OECD Guidance Document
for the Diagnosis of Endocrine-Related Histopathology of Fish Gonads

Dr. Rodney Johnson
Dr. Jeffrey Wolf
Dr. Thomas Braunbeck

Draft of January 7, 2009

OECD
Acknowledgements

This document has been compiled on the basis of the following documents:


The following persons contributed time and effort toward the creation of this document:

Gerald Ankley, US Environmental Protection Agency (EPA), USA
Thomas Braunbeck, University of Heidelberg, Germany
Stephen Duffell, Syngenta, UK
John Fournie, US Environmental Protection Agency (EPA), USA
Christiana Grim, US Environmental Protection Agency (EPA), USA
Anne Gourmelon, OECD
Narisato Hirai, National Institute for Environmental Studies (NIES), Japan
Rodney Johnson, US Environmental Protection Agency (EPA), USA
Susanne Knörr, Heidelberg, Germany
Gerd Maack, UBA, Germany
Christine Rühl-Fehler, Bayer AG, Germany
Christoph Schäfers, Fraunhofer Inst. for Molecular Biology and Applied Ecology (IME), Germany
Helmut Segner, University of Bern, Switzerland
Masanori Seki, Chemicals Evaluation and Research Institute (CERI) Environment, Inc., Japan
Leslie Touart, US Environmental Protection Agency (EPA), USA
Leo van der Ven, National Inst. for Public Health and the Environment (RIVM), The Netherlands
Piet Wester, National Institute for Public Health and the Environment (RIVM), The Netherlands
Jeffrey Wolf, Experimental Pathology Laboratories (EPL), Inc., USA
Marilyn Wolfe, Experimental Pathology Laboratories (EPL), Inc., USA
## Contents

1. **Purpose of this document** ................................................................. 7
2. **Post-mortem and histotechnical procedures** ..................................... 8
  2.1 Fixation of tissue specimens for gonad histopathology .......................... 8
  2.2 Euthanasia and necropsy .................................................................... 10
  2.3 Tissue processing ................................................................................ 13
  2.4 Embedding ......................................................................................... 15
  2.5 Microtomy .......................................................................................... 16
  2.6 Staining, cover-slipping, and slide labeling ........................................... 18
3. **Gonadal histopathology glossary and diagnostic criteria** .................... 20
  3.1 General approach to reading studies .................................................... 20
  3.2 Diagnostic criteria ............................................................................... 20
    3.2.1 Primary criteria in the gonads of male fathead minnow, Japanese medaka and zebrafish .......................................................... 21
    3.2.2 Primary criteria in the gonads of female fathead minnow, Japanese medaka and zebrafish .......................................................... 22
    3.2.3 Secondary criteria in the gonads of male fathead minnow, Japanese medaka and zebrafish ......................................................... 23
    3.2.4 Secondary criteria in the gonads of female fathead minnow, Japanese medaka and zebrafish ......................................................... 24
  3.3 Severity grading .................................................................................. 24
  3.4 Data recording ...................................................................................... 26
  3.5 Histopathology report format ............................................................... 26
4. **Glossary and diagnostic criteria** ........................................................ 27
  4.1 Normal gonadal architecture in fathead minnow, Japanese medaka and zebrafish .......................................................... 28
    4.1.1 Normal testicular architecture in male fathead minnow, Japanese medaka and zebrafish .......................................................... 28
    4.1.2 Normal ovarian architecture in female fathead minnow, Japanese medaka and zebrafish .......................................................... 34
  4.2 Primary diagnoses in fathead minnow, Japanese medaka and zebrafish following exposure to endocrine disruptors ......................................................... 43
    4.2.1 Primary diagnoses in male fathead minnow, Japanese medaka and zebrafish following exposure to endocrine disruptors ......................................................... 43
    4.2.2 Primary diagnoses in female fathead minnow, Japanese medaka and zebrafish following exposure to endocrine disruptors ......................................................... 55
  4.3 Secondary diagnoses in fathead minnow, Japanese medaka and zebrafish following exposure to endocrine disruptors ......................................................... 61
    4.3.1 Secondary diagnoses in male fathead minnow, Japanese medaka and zebrafish following exposure to endocrine disruptors ......................................................... 61
    4.3.2 Secondary in female fathead minnow, Japanese medaka and zebrafish following exposure to endocrine disruptors ......................................................... 66
  4.4 Additional diagnostic criteria and an illustrated glossary of microanatomical and diagnostic terms .......................................................... 69
5. **Gonadal staging criteria** .................................................................... 88
  5.1 Criteria for staging testes in fathead minnow and zebrafish .................. 89
  5.2 Criteria for staging testes in Japanese medaka ....................................... 91
  5.3 Criteria for staging ovaries in fathead minnow, Japanese medaka and zebrafish .......................................................... 92
6. **References** ....................................................................................... 95
Exposure of fathead minnow (*Pimephales promelas*), Japanese medaka (*Oryzias latipes*) and zebrafish (*Danio rerio*) to endocrine disrupting chemicals may result in histopathological alterations of gonadal structure and microarchitecture. The diagnosis of such histopathological changes may, therefore, assist in the detection of endocrine disrupting chemicals. Given the long historical background of piscine histopathology and diverse traditions in the terminology in different fish species and in different regions, there is a need for harmonization in order to facilitate non-biased comparisons of results from different laboratories and, thus, to optimize reproducibility and reliability of histopathological diagnoses. The purpose of this document is to provide guidance for the preparation and histopathological evaluation of gonads from fathead minnow, Japanese medaka and zebrafish following exposure to chemicals potentially disrupting the endocrine system.

Throughout this document, the proposed procedures were derived from consensus opinions of various fish pathologists, recommendations from the Bilthoven (2002), Paris (2003) and Heidelberg (2004) workshops, from information distilled from previous guidelines, and the scientific literature. Major sections of the guidance document comprise (1) post-mortem and histotechnical procedures (chapter 2), (2) primary, secondary and additional gonadal histopathology glossary and diagnostic criteria (chapters 3 and 4), and (3) gonadal staging criteria (chapter 5).

**Note:** It is the intention of the authors to make this document available on the internet, preferably on the OECD website. Further experience in the use of histopathological diagnoses of endocrine-disruptor-related changes in testes and ovaries of fathead minnow, Japanese medaka and zebrafish should be incorporated into future updates of this guidance document.
1. Purpose of this document

The purpose of this document is to provide guidelines for the preparation and histopathological evaluation of gonads from fathead minnow (*Pimephales promelas*), Japanese medaka (*Oryzias latipes*) and zebrafish (*Danio rerio*) following exposure to chemicals potentially disrupting the endocrine system.

The goals of these guidelines are

a) to harmonize techniques for the preparation of histological specimens for the diagnosis of endocrine-related histopathology in fish gonads,

b) to provide guidance for the diagnosis of histopathological alterations in fish exposed to potential endocrine-disrupting chemicals according to

   (1) the OECD “Fish Screening Assay” (FSA),

   (2) the US “Fathead Minnow 21-d Short-term Reproduction Assay”, and

   (3) any other assay designed for the detection of endocrine-disrupting chemicals and requiring histological analyses of the gonads.

c) to supply template text for laboratory protocols, and (d) to facilitate non-biased comparisons of results from different laboratories.

Throughout this document, the proposed procedures were derived from consensus opinions of various fish pathologists, recommendations from the Bilthoven (2002), Paris (2003) and Heidelberg (2004) workshops, information distilled from previous guidelines, and the scientific literature.

This guidance document is divided into three main sections:

   (1) post-mortem and histotechnical procedures (chapter 2),

   (2) gonadal histopathology glossary and diagnostic criteria (chapters 3 and 4),

   (3) gonadal staging criteria (chapter 5).

In order to facilitate interspecies comparisons, this document is not partitioned according to species; instead, for each diagnosis, all three fishes (fathead minnow, Japanese medaka, zebrafish) are considered simultaneously whenever possible.
2. Post-mortem and histotechnical procedures

The purpose of this section is to outline post-mortem steps and procedures that occur prior to evaluation of histological sections on glass slides including euthanasia, necropsy, tissue fixation, decalcification, tissue trimming, processing, embedding, microtomy, staining, cover-slippering, and slide labeling.

2.1 Fixation of tissue specimens for gonad histopathology

Techniques were selected that would most optimally

1. preserve the cellular structure of the gonads;
2. maximize the amount of gonad tissue available for analysis;
3. sample the gonads in a representative and consistent fashion; and
4. allow the pathologist to examine at least 3 step sections of both gonads on a single glass slide.

In all three fish species, the gonads are excised from the fish. In case very young individuals were used, the gonads may alternatively be fixed inside the carcass. However, in order, to allow optimal penetration of the fixative, head and tail portions should be removed by means of sharp razor blades.

Davidson's fixative is the recommended fixative. Compared to other common fixatives, such as 10% neutral buffered formalin or Bouin’s fixative, the advantages of Davidson’s fixative are as follows:

1. the morphologic appearance of gonad sections is generally considered to be comparable to sections fixed in Bouin’s fixative and superior to sections fixed in formalin;
2. compared to Bouin’s fixative, which contains picric acid, Davidson’s fixative is generally considered to be less noxious, less hazardous, and more easily disposed of;
3. there is anecdotal information which suggests that Bouin’s fixative may be difficult to obtain in the near future;
4. specimens fixed in Bouin’s fixative require multiple rinses prior to transfer to alcohol or formalin.

**Formulation of Davidson’s fixative** (1 L; Fournie et al., 2000):

- Formaldehyde (37 - 40 %): 200 ml
- Glycerol: 100 ml
- Glacial acetic acid: 100 ml
- Absolute alcohol: 300 ml
- Distilled water: 300 ml

**Formulation of Modified Davidson’s Fixative** (1 L):

- Formaldehyde (37 - 40 %): 220 ml
- Glacial acetic acid: 115 ml
- 95% Ethyl alcohol: 330 ml
- Distilled water: 335 ml
For a photographic comparison of specimens fixed in Davidson’s versus Bouin’s fixatives, see Fig. 1. Please be aware that different recipes and products that are designated as “Davidson’s fixative” may actually be modifications of the original formula (Appendix C); if a modified Davidson’s fixative is used, this should be noted by the laboratory. If necessary, a decalcification solution (EDTA, acetic acid) may be used. Factors that may affect the need for decalcification include the size of the individual fish, the length of time that the carcass was immersed in fixative, and the extent to which the abdominal cavity came into contact with the fixative.

Fig. 1. Fathead minnow, testis (a, b) and ovary (c, d) fixed in Bouin’s fixative (a, c) and modified Davidson’s fixative (b, d). Color contrast was slightly superior in testes fixed with Davidson’s fixative and was clearly superior in ovaries fixed with Bouin’s fixative. Either fixative is satisfactory for diagnostic purposes; however, Davidson’s fixative was eventually selected.
2.2  Euthanasia and necropsy

Objectives
(1)  Provide for the humane sacrifice of fish.
(2)  Obtain necessary body weights and measurements.
(3)  Obtain specimens for vitellogenin analysis.
(4)  Excise gonad specimens (not required for Japanese medaka and zebrafish).
(5)  Evaluate secondary sex characteristics (not required for Japanese medaka and zebrafish).
(6)  Provide for adequate fixation of the gonads and/within the carcass.

Materials
(1)  Fish transport container (~500 ml, contains water from the experimental tank or system reservoir).
(2)  Small dip net.
(3)  Euthanasia chamber (~500 ml vessel).
(4)  Euthanasia solution (see below).
(5)  Electronic slide caliper (minimum display: ± 0.1mm)
(6)  Electronic analytical balance (minimum display: ± 0.1mg) and tared vessels.
(7)  Stereoscopic microscope.
(8)  Pins and corkboard.
(9)  Small scissors (e.g., iris scissors).
(10) Small forceps.
(11) Microdissection forceps.
(12) Microdissection scissors.
(13) Gauze sponges.
(14) Davidson’s fixative (see 2.1).
(15) Plastic syringe (3 ml).
(16) Standard plastic tissue cassettes (one per fish).
(17) Fixation containers (100 ml, one per fish).

Euthanasia solution (MS-222; tricaine methanesulfonate)

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tricaine methanesulfonate</td>
<td>100 mg</td>
</tr>
<tr>
<td>Sodium bicarbonate</td>
<td>200 mg</td>
</tr>
<tr>
<td>Tank or reservoir water</td>
<td>1 L</td>
</tr>
</tbody>
</table>
**Procedures**

1. Fish should be sacrificed within one to two minutes prior to necropsy. Therefore, unless multiple prospectors are available, multiple fish should not be sacrificed simultaneously.

2. Using the small dip net, a single fish is removed from the experimental chamber and transported to the necropsy area in the transport container. For each test chamber, all male fish are sacrificed prior to the sacrifice of female fish; the sex of each fish is determined by external body characteristics (fathead minnow: presence or absence of nuptial tubercles, dorsal pad; medaka: shape of dorsal and anal fins; zebrafish: coloration, body shape).

3. The fish is placed into buffered MS-222 (tricaine methanesulfonate) solution. The fish is removed from the solution when there is cessation of respiration and the fish is unresponsive to external stimuli.

4. The fish is wet weighed, measured according to protocol, and a blood sample is obtained from the caudal artery/vein or heart.

5. The fish is placed on a corkboard on the stage of a dissecting microscope. Using iris scissors and small forceps, the abdomen is opened via a carefully made incision that extends along the ventral midline from the pectoral girdle to a point just cranial to the anus.

6. The fish is placed in dorsal recumbency and the opposing flaps of body wall are pinned laterally to expose the abdominal viscera (Figs. 2, 3).

7. Using the small forceps and small scissors, the abdominal viscera (liver, gastrointestinal tract, spleen, pancreatic tissue, and abdominal mesentery) are carefully removed en masse in the following manner:
   
a. The intestine is severed proximal to the anus.
   
b. A forceps is applied to the terminal portion of the intestine. Using gentle traction and taking care not to disturb the gonads, the viscera are dissected out of the abdominal cavity in a caudal to cranial direction.
   
c. The distal esophagus is severed just proximal to the liver.

8. **Fathead minnow:** Using a syringe, **approximately 0.5 ml of Davidson’s fixative is then gently applied to the gonads in situ.** Approximately 90 seconds following the application of fixative, the liquid fixative within the abdomen is removed with a gauze sponge, and the gonads are excised in a manner similar to the abdominal viscera:
   
a. Using the microdissection scissors, the spermatic ducts or oviducts are severed proximal to the genital pore.
   
b. Microdissection forceps are then applied to the spermatic ducts/oviducts. Using gentle traction, the gonads are dissected out of the abdominal cavity in a caudal to cranial direction, severing the mesorchial/mesovarial attachments as needed using the microdissection scissors. The left and right gonads may be excised individually or they may be excised simultaneously and subsequently divided at their caudal attachment.

   **Japanese medaka, zebrafish:** Using the microdissection scissors and forceps, the liver is dissected from the viscera and retained for vitellogenin analysis. **Go to step (11).**

9. **Fathead minnow:** The gonads (right and left) are placed into a pre-labeled plastic tissue cassette which is then placed into an individual container of Davidson’s fixative accompanied by the abdominal viscera. The volume of fixative in the container should be at least 10 times the approximated volume of the tissues. The fixative container is gently agitated for five seconds to dislodge air bubbles from the cassette.
(10) **Fathead minnow**: Using the carcass, the secondary sex characteristics are assessed (e.g., dorsal nape pad, nuptial tubercles). The carcass is then added to the fixative container. **Go to step (12).**

(11) **Japanese medaka, zebrafish**: The abdominal cavity is *gently* flushed with 0.5 ml of Davidson’s fixative, and then the carcass and abdominal viscera are placed in an individual container of Davidson’s fixative. The volume of fixative in the container should be at least 10 times the approximated volume of the tissues. **Go to step (12).**

(12) **All tissues remain in Davidson’s fixative overnight**, followed by transfer to individual containers of 10% neutral buffered formalin the next day. Containers with cassettes are gently agitated for 5 seconds to ensure adequate penetration of formalin into cassettes (it is not necessary to rinse with water or perform multiple changes in formalin).

---

**Fig. 2.** Fathead minnow, male: Excision of the testes during necropsy. (a) The abdominal wall is pinned laterally. (b) The terminal intestine is severed and retracted prior to removal. (c) The testes are grasped near the spermatic ducts. (d) Removal of the testes is complete.
Fig. 3. Fathead minnow, female: Excision of the ovaries during necropsy. (a) The abdominal wall is pinned laterally. (b) The terminal intestine is severed and retracted prior to removal. (c) The ovaries are grasped near the oviducts. (d) Removal of the ovaries is complete.

2.3 Tissue processing

Objectives

(1) **Japanese medaka and zebrafish**: Trim carcass for embedding and microtomy.
(2) Dehydrate tissue to provide for adequate penetration of paraffin.
(3) Impregnate the tissue with paraffin to maintain tissue integrity and create a firm surface for microtomy.

Materials

(1) Tissue processor.
(2) Paraffin heating pots.
(3) Processing unit oven.
(4) Activated charcoal.
(5) Paraffin (Paraplast™, or equivalent).
(6) 10 % neutral buffered formalin.
(7) Ethyl alcohol (absolute and dilutions as required).
(8) Proprietary clearing agent (Clear Rite-3™ or equivalent).
(9) Xylene.

Procedures

(0) **Japanese medaka, zebrafish**: Head and tail are severed and removed prior to embedding.

(1) Labeled tissue cassettes are removed from formalin storage and are washed in tap water.

(2) The cassettes are placed in the processing basket(s) in a single layer. The processing baskets are loaded into the tissue processor.

(3) The processing schedule is selected (see Appendix B, Schedule 1). The “Gonad Program” or equivalent is selected for fathead minnow.

(4) After the tissue processor has completed the processing cycle, the basket(s) may be transferred to the embedding station.

Schedule 1: Tissue Processing

<table>
<thead>
<tr>
<th>Station No.</th>
<th>Reagent</th>
<th>Pressure/Vacuum Cycle</th>
<th>Heat (°C)</th>
<th>GONAD program (min.)</th>
<th>WHOLE-FISH program (min.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>10 % NBFa</td>
<td>On</td>
<td>Ambient</td>
<td>40</td>
<td>60</td>
</tr>
<tr>
<td>2</td>
<td>70 % Ethanol</td>
<td>On</td>
<td>Ambient</td>
<td>40</td>
<td>60</td>
</tr>
<tr>
<td>3</td>
<td>80 % Ethanol</td>
<td>On</td>
<td>Ambient</td>
<td>40</td>
<td>60</td>
</tr>
<tr>
<td>4</td>
<td>95 % Ethanol</td>
<td>On</td>
<td>Ambient</td>
<td>40</td>
<td>60</td>
</tr>
<tr>
<td>5</td>
<td>95 % Ethanol</td>
<td>On</td>
<td>Ambient</td>
<td>40</td>
<td>60</td>
</tr>
<tr>
<td>6</td>
<td>100 % Ethanol</td>
<td>On</td>
<td>Ambient</td>
<td>40</td>
<td>60</td>
</tr>
<tr>
<td>7</td>
<td>100 % Ethanol</td>
<td>On</td>
<td>Ambient</td>
<td>40</td>
<td>60</td>
</tr>
<tr>
<td>8</td>
<td>100 % Ethanol</td>
<td>On</td>
<td>Ambient</td>
<td>40</td>
<td>60</td>
</tr>
<tr>
<td>9</td>
<td>Clear Rite 3</td>
<td>On</td>
<td>Ambient</td>
<td>60</td>
<td>80</td>
</tr>
<tr>
<td>10</td>
<td>Clear Rite 3</td>
<td>On</td>
<td>Ambient</td>
<td>60</td>
<td>80</td>
</tr>
<tr>
<td>11</td>
<td>Paraffin</td>
<td>On</td>
<td>60</td>
<td>45(60b)</td>
<td>75(100b)</td>
</tr>
<tr>
<td>12</td>
<td>Paraffin</td>
<td>On</td>
<td>60</td>
<td>45(60b)</td>
<td>75(100b)</td>
</tr>
<tr>
<td>13</td>
<td>Paraffin</td>
<td>On</td>
<td>60</td>
<td>45(60b)</td>
<td>75(100b)</td>
</tr>
<tr>
<td>14</td>
<td>Paraffin</td>
<td>On</td>
<td>60</td>
<td>45</td>
<td>75</td>
</tr>
</tbody>
</table>

* Neutral buffered formalin.

b Times are increased for processors that have three (versus four) final stations.

Automatic cleaning cycle to be run after removal of tissues from the processor. Time, temperature, and vacuum are preset by the manufacturer.

OECD Guidance Document for the Diagnosis of Endocrine-related Histopathology of Fish Gonads
2.4 Embedding

Objective

(1) Properly orient the tissue in solidified paraffin for microtomy.

Materials

(1) Embedding station (thermal, dispensing and cryo consoles).
(2) Paraffin heating pots.
(3) Paraffin transfer pots.
(4) Laboratory oven.
(5) Thermometer.
(6) Embedding molds.
(7) Block drawers.
(8) Forceps.
(9) Scraper.
(10) Standard paraffin.

Procedures

(1) The cryoconsole of the embedding station is turned on. (Power to the dispensing console and thermal console should remain on at all times.)

(2) The basket(s) of cassettes is/are removed from the processor and immersed in the paraffin-filled front chamber of the embedding station thermal console.

(3) The first cassette to be embedded is removed from the front chamber of the thermal console. The cassette lid is removed and discarded, and the cassette label is checked against the animal records to resolve potential discrepancies prior to embedding.

(4) An appropriately sized embedding mold is selected.

(5) The mold is held under the spout of the dispensing console and filled with molten paraffin.

(6) The gonads are removed from the base of the cassette and are placed in the molten paraffin in the mold. The two gonads (left and right) are oriented horizontal to their long axis in the mold to allow for longitudinal sectioning.

(7) The base of the cassette is placed on top of the mold. Additional paraffin is added to cover the bottom of the base.

(8) The mold with the cassette base is placed on the cooling plate of the cryo console.

(9) After the paraffin has solidified, the block (i.e., the hardened paraffin containing the tissues and the cassette base) is removed from the mold.

(10) Steps 3 through 10 are repeated for each cassette to be embedded.
2.5 Microtomy

Objective
(1) Create and mount histological sections for staining.

Materials
(1) Microtome.
(2) Disposable microtome knives.
(3) Lipshaw Pike® oil (or equivalent lightweight, machine oil).
(4) Temperature-controlled water bath.
(5) Ice.
(6) Microscope slides.
(7) Staining racks.
(8) Permanent slide marking pen.
(9) Forceps.
(10) Fine-tipped paint brush.
(11) Temporary labels.
(12) Slide warmer/oven.

Procedures
(1) The temperature in the water bath is allowed to stabilize so that ribbons cut from the tissue blocks will spread out uniformly on the surface without melting. This temperature assessment is a qualitative judgment made by the microtomist before and during microtomy.

(2) If necessary, a new blade is mounted onto the microtome and the microtome is lubricated with oil.

(3) The initial phase of microtomy is termed “facing” the block and is conducted as follows:
   a. The block is placed in the chuck of the microtome.
   b. The chuck is advanced by rotating the microtome wheel and thick sections are cut from the paraffin surface of the block until the knife reaches the embedded tissues. This process is referred to as “rough trimming” of the block.
   c. The section thickness on the microtome is set between 4 -10 microns. The chuck is advanced and multiple sections are cut from the block to remove any artifacts created on the cut surface of the tissue during rough trimming. This process is termed “fast trimming” of the block.
   d. The block is removed from the chuck and placed face down on ice to soak the tissue.
   e. Steps a. through d. are repeated until all blocks to be microtomed have been faced.
   f. If it is determined during facing that any block is not of acceptable quality for microtomy, it is returned for re-embedding before proceeding with microtomy.
g. Any extraneous pieces of paraffin are removed from the microtome and workstation periodically during facing and before proceeding with the next phase of microtomy.

h. **Japanese medaka:** The block is faced from the dorsum towards the ventrum (through the majority of the swim bladder) until the fused left and right gonads are reached.

**Zebrafish:** The block is faced from the ventrum and towards the dorsum until the level is reached at which the following anatomic landmarks are observed:

   i. **Testis:** disappearance of the liver, prior to the appearance of the rostral swim bladder. There is greater than 500 microns of tissue thickness at this level. Sections at this level do not contain intestine.

   ii. **Ovary:** the appearance of the swim bladder.

(4) The next phase of microtomy is final sectioning and mounting of tissue sections on slides. These procedures are conducted as follows:

   a. Macroscopic lesions (if any) that are reported in the records are noted. Care is taken to include any macroscopic lesions in the sections collected during final sectioning.

   b. The block is removed from the ice and placed in the chuck of the microtome.

   c. With the section thickness on the microtome set to 4 - 5 µm, the chuck is advanced by rotating the microtome wheel. Sections are cut from the block until a “ribbon” containing at least one acceptable section has been produced. As necessary during sectioning, the block may be removed from the chuck, placed on ice to soak the tissue, and replaced in the chuck.

   d. Each ribbon is floated flat on the surface of the water in the water bath. An attempt is made to obtain at least one section in the ribbon that contains no wrinkles and has no air bubbles trapped beneath it.

   e. A microscope slide is immersed beneath the best section in the floating ribbon. The section is lifted out of the water using the slide. This process is referred to as “mounting” the section on the slide.

   f. **A single slide is prepared for each fish. A total of three step sections (each section consisting of both the right and left gonad) are mounted on each slide. The first section is obtained at the point where approximately half of the gonad has been removed and the size of the section is maximized. For both the testis and the ovary, the second and third sections are taken at 50 micron intervals following the first section.**

   g. With a slide-marking pen, the block number from which the slide was produced is recorded on the slide.

   h. The slide is placed in a staining rack.

   i. The block is removed from the chuck and placed facedown for storage.

   j. Steps a. through h. are repeated for all blocks to be microtomed.
2.6 Staining, cover-slippping, and slide labeling

Objectives
(1) Differential staining of intra- and inter-cellular components of the gonads to facilitate diagnostic examination by bright field microscopy.
(2) Permanently seal mounted and stained tissues.
(3) Permanently identify stained sections in a manner that allows complete traceability.

Materials
(1) Automated slide stainer (optional).
(2) Robot cover-slippping machine (optional).
(3) Clarifier solution (Richard Allen or equivalent).
(4) Bluing reagent (Richard Allen or equivalent).
(5) Eosin-Y (Richard Allen or equivalent, Appendix C).
(6) Hematoxylin-2 (Richard Allen or equivalent, Appendix C).
(7) Xylene.
(8) Absolute ethyl alcohol (100% ETOH).
(9) 95% ETOH.
(10) 80% ETOH.
(11) Cover-slippping mountant (Permoun®️, DPex®️ or equivalent).
(12) Glass cover-slips, No. 1, 24 x 50 (or 60) mm.
(13) Slide flats.

Gill’s Hematoxylin Solution (Gill et al. 1974)
- Distilled water: 730 ml
- Ethylene glycol: 250 ml
- Hematoxylin, anhydrous: 2 g
- Sodium iodate: 0.2 g
- Aluminum sulfate: 17.6 g
- Glacial acetic acid: 20 ml

Eosin Solution
- Eosin Y (1 % aqueous solution): 100 ml
- Ethyl alcohol, 95%: 600 ml
- Glacial acetic acid: 4 ml
Procedures

(1) Staining
   a. Slides are routinely air-dried overnight before staining.
   b. An example H&E staining schedule for automated stainers is given below. A similar schedule can be adapted for manual staining.

(2) Cover-slipping
   a. Cover-slips can be applied manually or automatically.
   b. A slide is dipped in xylene, and the excess xylene is gently knocked off the slide.
   c. Approximately 0.1 ml of mounting medium is applied near the end of the slide opposite to the frosted end.
   d. A cover-slip is tilted at a shallow angle as it is applied to the slide.

(3) Labeling
   a. Each slide label should contain the following information:
      i. Laboratory name.
      ii. Species.
      iii. Specimen No./Slide No.
      iv. Chemical/Treatment group.
      v. Date (optional).

Schedule 2. Hematoxylin and eosin Staining

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Minutes in Reagent</th>
<th>Reagent Maintenance after 1st run</th>
<th>Reagent Maintenance after 2nd run</th>
</tr>
</thead>
<tbody>
<tr>
<td>Xylene</td>
<td>4</td>
<td>Remove</td>
<td>Remove</td>
</tr>
<tr>
<td>Absolute Alcohol</td>
<td>2</td>
<td>Remove</td>
<td>Remove</td>
</tr>
<tr>
<td>80% Alcohol</td>
<td>1</td>
<td>Renew</td>
<td>Renew</td>
</tr>
<tr>
<td>Water</td>
<td>1</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Hematoxylin</td>
<td>3</td>
<td>—</td>
<td>Remove</td>
</tr>
<tr>
<td>Water</td>
<td>2</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Clarifier</td>
<td>1</td>
<td>Renew</td>
<td>Renew</td>
</tr>
<tr>
<td>Water</td>
<td>1</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Bluing</td>
<td>1</td>
<td>Renew</td>
<td>Renew</td>
</tr>
<tr>
<td>Water</td>
<td>2</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>95% Alcohol</td>
<td>1</td>
<td>Renew</td>
<td>Renew</td>
</tr>
<tr>
<td>Eosin</td>
<td>1</td>
<td>—</td>
<td>Renew</td>
</tr>
<tr>
<td>Absolute Alcohol</td>
<td>4</td>
<td>Remove</td>
<td>Remove</td>
</tr>
<tr>
<td>Xylene</td>
<td>3</td>
<td>Remove</td>
<td>Remove</td>
</tr>
</tbody>
</table>
3. Gonadal histopathology glossary and diagnostic criteria

The purposes of this section are:

1. to provide general guidance for the light microscopic evaluation of tissue sections;
2. to promote a common awareness of various pathological findings that may be observed; and
3. to foster consistency in the use of diagnostic terminology.

3.1 General approach to reading studies

Studies are to be read by individuals experienced in reading toxicologic pathology studies, and who are familiar with normal small fish gonad histology, with gonadal physiology, and with general responses of the gonads to toxicologic insult. Pathologists may be board certified (e.g. American College of Veterinary Pathologists, The European Centre of Toxicologic Pathology, or other certifying organizations); however, certification is not a requirement as long as the pathologist has obtained sufficient experience with, and knowledge of, fish histology and toxicologic pathology. Technicians should not be used to conduct readings due to the subtle nature of some changes and the need for subjective judgments based on past experience.

It is recognized that there is a limited pool of pathologists with the necessary training and experience that are available to read the gonadal histopathology for the Fish Screening Assay (FSA) or the 21-day reproduction assay. If an individual has toxicological pathology experience and is familiar with gonadal histology in small fish species, he/she may be trained to read the fish assay(s). If pathologists with little experience are used to conduct the histopathological analysis, informal peer review may be necessary.

Pathologists are to read the studies non-blinded (i.e. with knowledge of the treatment group status of individual fish). This is because endocrine effects on histomorphology tend to be incremental, and subtle differences between exposed and unexposed animals may not be recognizable unless tissue sections from high dose animals can be knowingly compared to those from controls. Thus, the aim of the initial evaluation is to ensure that diagnoses are not missed (i.e., to avoid false-negative results). However, it is expected that any potential compound-related findings will be re-evaluated by the pathologist in a blinded manner prior to reporting such findings, when appropriate. Certain diagnostic criteria, such as relative increases or decreases in cell populations, cannot be read in a blinded manner due to the diagnostic dependence on control gonads. As a rule, treatment groups should be evaluated in the following order: control, high-dose, intermediate-dose, and low-dose.

It is suggested that the pathologists be provided with all available information related to the study prior to conducting their readings. Information regarding gross morphologic abnormalities, mortality rates, and general test population performance and health are useful for pathologists to provide comprehensive reports and to aid in the interpretation of findings. For a more comprehensive discussion of standard reading approaches for toxicologic pathology studies, please refer to the Society of Toxicologic Pathology Best Practices for reading toxicologic histopathology studies (Crissman et al., 2004).

3.2 Diagnostic criteria

Histopathology is a descriptive and interpretive science, and therefore somewhat subjective. However, histopathological evaluations of the same study by any qualified pathologist should identify the same treatment-related findings (Crissman et al., 2004). Therefore, we aim to define the diagnostic criteria...
that will likely be encountered during the histopathological analysis of the FSA or the 21-day reproduction assay in fathead minnow, medaka and/or zebrafish.

In the following text, a consolidated set of diagnostic criteria will be introduced. These criteria are based on pathologists’ experience with certain consistent histopathological changes that occur in fathead minnow, medaka and/or zebrafish gonads in response to chemical exposure; however, any additional/novel findings that are exposure-related shall also be reported.

The criteria below have been divided into two categories (Table 1):

1. primary criteria,
2. secondary criteria and
3. additional criteria.

The criteria are graded for severity on a numerical scale. Likewise, any novel findings are either graded on a numerical scale, or are qualitatively described.

Table 1. Primary and secondary diagnoses for histopathological changes in male and female fathead minnow, Japanese medaka and zebrafish after exposure to potential endocrine disruptors

<table>
<thead>
<tr>
<th>Primary Diagnoses</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>For males:</strong></td>
</tr>
<tr>
<td>1. Increased proportion of spermatogonia</td>
</tr>
<tr>
<td>2. Presence of testis-ova</td>
</tr>
<tr>
<td>3. Increased testicular degeneration</td>
</tr>
<tr>
<td>4. Interstitial (Leydig) cell hyperplasia/hypertrophy</td>
</tr>
<tr>
<td><strong>For females:</strong></td>
</tr>
<tr>
<td>1. Increased oocyte atresia</td>
</tr>
<tr>
<td>2. Perifollicular cell hyperplasia/hypertrophy</td>
</tr>
<tr>
<td>3. Decreased vitellogenesis</td>
</tr>
<tr>
<td>4. Changes in gonadal staging</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Secondary Diagnoses</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>For males:</strong></td>
</tr>
<tr>
<td>1. Decreased proportion of spermatogonia</td>
</tr>
<tr>
<td>2. Increased vascular or interstitial proteinaceous fluid</td>
</tr>
<tr>
<td>3. Asynchronous gonad development</td>
</tr>
<tr>
<td>4. Altered proportions of spermatozoa or spermatocytes</td>
</tr>
<tr>
<td>5. Gonadal staging</td>
</tr>
<tr>
<td>6. Granulomatous inflammation</td>
</tr>
<tr>
<td><strong>For females:</strong></td>
</tr>
<tr>
<td>1. Interstitial fibrosis</td>
</tr>
<tr>
<td>2. Egg debris in the oviduct</td>
</tr>
<tr>
<td>3. Granulomatous inflammation</td>
</tr>
<tr>
<td>4. Decreased post-ovulatory follicles</td>
</tr>
</tbody>
</table>

### 3.2.1 Primary criteria in the gonads of male fathead minnow, Japanese medaka and zebrafish

The following criteria have been defined in male fish as diagnoses of primary interest:

1. **Increased proportion of spermatogonia**: Increases in the proportion of spermatogonia are consequent of changes of the relative ratios of spermatogenic cells. This could be due to an increase in the number of spermatogonia, or a decrease in the number of other cell types, such as spermatocytes, spermatids, and spermatozoa. Because the diagnosis of increased proportion of spermatogonia is dependent on a comparison to controls, it is necessary to establish the normal range of the ratios of spermatogenic cells in control male fish testes prior to making determinations on relative proportions in dose groups.

2. **Presence of testis-ova**: The presence of one or more individualized or clustered oogenic cells within the testis. Oocytes within the testis may be determined to be perinucleolar, cortical alveolar, vitellogenic, or atretic. There is little or no evidence of ovarian architecture. Whenev-
er applicable, the term testis-ova should be used in preference to less precise terms such as “intersex” or “hermaphrodite”.

Terms such as hermaphroditism, hermaphroditism, intersex, mixed sex, sex reversal, ovotestis, ovatessis, ova-testis, testis-ova, testicular oocytes or testicular oogenesis abound in the lower vertebrate scientific literature. Three of these terms are listed in standard medical dictionaries (Dorland’s, 1981; Illustrated Stedman’s, 1982), and their consensus definitions are: (1) intersex: the presence of both male and female characteristics in the same individual; (2) hermaphroditism (= hermaphroditism): the presence of male and female gonadal elements within the same individual; (3) ovotestis: the presence of male and female gonadal elements within the same gonad. The validity of many other terms is largely based on traditional usage, which has not always been consistent.

(3) **Increased testicular degeneration**: Testicular degeneration is characterized by

a) individual or clustered apoptotic germ cells;

b) vacuolated germ cells; and/or

c) multinucleated (syncytial) cells in the germinal epithelium or testicular lumen.

Apoptotic germ cells are characterized by cell shrinkage, nuclear condensation, and fragmentation into spherical, membrane-bound bodies, which are often phagocytized by neighboring cells. There is no inflammation associated with these cells. If possible, testicular degeneration should be differentiated from necrosis, which is characterized morphologically by cytoplasmic coagulation or swelling, nuclear karyorrhexis (destructive nuclear fragmentation) or pyknosis (shrinkage of nuclei in conjunction with chromatin condensation of the chromatin to a central mass), associated inflammation, a locally extensive pattern of tissue involvement, and/or the involvement of different local tissue elements (e.g., both germinal and stromal tissues). Extensive testicular degeneration may lead to localized or generalized loss of the germinal epithelium.

(4) **Interstitial cell (Leydig cell) hyperplasia/hypertrophy**: An increase in number and/or size of the interstitial cells responsible for producing androgens. Interstitial cells may have larger, more rounded nuclei, and interstitial cell aggregates may occupy and expand some interstitial spaces.

### 3.2.2 Primary criteria in the gonads of female fathead minnow, Japanese medaka and zebra-fish

The following criteria have been defined in female fish as diagnoses of primary interest:

(1) **Increased oocyte atresia**: An increase in degradation and resorption of oocytes at any point in development. Atresia is characterized by clumping and perforation of the chorion, fragmentation of the nucleus, disorganization of the ooplasm, and/or the uptake of yolk materials by perifollicular cells.

(2) **Perifollicular cell hyperplasia/hypertrophy**: Increase in the size or number of granulosa, theca, and/or surface epithelium cells involved in a developing follicle. Abnormal perifollicular cell hypertrophy must be distinguished from the normally enlarged granulosa and theca cells of a post-ovulatory follicle.

(3) **Decreased yolk formation**: A decrease in the amount of vitellogenic/yolk material that is deposited in the developing oocyte. Decreased vitellogenesis is characterized by the presence of oocytes in which yolk material is not present despite their relatively large size. Note that oocytes may be affected to varying degrees. Some affected oocytes have extremely fine vitellogenic granules, and this is interpreted as ineffective vitellogenesis.
Changes in gonadal staging: Gonadal staging results are virtually meaningless in terms of individual fish (versus treatment groups). This is because considerable animal-to-animal variation in gonad cell proportions is to be expected, even among fish of the control groups, as a consequence of spawning cycle asynchrony. Consequently, following the gonadal staging of individual fish, each treatment group is assessed as a whole and compared to the appropriate control group to determine if a compound-related effect has occurred. Hence, gonadal staging cannot be performed in a blinded manner. Since the cell distribution pattern is likely to vary throughout a given tissue section, the gonad should be staged according to the predominant pattern in that section. Similarly, both gonads should be staged as a single organ according to the predominant pattern. Gonads that cannot be reasonably staged for various reasons (e.g., insufficient tissue, or extensive necrosis, inflammation, or artifact) should be recorded as UTS (unable to stage).

3.2.3 Secondary criteria in the gonads of male fathead minnow, Japanese medaka and zebrafish

The following criteria have been defined in male fish as diagnoses of secondary interest:

1. Decreased proportion of spermatogonia: Decreased relative proportion of spermatogonia to other spermatogenic cell types. This can be due to a decrease in the number of spermatogonia, or an increase in the number of other cell types, such as spermatocytes, spermatids, and spermatozoa. Because the diagnosis of decreased proportion of spermatogonia is dependent on a comparison to controls, it is necessary to establish the normal range of the ratios of spermatogenic cells in control male fish testes prior to making determinations on relative proportions in dose groups.

2. Increased vascular or interstitial proteinaceous fluid: Homogenous dark pink translucent material, presumably vitellogenin, within the testicular interstitium or blood vessels. The presence of this fluid may cause a thickening of interstitial areas that might be misinterpreted as “stromal proliferation”.

3. Asynchronous gonad development: The presence of more than one developmental phase of spermatogenic cells within a single spermatocyst enclosed by a Sertoli cell. For example, this term may be applied to a spermatocyst that contains a mixture of spermatocytes and spermatids, or a spermatocyst that contains more than one meiotic phase of primary spermatocytes (i.e., leptotene, pachytene, and/or zygotene). It also refers to the presence of distinctly different populations (i.e. developmental phases) of gametogenic cells in the right and left gonads.

4. Altered proportions of spermatocytes or spermatids: A change in the relative proportions of spermatocytes or spermatids to other spermatogenic cell types. Changes in relative ratios could be due to an increase in the number of spermatocytes or spermatids, or to a decrease in the number of other cell types. Relative changes may also occur between spermatocytes and spermatids.

5. Gonadal staging: Gonadal staging results are virtually meaningless in terms of individual fish (versus treatment groups). This is because considerable animal-to-animal variation in gonad cell proportions is to be expected, even among fish of the control groups, as a consequence of spawning cycle asynchrony. Consequently, following the gonadal staging of individual fish, each treatment group is assessed as a whole and compared to the appropriate control group to determine if a compound-related effect has occurred. Hence, gonadal staging cannot be performed in a blinded manner. Because the cell distribution pattern is likely to vary throughout a given tissue section, the gonad should be staged according to the predominant pattern in that section. Similarly, both gonads should be staged as a single organ according to the predominant pattern. Gonads that cannot be reasonably staged for
various reasons (e.g., insufficient tissue, or extensive necrosis, inflammation, or artifact) should be recorded as UTS (unable to stage).

(6) **Granulomatous inflammation:** This process is characterized by the presence of epithelioid macrophages that typically form sheets or nodules (granulomas) due to desmosome-like cytoplasmic attachments (Noga et al., 1989). When compared to histiocytic-type macrophages, epithelioid macrophages have larger, more open-faced, centralized nuclei and less abundant cytoplasm. During resolution of inflammation, the epithelioid macrophages may become flattened into fibrocyte-like cells. Lymphocytes, granulocytes, and multinucleated giant cells may also be components of granulomatous inflammation. Granulomatous inflammation is intrinsically a pathologic process that is often associated with reactions to infectious agents, foreign materials or the aftermath of necrosis; therefore, it is important to distinguish this, if possible, from the presence of histiocytic cells in the lumen of the testis.

3.2.4 Secondary criteria in the gonads of female fathead minnow, Japanese medaka and zebrafish

The following criteria have been defined in female fish as diagnoses of secondary interest:

1. **Interstitial fibrosis:** The presence of increased fibrous connective tissue (collagenous fibers and fibrocytes or fibroblasts) within the ovarian interstitium (stroma). Collagen may be difficult to appreciate in early phases of fibrosis.

2. **Egg debris in the oviduct:** The presence of inspissated-appearing, homogenous, irregular, dense pink material, presumed to be yolk, within the oviduct.

3. **Granulomatous inflammation:** This process is characterized by the presence of epithelioid macrophages that typically form sheets or nodules (granulomas) due to desmosome-like cytoplasmic attachments (Noga et al., 1989). When compared to histiocytic-type macrophages, epithelioid macrophages have larger, more open-faced, centralized nuclei and less abundant cytoplasm. During resolution of inflammation, the epithelioid macrophages may become flattened into fibrocyte-like cells. Lymphocytes, granulocytes, and multinucleated giant cells may also be components of granulomatous inflammation. Granulomatous inflammation is intrinsically a pathologic process that is often associated with reactions to infectious agents, foreign materials or the aftermath of necrosis; therefore, it is important to distinguish this, if possible, from the presence of macrophage aggregates in the ovary.

4. **Decreased post-ovulatory follicles:** A decrease in the number of collapsed perifollicular sheaths, or membranous structures lined by granulosa cells, theca cells and surface epithelium, following release of oocytes, in comparison to control fish. The granulosa cells are often hypertrophic, although this appears to be species dependent (Saidapur, 1982).

3.3 Severity grading

In toxicologic pathology, it is recognized that compounds may exert subtle effects on tissues that are not adequately represented by simple binary (positive or negative) responses. Severity grading involves a semi-quantitative estimation of the degree to which a particular histomorphological change is present in a tissue section (Shackelford et al., 2002). The purpose of severity grading is to provide an efficient, semi-objective mechanism for comparing changes (including potential compound-related effects) among animals, treatment groups, and studies.
Severity grading will employ the following system:

- **Not remarkable**
- **Grade 1 (minimal)**
- **Grade 2 (mild)**
- **Grade 3 (moderate)**
- **Grade 4 (severe)**

A grading system needs to be flexible enough to encompass a variety of different tissue changes. In theory, there are three broad categories of changes based on the intuitive manner in which people tend to quantify observations in tissue sections:

1. **Discrete**: these are changes that can be readily counted. Examples include atretic follicles, oocytes in the testis, and clusters of apoptotic cells.
2. **Spatial**: these are changes that can be quantified by area measurements. This includes lesions that are typically classified as focal, multifocal, coalescing, or diffuse. Specific examples include granulomatous inflammation and tissue necrosis.
3. **Global**: these are generalized changes that would usually require more sophisticated measurement techniques for quantification. Examples include increased hepatocyte basophilia, Sertoli cell/interstitial cell hypertrophy, or quantitative alterations in cell populations.

### General severity grading scale

- **Not Remarkable**: This grade is used if there are no findings associated with a particular diagnostic criterion.
- **Grade 1: Minimal**: Ranging from inconspicuous to barely noticeable but so minor, small, or infrequent as to warrant no more than the least assignable grade. For discrete changes, grade 1 is used when there are fewer than 2 occurrences per microscopic field, or 1 - 2 occurrences per section. For multifocal or diffusely-distributed alterations, this grade is used for processes where ≤ 20% of the tissue in the section is involved.
- **Grade 2: Mild**: A noticeable feature of the tissue. For discrete changes, grade 2 is used when there are 3 - 5 occurrences per microscopic field or per tissue section. For multifocal or diffusely-distributed alterations, this grade is used for processes where 20 - 50% of the tissue in the section are involved.
- **Grade 3: Moderate**: A dominant feature of the tissue. For discrete changes, grade 3 is used when there are 6 - 8 occurrences per microscopic field or per tissue section. For multifocal or diffusely-distributed alterations, this grade is used for processes where 50 - 80% of the tissue in the section are involved.
- **Grade 4: Severe**: An overwhelming feature of the tissue. For discrete changes, grade 4 is used when there are more than 9 occurrences per microscopic field or per tissue section. For multifocal or diffusely-distributed alterations, this grade is used for processes where > 80% of the tissue in the section are involved.

At least some of the histomorphological changes that have been associated with EDCs in fish are considered to be exacerbations of “normal”, physiologic findings (e.g., oocyte atresia [Nagahama, 1983; Tyler and Sumpter, 1996]). At the discretion of the pathologist, the severity of a given change should be scored according to one of the following two methods:

1. score compound-exposed animals relative to the severity of the same change in control animals, or
2. score all animals relative to “normal” as determined by the pathologist’s experience.
For each important (i.e., treatment-associated) finding, the method that was used should be stated in the Materials and Methods section of the pathology narrative report (see section 3.5: Histopathology report format). By convention, severity grading should not be influenced by the estimated physiological importance of the change, since this would add a further layer of subjectivity to the findings that complicates inter-laboratory results comparisons. For example, the presence of two oocytes in the testis should not be graded as “severe”, even if the pathologist considers this finding to be highly significant in terms of endocrine modulation.

3.4 Data recording

An Excel worksheet form has been created that includes worksheets for primary, secondary, and additional diagnoses to facilitate histopathology data collection. In this worksheet, each data entry cell represents an individual fish. Additional sheets are available for comments and additional findings. For each fish, the pathologist records a severity score associated with the diagnosis (see section 3: Severity Grading). Diagnostic criteria with non-remarkable findings shall be denoted using (−). If there is no reasonably appropriate diagnostic term for a particular finding, the pathologist can create a term that can be recorded in the “Additional diagnoses” worksheet. If insufficient tissue is available for diagnosis, this should be recorded as IT (insufficient tissue). If a target tissue is missing, this should be recorded as MT (missing tissue).

Adding a modifier term to a diagnosis may help to further describe or categorize a finding in terms of chronicity, spatial distribution, color, etc. In many instances, modifiers are superfluous or redundant (e.g., fibrosis is always chronic); therefore, the use of modifiers should be kept to a minimum. An occasionally important modifier for evaluating paired gonads is unilateral (UNI); unless specified in this manner, all gonad diagnoses are assumed to be bilateral. Other modifier codes can be created as needed by the pathologist.

3.5 Histopathology report format

Each histopathology narrative report should contain the following five sections:

- Introduction
- Materials and Methods
- Results
- Discussion
- Summary/Conclusions
  (References)

The Introduction section briefly outlines the experimental design. The Materials and Methods section describes any items or procedures that are essentially different from Section 1: Post-mortem and Histotechnical Procedures. As applicable, specific severity grading criteria (see Severity Grading) should also be listed in this section. The Results section should report findings that are: (1) compound-related; (2) potentially compound-related; (3) novel or unusual. Detailed histomorphological descriptions need only be included for findings that differ substantially from diagnoses presented in Section 4 (Glossary and diagnostic criteria). It is intended that the Results section should be as objective as possible (i.e., opinions and theories should be reserved for the Discussion section). The Discussion section, which contains subjective information, should address relevant findings that were reported in the Results section. Opinions and theories can be included in this section, preferably backed by references from peer-reviewed sources, but unsupported speculation should be avoided. The Summary/Conclusions section should encapsulate the most important information from the Results and Discussion sections.
4. Glossary and diagnostic criteria

The purposes of this section are:

(1) to provide photomicrographs of normal gonadal structure in fathead minnow, medaka and zebrafish,

(2) to provide a common technical “language” and

(3) to create a reference atlas of both microanatomical structures and potential pathological findings.

The information in this section is derived from a number of sources including scientific articles, conference proceedings, related guidelines, toxicologic pathology textbooks, medical dictionaries, and the personal experience of various fish pathologists. Regarding the last, opinions were solicited via a questionnaire that was circulated among conference participants following the October 2003 meeting of the histopathology subcommittee of the Fish Discussion Group in Paris. Consensus replies to this questionnaire form the basis for naming many of these terms. Additional guidance was provided by pathologists attending the 2nd OECD meeting of the Fish Pathologist Subcommittee at the University of Heidelberg, Germany, November 22 - 23, 2004. Other considerations include traditional usage and scientific precedence, and attributes such as clarity and brevity.

In addition to illustrating standard microanatomical features of Japanese medaka, fathead minnow, and zebrafish gonads, this document presents an array of histopathological diagnoses and their associated morphologic criteria. Whereas a few of these diagnoses have been consistently linked to endocrine disruptor exposure under certain specified laboratory conditions, many other findings are less well-established as EDC markers. It is also important to recognize that the utility of a given diagnosis for detecting EDC effects may depend greatly on the species and age of the test subjects, the effective dose of the test compound, and other factors such as husbandry practices. For example, juvenile and adult Japanese medaka seem to be relatively more susceptible to estrogen-induced testis-ova formation than either fathead minnow or zebrafish; however, another excellent response to estrogenic endocrine disruptors that has been primarily reported in juvenile fathead minnow and zebrafish is retained peritoneal attachments/gonadal duct feminization of the testis. On the other hand, while preliminary evidence indicates that decreased yolk formation may be a reliable marker for aromatase inhibitors and non-aromatizable androgens in reproductively active adult female Japanese medaka and zebrafish, this effect seems to be less robust in similarly-treated female fathead minnow. Conversely, adult male fathead minnow appear to be particularly well-suited for observing interstitial cell hyperplasia/hypertrophy of the testis as a reaction to the steroidogenesis inhibitor ketoconazole. One should also keep in mind that endocrine disruptors may induce different effects at high versus low exposure concentrations.

Chapter 4 is arranged as follows:

4.1 Normal testicular architecture
4.2 Primary diagnoses in males and females
4.3 Secondary diagnoses in males and females
4.4 Additional diagnostic criteria
4.1 Normal gonadal architecture in fathead minnow, Japanese medaka and zebrafish

4.1.1 Normal testicular architecture in male fathead minnow, Japanese medaka and zebrafish

Spermatogonia: The largest of the spermatogenic cells (~5 - 10 µm), spermatogonia generally have pale vesicular nuclei, prominent nucleoli, variably distinct nuclear membranes, perinuclear cytoplasmic granules, and moderate amounts of granular cytoplasm. Spermatogonia B are smaller than spermatogonia A, and spermatogonia B are usually present in larger clusters (e.g., > 4 cells). If at all possible, an attempt should be made to classify these cells as spermatogonia rather than to label them with a non-specific term such as “pale cells” or “light cells”.

Fig. 4. Spermatogonia A in male fathead minnow (a; GMA, H&E), Japanese medaka (b; paraffin, H&E) and zebrafish (c; paraffin, H&E)
**Spermatocytes**: Derived from spermatogonia, spermatocytes are of intermediate cell size (~ 4 - 6 µm), and have comparatively dense nuclei and minimal to moderate amounts of indistinct cytoplasm. Spermatocyte nuclei are usually evident in one of three meiosis phases: pachytene, leptotene, or zygotene. Primary spermatocytes are larger than secondary spermatocytes, and the latter are derived from primary spermatocytes following the first meiotic division. Spermatocytes are usually one of the most abundant spermatogenic cells, and they tend to contribute to the largest spermatocysts.

Fig. 5. Spermatocytes A: (a) male fathead minnow (GMA, H&E); (b) male Japanese medaka (paraffin, H&E), and (c) in the leptotene, zygotene, and pachytene phases, respectively, during the first meiosis in the zebrafish testis (paraffin, H&E).
**Spermatids**: Derived from spermatocytes following the second meiotic division, these cells have dense nuclei and narrow rims of eosinophilic cytoplasm. They are the smallest cells within the germinal epithelium (~2 - 3 μm), and the cells lose their cytoplasmic attachments to one another during spermogenesis.

Fig. 6. (a) Spermatids in male fathead minnow: Intercellular attachments are lost just prior to rupture of the spermatocyst and release of these cells as spermatozoa (GMA, H&E). (b) Spermatids in a Japanese medaka testis (paraffin, H&E). (c) Spermatids in a zebrafish (paraffin, H&E).
**Spermatozoa:** These cells have dark, round nuclei and minimal or no apparent cytoplasm. Tails are generally not apparent in histological sections. Spermatozoa are the smallest spermatogenic cells (~2 µm), and they exist as scattered individual cells within tubular lumen.

Fig. 7. (a) Spermatozoa in male fathead minnow (GMA, H&E). (b) Spermatozoa in a Japanese medaka testis (paraffin, H&E). (c) Spermatozoa in a zebrafish (paraffin, H&E).
Sertoli cells: These cells tend to have sharply-defined elongated or triangular nuclei, variably evident nucleoli, and cytoplasm that is often indistinct. The cytoplasmic arms of a Sertoli cell encircle a clonal group of spermatogenic cells, forming a spermatocyst. Compared to germinal cells, Sertoli cells are usually present in low numbers, usually as single cells located adjacent to lobular septa. In some instances, hypertrophic (enlarged, swollen) Sertoli cells may resemble spermatogonia.

Fig. 8. In all three species, Sertoli cells are flattened. (a) Fathead minnow testis (GMA, H&E), (b) Japanese medaka testis (paraffin, H&E), and (c) zebrafish testis (paraffin, H&E).
Interstitial (Leydig) cells: These cells have dense, dark round or oval nuclei with little detail and moderate amounts of variably-evident, faintly vacuolated cytoplasm. Compared to germinal cells, interstitial cells are usually present in low numbers, usually as single cells or small aggregates, within the interlobular interstitium. Although they may resemble spermatocytes, interstitial cells are only present in interlobular areas.

Spermatocyst: The functional unit of the testis, this structure consists of a clonal group of spermatogenic cells (spermatogonia, spermatocytes, or spermatids) that are surrounded by the cytoplasmic arms of (usually) one Sertoli cell. Cells within spermatocysts exist as syncytia, maintained by intercellular attachments (cytoplasmic bridges), until final maturation and release of spermatozoa occur (spermiogenesis; Grier, 1976).

Fig. 9. Interstitial cells (→) in male fathead minnow are only found in interlobular areas. Note the resemblance between these cells and spermatocytes (►) (GMA, H&E).

Fig. 10. (a) Spermatocyst in male fathead minnow: A group of dissociated spermatids are surrounded by the cytoplasmic “arms” of a single Sertoli cell (→). This arrangement is usually not as obvious as it is in this photograph. The nucleus of this particular Sertoli cell appears enlarged (hypertrophic; GMA, H&E). (b) Testis from adult male Japanese medaka: Each packet of cells (spermatocyst) represents a cohort of germ cells in approximately the same developmental phase (paraffin, H&E, bar = 10 µm).
Fig. 11. Spermatocysts in adult male fathead minnow are outlined in red and green contain spermatocytes and spermatids, respectively. Spermatogonia (black arrows) and spermatozoa within tubular lumina (blue arrows) are also indicated (GMA, H&E)

**Male germinal epithelium:** The germinative intratubular (intralobular) parenchyma of the testis, this membrane-bound structure consists of multiple spermatocysts in various phases of development. For FHM, boundaries of the germinal epithelium at various locations throughout the testis include the interlobular interstitium, the lobular lumina, collecting ducts, and the tunica albuginea.

Fig. 12. Male germinal epithelium in the testis of normal fathead minnow: Double arrow indicates width of germinal epithelium, which extends from the interlobular interstitium to the lobular lumen (GMA, H&E, bar = 25 µm).
4.1.2 Normal ovarian architecture in female fathead minnow, Japanese medaka and zebrafish

**Oogonia**: These cells represent the replicative pool of the ovary. Unlike mammalian oogonia (although this dogma may soon change based on recent data from rodent research), piscine oogonia continue to divide in juveniles and adults. The smallest of the oocytic cells, oogonia reside within the ovarian germinal epithelium, usually in comparatively low numbers. Oogonia are characterized by a relatively large nucleus with small or inapparent nucleolus, and minimal amounts of cytoplasm.

Fig. 13. Oogonia in fathead minnow: A small cluster of oogonia reside within a portion of germinal epithelium; the nucleus of only one oogonium is visible (small arrow). The oogonia are dwarfed by a perinucleolar oocyte (*; paraffin, H&E, bar = 10 µm).

**Chromatin nucleolar oocytes**: Slightly larger than an oogonium, this oocyte is formed when an oogonium becomes surrounded by prefollicle cells (presumptive granulosa cells), and the resulting complex buds from the germinal nest as a primordial follicle. The chromatin nucleolar oocyte has a relatively large nucleus that contains a single, large nucleolus. Compared to an oogonium, there is more cytoplasm which is slightly denser and finely granular.

Fig. 14. Chromatin nucleolar oocyte in fathead minnow: A single chromatin nucleolar oocyte protrudes from the germinal epithelium (paraffin, H&E, bar = 10 µm).
Perinucleolar oocytes: Concomitant with oocyte growth, the nucleus (germinal vesicle) increases in size and multiple nucleoli appear, generally at the periphery of the nucleus. The cytoplasm stains uniformly dark, although late perinucleolar oocytes may have small clear or amphophilic vacuoles in the cytoplasm. These cells tend to be abundant in normal adult ovaries.

Fig. 15. (a) Perinucleolar oocytes in fathead minnow: Arrows (→) indicate nucleoli at the periphery of the germinal vesicle (paraffin, H&E, bar = 10 µm). (b) Numerous perinucleolar oocytes in a Japanese medaka ovary appear virtually identical to perinucleolar oocytes of fathead minnow (cf. Fig. a; paraffin, H&E). (c) Perinucleolar oocytes in a zebrafish ovary (paraffin, H&E, original mag.: 20 ×).
Cortical alveolar oocytes: Generally larger than perinucleolar oocytes, cortical alveolar oocytes are characterized by the appearance of cortical alveoli (yolk vesicles) within the ooplasm. The cortical alveoli are technically not yolk, as they do not provide nourishment for the embryo (Selman and Wallace, 1989). The chorion becomes distinctly evident in this phase, and the perifollicular cells are more easily visualized.

Fig. 16. (a) Cortical alveolar oocytes in fathead minnow ovary: The cytoplasm is predominately filled by numerous cortical alveoli, which are amphiphilic within this preparation. Evident are oocytes in transition from the perinucleolar to cortical alveolar phase (→), and from the cortical alveolar to early vitellogenic phase (▶; paraffin, H&E, bar = 100 µm). (b) Japanese medaka ovary with a single large cortical alveolar oocyte adjacent to several smaller perinucleolar oocytes. Compared to fathead minnow, the cortical alveoli are fewer, larger, and are relatively clear in this preparation (paraffin, H&E). (c) Cortical alveolar oocyte in a zebrafish ovary (paraffin, H&E, original mag. 40 ×).
**Early vitellogenic oocytes:** Larger than cortical alveolar oocytes, these cells are characterized by the centralized appearance of spherical, eosinophilic, vitellogenic yolk granules/globules. In H&E sections, accumulations of fine yolk granules in the central region of the oocyte may somewhat resemble (and thus be confused with) the reddish nucleus.

Fig. 17. Early vitellogenic oocytes in fathead minnow ovary: Numerous fine pale pink granules (►) and a few larger dark red granules (→) are evident in the central region of an early vitellogenic oocyte. Although nuclei are present they are not apparent in every oocyte due to the comparatively vast amount of cytoplasm (paraffin, H&E, bar = 100 µm). (b) A few eosinophilic vitellogenic granules (arrow) are present in the ooplasm of this oocyte in a Japanese medaka ovary (paraffin, H&E).

**Late vitellogenic oocytes:** These cells are characterized by an increased accumulation of vitellogenic granules that displace the cortical alveolar material to the periphery of the cytoplasm. It is during this stage that the nucleus begins to migrate toward the periphery of the cell.

Fig. 18. (a) Late vitellogenic oocytes in fathead minnow: The yolk granules almost fill the ooplasm. The nucleus has not yet begun to migrate to the periphery (paraffin, H&E, bar = 100 µm). (b) Late vitellogenic oocyte in Japanese medaka: Yolk granules are fused into a central liquid yolk mass (paraffin, H&E). (c) Late vitellogenic oocytes in zebrafish close to the mature/spawning phase (paraffin, H&E, original mag. 10 ×).
**Mature/spawning oocytes:** In this phase of development, vitellogenesis has reached its peak, the cell has become larger and more hydrated, and the nucleus has migrated toward the periphery of the cell and is in the process of dissolution. The loss of nucleus is not a very helpful diagnostic feature, however, as the nucleus is often not visible in larger oocytes due to the plane sectioning. Because of the transient nature of these cells in fractional spawning fish, mature/spawning oocytes are uncommonly observed.

Fig. 19. (a) Mature/spawning oocytes in fathead minnow ovary: The oocytes and the yolk granules have attained their maximum size just prior to spawning, and the nucleus is not evident (paraffin, H&E, bar = 100 µm). (b) The yolk mass nearly fills the cytoplasm of the mature/spawning oocyte in this Japanese medaka ovary (paraffin, H&E). (c) Two large mature/spawning oocytes in a zebrafish ovary. The yolk granules have congealed into large globules (paraffin, H&E).
**Ovarian follicle:** The functional unit of the ovary, this term generally refers to an oocyte plus its surrounding sheath of perifollicular cells (granulosa cells, theca cells, and surface epithelium cells; Tyler and Sumpter, 1996). However, there are subtypes of follicles in which the oocyte is not present or may be difficult to appreciate; these include post-ovulatory (spent), empty, and atretic follicles. A *post-ovulatory follicle* (the follicle has ruptured to release an oocyte during spawning) is collapsed and often has enlarged (hypertrophic) granulosa and theca cells. Conversely, an empty follicle (in which the oocyte has been dislodged from the histological section as a post-mortem artifact) generally retains the shape of the oocyte and may or may not have enlarged granulosa and theca cells. An *atretic follicle* must be distinguished from both spent follicles and empty follicles; the presence of at least some ooplasmic material (often heterochromatic) within a follicle indicates that it contains an atretic oocyte (see also: oocyte atresia).

Fig. 20. Diagram of an ovarian follicle (from Tyler and Sumpter, 1996).

**Perifollicular cells:** These cells form a three-layered sheath around each oocyte, and combined with the oocyte itself comprise the ovarian follicle. These layers are more easily visualized as the oocyte matures. The innermost layer consists of the granulosa cells, the middle layer consists of the theca cells, and the outermost layer consists of the surface epithelial cells. The granulosa cells especially may become enlarged and vacuolated following ovulation or during oocyte atresia. The perifollicular sheath should not be confused with folds of the ovarian wall epithelium.

Fig. 21. Perifollicular cells in adult fathead minnow ovary, as compared to the cells of the ovarian wall epithelium, which contain dark brown (melanin) pigment (→) and are comprised of ciliated columnar cells (paraffin, H&E, bar = 25 µm).
Chorion: The chorion usually stains pale to dark eosinophilic and refractile, the chorion is the thick external layer of an oocyte that surrounds the ooplasm. The terms zona radiata and vitelline envelope have been used synonymously. In mature, unspent follicles, the chorion is noticeably surrounded by perifollicular cells (granulosa cells, theca cells, and surface epithelial cells). As viewed by light microscope, the chorion is often minimally apparent or inapparent prior to the cortical alveolar phase of oocyte development.

Fig. 22. The chorions (►) of two oocytes in an adult fathead minnow ovary: A smaller arrow (→) denotes a post-ovulatatory follicle (paraffin, H&E, bar = 25 µm).

Fig. 23. Chorion in the Japanese medaka ovary: Note the vast difference in thickness between the chorion of a cortical alveolar oocyte (→) and the chorion of a mature vitellogenic oocyte (►; paraffin, H&E, bar = 25 µm).

Fig. 24. Chorion in the zebrafish ovary: Three vitellogenic oocytes (V) have smooth, intact chorions. In contrast, this “membrane” is just beginning to form around a perinucleolar oocyte (P), and the chorion surrounding an atretic oocyte (A) has become thickened and fragmented (paraffin, H&E, bar = 25 µm).
**Post-ovulatory follicle:** A collapsed perifollicular sheath following release of the oocyte; this is a membranous structure lined by granulosa cells, theca cells, and surface epithelium. The granulosa cells are often hypertrophic, although this appears to be species dependent (Saidapur, 1982). Mammalian terms such as “corpus lutea” and “Graafian follicles”, are probably less desirable, due to structural and functional differences between these entities and piscine post-ovulatory follicles. Whenever possible, post-ovulatory follicles should be differentiated from atretic follicles, the latter of which contains oocyte debris.

Fig. 25. Post-ovulatory follicle in an adult fathead minnow ovary: Situated between three oocyte-containing follicles is a collapsed follicle, which does not contain oocyte remnants (→; paraffin, H&E, bar = 25 µm). (b) Post-ovulatory follicle in an adult Japanese medaka ovary: A number of post-ovulatory follicles (→), indicating recent spawning, are evident in this ovary (paraffin, H&E, bar = 250 µm). (c) Ovary from an adult female fathead minnow. These oocytes (→) are actually *not post-ovulatory follicles*. The presence of yolk material indicates that they are late phase atretic follicles. (d) Ovary from an adult female zebrafish: The solid arrow indicates a post-ovulatory follicle, whereas the open arrow points to a late phase atretic follicle (paraffin, H&E).
4.2 Primary diagnoses in fathead minnow, Japanese medaka and zebrafish following exposure to endocrine disruptors

4.2.1 Primary diagnoses in male fathead minnow, Japanese medaka and zebrafish following exposure to endocrine disruptors

**Increased proportion of spermatogonia in male fathead minnow**

*Increased proportion of spermatogonia:* It is recognized that endocrine active compounds may alter the proportional distribution of gametogenic cell types in the testis (or ovary). Certain types of alterations (for example, the proliferation or absence of single cell population) may not be adequately documented by gonadal staging. This diagnostic term provides a mechanism for documenting such changes.

Quantitative alterations are:

1. relative to other cell types in the gonad;
2. relative to cell numbers in control animals; and
3. estimates only, versus actual cell counts.

A proportional increase in spermatogonia was observed consistently in the testes of fathead minnow and zebrafish (and less dependably in Japanese medaka) as an exposure response to, e.g., the estrogenic compound 4-tert-pentylphenol. Other experiments have also linked exposure to estrogens, or substances with estrogenic activity, to increases in spermatogonia (Condeça and Canario, 1999; Sohoni et al., 2001; Wester et al., 2003). Proportional increases in spermatogonia are often associated with increases in the thickness of the testicular germinal epithelium, but this is not always the case. Testes exhibiting Grade 4 severity for this change may be difficult, if not impossible, to distinguish from juvenile testes, and in truth, the distinction may not be important. Other potential ruleouts for gonads that contain large numbers of immature gonial cells would include germ cell neoplasms such as seminomas and dysgerminomas.

---

Fig. 26. Increased cells (spermatogonia) in fathead minnow testis: (a) Testis from adult male FHM negative control. (b) Spermatogonia dominate the germinal epithelium in this testis from adult male FHM exposed to 10 nM 17β-estradiol for 10 days. Other diagnoses for this section include “decreased cells, spermatocytes” and “decreased cells, spermatids” (GMA, H&E, bars = 25 µm).
Fig. 27. Testis from an adult male fathead minnow: There is a minimal (Grade 1) increase in the proportion of spermatogonia (→; GMA, H&E).

Fig. 28. Testis from an adult male fathead minnow: There is a slight/mild (Grade 2) increase in the proportion of spermatogonia throughout the germinal epithelium (→; GMA, H&E).

Fig. 29. Testis from an adult male fathead minnow: There is a moderate (Grade 3) increase in the proportion of spermatogonia (→; GMA, H&E).

Fig. 30. Testis from an adult male fathead minnow: There is a severe (Grade 4) increase in the proportion of spermatogonia (GMA, H&E).
Fig. 31. Spermatogonia, increased. (a) Testis of an adult control fathead minnow. (b) Testis of an adult male fathead minnow exposed to 1000 µg/L flutamide. In this minimally affected testis, spermatogonia dominate the germinal epithelium, but many spermatocytes are evident also. (c) Testis of an adult male fathead minnow exposed to 100 ng/L estradiol. Multiple layers of spermatogonia surround seminiferous tubules in this mildly affected testis. (d) Moderately affected testis. (e) Severely affected testis: Due to their longevity, spermatozoa may be abundant despite the lack of intermediate germ cell phases (i.e., spermatocytes and spermatids; paraffin, H&E, bars = 25 µm)
Fig. 32. Spermatogonia, increased: A proportional increase in spermatogonia has been observed in the testes of male zebrafish exposed to 17ß-estradiol. (a) Testis from adult male zebrafish (negative control). (b) Testis from adult male zebrafish exposed to 17ß-estradiol for 3 weeks. Again, spermatogonia are present in greater numbers compared to control fish (paraffin, H&E)
Presence of testis-ova in male fish

The presence of testis-ova defines a state in which fully formed male and female gonad tissues are present in the same individual. The phrase “fully formed” indicates that (1) the male and female gonadal tissues are in discrete compartments; (2) the organizational architecture of the gonads is maintained; and/or (3) there is visible evidence of supportive structures (e.g., tunica albuginea, ducts) in addition to germinal cells. The testicular and ovarian tissues may be present within confines of the same gonad (ovotestis) as defined by the tunica albuginea, or they may exist as completely separate organs (e.g., left and right, rostral and caudal).

Fig. 33. Hermaphroditism in young adult fathead minnow: This very rare finding occurred spontaneously in an untreated fish. The small arrows (→) indicate perinucleolar stage oocytes. This case should not be diagnosed merely as “testicular oocytes”, because this gonad has additional ovarian features, such as the presence of an ovarian cavity (OV) that is lined by an ovarian wall-type epithelium (open arrow). As further evidence for hermaphroditism, the contralateral gonad (not shown) was a normally appearing ovary (whole-body cross section, paraffin, H&E, bar = 500 µm).

Fig. 34. Hermaphroditism in young adult fathead minnow: example of spontaneous testicular oocytes. Whether spontaneous or induced, testicular oocytes are a rare finding in fathead minnow (paraffin, H&E, bar = 100 µm).
Fig. 35. Testis-ova in adult male Japanese medaka: Testicular oocytes (paraffin, H&E). Photomicrographs from four different negative control fish from several studies. The large arrows indicate perinucleolar oocytes, whereas the small arrows are chromatin nucleolar oocytes (paraffin, H&E).

Fig. 36. Testicular oocytes in adult male Japanese medaka: Red arrows indicate oocytes in the testis of this fish treated with 17ß-estradiol (paraffin, H&E, bar = 30 μm).
Fig. 37. Examples of testis-ova severity grading in male Japanese medaka: It is important to note that the severity grade can vary dramatically from section to section within the same testis; therefore, the value of severity grading becomes questionable when it is based on the evaluation of only one or two sections per fish. (a) A minimally affected testis with a single perinucleolar oocyte (→) in one section. (b) Mildly affected fish with a number of scattered, individual perinucleolar oocytes (→). (c) In this moderately affected testis the oocytes occur within large lobular nests that also contain oogonia. (d) Small remnants of spermatogenic tissue and the bilobed configuration of this gonad are evidence that this is a testis rather than an ovary. Another potential ruleout for this finding would be dysgerminoma. This is an 8 weeks old genetically male fish that was exposed to 100 ng/L 17β-estradiol for approximately eight weeks (all images are paraffin, H&E, bar = 100 µm).
Fig. 38. Hermaphroditism in zebrafish: In these sections from two different adult fish, each section contains fully-formed male and female gonad tissues. According to a convention established herein, (see Gender), the fish in image “a” would be designated as a male, whereas the fish in “b” would be designated as a female. (whole-body cross-section, H&E; bar = 500 µm).

Fig. 39. Testicular oocytes in adult male zebrafish: Red arrows indicate oocytes. Note that the fish on the right is a control animal. The testis in the lower image would be considered mild (Grade 2) severity. (paraffin, H&E).
**Increased testicular degeneration in male fathead minnow**

**Testicular degeneration**: Examples of degenerative findings in the testis include:

1. Individual or clustered apoptotic germ cells;
2. Vacuolated germ cells;
3. Multinucleated (syncytial) cells in the germinal epithelium or testicular lumen.

These diagnoses may be “lumped” together under the term testicular degeneration. Apoptotic germ cells are characterized by cell shrinkage, nuclear condensation, and fragmentation into spherical, membrane-bound bodies, which are often phagocytized by neighboring cells. There is no inflammation associated with these cells. If possible, testicular degeneration should be differentiated from necrosis, which is characterized morphologically by cytoplasmic coagulation or swelling, nuclear karyorrhexis or pyknosis, associated inflammation, a locally extensive pattern of tissue involvement, and/or the involvement of different local tissue elements (e.g., both germinal and stromal tissues). Extensive testicular degeneration may lead to localized or generalized loss of the germinal epithelium.

![Fig. 40. Testicular degeneration in adult male fathead minnow: (a) Multiple clusters of apoptotic germ cells (→) and vacuolated germ cells (red →) within the germinal epithelium. (b) Another mildly-affected testis with three large germ cell syncytia (→; GMA, H&E, bars: 25 µm).](image-url)
Fig. 41. Increased testicular degeneration in adult male fathead minnow: There are low numbers of germ cell syncytia (→) in this minimally affected testis. This degree of degeneration is occasionally seen in control males; therefore, this diagnosis should be made relative to the degree of degeneration evident among the majority of the concurrent controls (paraffin, H&E, bar = 100 µm).

Fig. 42. Increased testicular degeneration in adult male Japanese medaka: (a) In most studies, an occasional degenerating cell, like this multinucleated germ cell (arrow), would be below the threshold for “increased” and therefore would be considered non-remarkable. (b) There are several clusters of apoptotic cells (arrows) in a relatively small area of this mildly affected testis (paraffin, H&E, bar = 25 µm).
Interstitial cell (Leydig cell) hyperplasia/hypertrophy in male fish

Fig. 43. Testis from an adult male fathead minnow: Interstitial areas contain small aggregates of interstitial (Leydig) cells (→). Most interstitial cells have wispy, pale cytoplasm (GMA, H&E).

Fig. 44. Testis from an adult male fathead minnow: Interstitial cell aggregates (→) in the testis of this fish are larger than in control fish, and the cytoplasm of these cells is slightly denser. This was diagnosed as “Increased Cells, Interstitial Cells”, Grade 1 (minimal) severity (GMA, H&E).

Fig. 45. Testis from an adult male fathead minnow: Interstitial cell aggregates (→) in the testis of this fish are larger than in control fish, and the cells tend to fill and expand the interstitial spaces. This was diagnosed as “Increased Cells, Interstitial Cells”, Grade 2 (mild) severity (GMA, H&E).
Fig. 46. Interstitial (Leydig) cell hyperplasia in adult male fathead minnow: (a) Unaffected testis from control fathead minnow. Arrowheads (►) indicate occasional interstitial cells as single cells or small clusters. (b) Testis with a minimal increase in the number of interstitial cells. Interstitial cells are present in small to medium-sized aggregates (►). Note that the cell nuclei are small and condensed, and the cytoplasm of these cells is clear. (c) Testis with mild interstitial cell (IC) hyperplasia. (d) Testis with moderate interstitial cell hyperplasia/hypertrophy (►) associated with ketoconazole exposure (unpublished data). There are relatively large aggregates of interstitial cells, and these cells have rounded nuclei and dense cytoplasm; paraffin, H&E, bar = 25 µm).
4.2.2 Primary diagnoses in female fathead minnow, Japanese medaka and zebrafish following exposure to endocrine disruptors

Increased oocyte atresia

Oocyte atresia, increased, immature/mature: Degradation and resorption of an oocyte at any point in development. Histopathologically, atresia is often characterized by clumping and perforation of the chorion, fragmentation of the nucleus, disorganization of the ooplasm, and/or the uptake of yolk materials by perifollicular cells. Separate diagnoses and severity grades can be given to atretic oocytes that are mature (“oocyte atresia, increased, mature”) versus immature (“oocyte atresia, increased, immature”). In this context, oocytes will be considered “mature”, if they appear to have been interrupted in either the late vitellogenic oocyte phase or the mature/spawning phase of development.

Fig. 48. Ovary from an adult female fathead minnow: Oocyte atresia, mature oocytes – Note clumping and pore-formation in the vitelline envelope (chorion) of the early atretic oocyte (large →), and the vacuolar hypertrophy of its surrounding granulosa cells (small →). Compare these with granulosa cells that surround a non-atretic late vitellogenic oocyte (►). In FHM, granulosa cells of atretic oocytes often appear to contain phagocytized material, whereas the granulosa cells of non-atretic oocytes do not (paraffin, H&E).
Fig. 49. Ovary from a normal (control) adult female fathead minnow (paraffin, H&E).

Fig. 50. Ovary from an adult female fathead minnow: Numerous atretic oocytes are evident – Grade 3 (→; paraffin, H&E).

Fig. 51. Stage 4 ovary from an adult female fathead minnow as characterized by severe oocyte atresia. Asterisks (*) indicate the relatively few non-atretic oocytes (paraffin, H&E).
Fig. 52. Oocyte atresia in ovaries from adult female fathead minnow: (a) Although atresia is most often observed in vitellogenic oocytes, it can occur at any phase of development, such as in this cortical alveolar oocyte (→). (b) Although not usually diagnosed as atresia, increased apoptotic-type death of early germ cells (→) could be a finding in some studies (paraffin, H&E).

Fig. 53. Moderately atretic zebrafish ovary (paraffin, H&E, original mag. 10 ×).
Perifollicular cell hyperplasia/hypertrophy

Exposure to aromatase inhibitors (e.g., fadrozole, prochloraz) has been associated with these perifollicular cell changes in Japanese medaka. A similar effect has also been linked with exposure to the non-aromatizable androgen, trenbolone (unpublished data). This finding is characterized by an increase in the height and number of granulosa cells, which gives this cell layer a “pseudostratified” appearance in extreme cases. A common coexisting change in affected Japanese medaka was decreased yolk formation. Since perifollicular cells (i.e., granulosa cells) are thought to be involved with aromatase production in fish (Devlin and Nagahama, 2002; Nagahama, 1987), it is possible that the increased number and size of these cells is a compensatory mechanism aimed at restoring aromatase to levels required for vitellogenesis. It is important to note that (1) normal perifollicular cells may appear hypertrophic in tangentially-sectioned oocytes, and (2) perifollicular cell changes are best identified by comparisons made with concurrent control fish.

Fig. 54. Perifollicular cell hyperplasia/hypertrophy: (a) Ovary from an untreated adult female Japanese medaka: Perifollicular cells (→) are cuboidal and have small, condensed nuclei. (b) Ovary from female Japanese medaka exposed to 3 µg/L fadrozole illustrating minimal perifollicular cell hyperplasia: Perifollicular cells surrounding some oocytes are columnar. It should be noted that in certain studies the control animals can look like this. (c) Ovary from female Japanese medaka exposed to 100 µg/L fadrozole: Ovaries graded as mildly affected, such as this, should be distinctly different from concurrent controls. (d) Ovary from female Japanese medaka exposed to 300 µg/L prochloraz: Moderately hyperplastic perifollicular cells (→) have a pseudostratified columnar appearance, and relatively enlarged oval nuclei (paraffin, H&E, bars = 25 µm).
Decreased yolk formation

Fig. 55. Stage 3 ovary from a normal (control) adult female fathead minnow: A single atretic ovary is evident (→; paraffin, H&E).

Fig. 56. Ovary from an adult female fathead minnow: Decreased yolk formation is characterized by the presence of oocytes in which yolk material is not present despite their relatively large size (large →). Note that oocytes are affected to varying degrees. Some affected oocytes have extremely fine vitellogenic granules (small →), and this is interpreted as ineffective yolk formation and deposition (paraffin, H&E, bar = 250 μm).

Fig. 57. Ovary from an adult female zebrafish: A striking example (i.e., severity Grade 4) of decreased yolk formation in an adult female zebrafish exposed to fadrozole (paraffin, H&E, bar = 500 μm).
Fig. 58. Decreased yolk formation in the ovaries of Japanese medaka: (a) Ovary from a control medaka. (b) Ovary from a female medaka exposed to 150 ng/L trenbolone (non-aromatizable androgen) shows minimal decreased yolk formation; many vitellogenic oocytes have pale, watery yolk (→). (c) Ovary from a female medaka exposed to 500 ng/L trenbolone: mildly decreased yolk formation as indicated by inadequate amounts of yolk that is excessively vacuolated. (d) Ovary from female medaka exposed to 500 ng/L trenbolone: moderately decreased yolk formation characterized by the presence of scanty yolk in relatively few oocytes. (e) Ovary from female medaka exposed to 5 µg/L trenbolone: yolk granules are absent, cortical alveoli (A) are disrupted, and there is dramatic perifollicular cell hypertrophy/hyperplasia. (f) Severely-affected ovary from a female medaka exposed to 300 µg/L prochloraz (aromatase inhibitor). All images: paraffin, H&E, bar = 500 µm except for (f): bar = 250 µm.
Changes in ovarian staging

See Chapter 5.

4.3 Secondary diagnoses in fathead minnow, Japanese medaka and zebrafish following exposure to endocrine disruptors

4.3.1 Secondary diagnoses in male fathead minnow, Japanese medaka and zebrafish following exposure to endocrine disruptors

Decreased proportion of spermatogonia

Care for Chapter 4.2.1 (Primary diagnoses in male fathead minnow, Japanese medaka and zebrafish following exposure to endocrine disruptors).

Increased vascular or interstitial proteinaceous fluid

“Increased vascular or interstitial proteinaceous fluid” indicates homogenous dark pink translucent material, presumably vitellogenin, within the testicular interstitium or blood vessels. The presence of this fluid may cause a thickening of interstitial areas that might be misinterpreted as “stromal proliferation”.

Proteinaceous fluid, interstitial (male or female)

“Interstitial proteinaceous fluid” indicates homogenous dark pink translucent material, presumably vitellogenin, within the testicular or ovarian interstitium. In male fish especially, this finding has been associated with exposure to estrogenic substances. The presence of this fluid may cause a thickening of interstitial areas that might be misinterpreted as “stromal proliferation”.

Fig. 59. Proteinaceous fluid in the interstitium of ovaries of adult female fathead minnow: There is homogenous dark pink material in interstitial spaces (→; paraffin, H&E, bar = 50 µm).
**Proteinaceous fluid, intravascular (male or female)**

“Proteinaceous fluid in intravascular spaces” of male or female fish indicates homogenous dark pink translucent materials, presumably vitellogenin, within testicular or ovarian blood vessels. In male fish especially, this finding has been associated with exposure to estrogenic substances.

**Fig. 60.** Proteinaceous fluid in blood vessels in the testis of adult male fathead minnow as homogenous dark pink material (→; GMA, H&E, bar = 25 µm).

**Fig. 61.** Intravascular proteinaceous fluid (→) in (a) the testis of adult male Japanese medaka exposed to 100 ng/L 17 β-estradiol for 4 weeks (paraffin, H&E, bar = 25 µm). (b) Ovary of 8 weeks old Japanese medaka female after 8 week exposure to 90 µg/L 4-tert-octylphenol displays proteinaceous fluid (→; paraffin, H&E, bar = 50 µm). (c) Proteinaceous fluid is evident in zebrafish testicular and hepatic blood vessels (→) and also liver sinusoids (►). (d) Proteinaceous fluid in ovarian and pancreatic blood vessels (→) and also within the ovarian interstitium (►) of zebrafish (paraffin, H&E).
Asynchronous gonad development

“Asynchronous gonad development” indicates the presence of more than one developmental phase of spermatogenic cell within a single spermatocyst enclosed by a Sertoli cell. For example, this term may be applied to a spermatocyst that contains a mixture of spermatocytes and spermatids, or a spermatocyst that contains more than one meiotic phase of primary spermatocytes (i.e., leptotene, pachytene, and/or zygotene). It also refers to the presence of distinctly different populations (i.e. developmental phases) of gametogenic cells in the right and left gonads.
**Asynchronous development, gonad (male or female)**

“Asynchronous development” indicates the presence of distinctly different populations (i.e., range of developmental phases) of gametogenic cells in different regions of a gonad.

Fig. 63. Asynchronous development in adult fathead minnow: With respect to developmental stage, there is a fairly distinct difference between the lower left region of testis (resembles Stage 1) and the upper right region (resembles Stage 3; GMA, H&E, bar = 100 µm).

Fig. 64. Asynchronous development in 8 weeks old Japanese medaka: This animal had been exposed for approximately eight weeks to 27 µg/L 4-tert-octylphenol. In addition to the presence of numerous testis-ova, the efferent duct system is abnormally irregular, and spermatogonium-containing spermatocysts (→) are located in an atypical position adjacent to the ducts (asynchronous development; H&E, bar = 100 µm).
Asynchronous development, spermatocyst (male)

“Asynchronous development in spermatocyst” indicates the presence of more than one developmental phase of spermatogenic cell within a single spermatocyst. For example, this term may be applied to a spermatocyst that contains a mixture of spermatocytes and spermatids, or a spermatocyst that contains more than one meiotic phase of primary spermatocyte (i.e., leptotene, pachytene, and/or zygotene).

Fig. 65. Asynchronous development within a spermatocyst in the testis of an adult male zebrafish exposed to 320 µg/L tamoxifen: Each spermatocyst contains multiple developmental phases of spermatogenic cells (l = leptotene, z = zygotene, p = pachytene, sc = spermatocytes, st = spermatids; paraffin, H&E).

Asynchronous development, right and left gonads (male or female)

“Asynchronous development between right and left gonads” indicates the presence of distinctly different populations (i.e., developmental phases) of gametogenic cells in the right and left gonads.

Fig. 66. Asynchronous development between right and left ovaries of an adult female zebrafish: Images (a) and (b) are contralateral ovaries from a single fish. (a) Stage 2 ovary. (b) Stage 4 ovary (paraffin, H&E).
**Altered proportions of spermatocytes or spermatids**

“Altered proportions of spermatocytes or spermatids” indicates a change in the relative proportions of spermatocytes or spermatids to other spermatogenic cell types. Changes in relative ratios could be due to an increase in the number of spermatocytes or spermatids, or to a decrease in the number of other cell types. Relative changes may also occur between spermatocytes and spermatids.

**Gonadal staging**

See Chapter 5.

**4.3.2 Secondary in female fathead minnow, Japanese medaka and zebrafish following exposure to endocrine disruptors**

**Interstitial fibrosis (male or female)**

“Interstitial fibrosis” indicates the presence of increased fibrous connective tissue (collagenous fibers and fibrocytes or fibroblasts) within the testicular or ovarian interstitium (stroma). Collagen may be difficult to appreciate in early phases of fibrosis. In most cases, this term should be used in preference to terms such as “stromal hyperplasia.”

![Fig. 67. Interstitial fibrosis in adult male Japanese medaka: Green arrows indicate focally extensive interstitial fibrosis, whereas red arrows indicate focal atrophy of the germinal epithelium (paraffin, H&E).](image1)

![Fig. 68. Interstitial fibrosis in adult female zebrafish: Arrows (→) point to bundles of fibrous connective tissue and possibly increased perivascular smooth muscle (paraffin, H&E; original mag. 20 ×).](image2)
**Egg debris in the oviduct**

“Egg debris in the oviduct” indicates the presence of inspissated-appearing, homogenous, irregular, dense pink material, presumed to be yolk, within the oviduct.

![Fig. 69. Egg debris in the oviduct of an adult female fathead minnow (paraffin, H&E).](image)

**Granulomatous inflammation**

In the early stages of inflammation, this process is characterized by the presence of epithelioid macrophages that typically form sheets or nodules (granulomas) due to desmosome-like cytoplasmic attachments (Noga et al., 1989). When compared to histiocytic-type macrophages, epithelioid macrophages have larger, more open-faced, centralized nuclei and less abundant cytoplasm. During resolution of inflammation, the epithelioid macrophages may become flattened into fibrocyte-like cells. Lymphocytes, granulocytes, and multinucleated giant cells may also be components of granulomatous inflammation. Granulomatous inflammation is intrinsically a pathologic process that is often associated with reactions to infectious agents, foreign materials, or the aftermath of necrosis; therefore, it is important to distinguish this, if possible, from the presence of macrophage aggregates in the ovary or histiocytic cells in the lumen of the testis.

![Fig. 70. Egg debris in the oviduct of an adult female zebrafish: The oviduct is obstructed by a large yolk plug (→; paraffin, H&E).](image)

---

OECD Guidance Document for the Diagnosis of Endocrine-related Histopathology of Fish Gonads

Page 67/96
Fig. 71. Granulomatous inflammation. (a) Sheets of macrophages and other inflammatory cells eclipse much of the germinative tissue in the testis of this adult male fathead minnow. (b) Relatively few viable-appearing oocytes remain in this ovary of an adult female fathead minnow. As in the testis photo, the inciting cause of the inflammation is not evident at this magnification (paraffin, H&E).

Fig. 72. (a) Granulomatous inflammation in the ovary of an adult female zebrafish featuring multinucleated giant cells, a hallmark of granulomatous inflammation (H&E, original mag. 20 ×). (b) Ovary from an adult female zebrafish showing lightly pigmented sheets of histiocytic macrophages, which aid in the resorption of atretic oocyte material (see: “pigmented macrophage aggregates”). This is actually not granulomatous inflammation, and it is usually not necessary to record this as a pathologic finding. It may be difficult to distinguish these two types of responses in some instances. To further complicate matters, granulomatous inflammation may initiate from, and merge with, pigmented macrophage aggregates. (paraffin, H&E).
“Decreased post-ovulatory follicles” indicates a decrease in the number of collapsed perifollicular sheaths, or membranous structures lined by granulosa cells, theca cells and surface epithelium, following release of oocytes, in comparison to control fish. The granulosa cells are often hypertrophic, although this appears to be species dependent (Saidapur, 1982).

4.4 Additional diagnostic criteria and an illustrated glossary of microanatomical and diagnostic terms

In this section, a number of additional terms and diagnoses are listed alphabetically followed by working definitions or descriptions.

Gender
Since the genetic sex of a fish cannot be determined within the context of a screening assay, and because the external phenotypic sex may be an unreliable indicator and/or is not easily determined in some species, by convention the gender of a fish will be assigned according to the most abundant mature cell type that is present in the gonad.

Germ Cell Neoplasms
As spontaneous findings, germ cell neoplasms such as seminomas and dysgerminomas are rare in fathead minnow and Japanese medaka, although both spontaneous and carcinogen-induced seminomas may be relatively more common in zebrafish. There is currently little evidence to support the idea that such tumors are linked to endocrine disruptor exposure, and control animals seem to be affected as often as chemically-exposed individuals. In many fishes, distinguishing features of germ cell neoplasms include haphazard anatomic organization and progression of cell development, and a tendency to form mass-like lesions that distort the gonad architecture. In early life stage studies in which fish are exposed to potent hermaphroditic chemicals such as 17 β-estradiol or 17α-ethynylestradiol, it may be difficult to distinguish germ cell neoplasms from malformed intersex gonads.

Fig. 73. Germ cell neoplasms (following page): (a) Seminoma in a fathead minnow testis: This particularly well-differentiated tumor presents as a mass (→) that markedly deforms the contour of the testis (paraffin, H&E, bar = 250 μm). (b) Higher magnification of the tumor in Fig. a: Arrows (→) indicate the margin of the neoplasm (lower portion of image) relative to the non-neoplastic testis. As is the case for many fish seminomas, this tumor is “spermatocytic”; i.e., there is at least some developmental progression toward more mature cell types within the tumor (paraffin, H&E, bar = 25 μm) (c) Dysgerminoma in Japanese medaka testis: The caudal pole of the testis is effaced by a mass (→) consisting of oogenic tissue (paraffin, H&E, bar = 250 μm). (d) Higher magnification of the tumor in Fig. c: The disorganization of the oogenic tissue is apparent. It is important to distinguish this neoplasm from other findings such as: asynchronous development, gonad (in which different areas of the gonad are in different stages of development that blend almost imperceptibly and do not form a mass); testis-ova formation (in which the scattered oocytes do not form a mass capable of distorting the gonad); and possibly from hermaphroditism (in which the anatomic arrangement and developmental progression of the aberrant tissue is orderly and essentially resembles the normal gonad; paraffin, H&E, bar = 50 μm). (e) Well-differentiated seminoma in a zebrafish testis. The large neoplastic mass expands the abdomen and displaces other organs (paraffin, H&E, bar = 800 μm). (f) Higher magnification of the tumor in Fig. e: In this particular case, the neoplastic tissue closely resembles normal testis in terms of organization and cellular constituency, at least in some areas of the mass (paraffin, H&E, bar = 25 μm). (g) Poorly-differentiated seminoma in a zebrafish testis (paraffin, H&E, bar = 800 μm). (h) Higher magnification of Fig. g: There is loss of the normal testicular architecture (paraffin, H&E, bar = 25 μm).
Germinal epithelium (female)

The germinative parenchyma (epithelium) of the ovary is a membrane-bound structure and constitutively contains oogonia, prefollicular and prethecal cells, epithelial cells, and occasionally small chromatin nucleolar (primary growth) oocytes (Norberg et al., 1999; Parenti and Grier, 2003). The germinal epithelium separates the ovarian lumen from the stroma, the latter of which often contains perinucleolar, cortical alveolar, and vitellogenic follicles within a variably-apparent extravascular space.

Germinal epithelium (male)

The germinative intratubular (intralobular) parenchyma of the testis is a membrane-bound structure and consists of multiple spermatocysts in various phases of development. For fathead minnow and zebrafish, boundaries of the germinal epithelium at various locations throughout the testis include the interlobular interstitium, the lobular lumina, collecting ducts, and the tunica albuginea. For Japanese medaka, boundaries include the interlobular interstitium, efferent ducts, and the tunica albuginea.

Fig. 74. The germinal epithelium (→) in the ovary of a normal adult Japanese medaka is a membranous structure that separates the ovarian lumen (L) from the extravascular space (EVS) of the ovarian stroma. (paraffin, H&E).

Fig. 75. Germinal epithelium, testis. (a) In the normal testis from an adult fathead minnow, the germinal epithelium (↔) extends from the interlobular interstitium to the lobular lumen (GMA, H&E, bar = 25 µm). (b) In the testis of normal adult Japanese medaka, the germinal epithelium (↔) extends from the tunica albuginea to the efferent duct (paraffin, H&E).
**Germinal epithelium, atrophy/hypoplasia (male)**

“Germinal epithelium, atrophy/hypoplasia” indicates loss or underdevelopment of germinal epithelium, respectively, this condition may be associated with interstitial fibrosis and increased prominence of interstitial cells in affected areas of the testis. It may be difficult to distinguish atrophy from hypoplasia. Care should be taken to avoid mistaking areas of collecting ducts for atrophy. Severity of this finding can vary from Grade 1 (minimum, focal) to Grade 4 (severe, diffuse). If thinning of the epithelium appears to be caused by degenerative changes that are obvious in the section, the diagnostic term *testicular degeneration* should be used instead.

![Fig. 76](image)

(a) Atrophy of germinal epithelium in the testis of an adult fathead minnow; also note the prominence of interstitial (Leydig) cells (red circles) and interstitial fibrosis. (b) Normal fathead minnow testis. (c) Normal collecting duct region in an adult male fathead minnow testis (GMA, H&E, bars = 25 µm). The presence of pigment in the duct walls and the lack of interstitial cells are distinguishing features.
Fig. 77. Hypoplasia of the germinal epithelium: (a) Normal testis from an 8 weeks old control Japanese medaka. (b) Higher magnification of the testis in Fig. a. (c) In the testis from an 8 weeks old male Japanese medaka exposed to 450 mg/L 4-n-aminylaniline for approx. 8 weeks, the hypoplastic testis (→) is small and poorly formed. (d) Higher magnification of the testis in Fig. c: Clusters of primarily spermatogonia (S) are haphazardly arranged and are surrounded by mononuclear cell infiltrates. Spermatozoa are present in a lumen-like space (paraffin, H&E, bars = 250 µm for Figs. A and c, bars = 25 µm for Figs. b and d).


**Gonadal stromal tumors**

Among the three fish species, gonadal stromal tumors seem to be even less common than germ cell neoplasms. Examples include Sertoli cell tumors, granulosa cell tumors, and teratomas.

Fig. 78. Gonadal stromal tumors: (a, b) Teratoma in a fathead minnow ovary: Disparate elements such as developing bone, CNS-type neural tissue, and ocular pigment cells are evident. (c) Teratoma in a Japanese medaka ovary represent generally uncommon, although not exceedingly rare, incidental findings. (d) In another tissue section from the same ovary as in Fig. c, the ocular tissue is remarkably well-formed (paraffin, H&E, bars in (a) = 100 µm, (b) = 25 µm, (c) = 100 µm, (d) = 50 µm)
**Hepatocyte basophilia, increased/decreased**

A generally diffuse increase in hepatocellular cytoplasmic basophilia has been observed in male fish that have been exposed to compounds that are able to interact with hepatic estrogen receptors, including 17β-estradiol and 17α-methyl-dihydrotestosterone (Wester et al., 2003). This increase in basophilia, which is correlated with increased vitellogenin production, presumably mimics the heightened metabolic state (e.g., increased endoplasmic reticulum) that is required for the production of vitellogenin in the reproductively-active female fish.

![Fig. 79. Hepatocyte basophilia: (a) Liver from a control adult male fathead minnow: In addition to the overall coloration, note the hepatocellular cytoplasmic vacuolization as indicated by the arrows. (b) Liver from an adult male fathead minnow that was exposed to a compound with estrogenic activity: There is a diffuse increase in hepatocellular basophilia, a loss of cytoplasmic vacuolization, and hepatic blood vessels contain proteinaceous fluid (→; paraffin, H&E). (c) Liver from an adult control Japanese medaka. (d) Liver from an adult Japanese medaka exposed to 100 µg/L 4-tert-octylphenol, an estrogenic substance (paraffin, H&E, bars = 25 µm).](image)
**Histiocytic cells (male)**

“Histiocytic cells” indicates the presence of individual or clustered cells with small eccentric nuclei and moderate to abundant, pale or vacuolated cytoplasm within the testicular lumen, germinal epithelium, efferent ducts and/or ductus deferens. Such cells may contain intracytoplasmic cellular debris (presumably phagocytized). The origin of the histiocytic cells in each particular case may not be clear; for example, they may be hematogenous macrophages or Sertoli cells. Histiocytic cells should be differentiated from *macrophage aggregates* (these variably pigmented cells are primarily interstitial) and *granulomatous inflammation* (which is predominately comprised of “epithelioid” macrophages and/or flattened, fibrocytic cells).

Fig. 80. Histiocytic cells in adult male fathead minnow: (a) Cells with small peripheral nuclei and abundant vacuolated cytoplasm are present within the germinal epithelium and are scattered throughout the tubule lumen (→). Some of these cells contain phagocytized cellular debris. (b) Similar cells (→) are evident within the lumen of the collecting duct (paraffin, H&E).
**Increased/decreased cells, [insert cell type] (testis or ovary)**

It is recognized that endocrine active compounds may alter the proportional distribution of gametogenic and supportive cell types in the testis or ovary. Certain types of alterations (for example, the proliferation or absence of single cell population) may not be adequately documented by gonadal staging. This diagnostic term provides a mechanism for documenting such changes. For consistency, the pathologist should presume that these semi-quantitative changes are: (1) relative to other cell types in the gonad; (2) relative to cell numbers in control animals; and (3) estimates only, versus actual cell counts. See example of increased spermatozoa below.

![Figure 81](image1)

**Fig. 81.** Increased cells (*spermatozoa* in this case). (a) Testis from an adult control male fathead minnow. (b) Testis from an adult male fathead minnow exposed to 300 µg/L prochloraz: Spermatozoa density is increased relative to control. Ideally, comparisons should be made between same stage gonads (testis in (a) is Stage 2, whereas testis in (b) is Stage 3) (paraffin, H&E, bar = 25 µm).

![Figure 82](image2)

**Fig. 82.** Testis from an adult male zebra-fish: increase in the proportion of spermatocytes (paraffin, H&E).
(Pigmented) Macrophage aggregates

(Pigmented) Macrophage aggregates are cell clusters constitutively present in the interstitium of the ovary primarily, although they may also be found in the fish testis (unusual for, e.g., tank-raised fathead minnow). These phagocytes usually have small condensed eccentric or peripheralized nuclei and various brown, yellow, red, or gold pigmented granules (lipofuscin, ceroid, hemosiderin, and/or melanin) that often impart a slightly crystalline appearance to their comparatively abundant pale cytoplasm. In the normal ovary, macrophage aggregates are thought to be involved in the processing of breakdown products associated with atresia of non-spawned oocytes. It has been demonstrated that macrophage aggregates may become larger and/or more numerous following exposure to certain toxicants or infectious agents (Blazer et al., 1987). Whenever possible, macrophage aggregates should be distinguished from granulomatous inflammation, which is characterized by the presence of epithelioid macrophages. This is not always easy, as macrophage aggregates often proliferate with, and become incorporated into, granulomatous inflammation.

Fig. 83. (a) Macrophage aggregates in adult female fathead minnow: Arrows (→) indicate multiple aggregates within the ovarian interstitium. (b) Large macrophage aggregate (→) in a Japanese medaka ovary (paraffin, H&E).
**Mineralization (male or female)**

In both the testis and ovary, mineralization may be evident in the germinal parenchyma, luminal areas, or duct system. In H&E-stained sections it usually manifests in one of several forms:

1. dark purple, laminated, spherical concretions;
2. purple or dark blue, finely-granular deposits;
3. purple-discolored, fragmented, gonadal structures; or
4. irregular spicules of dystrophic mineralization in areas of granulomatous inflammation.

The last form, when observed within ovaries, is often associated with resorption of atretic oocytes. In the testis, mineralization must be distinguished from testis-ova formation; on occasion, this may be difficult, as degenerating testis-ova may also become mineralized. Rarely, mineralization may be seen as a presumably non-endocrine treatment effect, possibly due to vehicle-induced changes in the exposure water chemistry.

Fig. 84. Mineralization: (a) Testis from an adult male fathead minnow showing numerous foci of angular, roughly spherical, mineralization in the collecting ducts. Surprisingly, even this profound degree of mineralization is not usually associated with morphologic evidence of obstructive dysfunction. (b) Ovary from an adult male zebrafish: Although difficult to say with certainty, this may represent mineralization of atretic oocyte material (paraffin, H&E, bars = 25 and 100 μm, respectively).
Nephropathy

“Nephropathy” indicates degenerative renal disease has been observed in a variety of fishes that have been exposed to compounds with estrogenic activity (Herman & Kincaid, 1988; Palace et al., 2002; Zillioux et al., 2001). Renal impairment presumably occurs due to increased production of vitellogenin (especially in males) that stresses the kidney via protein overload. Microscopic lesions may include swelling of tubular epithelial cells, tubular necrosis, dilation of Bowman’s capsule, interstitial fibrosis, casts, and hyaline droplets in tubules or glomeruli.

Fig. 85. Nephropathy: In contrast to the kidney from an untreated adult male fathead minnow (a), the kidney from an adult male fathead minnow exposed to a compound with estrogenic activity (b) shows glomerular epithelial cell hypertrophy, vacuolar swelling and necrosis of the tubular epithelium, and hyaline droplets within glomerular and tubular epithelia (paraffin, H&E, bars = 25 µm).

Fig. 86. Nephropathy: In contrast to the kidney from an untreated adult male Japanese medaka (a), the kidney from an adult female Japanese medaka exposed to 1 mg/L 4-tert-pentylphenol shows changes similar to those in fathead minnow in Fig. 85b. Arrow (→) indicates large hyaline droplet in glomerular capillary (paraffin, H&E, bars = 25 µm).
Oocyte membrane folding

“Oocyte membrane folding” indicates a compound-associated, presumably degenerative (atretic) process characterized by abrupt, usually multiple, invaginations of the chorion. As an example, this particular lesion was observed in ZBF following exposure of adult females to the anti-estrogen tamoxifen (Wester, et al., 2003).

Fig. 87. Membrane folding in adult female fathead minnow: two cortical alveolar oocytes show folding of the oocyte membrane (→; paraffin, H&E).

Fig. 88. Membrane folding in adult female fathead minnow: This is actually not an example of membrane folding. The arrow (→) indicates the micropyle, which is a normal funnel-shaped structure located at the animal pole of the oocyte. The inset demonstrates this structure at a higher magnification and a different angle (paraffin, H&E).

Fig. 89. This high magnification image illustrates abrupt folding of the chorion with incorporation of the granulosa cells in this oocyte from an adult female zebrafish exposed to 320 mg/L tamoxifen (→; paraffin, H&E).
Ovarian cysts

Although ovarian cysts are relatively common in the ovaries of adult female fathead minnow, no references to these structures were found in the scientific literature. Large cysts may be grossly visible. By light microscopy they present as single or multiple, thin-walled, spherical to irregular structures (~50 - 500 µm) that are lined by a mildly vacuolated, ciliated, cuboidal epithelium that rests on a very fine basement membrane. On occasion the cyst lumen may contain uniform pale eosinophilic material, and/or a few exfoliated cells, but usually it appears completely empty. In rare cases, these cysts may comprise a large percentage of the ovary and crowd out much of the oogenic tissue. To date there have been no compound exposure effects associated with ovarian cysts. Their developmental origin has not been determined, although it is suspected that they most likely represent spontaneous congenital or developmental anomalies. Ovarian cysts should not be confused with post-ovulatory follicles, which usually have a thicker wall, are often collapsed, and are never lined by ciliated cells.

![Fig. 90. Ovarian cyst](image)

(a) Low magnification view of three ovarian cysts in adult female fathead minnow (paraffin, H&E, bar = 100 µm). (b) Higher magnification of two cyst walls from Fig. a to demonstrate the ciliated epithelium (bar = 25 µm).
**Ovarian spermatogenesis**

“Ovarian spermatogenesis” indicates the presence of non-neoplastic spermatogenic cells, usually immature, within the ovary. There is little or no evidence of lobular or tubular testicular architecture. Care should be taken to distinguish ovarian spermatogenesis from mitotically dividing oogonia; a key feature of ovarian spermatogenesis is the presence of multiple spermatogenic phases.

---

**Fig. 91.** Oogonia dividing mitotically in the ovaries of adult female fathead minnow: Groups of cells resembling spermatocytes (→) are situated between perinucleolar and cortical alveolar oocytes. This should not be mistaken for spermatogenesis (paraffin, H&E).

---

**Fig. 92.** Ovarian spermatogenesis in adult female Japanese medaka: Aggregates of spermatogenic cells including spermatogonia (large arrow), spermatocytes (small arrow), and spermatzoa (►) are located between ovarian follicles (paraffin, H&E).
**Retained peritoneal attachments/gonadal duct feminization (male)**

“Retained peritoneal attachments/gonadal duct feminization” has been recognized in male zebrafish that have been exposed to estrogen (17β-estradiol; van der Ven and Wester, 2004). In normal (control) fish, the cranial attachment of the testis to the peritoneum is lost prior to reproductive maturity. Conversely, the testes of affected fish retain both of their cranial and caudal peritoneal attachments, thus forming an ovarian-like cavity. A similar response to compounds with estrogenic activity has also been reported for juvenile fathead minnow (Panter et al., 2006; Van Aerle et al., 2002). In addition to retention of the membranous peritoneal attachments, the lining of these membranes may become transformed into a ciliated simple or pseudostratified columnar epithelium that strongly resembles oviduct epithelium (unpublished data).

![Fig. 93. Retained peritoneal attachments: (a) Peritoneal cavity and testes of an untreated 6 weeks old male zebrafish: Arrows (→) indicate normal loss of cranial peritoneal attachments (paraffin, H&E, coronal section). (b) This 6 weeks old male zebrafish was exposed (lifetime) to 1 nM 17β-estradiol, and has retained its cranial peritoneal attachment (→; paraffin, H&E, coronal section).](image)
Fig. 94. Feminization of gonadal ducts: (a) Testis from a control male fathead minnow 120 days post-hatch (dph): Note the single attachment site of the gonadal duct (→) that is appropriate for the testis. (b) Testis from a male fathead minnow 120 dph exposed to a xenoestrogen from early embryo stage: The two attachment sites (→) form an intervening ovarian-like cavity (*) that is lined by oviduct-type epithelium. In this instance, the cavity contains spermatozoa, exfoliated cells, and cellular debris. (c) Testis from another male fathead minnow 120 dph exposed to a xenoestrogen: Similar appearance to Fig. b, and, in addition, with a single oocyte in the testicular parenchyma. (d) Ovary from a control female fathead minnow 120 dph, which was included to illustrate normal oviduct epithelium for comparison with Figs. b and c (paraffin, H&E, transverse sections).
**Sertoli cell hypertrophy**

Exposure of male fish to estrogen-active compounds has been reported to cause enlargement of Sertoli cells, with or without Sertoli cell proliferation (Kinnberg et al., 2000; Miles-Richardson et al., 1999a, b; van der Ven et al., 2003). In the scientific literature, the light microscopic appearance of hypertrophic Sertoli cells tends to be ambiguous, as Sertoli cells resemble spermatogonia in some descriptions and images.

![Fig. 95. Nuclear enlargement of Sertoli cells (large arrows) is evident in the testes of adult male zebrafish after exposure to 17α-methyltdihydrotestosterone for 10 days (b, c) as compared to the Sertoli cell nuclei of an adult male control (a; small arrows; paraffin, H&E).](image)

**Sperm necrosis**

“Sperm necrosis” has been observed occasionally as a focal, or more often multifocal, lesion in fathead minnow testes that are otherwise well-preserved. It is characterized by patchy areas of increased pallor and fragmentation +/- swelling of spermatozoa within testicular lumina. To date, sperm necrosis has not been related to compound exposure. Sperm necrosis can be seen concurrent with overt testicular mineralization, and it may well be a more subtle manifestation of mineralization.

![Fig. 96. Sperm necrosis in adult male fathead minnow: Within testicular lumen, spermatozoa are pale and fragmented (paraffin, H&E, bar = 25 µm).](image)
**Vitellogenic oocyte**

A vitellogenic oocyte is an oocyte that contains microscopically visible yolk material. Generally, such material is strongly eosinophilic and slightly refractile in hematoxylin- and eosin-stained sections. This material may be present in the form of spherical, globular, yolk granules (fathead minnow, zebrafish). In some scholarly sources (e.g., Iwamatsu, et al., 1998), the term “vitellogenic” has been applied to cortical alveolar oocytes, which lack eosinophilic yolk granules/globules (although their amphophilic or clear cortical alveoli are also known as yolk vesicles).
5. Gonadal staging criteria

The goal of gonadal staging is to determine if the administration of a particular endocrine-active substance affects the reproductive cycle status of adult male and female fish. The purpose of this section is to describe a rapid, semi-quantitative method for assessing the proportions of various gametogenic cell types (gonadal staging) based on the light microscopic examination of hematoxylin and eosin-(H&E-)stained histological sections.

Semi-quantitative gonadal staging has been proposed for, or employed in, studies involving fathead minnow and other fish species (Ankley et al., 2002; Jensen et al., 2001; Miles-Richardson et al., 1999a; Nichols et al., 2001; US Environmental Protection Agency, 2002). Although such studies generally included excellent descriptions of the different gametogenic maturation stages (e.g., spermatogonium through spermatozoa for the testis), they did not incorporate pre-defined categorical guidelines for evaluating and reporting the reproductive cycle status of an individual fish. To maintain scientific integrity across the board in a program that involves multiple studies, multiple laboratories, and large numbers of animals, it is essential that observations are recorded on a fish-by-fish basis. The use of a categorization system can improve the consistency and objectivity of reported observations within and among experiments; consequently, comparisons of the results are more meaningful.

Categorization systems also have some drawbacks and limitations, the most significant of which are (1) the potential loss of discriminatory data when similar, but not identical, types of observations are combined (binned) into a single class; (2) the questionable biological relevance of the classification criteria in some cases; and (3) the inability of any single classification system to address every type of observation (either predicted or unforeseen).

To address this last limitation, gonadal staging is accompanied by a complete histopathological evaluation of the gonads; in this manner, the loss or overabundance of a specific gametogenic cell type, for example, can be documented.

The semi-quantitative gonadal staging scheme selected for analysis of fish gonads is a modification of a system adopted by the United States Department of the Interior, U.S. Geological Survey, Biological Resources Division as part of the “U.S. Biomonitoring of Environmental Status and Trends (BEST) Program” (McDonald et al., 2000). The authors of the BEST system credit previous work by Treasurer and Holiday (1981), Nagahama (1983), Rodriguez et al. (1995), and Goodbred et al. (1997). The foremost benefits of this system are speed and ease of use, especially when compared to fully-quantitative staging. The basis of the BEST system is a visual assessment of the density of gametogenic precursors as compared to mature gametocytes in one or more gonad sections. Accordingly, the stage numbers (testis: Stages 0 to 4; ovary: Stages 0 to 5) increase in direct relationship to the relative proportion of mature cells.

Although the BEST system was initially developed to assess reproductive function in seasonal spawners such as carp (Cyprinidae) and black basses (Centrarchidae), the same stage categories can be applied to fractional spawners such as fathead minnow. This was demonstrated at the October 2003 meeting of the histopathology subcommittee of the Fish Drafting Group at the Paris meeting, at which the participants were asked to briefly evaluate the applicability of a modified BEST system using actual histological specimens of the three species fathead minnow, medaka and zebrafish. In general, the participants agreed that they could readily recognize the various gonadal stages as defined by modified BEST criteria. The participants acknowledged that the terminal stages of the system (testis: Stage 4, “Spent”; ovary: Stage 5, “Post-ovulatory”) would be minimally present (or possibly not present at all) among reproductively-active adult fathead minnow, medaka and zebrafish, because they are not seasonal spawners. Similarly, it was thought that Stage 4, “Late development/hydrated”, would be rarely observed in the test fishes, due to the very transient nature of this stage in fractional spawners.

A few modifications have been made to the BEST system to adapt it for use. For example, there is currently no provision in the system for gonads that are comprised entirely of spermatogonia or oogonia. Although it is intended that reproductively mature fish are used, it is possible that an occasional
animal may not attain sexual maturity by the time the experiment is terminated, or that certain test compounds might cause reversion of the gonads to a juvenile phenotype. Therefore, a pre-staging category called “juvenile” has been added for both male and female fish as one modification of the BEST system. Another modification to the system involves an apparent discrepancy between the BEST system and Goodbred et al. (1997) concerning the thickness of the testicular germinal epithelium as a staging criterion. As indicated by Goodbred et al. (1997), the germinal epithelium becomes thinner as the testis stage increases, whereas the reverse occurs according to the BEST system (as presented in McDonald et al., 2000). Although it is difficult to find corroborating statements in the scientific literature, empirical evidence indicates that Goodbred et al. (1997) is correct on this point. A third modification to the system is the option to subdivide a stage into two subordinate stages (e.g., Stages 3A and 3B), if the pathologist believes that this tactic would reveal a subtle, compound-related effect that might otherwise be missed. Other modifications to the system are relatively minor and primarily involve rewording for clarification.

The cell distribution pattern is likely to vary throughout a given tissue section, the gonad should be staged according to the predominant pattern in that section. Both gonads should be staged as a single organ according to the predominant pattern. Gonads that cannot be reasonably staged for various reasons (e.g., insufficient tissue, or extensive necrosis, inflammation, or artifact) should be recorded as UTS (unable to stage).

5.1 Criteria for staging testes in fathead minnow and zebrafish

- **Juvenile**: gonad consists of spermatogonia exclusively; it may be difficult or impossible to confirm the sex of these individuals.
- **Stage 0 – Undeveloped**: entirely immature phases (spermatogonia to spermatids) with no spermatozoa.
- **Stage 1 – Early spermatogenic**: immature phases predominate, but spermatozoa may also be observed; the germinal epithelium is thinner than it is during Stage 2.
- **Stage 2 – Mid-spermatogenic**: spermatocytes, spermatids, and spermatozoa are present in roughly equal proportions; the germinal epithelium is thinner than Stage 1, but thicker than Stage 3.
- **Stage 3 – Late spermatogenic**: all stages may be observed, however, mature sperm predominate; the germinal epithelium is thinner than it is during Stage 2.
- **Stage 4 – Spent**: loose connective tissue with some remnant sperm.
Fig. 97. Examples of staging system applied to four testes from fathead minnow: There is progressive thinning of the germinal epithelium and expansion of the lobular lumen with each increase in stage. Note that no spermatozoa are present as in the Stage 0 image (GMA, H&E).

Fig. 98. Examples of staging system applied to testes from zebrafish: The zebrafish testis is essentially similar to the fathead minnow testis in terms of staging. For some reason, spermatozoa are somewhat sparse in the illustrated testes (paraffin, H&E).
5.2 Criteria for staging testes in Japanese medaka

To derive each stage score, the estimated width of the germinal epithelium (EWG) can be compared to the estimated width of the testis (EWT) in the following manner:

- **Stage 1**: EWG > ⅔ EWT
- **Stage 2**: EWG ⅔ to > ½ EWT
- **Stage 3**: EWG ½ to > ¼ EWT
- **Stage 4**: EWG < ¼ EWT

Fig. 99. Examples of staging system applied to testes from Japanese medaka: Black arrows (↔) represent the estimated width of the testis (EWT), and red arrows (↔) represent the estimated width of the germinal epithelium (EWG; measurements are illustrated unilaterally for simplicity). In order to obtain comparable sections, it is imperative that each section contains a portion of the central duct (CD), preferably at its widest and longest extent (paraffin, H&E).
5.3 Criteria for staging ovaries in fathead minnow, Japanese medaka and zebrafish

The following are morphologic criteria for staging female fathead minnow, Japanese medaka and zebrafish:

- **Juvenile**: gonad consists of oogonia exclusively; it may be difficult or impossible to confirm the sex of these individuals.
- **Stage 0 – undeveloped**: entirely immature phases (oogonia to perinucleolar oocytes); no cortical alveoli
- **Stage 1 – early spermatogenic**: vast majority (e.g., > 90%) are pre-vitellogenic follicles, predominantly perinucleolar through cortical alveolar.
- **Stage 2 – mid-development**: at least half of observed follicles are early and mid-vitellogenic.
- **Stage 3 – late development**: majority of developing follicles are late vitellogenic.
- **Stage 4 – late development/hydrated**: majority of follicles are late vitellogenic and mature/spawning follicles; follicles are larger as compared to Stage 3.
- **Stage 5 – post-ovulatory**: predominately spent follicles, remnants of theca externa and granulosa.

Fig. 100. Examples of the staging system applied to the ovaries of the fathead minnow: Due to its transient nature in fathead minnow, Stage 4 is not often observed (paraffin, H&E).
Fig. 101. Examples of the staging system applied to the ovaries of adult Japanese medaka (paraffin, H&E).
Fig. 102. Examples of the staging system applied to the ovaries of adult zebrafish (paraffin, H&E).
6. References


OECD Guidance Document for the Diagnosis of Endocrine-related Histopathology of Fish Gonads


