## CHAPTER 2

# Sample Collection and Submission 

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## I. Introduction

The purpose of this section is to provide guidance to laboratory staff and user groups regarding the proper sample collection and submission procedures. This is an absolute necessity to assure that samples received by the Centers are acceptable specimens for definitive pathogen identification. Each sample submission must include a NWFHS Submission Form (Appendix B). Specimens will be stored, maintained and processed in accordance with protocols described in subsequent chapters of this Manual. Sampling procedures not found in this Manual will follow diagnostic procedures outlined in the AFS-FHS Bluebook titled Suggested Procedures for the Detection and Identification of Certain Finfish and Shellfish Pathogens, 2004, 6th edition. American Fisheries Society, Fish Health Section.

Additional, detailed information regarding sample collection and processing can be found for specific assays in the following chapters:

| Chapter | Sample Receipt and Laboratory Tracking |
| :---: | :---: |
| Chapter 4 | Standard Necropsy Procedures for Finfish |
| Chapter 5 | Bacteriology (Aeromonas salmonicida, Yersinia ruckeri, Edwardsiella ictaluri) |
| Chapter | ELISA for Detection of Renibacterium salmoninarum Antigen in Fish Tissue |
| Chapter | Corroboration Testing of Renibacterium salmoninarum by PCR |
| Chapter | Parasitology (Myxobolus cerebralis, Bothriocephalus acheilognathi, Ceratomyxa shasta) |
| Chapter | Corroboration Testing for Parasites by PCR |
| Chapter 10- | Tissue Culture of Fish Cell Lines |
| Chapter 11- | Virology |
| Chapter 12- | Corroborative Testing of Viral Isolates |
| Chapter 13- | Histology of Finfish |
| Chapter 14- | Non-Lethal Methodology for Detection of Fish Pathogens |
| Chapter 15- | Procedure for Revision to the National Wild Fish Health Survey Laboratory Procedures Manual |

## II. Considerations for Sampling Fish Populations

Disease Recognition and Action - The majority of sampling conducted under the Survey will occur when no external signs of disease exist in wild fish populations. However, the Survey may also be helpful in determining the cause of fish kills or monitoring wild populations when abnormal behavior patterns, external abnormalities, or high mortality are reported for natural fish populations. In these cases, an immediate response is needed to determine the cause of mortality and determine if infectious agents are present or if adverse environmental conditions exist (low dissolved oxygen, elevated temperatures, toxic algal bloom, water contaminants, etc.). The following offers guidelines for sampling fish under various scenarios:

- In clinical cases of disease ( $\geq 0.5 \%$ mortality/day) 10 moribund fish are generally sufficient to detect fish pathogens and make a disease diagnosis.
- In survey or monitoring situations where no excessive mortality or clinical disease is apparent, a larger sample size of 60 animals may be necessary to detect infection rates below $5 \%$. However, depending upon individual circumstances, sample sizes may vary between 10 and 60 fish. Samples should be examined from several stretches of a stream or body of water to ensure representation of the entire population. It is also extremely important that sampling techniques are optimal and laboratory assays as sensitive as possible to allow detection of fish pathogens, especially when sample numbers are small (<60 fish).
- Donor populations - in some cases, natural populations will be used as donor broodstock to provide gametes to a hatchery program involved with a captive broodstock program or restoration activities. When a disease history for natural population is needed, a minimum of 60 samples from spawning adults is required to detect pathogens at a $5 \%$ prevalence and a confidence interval of $95 \%$. Samples of choice are from spawning or post spawning individual female fish consisting of ovarian fluid and kidney/spleen tissue.

The priority for acceptable samples submitted for Wild Fish Survey testing is as follows:

- Live specimens or samples taken on site
- Iced specimens or tissues
- Preserved specimens or tissues


## III. Sample Collection

Prior to collecting samples, the Center will contact sampling personnel with instructions on the appropriate types of samples and numbers of fish needed. Partners collecting specimens need to provide at least one-week advance notice of sample collection. If advance notice is not given, lab personnel may not be available to receive and process the submission after it is collected and shipped. Samples that are not in an acceptable condition (either substandard or improperly packaged) upon arrival cannot be processed due to the poor sample quality and unreliability of pathogen testing.

The following instructions are general guidelines. User groups and or individual collectors should be properly trained in the use of these sampling procedures by a fish health specialist, pathologist, or technician prior to sampling fish. Different procedures are followed in bacteriology, virology, parasitology, ELISA, PCR, and histological analyses.

Further details regarding the procedures below are addressed in the appropriate chapters of this Manual.

- Live fish are preferred, and should be sampled immediately upon removal from the water.

If this is not possible, fish should be held on ice and processed within 1-2 hours after collection. Whole fish can be packaged live, or freshly killed (iced) for shipment to the Centers for processing within 24 hours. When individual tissues are collected, these should be kept cold with ice or artificial icepacks but should not be allowed to freeze (insulated from direct contact with ice).

- Samples for virology should be processed within 48 hours and inoculated into cell culture within 72 hours of sample collection. Upon specimen arrival determine the freshness of the fish. Criteria for this freshness are: smell, appearance of eyes, gills, internal organs and presence of postmortem stiffness (rigor mortis). If shipping is delayed beyond 24 hours, some tissues may not be suitable for processing.
- Euthanize the fish or group of fish. Use clean sterile dissecting instruments. Clean dissecting tools with soap and water and disinfect with alcohol between the sampling of each individual fish (species) or species group. The use of disposable tools is recommended for ELISA samples, as Rs antigen is difficult to remove by standard disinfection methods. Disinfect hands between the sampling of each fish (species) or species group. Before taking tissue samples, wipe clean any mucus or debris from the fish and then disinfect the outer surface of the fish by flooding with $70 \%$ alcohol. Proceed with necropsy of fish tissues.


## IV. Bacteriology

Bacteriological samples should be taken first to reduce the chance of contamination. If open sores or lesions are present swab these areas and streak the sample onto a BHIA agar slant. Discard swab and re-cap slant. The abdominal cavity is entered by cutting into the abdominal wall at the base of the pectoral fin with a pair of small sterile scissors or scalpel. The cut is continued dorsally to just below the lateral line. Start again at the base of the pectoral fin and continue the incision towards the posterior of the fish along the ventral abdominal wall to the vent. Stay slightly above the intestinal tract when making the incision so that it is not punctured, thereby contaminating the abdominal cavity and target tissues.

Use the butt end of a sterile inoculating loop or forceps to pull back the internal viscera and air bladder to expose the kidney. Stab the posterior kidney with a sterile forceps and collect a large piece of kidney tissue and streak it directly onto a BHIA agar slant or plate. The struck kidney tissue can be used for the virology or ELISA sample as well. Record sample number and date on tube or plate. Sample numbers can be any logical group of letters and numbers in order (RBT-1, RBT-2). Store tubes in a cool place such as cooler or cool room $\left(15-20^{\circ} \mathrm{C}\right)$.

## V. Virology

Kidney/Spleen: Dissect approximately 0.5 gm piece of kidney and/or spleen and place into a small WHIRL- PAK ${ }^{\mathrm{TM}}$ bag or snap-cap tube with a small amount of HBSS (Hank's Balance Salt Solution) to cover sample. Keep the HBSS cold at all times. Label bag with sample number, K/S and date. Up to 5 fish samples may be pooled in one bag or tube if appropriate. Keep samples on ice while in the field. Samples must be kept cold $\left(5-18^{\circ} \mathrm{C}\right)$, but do not freeze, during shipment to the Center.

Ovarian Fluid: Sexually mature females only. Remove a small amount of ovarian fluid from the oviduct using a pipettor or express ovarian fluid into a paper cup (approximately 1 mL per fish) Place the fluid in a small snap-cap or transport tube. Up to five (5) fish may be pooled into one sample. Label tube with sample number (OV-1, OV-2, etc.) and date. Keep samples cold ( $4{ }^{\circ} \mathrm{C}$ ) with pre-packaged ice packs, but do not freeze during collection or transport. Sample tubes should be placed within a WHIRL-PAK ${ }^{\mathrm{TM}}$ or zip- lock bag which is labeled with the stream or lake name, fish species, sample type, etc.

## VI. ELISA for Renibacterium salmoninarum (BKD)

Remove the remaining amount of kidney or sizable portion using individual, or disposable instruments. Place the kidney tissue into a snap-cap tube or small WHIRL-PAK ${ }^{\mathrm{TM}}$ bag. Up to fish five can be pooled into one sample if necessary to obtain a minimum tissue weight of 0.08 g per sample ( 0.08 g diluted $1: 8$ during processing will yield $560 \mu \mathrm{~L}$ for the ELISA assay). Label with sample number, KD-ELISA and date (similar numbering scheme as with bacteriology slants). Keep samples on ice for transport to the lab. Samples can be frozen at $-20^{\circ} \mathrm{C}$ for delayed processing, however if long-term storage is needed, $-70^{\circ} \mathrm{C}$ is recommended.

## VII. Parasitology

Myxobolus cerebralis (Whirling Disease):
Salmonid Fishes Only. For small fish remove the entire head and gill arches. For larger fish, take a cranial core sample and/or gill arches. If taking a core sample, see Chapter 9 Corroborative Testing of Parasites by PCR, for the target tissue site for Myxobolus cerebralis. Up to (5) fish may be pooled into one sample. Place samples in a WHIRL-PAK ${ }^{\mathrm{TM}}$ or zip-lock bag labeled with sample information. Pack samples on ice for transport to the CENTER.

Either during sample collection, or during processing in the laboratory, the head tissue is halved into two separate pieces to allow testing of one half by Pepsin-Trypsin Digest (PTD) and archiving of the second specimen for corroborative testing by PCR or histology. If tissue is halved during collection, label each half to allow correlation between archive and tissue to be used for PTD tissues. Recommend a notation of "PTD" and "ARC", as well as FISH ID

NUMBER on each 5-pool sample (e.g., PTD 1-5, PTD 6-10, ARC1-5, ARC 6-10, etc). Tracking the fish identification for each sample pool will facilitate corroborative testing by PCR. In this way, only the PTD positive sample pool will require corroborative testing of the archive sample by PCR, rather than the entire sample set.

Bothriocephalus acheilognathi (Asian Tapeworm) and Ceratomyxa shasta (salmonid ceratomyxosis):

Remove the GI track of the fish from the esophagus to the anus. Place GI track into WHIRL-PAK ${ }^{\text {TM }}$ or zip-lock bags. Small fish of the same species can be pooled ( 5 fish) if applicable. Label samples with appropriate information. Pack samples on ice for transport to the Center.

## VIII. Histological Samples

Histological samples should be fixed in Bouin's solution, $10 \%$ buffered formalin, Davidson's fixative, or Prefer fixative. Fix live fish after anesthetizing. Fix tissues within 2 minutes of removal from water and/or time of death. Fish rapidly autolyze (especially gill) and only freshly fixed tissues are worth processing for histological analysis. The volume of fixative must be at least 10 times the volume of tissue. For fish longer than 6 cm , slit the abdomen, detach the intestine at the anus, and pull the internal organ mass out slightly to allow penetration of fixative within the body cavity. For larger fish, send only specified organs in fixative. Cut tissues with a sharp blade or scissors - don't tear as this action creates artifacts.

## SAFETY NOTE! formalin-based fixatives are toxic and strong irritants.

 Read the entire Material Safety Data Sheet. Avoid contact with skin and eyes by wearing gloves and a face shield. Use only in well-ventilated areas (outdoors or under a fume hood).HAZARDOUS MATERIALS - Fixatives and alcohol require special shipping procedures as Dangerous Goods, or Hazardous Materials. Refer to your local regulatory agency and commercial carrier for requirements for shipping these materials.

Place a paper penciled label (location, date, species, tissue type, and initials) inside the fixative container (alcohol and fixatives tend to wash off pen marks on the container).

Prevent spills during transport. Tightly cap the container, then wrap the cap with several layers of parafilm, and place the containers inside Ziploc bag(s). If samples fixed in Bouin's or Davidson's Fixative cannot reach the laboratory in 48 hrs , it will be necessary to transfer fixed tissues to 70\% ethanol after 24-48 hours and then transport the tissues to the Center (Samples fixed in $10 \%$ Buffered Formalin or Prefer fixative can be held in fixative indefinitely).

## IX. Non-lethal Collection of Tissue Samples

Compliance with the Endangered Species Act (ESA) of 1973 requires special consideration regarding take of threatened or endangered (T\&E) species if they occur in a proposed sampling site or watershed. All Centers and Partners in the Survey are responsible for obtaining appropriate collection permits, coordinating sample collection with Federal, State and local regulatory agencies, and fully complying with the regulatory statutes of the Endangered Species Act.

When lethal sampling of T\&E species is prohibited, non-lethal sampling techniques should be considered. While non-lethal sampling methods are less sensitive than standard detection methods, they may provide limited fish health information when no other alternative exists. Refer to Chapter 14 -Non Lethal Methodology for Detection of Fish Pathogens for specific protocols for non-lethal sampling.

## Bacteriology

Blood Samples
Obtain blood via heart puncture or caudal vein or artery using a needle and syringe. Streak blood onto BHIA slant with sterile loop. Discharge appropriate amount of blood directly into the appropriate volume of PBS-T20 for ELISA. Heparinized blood may be used for delayed transport and processing.

## Ovarian Fluid Samples

If the fish is a sexually mature female, remove a small amount of ovarian fluid and inoculate a BHIA plate for bacterial growth. Also, the same sample can be placed into a centrifuge tube. Up to five (5) fish may be pooled into one sample. Keep samples cold for transport, but do not freeze. Following centrifugation and processing for virology, the Ovarian Fluid Pellet can be used to screen for Renibacterium salmoninarum by FAT.

## Mucus Samples

Pass a sterile swab along the lateral surface of the fish. Streak the sample onto a BHIA agar slant.

## Vent Samples

Place a sterile swab or loop approximately $1 / 2$ to 1 inch into the anal vent and remove. Streak the sample onto a BHIA agar slant. (Discard the swab or loop).

## External Lesions

Pass a sterile swab along the surface of an external lesion. Streak the sample on selective media for the targeted bacteria. Plate serial dilutions of the inoculum to decrease the number of interfering bacteria and fungi that are likely to be present in this type of sample.

## Virology

## Blood Samples

Obtain blood via heart puncture or caudal vein or artery using a needle and syringe. Discharge appropriate amount of blood directly into the appropriate volume of antibioticantimycotic incubation (anti-inc) solution. Keep samples cold for transport, but do not freeze. Follow normal processing protocol.

Ovarian Fluid Samples
If the fish is a sexually mature female, remove a small amount of ovarian fluid and place into a centrifuge tube. Up to five (5) fish may be pooled into one sample. Keep samples cold for transport, but do not freeze. Follow normal processing protocol.

## Mucus Samples

Collect mucus by passing a blunt edge instrument along the lateral surface of the fish, head to tail. This is easily accomplished when the fish is removed from the water, holding the fish head to tail in a vertical position. Mucus is placed in a $15(\mathrm{~mL})$ or smaller graduated centrifuge tube with antibiotics used in viral sample processing. Keep the samples cold for transport to the Center, but do not freeze. In the lab, samples should be vortexed, and then a low dilution scheme ( $1: 2,1: 5,1: 10$ ) is set up using Hanks Balanced Salt Solution (HBSS). Samples are centrifuges at low speed, 2000-3000 rpm for ten to pellet cellular debris. Supernatant is inoculated into cell culture.

## Fecal samples

Feces are collected by aspiration with a syringe and small tubing catheter. Samples are placed into small collection tubes with antibiotics (may need to increase concentrations of antibiotics by $25-50 \%$ for fecal samples). Keep the samples cold for transport, but do not freeze. Follow processing as in mucus samples (may require higher dilutions to avoid toxicity to cell lines).

## External Lesions

Pass a sterile swab along the surface of an external lesion. Place the swab in anti-inc solution pressing, or rolling the swab against the interior of the tube to release the material into solution. Keep samples cold for transport, but do not freeze. Follow normal processing protocol.

## Parasites

## Blood Samples

Obtain blood via heart puncture or caudal vein or artery using a needle and syringe. Discharge appropriate amount of blood directly into a slide and prepare a thin blood film (Chapter 4, page 11 ). Blood films are air dried, and fixed in absolute methanol for 10 minutes. Blood parasites can be viewed following staining with DiffQuick or a Gram stain (Chapter 4, page 10).

Mucus Samples
Collect mucus by passing a blunt edge instrument along the lateral surface of the fish,
head to tail. This is easily accomplished when the fish is removed from the water, holding the fish head to tail in a vertical position. Mucus is placed in a 15 (mL)
or smaller graduated centrifuge tube. The tissue can be examined directly under microscopy for parasites or kept cold for transport and examined in the laboratory.

## Fecal samples

Feces are collected by aspiration with a syringe and small tubing catheter. Samples are placed into small collection tubes. The tissue can be examined directly under microscopy for parasites or kept cold for transport and examined in the laboratory.

## External Lesions

Pass a sterile swab along the surface of an external lesion. Place the swab in normal saline or PBS solution pressing, or rolling the swab against the interior of the tube to release the material into solution. The tissue can be examined directly under microscopy for parasites or kept cold for transport and examined in the laboratory.

## Tissue Biopsy

Gill filament can be removed from anesthetized fish with little injury to the fish. The tissue can be examined directly under microscopy for parasites, preserved for histology, or frozen for examination using other diagnostic methods. See Chapter 14 for more a detailed protocol for gill biopsy.

Water and sediments sampling for virus, bacteria and parasites is also discussed in Chapter 14 -Non-Lethal Methodology for Fish Pathogens.

## X. Shipping Samples - See Appendix A for shipping addresses of Centers

A. Ship samples in small/medium (heavy duty) mailing cartons or plastic coolers lined with a plastic trash bag. Group the samples by type in separate Ziploc bags or racks that are labeled with the number of samples, location, species, and date. Position the samples upright and use packing material to hold samples in place.
B. Place an adequate amount of ice (or artificial gel packs) around the insulating layer of packing material. Seal the outer trash bag to prevent leakage.
C. Complete the NWFHS Submission Form (Appendix B) for each species and enclose in a waterproof plastic bag within the cooler.
D. Close, seal and label the ice chest with laboratory address, be sure to include the Center contact's NAME AND PHONE NUMBER).
E. Appropriate shipping labels should also be affixed to ensure proper handling during shipment and upon receipt of containers.
"Live Fish - Do Not Freeze" ...........for live samples
"Keep Frozen" ...........for ELISA and/or Head tissue
"Refrigerate but DO NOT FREEZE"...........for virology or sets of samples that include both Bacteriology and Virology samples.

NOTE: Temperature indicator strips can be placed with the samples.
These strips will change color if $20^{\circ} \mathrm{C}$ has been exceeding during shipment.
F. Transport within 24 hours via overnight U.S. mail or Federal Express.

NOTE: Collectors should always include a Submission Form with samples to provide the collection information for the laboratory tracking and database entry. Samples submitted without this information may be refused or at least delayed for processing. When in doubt regarding collection and shipping instructions, consult the Center contact for the Survey.

Centers are prepared to provide all supplies needed for field sampling and training in sample collection. Transportation costs, personnel to assist with sampling and shipping costs may be provided if possible.

## XI. Bibliography

AFS-FHS. 2004. Procedures for the detection and identification of certain finfish and shellfish pathogens. $6^{\text {th }}$ edition. Fish Health Section, American Fisheries Society.

CA-NV FHC, 1997. Histological sampling of fish tissues. S.O.P., California-Nevada Fish Health Center. Anderson, CA. 2 pp.

Lasee, B.A., editor. 1995. Introduction to fish health management. $2^{\text {nd }}$ edition. U.S. Fish and Wildlife Service, La Crosse Fish Health Center. Onalaska, WI. 139 pp.

Meyers, Theodore R., editor 1997. Fish pathology section laboratory manual, special publication No. 12, Alaska Department of Fish and Game.

Mitchell, A.J., and G.L. Hoffman. Submitting samples for fish disease diagnosis. U.S. Fish and Wildlife Service, Fish Farming Experimental Station, Stuttgart, AR. 15 pp.

Plumb, J.A., and P.R. Bowser. 1983. Microbial fish disease laboratory manual. Alabama Agricultural Experimental Station, Auburn University, AL. 95 pp.

Thoesen, J.C., (ed). 1994. Suggested procedures for the detection and identification of certain finfish and shellfish pathogens (4 $4^{\text {th }}$ Edition). Fish Health Section. American Fisheries Society, Bethesda, MD.

## Appendix 2.A - <br> Shipping Addresses and Contacts for Fish Health Centers

| Bozeman Fish Health Center | Crystal Hudson <br> Kenneth Peters |
| :--- | :--- |
| Bozeman, MT 59718, Suite G | $406-582-8656$ |
|  | $406-587-3998$ (fax) |
|  |  |
|  |  |
| California-Nevada Fish Health Center | Scott Foott |
| 24411 Coleman Hatchery Road | Kimberly True |
| Anderson, CA 96007 | $530-365-4271$ |
|  | $530-365-7150$ (fax) |

Idaho Fish Health Center Kathy Clemens
P.O. Box 272

Orofino, ID 83520
Laura Kessel
FOR SHIPPING:
208-476-9500
4447 Ahsahka
Ahsahka, ID 83520

La Crosse Fish Health Center 555 Lester Avenue, Suite 100 Onalaska, WI 54650-8552

Lamar Fish Health Center
P.O. Box 155

Lamar, PA 16848

FOR SHIPPING:
Hatchery Road
Lamar, PA 16848

Rick Nelson
Becky Lasee

608-783-8444
608-783-8450 (fax)

John Coll
Patricia Barbash

570-726-6611
570-726-7379 (fax)

| Lower Columbia Fish Health Center 61552 SR14 <br> Underwood, WA 98651 | Susan Gutenberger Ken Lujan |
| :---: | :---: |
|  | 509-493-3156 |
|  | 509-493-2748 (fax) |
| Olympia Fish Health Center 3859 Martin Way E., Suite Olympia, WA 98506 | Ray Brunson |
|  | Chris Patterson |
|  | 360-753-9046 |
|  | 360-753-9403 (fax) |
| Pinetop Fish Health Center P.O. Box 160 Pinetop, AZ 85935 | John C. Thoesen |
|  | Phil Hines |
|  |  |
|  | 928-367-1902 |
| FOR SHIPPING: | 928-367-1957 (fax) |
| 1684 East White Mountain Blvd., Suite 7 Pinetop, AZ 85935 |  |
|  |  |
| Warm Springs Fish Health Center 5151 Spring Street Warm Springs, GA 31830 | Norm P. Heil Robert Bakal |
|  |  |
|  | 706-655-3382 Ext. 233 |
|  | 706-655-3389 (fax) |

## Appendix 2.B - NWFHS Submission Form



## Species and Sample Information

| SPECIES: | No. FISH | No. Samples Pool Size | Sample ID Numbers |
| :--- | :--- | :--- | :--- |
| Viral Tissue: |  |  |  |
| Bact Cultures: |  |  |  |
| R.sal Tissues (KD): |  |  |  |
| Parasite Tissues: <br> Type: |  |  |  |
| Type: |  |  |  |
| Comments: |  |  |  |


| SPECIES: | No. FISH | No. Samples Pool Size | Sample ID Numbers |
| :--- | :--- | :--- | :--- |
| Viral Tissue: |  |  |  |
| Bacti Cultures: |  |  |  |
| R.sal Tissues (KD): |  |  |  |
| Parasite Tissues: <br> Type: |  |  |  |
| Type: |  |  |  |
| Comments: |  |  |  |


| SPECIES: | No. FISH | No. Samples Pool Size | Sample ID Numbers |
| :--- | :--- | :--- | :--- |
| Viral Tissue: |  |  |  |
| Bacti Cultures: |  |  |  |
| R.sal Tissues (KD): |  |  |  |
| Parasite Tissues: <br> Type: |  |  |  |
| Type: |  |  |  |
| Comments: |  |  |  |

## National Wild Fish Health Survey Submission Form: Sample Tracking

Location:
Date:

| Fish <br> No. | Length (mm) | Weight $\qquad$ | $\begin{gathered} \text { Sex } \\ (\mathrm{M} / \mathrm{F}) \end{gathered}$ | BHIA \# | Kidney \# (ELISA) | Virology sample \# | Head (pool \#) | other | other | Comments |
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| Total \# | mples (p | Is) sub | itted |  |  |  |  |  |  |  |

## NOTICE

In accordance with the Paperwork Reduction Act (44 U.S.C. 3501), please be advised that:

1. The gathering of information is authorized by the Fish and Wildlife Act of 1956 (16 U.S.C. 742f), the Wildlife Coordination Act (16 U.S.C. 661-666c), and the Anadromous Fish Conservation Act (16 U.S.C. 757a - 757g).
2. Failure to provide all of the requested information is sufficient cause for the U.S. Fish and Wildlife Service to deny your request for Aquatic Animal Health Inspection under 713 FW 4.
3. You are not required to respond to a collection of information unless it displays a currently valid OMB control number.
4. This information collection has been approved by OMB and assigned clearance number 1018-XXXX.
5. The requested information may be subject to disclosure under provisions of the Freedom of Information Act (5 U.S.C. 552).

The public reporting burden for the information collected on this form is 15 minutes. This burden estimate includes time for reviewing instructions, gathering data, and completing and reviewing the form. Comments on this form should be directed to the Information Collection Officer, Mail Stop 222, Arlington Square, U.S. Fish and Wildlife Service, Washington, DC 20240.

