in the taped mold. (D) 473 Agarose block is removed from the mold after solidification by peeling the gel downwards. D. 474 magna samples laid on their sides with a swimming antenna in the wells (E), the rostra facing the 475 same direction for sagittal plane sectioning and positioned in the wells on their back (F) for coronal and transverse plane sectioning. (G) Histological section showing the position of 476 477 samples at sagittal plane.

478

479 In summary, to date, *D. magna* lacked comprehensive and annotated histology resources, despite

- 480 its regulatory importance as a sentinel of environmental hazards and as a model in ecology, and
- 481 evolution. Our open-source visualization platform for *D. magna* histology with its user-friendly
- 482 interface that will help the utilization of this tool by non-specialists and students, contributing to

483 the growing resources for ecotoxicogenomics, biology and comparative medicine. Our atlas 484 provides detailed high-resolution information on tissue-specific phenotypes thereby offering 485 unprecedented opportunities to identify tissue-specific toxicity and genotype-phenotype 486 associations. The coupling of histopathological data with tissue-specific biomarkers (e.g., single 487 cell and spatial transcriptomics) has the potential to revolutionize the assessment of hazardous 488 substances in non-targeted species based on knowledge of the chemical modes of action. Using 489 functional conservation of gene and metabolite network across species, the atlas can assist in 490 revealing tissues target of chemical toxicity across species, including humans (94,95). The 491 pipeline of our atlas was designed with the capability to expand the ontology as more cell or 492 tissue being identified and labeled, and with the intent to extend its features to the other model 493 species. Our ambition is to contribute to the call for cross-species atlas which will be a key 494 resource for the assessment of phenotypes and diseases (96).

495

496 Supporting Information

497 Supplementary Table 1. Fixatives and fixation parameters tested for best preservation of whole498 *D. magna* samples

- Supplementary Table 2. Tissue processing steps for serial dehydration and infiltration of *D*.
 magna samples with Formula R paraffin in tissue processor
- 501 Supplementary Table 3. Automated steps for staining D. magna 5-µm sections with Harris'
- 502 hematoxylin and eosin in an auto-stainer
- 503 Supplementary Figure 1. Anatomy of adult male and female D. magna
- 504 Supplementary Figure 2. "Ballooning" artifact observed in *D. magna* sample fixed with 4% PFA 505 and 10% NBF
- 506 Supplementary Figure 3. 3D-SLA printed casting molds of different teeth designs that were
- 507 tested for orientation and positioning of *Daphnia* samples
- 508 Supplementary Text 1. Daphnia anatomy glossary
- 509 Supplementary File 1. Anatomical ontology for DaHRA
- 510 Supplementary File 2. STL file of casting mold with triangular teeth
- 511

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526

527 **Competing Interests**

528 The authors declare no competing interest.

529

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