



**Standard Operating Procedure**  
**Laboratory Methods: NEON-Zooplankton Version 1.0**

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## Contact

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# **STANDARD OPERATING PROCEDURE**

## **LABORATORY METHODS: NEON-ZOOPLANKTON**

### **Scope and Applicability**

This standard operating procedure (SOP) is used to establish a uniform format for sample handling and analysis for samples from the National Ecological Observatory Network (NEON). This method is applicable to all samples received from NEON. All staff at Limnopro Aquatic Science, Inc., Saint Cloud, MN, performing zooplankton identification and measurements associated with NEON contracts must follow this SOP without deviation.

### **Sample Receiving**

1. To receive samples, collect the delivered box as soon as possible, and store them at room temperature until sample check-in can be performed. Proceed to download Battelle-generated chain of custody (COC) and associated electronic files.
2. To collect the COC, within 72 hours of shipment receipt, search email for the electronic chain of custody (COC) that will come from [CLANotify@battelleecology.org](mailto:CLANotify@battelleecology.org) email address. Each COC contains a sample receipt file (file name prefix "receipt\_form") that serves as the electronic version of the COC (e-COC). Check the electronic file into the laboratory information management system (LIMS). Proceed to unpack sample bottles and check for their condition.
3. To unpack samples and check for their condition, remove each individual sample bottle from the box and line them up on the processing table to ensure each is in good condition. Specifically, check each sample for container damage, preservative leakage, and for labels being present, complete and legible. Any leakage or damage, and any discrepancies between sample labeling and the e-COC document must be noted. For any samples received in compromised condition, take a digital photograph of the sample, record the nature of the compromise on the e-COC. If possible, rectify the compromised samples by adding preservative (40% ethanol), moving sample to new bottle, and/or rewriting labels. Create a notation on the e-COC for any rectifying activity on a sample bottle. Email Battelle to report any problems with the shipment and include digital photograph of compromised samples within 72 hours. After unpacking samples and checking condition, immediately check in samples.
4. To check in samples, the label of each sample container must be checked against the e-COC and marked off as they are identified. For each bottle identifier that matches the sample code in the receipt\_form, type in "Y" in the "sampleReceived" field. For any discrepancies between sample labels and e-COC, take a digital photograph that clearly shows the discrepancy. Email Battelle to report any problems with the shipment and include a digital photograph of label mismatch within 72 hours. Once all samples have been checked off against the e-COC as

instructed by the COC email, upload the completed e-COC to the NEON data ingest portal, which is to be accessed at [data.neonscience.org/web/external-lab-ingest](http://data.neonscience.org/web/external-lab-ingest). Place a round green sticker on the sample bottle indicating the sample bottle has been checked in and is ready for processing. Repack samples for storage.

5. To repack samples for storage, ensure caps are tightly sealed on storage bottles, that they contain ethanol by noting the mark on the label or checking for odor if not and store them at room temperature until sample processing occurs.

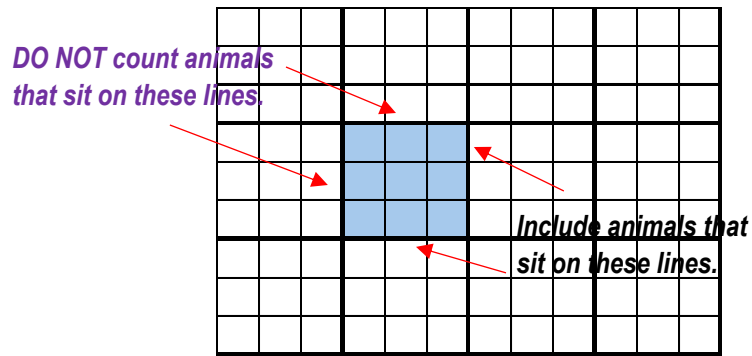
### **Sample Processing**

1. To prepare a sample for processing, randomly select a bottle from a shipment in the order shipment was received. Open the bottle and observe the liquid to make a judgement about whether the sample is dilute and requires concentration. The majority of samples will require concentration, but occasionally, abundant zooplankton observed will indicate sample can be processed without. If no concentration is required, advance to step 3 below.
2. To concentrate sample, pour the entire volume of the bottle as sent through a 37-micron mesh net (i.e., “concentration net”) and collect the filtrate in a 1L glass wide-mouth jar that has been tripled rinsed with tap water. Once the sample has filtered through the concentration net, pull the concentration net away from the 1L glass jar filtrate reservoir. Fill up the original empty sample bottle approximately one third of way with tap water, cap and invert several times. Uncap and pour the water with residual animals through the concentration net allowing the water to pass through and drain down the sink. Repeat this process of rinsing the original bottle out three times. If the sample contains a separate mesh net that was packed with the sample prior to shipping (i.e., “sampling net”), pull it out, and holding the 37-micron concentration net that contains all the animals over the sink, use a rinse bottle with a wide-bore tip filled with tap water to rinse any animals that might be adhered to the sampling net into the concentration net. Place the sampling net back into the original sample container. Using the same rinse bottle with tap water, concentrate animals to the apex of the concentration net. Place a funnel into the 50 ml Falcon tube. Flip over the concentration net mesh and rinse the sample through a funnel into the Falcon tube using 40% ethanol rinse bottle filled with the rinsate collected from the 1L glass wide-mouth jar. Fill the tube to between 25-50 ml of the ethanol, as much as is required, to rinse the animals off the concentration mesh. If animals are observed to be highly concentrated, fill the Falcon tube all the way to 50 ml with additional 40% ethanol. Add one drop of DAWN dish detergent to the Falcon tube to help with the clumping of organisms. Using the LIMS, create, print off, and adhere a 1 x 1.5 inch label for a 50 ml Falcon tube. Proceed to sample concentration. Pour the filtrate from the glass-mouth jar and the 40% ethanol rinse bottle, and place the sampling net (if present), back into the original sample jar. Replace the green round sticker on the bottle with a yellow sticker to indicate the sample is currently being processed. Proceed to prepare a bench sheet for identifications and counting.

3. To prepare a bench sheet for identification and counting, obtain a bench sheet and record the sample code, sample site/station, sample date, volume of original sample, volume of the sample as concentrated into the Falcon tubes, the date for start of identification of sample and the taxonomist name on the appropriate line. Proceed to subsample from the Falcon tube.
  
4. To subsample, target a minimum of 300 animals per sample. Rotate the concentrated sample in the 50-ml Falcon tube gently 10 times to mix the contents. Immediately, insert a pipette with a disposable 1 mL wide-bore pipette tip into the sample and further homogenize, mixing it in a random fashion by pumping the pipette up and down several times. A 1 mL subsample is captured during this mixing process to avoid bias due to sinking of heavier planktonic organisms. Add subsampled contents to a gridded Sedgewick Rafter Cell. Allow animals to sink and settle and stop swirling for 5 – 10 minutes. Identify animals to the following target resolution or lowest practical level:

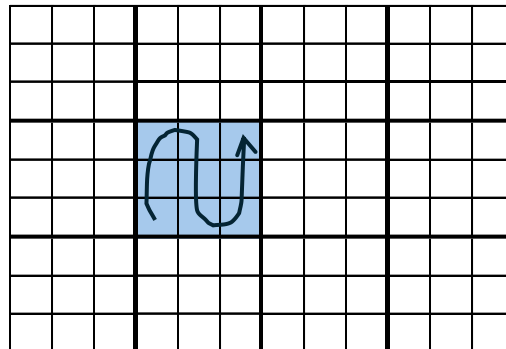
<b>Taxonomic group</b>	<b>Target</b>
Cladocera	Genus
Cyclopoida	Order
Calanoida	Order
Anostraca	Genus
Rotifera	Genus
Hydracarina	Genus
Diptera	Family
Harpacticoida	Order
Ostracoda	Class

Identify and count all zooplankton under a compound microscope using 40X magnification by moving up and down columns of 9 grid cells (3 x 3) per field. Count and identify animals that overlap on the right and bottom grid lines but not on the left or top as viewing the slide. Monitor liquid levels on the slide under the microscope as the ethanol will evaporate quickly. Add distilled water as necessary to keep organisms from drying out on the side.



**Illustration 1.** Count all zooplankton within the nine-grided blue area of a Sedgewick Rafter slide and then move up to the next 9 grid blue area and so on until all slide has been counted. A nine-grided field is an ideal field under a compound scope at 40X magnification. Each individual grid is 1 mm<sup>2</sup>.

If higher magnification is required to identify smaller rotifers or where animals are highly concentrated, move through individual grids one at a time through the 3 x 3 grid.



**Illustration 2.** If animals are small or densely packed on the slide, it may be necessary to use higher magnification and scan each of the nine-grids individually as shown.

Once the entire Sedgewick Rafter slide has been processed, determine if a minimum of 300 animals has been counted. If not, add another 1 mL of the concentrated sample and repeat the process until 300 animals have been counted. Record the final volume counted to the top of the bench sheet in the appropriate space. Note that the goal is to process between 1.0 – 3.0 ml of the concentrate. If after processing the first slide, there are fewer than 100 animals, the subsample is too dilute. If this is the case, repeat above “Sample Processing: Step 2” but decrease the volume used in the concentration step in the Falcon tube to a value < 25 ml that will yield 300 plus organisms in 1 ml subsample. Once enough animals have been identified and counted, proceed to measure.

5. To measure animals, scan through the sample from the beginning of the Sedgewick Rafter Slide. Use the microscope camera to photograph 15 individuals selected as first encountered for each taxonomic unit. Use the software to measure the length and width of each rotifer genera or length only for all other taxonomic units. All measurements are linear and made as follows:
  - Copepod measurements are total length including the caudal rami, excluding antennae and terminal setae of caudal rami.
  - Cladoceran measurements are total length, including helmets and excluding tail spines/mucros.
  - Leptodora lengths are taken from the distal end of the ocular lobe to the posterior end of the abdomen, excluding the furcae. If specimens are bent, usually two measurements are taken and summed.
  - Chaoborus length measurements are from the anterior point of the head capsule to the posterior end of the abdomen. If specimens are bent, usually two measurements are taken and summed.
  - Rotifer lengths exclude anterior and posterior spines (posterior spine is included in single-spine Keratella, e.g.- Keratella cochlearis). Widths taken are maximum widths of the body.

Prior to each series of measurements, calibration of the software will be checked using a 0.01 mm micrometer. Measurements are binned to the nearest 0.01mm for rotifers and 0.1mm for all other taxonomic units. Body fragments should not be counted unless the head is attached. If damaged, the record may be recorded as “Sample condition” = “Damaged, affecting measurement”. Record measurements to the bench sheet. Proceed to create a voucher (5a) if a new genus is encountered not currently part of the NEON taxonomic database. If no new genera are encountered, proceed directly to repacking samples for return (5b).

6. To create a voucher and/or repack samples, do as follows:
  - a. To create a voucher for a new taxonomic unit not already in the reference collection, create a slide mount and digital image of the individual(s). Record any individuals removed from the final processed sample for the reference collection or slide mounts. Save image files labeled with the original NEON sample ID and taxon abbreviation or scientific name to the Biorepository’s Dropbox folder for inclusion into the Symbiota portal. File names should not have spaces. Create a supplemental datasheet using client provided format detailing image file names, scientific names, identified by, etc. to be included in the Dropbox. Proceed to repack sample for return.
  - b. To repack samples for return for archiving at the NEON biorepository, first return the remaining concentrated sample in the 50 ml Falcon tube back into the original container. Remove some of the filtrate from the original bottles and pouring the contents of the Falcon tube back into the original bottles. The emptied Falcon tubes

will be triple rinsed using a small amount of 40% ethanol back into the original bottles. Once emptied, the Falcon tubes will be discarded. Once the contents have been returned to the original bottles, the bottles will be brought back to full using 40% ethanol. At that point, the yellow round sticker on the original bottle will be replaced with a red round sticker, indicating the sample has been processed and complete. Store the sample at room temperature until return to the client. Proceed to enter bench sheet data to LIMS.

### **Quality Control**

Quality Control (QC) requires the recount and reidentification of 10% of all samples in their entirety from a taxonomist different than the original taxonomist. A random number generator will select a minimum of 10% of samples. Samples selected for QC will be done at the same date and sequentially in time by two different taxonomists. Once QC samples have been distributed to the Sedgewick Rafter slide, taxonomist 1 will proceed to count and identify all zooplankton as described above and report data to a bench sheet. Following this, taxonomist 2 will do identifications and counts on the same slide and record data to their own bench sheet. If after the first distribution of animals is counted on the slide does not reach 300 organisms at a minimum, the slide will be rinsed and an additional distribution from the samples will be made to the slide and the process will continue until a minimum of 300 animals are counted.

Once a minimum of 300 animals are counted for the original count, the following will be used in order to determine both the Percent Taxonomic Difference (PTD) and Percent Difference in Enumeration (PDE) (Stribling et al. 2008) will be estimated as described below.

1. Percent Taxonomic Differences (PTD): Percent taxonomic difference must be  $\leq 15\%$  between the sample identified by Taxonomist 1 and the re-identified (quality checked) sample identified by Taxonomist 2 (Stribling et al. 2008).

$$PTD = \left( 1 - \left[ \frac{a}{N} \right] \right) 100$$

where  $a$  is the number of matches between taxonomist 1 and taxonomist 2 for a single sample and  $N$  is the total number of individuals identified in the larger of the 2 counts for the sample.

If PTD is  $> 15\%$ , the taxonomist 2 will update taxonomic identifications and reconcile any other data that have been affected. The QC results will be presented as numerical values in the required datasheet and an indication that QC was performed for a given sample will be recorded.

2. Percent Different in Enumeration (PDE): Percent difference in enumeration must be  $< 5\%$  between the sample enumerated by Taxonomist 1 and the re-identified (quality-checked) sample enumerated by Taxonomist 2.



$$PDE = \left( \frac{|n_1 - n_2|}{n_1 + n_2} \right) 100$$

where  $n_1$  is the number of individuals counted by taxonomist 1 and  $n_2$  is the number of individuals counted by taxonomist 2.

If PDE is >5% taxonomist 2 will update counts per taxon and reconcile any other data that have been affected. The QC results will be presented as numerical values in the required datasheet and an indication that QC was performed for a given sample will be recorded.

After QC is performed, values will be entered on NEON standard datasheets. Proceed to sample archiving.

### **Zooplankton Sample Archive Preparation**

All completed processed samples will be stored at room temperature preserved in 40% ethanol for up to six months following data return to allow NEON to perform a post-analytical review of the data. After the six-month period, Limnopro will ship the archive samples to the NEON Biorepository, located at Arizona State University in Tempe. Every sample/subsample sent to the Biorepository for archiving will get a unique identifier and QR Barcode that is specified on the labels and entered/submitted in the respective Biorepository shipment datasheets. Container size and labeling requirements will be matched to NEON instructions.

### **Zooplankton Data Return**

Data from these samples will be uploaded to the NEON portal prior to shipping to the Biorepository based on the following schedule

<b>Data Ingest Template</b>	<b>Type of Data Return</b>	<b>Upload Frequency</b>
<b>General Data Return</b>		
Receipt form – provided as an email attachment with each automated shipping notification, used to report condition of each sample/shipment	'receipt' load group on NEON Data Portal	Upload after the receipt of each sample shipment
<b>Zooplankton Taxonomy Data Return</b>		
Attachment2g_zooplankton_perSample	Zooplankton_Lab_Data' load group on NEON Data Portal	Each completed analytical batch
Attachment2h_zooplankton_perTaxon	Zooplankton_Lab_Data' load group on NEON Data Portal	Each completed analytical batch
Attachment2i_zooplankton_perVial	Zooplankton_Lab_Data' load group on NEON Data Portal	Each completed analytical batch
<b>Taxonomic Table Update Request</b>		

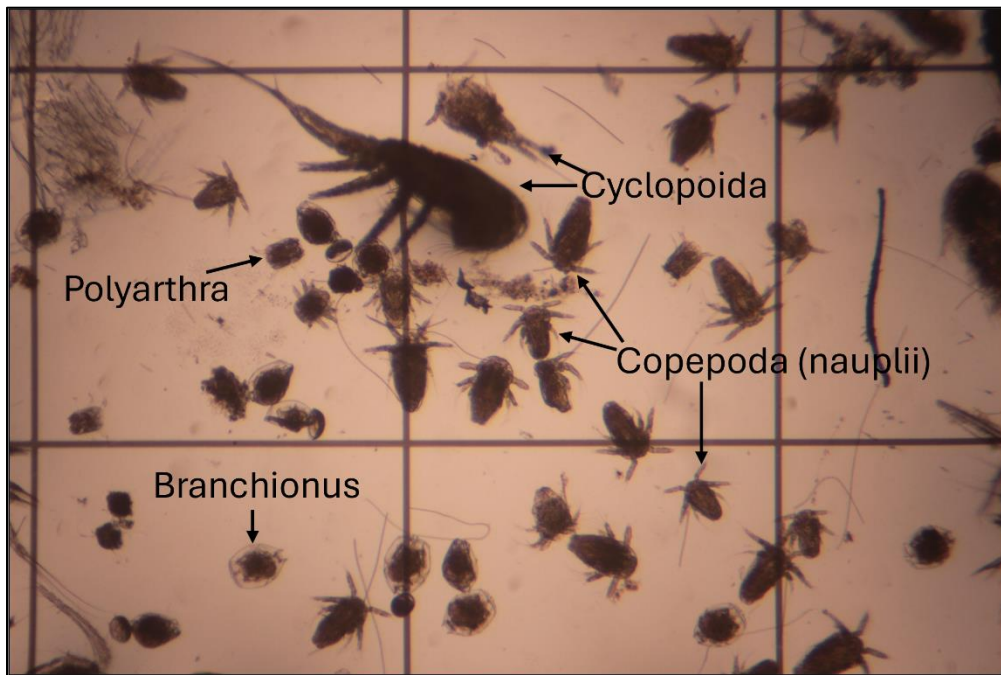
Attachment2k_taxon_request_template	Email to NEON technical representative	Up to 1 request per month
<b>Biorepository Shipments</b>		
Attachment3a_template_receipt_form_for_Biorepository shipments	shipment' load group on NEON Data Portal	Per Shipment to Biorepository

## Processing Equipment



Processing supplies including 50 ml Falcon tubes, original sample bottles, 1 L glass jar with attached funnel and 37 micron net, pipette with wide bore 1 mL tips and Sedgewick Rafter slide.

## Example Slide



Example camera view at 40X of diversity of zooplankton on Sedgewick Rafter slide with grids measuring 1 mm<sup>2</sup>.