Video Article

Necropsy-based Wild Fish Health Assessment

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Abstract

Anthropogenic influences from increased nutrients and chemical contaminants, to habitat alterations and climate change, can have significant effects on fish populations. Adverse effects monitoring, utilizing biomarkers from the organismal to the molecular level, can be used to assess the cumulative effects on fishes and other organisms. Fish health has been used worldwide as an indicator of aquatic ecosystem health. The necropsy-based fish health assessment provides data on visible abnormalities and lesions, parasites, condition and organosomatic indices. These can be compared by site, season and sex, as well as temporally, to document change over time. Severity ratings can be assigned to various observations to calculate a fish health index for more quantitative assessment. A drawback of the necropsy-based assessment is that it is based on visual observations and condition factors, which are not as sensitive as tissue and subcellular biomarkers for sublethal effects. Additionally, it is rarely possible to identify causes or risk factors associated with observed abnormalities. So, for instance a raised lesion or "tumor" on the fins, lips or body surface may be a neoplasm. However, it could also be a response to a parasite, chronic inflammation or hyperplasia of normal cells in response to an irritant. Conversely, neoplasms, certain parasites, other infectious agents and many tissue changes are not visible and so may be underestimated. However, during the necropsy-based assessment, blood (plasma), tissues for histopathology (microscopic pathology), genomics and other molecular analyses, and ooliths for aging can be collected. These downstream analyses, together with geospatial analyses, habitat assessments, water quality and contaminant analyses can all be important in comprehensive ecosystem evaluations.

Video Link

The video component of this article can be found at https://www.jove.com/video/57946/

Introduction

Human activities have numerous adverse effects on aquatic environments. Fish inhabit various water bodies that the human population recreates in and often uses as a drinking water source and hence are important indicators of the health of the aquatic environment. Wild fish that live and reproduce in a particular habitat are exposed throughout their lives to various stressors including pathogens, parasites, poor water quality and chemical contaminants. Thousands of chemicals enter our waterways through industrial and human wastewater, suburban/urban stormwater and agricultural runoff. These complex mixtures of chemicals can have additive, synergistic or antagonistic effects on exposed organisms¹,²,³. In addition, other environmental stressors such as elevated nutrients, elevated temperature, low dissolved oxygen or fluctuating pH can exacerbate the effects of chemical contaminants¹,³. Environmental stressors can also influence infectious disease outcomes directly by increasing the number of infectious agents, increasing the virulence of opportunistic pathogens, or suppressing the immune response and disease resistance of the host⁴,⁵,⁶. For these reasons, there is increasing interest in biological or adverse effects monitoring¹,⁵,⁶,⁷, utilizing fish and other aquatic organisms to identify populations and ecosystems at risk.

Adverse effects monitoring utilizes biomarkers at various levels of organization, from the organismal to the subcellular or molecular, to identify sublethal effects which may influence populations and be indicative of exposure to various stressors. Indicators at the organismal level include visible abnormalities and conditions. Condition indices based on length and weight are calculated to evaluate the well-being or fitness of fish populations. The most common is Fulton's condition factors (K) = (weight/length⁴)⁵, Another indicator is the presence of visible abnormalities. A variety of methods have been used in individual studies and monitoring programs to assess, document, and evaluate visible abnormalities. Assessment based only on external abnormalities, i.e., the proportion of individuals with disease, fin damage, tumors and skeletal anomalies, is one of the metrics for the index of biotic integrity (IBI) that evaluates community health⁴. A similar assessment termed DELTs (deformities, erosions, lesions, tumors) has also been used to evaluate the health of fish communities. However, these methods only assess external visual abnormalities and not internal lesions or early sublethal indicators.

Necropsy-based assessments include external and internal observations and allow for the measurement of additional condition indices. Hepatosomatic index (liver weight/total body weight) has also been used as an indicator of fitness or energy reserves⁸ for which a higher index value indicates healthier fish. However, a number of studies have shown that hypertrophy or an increase in liver size occurs due to exposure to various contaminants metabolized by the liver⁹,¹⁰. In this case a higher index would be indicative of exposure to certain chemical classes. The gonadosomatic index (gonad weight/total body weight) is another condition index directed toward reproductive health¹¹. Observations made...
The standardized necropsy-based assessment described here can be used to augment the grossly visible assessment in multiple ways depending on the question(s) to be answered, expertise and other available resources. Our routine approach is to collect biometric data (length, weight, liver weight, gonad weight), blood for plasma/serum analyses, document external and internal visible abnormalities, preserve pieces of organs for microscopic analyses and collect otoliths for age analyses. The necropsy-based assessment plus age analysis and histopathology of various organs, allows for the calculation and comparison of various condition indices, prevalence of visible abnormalities, as well as microscopic tissue changes, by sex, age, site and sampling period. Additional tissue collections can be made for many other analyses including electron microscopy, bacteriology, virology, parasitology and chemical concentrations. These methods can also be part of more in-depth analyses used to diagnose the cause of fish kills or mortalities of captive fishes. Methods for collection of tissue for two additional analyses, gene expression and functional immune analyses are illustrated.

### Protocol

Methods described here have been approved by the Leetown Science Center’s Institutional Animal Care and Use Committee.

1. **Fish Collection**

   1. Collect live fish with a minimum of stress. Use boat or backpack electrofishing, hook and line or nets.
   2. Hold fish in live wells or aerated containers until sampling.

   NOTE: The American Fisheries Society has published a number of guides for fish collection, handling and anesthesia/euthanasia. Wear gloves when handling fish.

2. **Fish Necropsy**

   1. **Euthanize a fish.**
   2. Measure biometric characteristics.
      1. Weigh the fish to the nearest gram.
      2. Measure the fish length to the nearest millimeter.
         1. Measure the total length from the tip of the snout with the mouth closed to the end of the tail when pinched together.
         2. Measure the fork length from the fork in the tail to the tip of the snout, and the standard length from the tip of the snout to the end of the body (beginning of the tail).
   3. Calculate the condition factor using the following formula:
      
      \[
      \text{Condition factor} = \frac{\text{total body weight} - \text{gonad weight}}{\text{total length}^3}
      \]
      
   NOTE: Gonad weight is subtracted from the total body weight since gonads can contribute significantly to the total body weight, particularly in prespawn female fish.

   3. Obtain a blood sample.
      
      NOTE: Blood is most commonly taken from the caudal vein but can also be withdrawn from the dorsal aorta or by cardiac puncture.
      1. Extract a peripheral blood sample from the caudal vein with a 22 or 23 G needle on a 1 to 5 mL syringe, depending on size of the fish. Insert the needle anterior to the caudal area below the lateral line (Figure 1A and 1B). Angle it upwards until hitting the spine and then withdraw slightly. The vein is ventral to the overlying spine.
      
      NOTE: If blood smears will be made or serum is required, no anticoagulant is used. In most cases, plasma will be collected and, hence, an anticoagulant such as sodium heparin, EDTA or lithium is used to coat the needle and syringe and is also in the blood collection tube (e.g., vacutainer).
      2. Remove the needle and place into a sharps disposal container prior to putting the blood into the collection tube.
      
      NOTE: Blood can be held on ice but depending on subsequent analyses should be centrifuged as soon as possible.
      3. If nuclear abnormalities or differential blood counts will be evaluated, immediately place a drop of blood on duplicate clean glass microscope slides. Back a second slide at a 45° angle into the drop, which is then drawn across the surface by capillary action. Allow to air-dry.
      4. Centrifuge blood at 1,500–2,500 x g for 15 min to sediment the cells. Remove plasma/serum with a sterile transfer pipet, aliquot into cryogenic vials, and store at -80 °C.
Figure 1: Obtaining a blood sample from a fish. (A) A recently euthanized fish is laid on its side and the lateral line located. (B) A needle is inserted ventral to lateral line (arrow), angled upward until needle touches the backbone. It is then slightly withdrawn, and suction initiated to withdraw blood. Please click here to view a larger version of this figure.

4. **Conduct a necropsy-based health assessment on each fish.**

   NOTE: A number of publications illustrating and describing lesions and abnormalities are available.

   1. Document external abnormalities including lesions on body surface and fins (Figure 2), eyes and gills (Figure 3), external parasites such as leeches (Figure 2D), grubs or trematode metacercarial cysts (Figure 2D, 3B) and gill parasites (Figure 3D). Document type, location and size of observed abnormalities on data sheets, as well as photographically, if possible.

   2. Open the abdominal cavity (Figure 4A) using a scissors by cutting from the anal area to the operculum and then removing the flap of muscle to expose the internal organs.

   **NOTE:** If anterior kidney will be collected for immune function (see step 5 below) or samples collected for bacteriology or virology, the external body surface should be disinfected with 70% alcohol and those samples should be obtained before the necropsy performed. If tissues are only being used for visual observations, plasma analyses and histopathology sterile technique is not necessary.

   3. Document internal abnormalities (Figure 4) including general or focal discolorations of the various organs (Figure 4B-4D), presence of raised areas (Figure 4E), cysts, parasites, and size abnormalities (enlarged, atrophied).
Figure 2: Examples of visible lesions observed on body surface and fins of fish. (A) A small, slightly eroded lesion (arrow) on the lateral body surface. (B) A large reddened area (arrow) involving the caudal body surface. (C) Raised, black lesions (arrows) on the body surface and fins. (D) Leeches (white arrow) and small black spots (black arrows) on the fin. Scale bar = 3 mm. (E) A raised, multilobed, pale lesion (arrow) on the body surface. Please click here to view a larger version of this figure.
Figure 3: Examples of visible lesions of the gills and eyes of fish. (A) A pale area (arrow) within the lens of an eye. Scale bar = 5 mm. (B) White cysts (white arrows) and small black spots (black arrows) caused by trematode parasites on the operculum covering the gills (a). Scale bar = 1 cm. (C) A pale, eroded area (arrow) on the gill (a). Scale bar = 5 mm. (D) A gill that has been removed showing parasites (arrows) attached to the gill filaments. Scale bar = 2 mm. Please click here to view a larger version of this figure.
Figure 4: Examples of a necropsy and internal abnormalities of fish. (A) During a necropsy the fish is cut open (along the white arrow) and a flap of muscle (black arrow) removed to expose the gonad (a) and the spleen, being held by forceps and scissors. (B) Mottled liver (a), testes (b), intestine surrounded by adipose fat (c) and stomach (d). Scale bar = 5 mm. (C) Liver (a) with a dark red area (arrow), ovary (b) and intestines (c). Scale bar = 5 mm. (D) Liver with greenish discolored areas (arrows). Scale bar = 1 cm. (E) Example of a normal (a) and abnormal (b) testes with raised nodules. Scale bar = 1 cm. Please click here to view a larger version of this figure.

5. Obtain hepatosomatic index (HSI).
   1. Remove liver by severing the hepatic artery and the connective tissue of the anterior end. Gently lift out while trimming adhesions and other connections to intestine and adipose fat. Take care to not puncture the gallbladder. Weigh the liver.
   2. Calculate hepatosomatic index (HSI) using the formula:
   
   \[ \text{HSI} = \frac{\text{liver weight}}{\text{total body weight}} \]

6. Calculate the gonadosomatic index.
   1. Remove the gonads and weigh it.
   2. Calculate gonadosomatic index (GSI) using the formula:
   
   \[ \text{GSI} = \frac{\text{gonad weight}}{\text{total body weight}} \]
3. Preserve Tissues for Microscopic Pathology

NOTE: A number of fixatives including 10% neutral buffered formalin and Z-fix, a formalin-based fixative with zinc, can be used for preservation of tissue in the field. The latter is preferred if methods such as in situ hybridization or fluorescent antibody staining may be used.

1. Carefully cut but do not pull out tissue samples. Keep individual tissue pieces <2 cm in size and <5 mm thick for proper fixation. As a rule of thumb, use approximately 10x more fixative by volume than the tissue for proper preservation. Place all tissue samples from one fish in the same leak-proof container of the appropriate size, depending on the size of fish being sampled.
2. Place pieces of any external abnormalities in the fixative container. Also, include an adjacent piece of normal tissue.
3. Cut at least five 3–4 mm thick pieces of liver from various regions and place into the fixative container. Include normal and abnormal areas, if observed.
4. Depending on size, place a whole gonad or multiple pieces along one gonad into the fixative container.
5. Place either whole organs, if small, or pieces of all other organs (spleen, anterior and posterior kidney, gills, heart, intestine and stomach) in the fixative container. If abnormal tissue is observed, preserve an adjacent piece of normal tissue as well.

4. Remove the Otoliths for Age Analyses

NOTE: Age can be an important variable in fish disease/fish health studies. While a number of structures, including scales and spines, have been used for age determination, most studies comparing structures have found the otoliths to give the best results. Teleost fishes have three pairs of otoliths - lapillus, sagitta and asteriscus. Generally, the sagittal or lapillus otoliths are collected for aging although that may vary by species. Removal and aging techniques have been previously described.

1. Cut through the gill isthmus, and bend the head back. Strip away connective and muscular tissue around the inferior portions of the neurocranium to locate prootic bullae, a raised bony area.
2. Score or cut with bone cutters and crack to expose the otoliths. They can be seen with the naked eye.
3. Place otoliths in a labelled vial or a coin envelope and store at room temperature until analyzed for age by counting the rings or increments. If placing in a vial, open cap once returned to the laboratory and allow to thoroughly dry prior to storage.

5. Obtain Tissue for Immune Function Assays

NOTE: The anterior kidney is the major hematopoietic organ, the source of lymphocytes and macrophages for functional assays, and must be removed aseptically if cells will be cultured for functional assays, such as mitogenesis, phagocytic and killing ability of macrophages.
1. Spray the external surface of the fish with 70% ethanol. Use sterile scissors, a scalpel and forceps to open the abdominal cavity and remove the anterior kidney tissue, which is a dark red organ located anterior to the swim bladder.

2. Place the anterior kidney sample in media (e.g., Leibovitz's L-15) to keep the cells alive. Homogenize kidney samples with a sterile hand-held tissue grinder (e.g., Tenbroeck tissue grinder) into single cell suspensions. Hold on wet ice until returned to the laboratory.

6. Preserve Tissue for Nucleic Acid Analyses

NOTE: If downstream molecular analysis will be conducted, such as gene expression using transcript abundance or quantitative PCR (polymerase chain reaction), place the pieces of tissue to be assessed in an appropriate preservative (e.g., RNAlater stabilization solution) as soon as possible.

1. For RNA preservation, place two to three small (2–3 mm) pieces in the appropriate preservative at a 10:1 ratio of preservative volume to tissue.

NOTE: Samples should be shielded from sunlight or excessive heat and transported on wet ice.

2. For DNA preservation, place two to three small pieces of tissue into 95% ethanol (10:1 ethanol to tissue by volume). Then hold the samples on wet ice and then store at -20 °C.

Representative Results

Great Lakes Areas of Concern (AOC) are geographic areas that were designated due to impairments of various beneficial uses. One of the beneficial use impairments (BUIs) at many AOC is the fish tumors or other deformities. Millions of dollars have been spent for remediation and restoration of each of these areas in order to delist the various BUIs and ultimately the AOC. The criteria for delisting the fish tumor BUI differs from state to state (see epa.ohio.gov/portals/35/lakeerie/ohio_AOC_delisting_guidance.pdf and dnr.wi.gov/topic/GreatLakes/documents/SheboyganRiverFinalReport2008.pdf); however, as noted in the delisting documents, there is a requirement to determine the prevalence of liver tumors and in some cases skin tumors. In many cases, the prevalence is compared to a non-AOC reference site.

The fish tumor BUI was evaluated at three AOCs (St. Louis River, Milwaukee River and Sheboygan River) and a non-AOC reference site (Kewaunee River) on Lakes Superior and Michigan, utilizing a necropsy-based assessment of white sucker (Catostomus commersonii), followed by microscopic pathology of skin and liver tissue. Fish were collected from the Milwaukee, Sheboygan and Kewaunee rivers in 2012 and 2013 and from the St. Louis River in 2015 (unpublished data). Two hundred white suckers were assessed from Milwaukee, Kewaunee and St. Louis, and 193 from Sheboygan.

By definition, a tumor can be any swelling or raised area, although it is generally considered that a swelling caused by an abnormal growth of tissue with abnormal cells is either a benign or malignant neoplasm. White sucker collected from all sites exhibited a variety of external raised lesions including small, discrete white spots, larger white areas, slightly raised mucoid lesions and multilobed raised areas on the body surface and lips (Figure 6). Fish were weighed and measured to obtain a condition factor, external and internal abnormalities were documented, and skin and liver tissue was collected for histopathology.

Figure 6: Raised skin lesions observed on white sucker from the Great Lakes. (A) A discrete white spot on the body surface. Scale bar = 5 mm. (B) A slightly raised mucoid (arrows) and multilobed lesions (a) on the posterior body surface. Scale bar = 1 cm. (C) A large, multilobed lesion on the body surface. Scale bar = 1 cm. (D) Numerous multi-lobed lesions on the lips. Please click here to view a larger version of this figure.
The percent of fish with external tumors or raised discolored areas ranged from 15.5% at the St. Louis AOC to 58.0% at the Milwaukee AOC. In general, the discrete white spots were the least common visual lesion while the multilobed lip and body surface lesions were most common. The number of fish with observable liver nodules was low, ranging from 1.5% at Kewaunee and St. Louis to 2.5% at Milwaukee (Table 1).

### Table 1: Necropsy-based Observations of White Sucker Collected at Great Lakes Areas of Concern and a Reference Site (Kewaunee River), Presented as a Percentage.

| Visible Lesions          | Rivers and Year Sampled |
|--------------------------|-------------------------|
|                         | Kewaunee 2013 | St. Louis 2015 | Sheboygan 2012 | Milwaukee 2013 |
| Discrete white spots     | 16           | 3             | 3.1           | 5             |
| Mucoid                   | 20           | 9.5           | 9.8           | 30.5          |
| Multilobed               | 22.5         | 3             | 29.5          | 40            |
| Total Raised Skin Abnormalities<sup>a</sup> | 46           | 15.5          | 38.3          | 58            |
| Visible liver nodules    | 1.5          | 1.5           | 1.6           | 2.5           |

<sup>a</sup>Total number of fish with raised lesions. Some fish had multiple types of abnormalities.

Visual examination can be used to document the percent of fish with various abnormalities. However, to definitively diagnose the presence and type of neoplasia, tissues must be examined microscopically (histopathology). Upon microscopic examination, it was found that not all of the raised lesions were neoplastic. Many of the discrete white spots and the mucoid lesions, particularly at Kewaunee, were hyperplastic lesions rather than neoplasia (Table 2). Additionally, at Kewaunee and St. Louis, all of the skin tumors observed were benign papillomas. At Sheboygan and Milwaukee both papillomas and squamous cell carcinomas, malignant skin tumors, were observed (Table 2).

### Table 2: Microscopically Verified Neoplastic Lesions of White Sucker Collected at Great Lakes Areas of Concern and a Reference Site (Kewaunee River), Presented as a Percentage.

| Neoplasm Type                  | Rivers Sampled |
|--------------------------------|----------------|
|                               | Kewaunee 2013 | St. Louis 2015 | Sheboygan 2012 | Milwaukee 2013 |
| Papilloma                      | 21            | 5.2            | 30.5           | 37.5          |
| Squamous cell carcinoma        | 0             | 0              | 2.1            | 10.5          |
| Total skin neoplasms          | 21            | 5.2            | 32.6           | 48            |
| Bile duct neoplasms<sup>a</sup>| 2.5           | 4              | 6.2            | 9.5           |
| Hepatic cell neoplasms<sup>b</sup>| 1             | 0              | 2.1            | 8             |
| Total liver neoplasms         | 3.5           | 4              | 8.3            | 15.0<sup>c</sup> |

<sup>a</sup>Includes cholangioma and cholangiocarcinoma

<sup>b</sup>Includes hepatic cell adenoma and hepatic cell carcinoma

<sup>c</sup>Some fish had both bile duct and hepatic neoplasms

### Discussion

The necropsy-based assessment of fish health can be utilized on any fish species for which the investigator has an understanding of the normal appearance of both external and internal structures. Using a standardized approach allows for comparisons between sites and species as well as seasonal and temporal changes in a population. The findings can be used to identify effects associated with point and nonpoint sources of contaminants and to inform management actions. It can also be used to track improvements once management actions are initiated. The methodology can be modified to augment the documentation of visual external abnormalities in a variety of ways. Assessments, based only on visual observations, can be non-lethal, relatively inexpensive and data can be generated quickly for a large number of individuals. Consequently, they can be useful for exploratory or initial assessments, to monitor change over time or in combination with other indicators. If the length and weight of fish are measured during visual observations, the condition factor can also be calculated. Although assessments based only on visual observation do not provide information on cause or associated risk factors, long term trends of certain skin abnormalities<sup>46</sup> and biometric parameters<sup>46</sup> have indicated improvement in some areas associated with water quality improvements.

The necropsy-based assessment provides more information as internal organs are also examined and other condition factors such as hepatosomatic index and gonadosomatic index can be calculated. Goede and Barton<sup>23</sup> developed a field necropsy method that included blood parameters, biometric factors, the percentage of abnormalities, and index values for specific abnormalities. A refinement of the method included a severity rating for some variables that allowed for calculation of a health assessment index that could be compared statistically<sup>23</sup>. This health
assessment index has been used in regional site comparisons, and in combination with other biological indicators including plasma and histopathological analyses in the U.S. Geological Survey's Biomonitoring of Environmental Status and Trends Program evaluating potential effects of contaminant exposure in large rivers nationwide. A Fish Disease Index based on externally visible diseases and parasites, visible liver neoplasms and other histopathologically detected liver lesions has been developed and used extensively in the North Sea, Baltic Sea, and off Iceland. This index was found to be an important tool as an ecosystem health indicator.

There are some critical factors in conducting the necropsy-based assessment on fish. First, assessments must be conducted on fish immediately after death. Changes in organ color and consistency can occur fairly rapidly after death. Additionally, some parasites may leave the host soon after death. Second, it is important to know what is normal for the species of interest. For instance, some fish normally have fatty and consequently, pale livers, while for most species a pale liver would be abnormal. It is also important to recognize seasonal changes that naturally occur. Some fish will have color changes or develop breeding tubercles during the spawning season.

The limitations of the necropsy-based assessment as a method for fish health assessment include the inability to 1) consistently identify the "cause" of specific lesions and 2) identify effects that may not be visible to the naked eye. These drawbacks can be overcome with the addition of histopathology, molecular or cultural identification of pathogens and parasites, and gene expression. For instance, a "tumor" or raised lesion (swelling) may be actual neoplasia or it may be a parasite, inflammation, edema or hyperplasia (increase in number of normal cells), caused by chemical exposure, infectious agents or other irritants. As shown in the representative results, definitive tumor or neoplasia diagnosis requires microscopic pathalogy to identify the lesion type and severity (i.e., benign or malignant). Assessment of white sucker external "tumors" by visual observation overestimated the prevalence, particularly at the reference site. Many of the raised lesions were not neoplasms but rather hyperplastic lesions. It is currently not known whether these hyperplastic lesions are pre-neoplastic. Conversely, the observation of raised nodules in the liver significantly underestimated the prevalence of liver neoplasms. Hence, collection of tissue for microscopicpathalogy was necessary to adequately address the potential for delisting.

Disclosures

The authors have nothing to disclose.

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