

Mosquitoes sampled from CO2 traps (DP1.10043.001)

Measurement

Mass, taxonomic identification, and sex of mosquitoes collected in CO2 traps.

Collection methodology

Traps consist of fans attached to insulated coolers filled with dry ice and hung from a Shepard's hook. A small hole in the cooler ensures that as the dry ice sublimates, the CO2 escapes. This is the lure for mosquitoes. The fan distributes the CO2 around the catch cup located under the cooler. Traps are set for day and night collection. The catch cup is tied off and stored on dry ice while the insulated cooler is refilled with dry ice. Mosquitoes are then stored in the -80°C freezer and obvious bycatch is removed. Samples are sent to the external laboratory in falcon tubes with tissue paper as padding to protect specimens.

For information about disturbances, land management activities, and other incidents that may impact data at NEON sites, see the Site management and event reporting (DP1.10111.001) data product.

Data package contents

mos_trapping: Mosquito trapping data from field operations

mos_barcoding: Mosquito barcoding sample data from external labs mos_archivepooling: Mosquito archival information from external labs

mos_expertTaxonomistIDRaw: Mosquito identifications by expert taxonomists - raw

mos_subsampling: Generation of mosquito samples for barcoding, archiving and pathogen testing mos_expertTaxonomistIDProcessed: Mosquito identifications by expert taxonomists - desynonomized

mos_sorting: Mosquito sorting data from external labs

 $mos_identification History: Mosquito\ identification\ history\ for\ records\ where\ identifications\ have\ changed$

variables: Description and units for each column of data in data tables

readme: Data product description, issue log, and other metadata about the data product validation: Description of data validation applied at the points of collection and ingest

Data quality

In the mos_trapping table, the fanStatus, catchCupStatus, and drylceStatus fields record any problems with the trapping apparatus noted at the time of sample collection. The sampleCondition field notes any issues with the sample itself, and is present in the mos_trapping, mos_sorting, and mos_expertTaxonomistID.



tables, tracking whether the condition of the sample changes as it travels through different facilities and analyses. In the mos_sorting table, differences between repeat measurements are noted for count, genus, and species; see variables file for details.

Standard calculations

For wrapper functions to download data from the API, and functions to merge tabular data files across sites and months, see the neonUtilities R package.

During the identification process any trap that appears to contain more than 200 mosquitoes is subsampled such that only a random subset of ~200 mosquitoes are identified. The laboratory reports the weight of the total sample, the subsample, and then the weight of all bycatch removed from the subsample. The individualCount reported for subsampled traps should be corrected for the proportion of the sample analyzed. To make this correction individualCount should be multiplied by totalWeight/subsampleWeight to estimate the full number of mosquitoes present in the trapped sample.

Table joining

| Table 1 | Table 2 | Join by field(s) |
|-------------------------------------|-------------------------------------|--|
| mos_trapping | mos_sorting | sampleID |
| mos_sorting | mos_subsampling | Requires intermediate table: join via mos_expertTaxonomistID table (either Processed or Raw) |
| mos_trapping | mos_subsampling | Requires intermediate table: join via mos_sorting table and mos_expertTaxonomistID table (either Processed or Raw) |
| mos_sorting | mos_expertTaxonomistIDProce ssed | subsampleID |
| mos_sorting | mos_expertTaxonomistIDRaw | subsampleID |
| mos_expertTaxonomistIDProce ssed | mos_archivepooling | Requires intermediate table: join via mos_subsampling table |
| mos_expertTaxonomistIDRaw | mos_archivepooling | Requires intermediate table: join via mos_subsampling table |



| Table 1 | Table 2 | Join by field(s) |
|-------------------------------------|-------------------------------------|---|
| mos_archivepooling | mos_barcoding | Full join not recommended: tables not related |
| mos_archivepooling | mos_sorting | Requires intermediate table: join via mos_sorting table and mos_expertTaxonomistID table (either Processed or Raw) and mos_subsampling table |
| mos_archivepooling | mos_subsampling | archiveID |
| mos_archivepooling | mos_trapping | Requires intermediate table: join via mos_sorting and mos_expertTaxonomistID table (either Processed or Raw) and mos_subsampling tables |
| mos_barcoding | mos_expertTaxonomistIDProce ssed | Not fully automatable: multiple individualIDs pooled into each individualIDList |
| mos_barcoding | mos_expertTaxonomistIDRaw | Not fully automatable: multiple individualIDs pooled into each individualIDList |
| mos_barcoding | mos_sorting | Not fully automatable: multiple individualIDs pooled into each individualIDList |
| mos_barcoding | mos_subsampling | Not fully automatable: multiple individualIDs pooled into each individualIDList |
| mos_barcoding | mos_trapping | Not fully automatable: multiple individualIDs pooled into each individualIDList |
| mos_expertTaxonomistIDProce ssed | mos_expertTaxonomistIDRaw | subsampleID,scientificName, sex |
| mos_expertTaxonomistIDProce ssed | mos_subsampling | subsampleID,scientificName, sex |



| Table 1 | Table 2 | Join by field(s) |
|-------------------------------------|-----------------|---|
| mos_expertTaxonomistIDProce ssed | mos_trapping | Requires intermediate table: join via mos_sorting table |
| mos_expertTaxonomistIDRaw | mos_subsampling | subsampleID,scientificName, sex |
| mos_expertTaxonomistIDRaw | mos_trapping | Requires intermediate table: join via mos_sorting table |
| mos_identificationHistory | Any other table | Full join not recommended: Previous identifications of the same individual can be linked by identificationHistoryID in the mos_expertTaxonomistID table (either Processed or Raw) |

Documentation







NEON User Guide to Mosquitos Sampled From CO2 Traps (DP1.10043.001) and Mosquito-borne Pathogen Status (DP1.10041.001)

NEON_mosquito_userGuide_vD | 246.7 KiB | PDF

Vector Disease Control International Denver Surveillance Laboratory Standard Operating Procedures [NEON Sample Processing Version, 2020]

VDCI_Mosquito_Identification_2020 | 701.4 KiB | PDF

For more information on data product documentation, see: https://data.neonscience.org/data-products/DP1.10043.001



Citation

To cite data from Mosquitoes sampled from CO2 traps (DP1.10043.001), see citation here: https://data.neonscience.org/data-products/DP1.10043.001

For general guidance in citing NEON data and documentation, see the citation guidelines page: https://www.neonscience.org/data-samples/guidelines-policies/citing