# Meier et al. 2022 - Supporting Information File 2: Example linking NEON TOS Coarse Downed Wood volume and Small Mammal abundance data at plot and site scales

Courtney L. Meier, Katherine M. Thibault, and David T. Barnett

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#### Introduction

The NEON Terrestrial Observation System (TOS) data products are frequently spatially integrated at multiple scales, from the site level to the level of relatively small 'sampling cells' that occur within plots (see Supporting Information File 1, Table 3). In addition, the frequency with which TOS data products are generated at each site varies widely, from multiple bouts per year to once every 5 years. This worked example is a companion piece to our manuscript, "Spatial and temporal sampling strategy connecting NEON Terrestrial Observation System protocols," and the purpose is to illustrate how a user of NEON data can draw scientific insight by working with two TOS data products collected at different spatial scales and at different temporal frequencies.

Here, we focus on the 'Coarse downed wood log survey' data product (DP1.10010.001; generated every 5-years) and the 'Small mammal box trapping' data product (DP1.10072.001; generated multiple times per year). Using R software (https://www.r-project.org/) to download data from the NEON API via the neonUtilities package (https://cran.r-project.org/web/packages/neonUtilities/), we show how to link data at both the plot and site spatial scales via the plotID. We also develop one method for comparing TOS data across products generated with very different temporal frequencies. For this example, we chose to investigate correlations between 'Coarse downed wood log survey' data and 'Small mammal box trapping' data based on the hypothesis that coarse downed wood (CDW) provides habitat for small mammals in forested systems, and increasing CDW volume may therefore be correlated with increasing small mammal abundance, both within and among sites.

# Load data from the NEON Data Portal

Note: When retrieving data from the NEON API using the neonUtilities functions (see code chunks below), use of the token argument allows authentication with a user-specific API token. Use of a token is

not required, but tokens provide faster download speeds and help NEON to measure data use for reporting to the NSF.

## Load Coarse Downed Wood Tally Data

At sites with sufficient logs >= 2 cm diameter, the 'Coarse downed wood log survey' data product, hereafter referred to as 'CDW Tally', is generated from both Distributed and Tower plots on a 5 year interval. However, sampling of Distributed and Tower plots is staggered through time such that each plot type is sampled every 2-3 years within a site. Distributed plots are allocated across each site in proportion to the area of dominant National Land Cover Database (NLCD) Vegetation Classes, and these plots allow creating statistically robust site-level estimates of both CDW volume (derived from tallies) and small mammal density. The Small Mammal (MAM) data product is generated annually from most TOS sites, and MAM data are collected from grids that are typically colocated with Distributed base plots, but not Tower plots, which also support CDW sampling. Thus, the first tasks for this example are to identify which NEON sites have produced CDW Tally data, identify several forested sites with recent Distributed plot data for the example, and then identify which Distributed plotIDs are colocated with MAM sampling.

The NEON 'Coarse downed wood log survey' data product citation is:

• NEON (National Ecological Observatory Network). Coarse downed wood log survey, RELEASE-2022 (DP1.10010.001). https://doi.org/10.48443/54dd-9407. Dataset accessed from https://data.neonscience.org on March 18, 2022

```
### Retrieve CDW Tally data for last 6 years from NEON Data Portal for all sites and
### dates
cdwDP <- neonUtilities::loadByProduct(</pre>
  dpID = "DP1.10010.001",
  site = "all",
  startdate = "2016-01",
  enddate = "2021-12",
  package = "basic",
  release = "RELEASE-2022",
 tabl = "all",
 check.size = FALSE,
  token = Sys.getenv('NEON_PAT')
  Extract the field tally data from the download list, and save for quicker read-in
cdw <- cdwDP$cdw_fieldtally</pre>
## Save local copy of data for faster subsequent work
# saveRDS(cdw, file = "cdw_fieldTally_2016-2021.RDS")
#
# #
      Read in data saved via neonUtilities code chunk above
# cdw <- readRDS(file = "cdw_fieldTally_2016-2021.RDS")</pre>
#
   Summarise data to identify 5-6 sites with recent CDW Tally data from Distributed
   plots that can be paired with MAM data; for demonstration purposes, goal is to
   choose a small number of forested sites that span a range of habitats.
summaryDist <- cdw %>%
  dplyr::filter(plotType=="distributed") %>%
  dplyr::group by(domainID, siteID, eventID, samplingImpractical) %>%
  dplyr::summarise(
```

```
plotCount = length(unique(plotID))
 )
  From 'summaryDist' output, select 6 sites with closed-canopy forest
theSites <- c("HARV", "JERC", "STEI", "UKFS", "RMNP", "ABBY")
  Filter CDW data to selected sites, filter to most recent eventID, remove
  unneeded columns
tempCDW <- cdw %>%
  dplyr::filter(
   plotType=="distributed",
   siteID %in% theSites
  dplyr::group_by(domainID, siteID) %>%
  dplyr::filter(
   yearBoutBegan==max(yearBoutBegan)
  dplyr::arrange(domainID, siteID, plotID) %>%
  dplyr::ungroup() %>%
  dplyr::select(-uid, -coordinateUncertainty, -elevationUncertainty, -publicationDate)
### Identify plotIDs in CDW dataset that are colocated with MAM sampling grids
  First, create MAM 'namedLocation' value for each CDW namedLocation value (e.g.,
#
   'HARV_001.mammalGrid.mam'); the goal is to check whether each CDW namedLocation
  has a colocated MAM namedLocation. The structure of the namedLocation value for
  any NEON data product can be determined by downloading data and checking the
  namedLocation\ field.
tempCDW <- tempCDW %>%
  dplyr::mutate(
   mamNamedLoc = stringr::str_replace(
     string = namedLocation,
     pattern = "basePlot.cdw",
     replacement = "mammalGrid.mam"
   ),
    .after = namedLocation
  ) %>%
  as.data.frame()
  Second, retrieve namedLocation data for each mamNamedLoc value that was constructed
  above; if a CDW namedLocation is also a MAM namedLocation, the geoNEON::getLocByName()
   function will return a value from the NEON API; if a CDW namedLocation is not also
  a MAM namedLocation, the function does not return a value. Note: The 'data' argument
  must be a data.frame() and CANNOT be a tibble (the latter is commonly returned by
   dplyr).
mamPlots <- geoNEON::getLocByName(</pre>
 data = tempCDW,
 locCol = "mamNamedLoc",
 locOnly = TRUE,
 token = Sys.getenv('NEON_PAT')
```

```
## Save data locally for quicker subsequent read-in
# saveRDS(mamPlots, file = "cdw_mam_collocation.RDS")
# mamPlots <- readRDS(file = "cdw_mam_collocation.RDS")

# Third, join tempCDW data with mamPlots and create a 'subtype' column to identify
# colocated sampling
tempCDW <- tempCDW %>%
    dplyr::left_join(
    mamPlots %>%
        select(plotID, subtype),
    by = "plotID"
    ) %>%
    dplyr::select(-mamNamedLoc)
```

As an aside, rather than programmatically retrieving TOS Spatial Data via the API with the geoNEON functions, as is done in the code chunk above, these spatial data may also be manually downloaded:

- First, navigate to https://data.neonscience.org/documents
- Click on 'Spatial Data' and click on the 'All\_NEON\_TOS\_Plots\_VX' link to retrieve a zip that contains a .csv with plot data.
- If this approached is used, one can then read in the included .csv file.
  - For example: tosPlots <- read.csv(file = "All\_NEON\_TOS\_Plot\_Centroids\_V8.csv",
    header = TRUE)</pre>

#### Load Small Mammal Data

Small Mammal sampling grids (n=3 to n=8 per site) are colocated with Distributed base plots that support multiple TOS protocols (including Coarse Downed Wood), and grids are sampled one to several times per year (ideally, 4-6 times per year). Similar to Distributed base plots, mammal grids are allocated using a stratified random design informed by NLCD Vegetation Class.

The NEON 'Small mammal box trapping' data product citation is:

• NEON (National Ecological Observatory Network). Small mammal box trapping, RELEASE-2022 (DP1.10072.001). https://doi.org/10.48443/h3dk-3a71. Dataset accessed from https://data.neonscience.org on March 18, 2022

```
### Retrieve small mammal box trapping data for same sites and date range as retrieved
### for CDW
mamDP <- neonUtilities::loadByProduct(
    dpID = "DP1.10072.001",
    site = theSites, #--> same sites identified for CDW
    startdate = "2016-01",
    enddate = "2021-12",
    package = "basic",
    release = "RELEASE-2022",
    tabl = "all",
    check.size = FALSE,
    token = Sys.getenv('NEON_PAT')
)
## Save data locally for quicker subsequent read-in
```

```
# saveRDS(mamDP, file = "mam_allTables_2016-2021.RDS")
#
# #
      Read in stored MAM data
# mamDP <- readRDS(file = "mam_allTables_2016-2021.RDS")</pre>
    Convert list tables to dataframes in the Global environment
list2env(mamDP, envir=.GlobalEnv)
### Read in master SMALL MAMMAL taxon table via the NEON API
    https://www.neonscience.org/resources/learning-hub/tutorials/neon-api-usage
    using the verbose option to get the taxonProtocolCategory field
mam.req <- httr::GET(</pre>
  paste(
    "https://data.neonscience.org/api/v0/taxonomy/?taxonTypeCode=SMALL_MAMMAL",
    "verbose=true",
    "offset=0",
    "limit=1000",
    sep = "&"
  )
mam.list <- jsonlite::fromJSON(httr::content(mam.req, as="text"))</pre>
```

# Data QC Checks

#### Coarse Downed Wood Tally QC Checks

Note: The word 'log' is used frequently in the Coarse Downed Wood dataset, and refers to a downed piece of wood, not a logarithm.

For CDW tally data, prior to analysis it is important to:

- Check for duplicates, and
- Remove any tallied logs with a logDistance greater than the protocol-specified transect length, and
- Verify that all tallied logs meet 'limiting distance' tally criteria based on reported equivalentLogDiameter, logDistance, and volumeFactor data (see Affleck 2010 for details).
  - The equivalentLogDiameter is used so that round and elliptical logs (typically highly decayed)
    can be assessed with the same criteria. Logs with logDistance greater than the limiting distance
    are removed from the dataset.
  - Because technicians sometimes use look-up tables to determine limiting distance from a subset of equivalentLogDiameter values (e.g., when digital ingest devices fail), and look-up tables require choosing the nearest equivalentLogDiameter rather than using the actual measured value, it is expected that some logs will have logDistance greater than the calculated limiting distance and will be removed.

```
### Check for individualID duplicates; duplicate checks are important because log
### counts are used to estimate CDW volume
# First, construct an individualID for logs < 10 cm diameter that are not tagged;
# smaller logs are not tagged and are given a unique temporary logID beginning with 'L'
# since they are less likely to persist over the 5-year CDW Tally measurement interval;
# second, construct a primary key from plotID, date, lidsAzimuth, and individualID.

tempCDW <- tempCDW %>%
```

```
dplyr::mutate(
   individualID = case_when(
      is.na(individualID) & targetTaxaPresent=="Y" ~ paste(
        "NEON.CDW", domainID, plotID, logID, sep = "."),
     TRUE ~ individualID
   ),
   key = paste(plotID, date, lidsAzimuth, individualID, sep = "_")
  Identify duplicates ('dups') based on 'key' value
cdwDupsKey <- tempCDW %>%
  dplyr::filter(duplicated(tempCDW$key)) %>%
  dplyr::select(key)
cdwDups <- tempCDW %>%
  dplyr::filter(key %in% cdwDupsKey$key)
# Log data assessment for dups: Only first one listed appears to be a real duplicate
# based on examination of other data fields; ABBY records are mis-indentified by the
# function as 'dups' because transect was reflected (i.e., not real duplicates).
    Remove second instance of cdwDups$key=="STEI_002_2016-05-26_320_NEON.CDW.D05.00375"
cdwRemoveDup <- cdwDups %>%
  dplyr::distinct(key)
tempCDW <- tempCDW %>%
  dplyr::filter(!key %in% cdwRemoveDup$key[1]) %>%
  dplyr::bind rows(cdwDups[1,]) %>%
  dplyr::arrange(domainID, siteID, plotID, lidsAzimuth, individualID)
### Perform QC checks for volumeFactor, logDistance, and equivalentLogDiameter
## Volume Factor: Check value matches protocol and flag if no match
  Read in CDW volumeFactor and transectLength look-up table derived from protocol
  (NEON.DOC.001711). Note that RMNP has different F-values and transect lengths for
# Distributed and Tower plots due to distant parcels of land used for each plot type
  and different forest types. The RMNP values in this table are specific to
  Distributed plots since we wish to compare data with Small Mammal data.
cdwParam <- read.csv(file = "cdw_siteParamLookup.csv", header = TRUE)</pre>
    Join look-up table values with CDW data and flag records with incorrect volume
   factor (i.e., f-value)
tempCDW <- tempCDW %>%
  dplyr::left join(
    cdwParam %>% dplyr::select(siteID, fValue, transectLength),
   by = "siteID"
  ) %>%
  dplyr::mutate(
   fValQF = case_when(
     is.na(volumeFactor) ~ 1,
      volumeFactor==fValue ~ 0,
      TRUE ~ 1
   )
  )
```

```
# Check records with volumeFactor problems and evaluate
fCheck <- tempCDW %>%
  dplyr::filter(fValQF==1) %>%
  dplyr::select(domainID, siteID, volumeFactor, fValue, fValQF)
   volumeFactor problems identified for HARV and STEI; volumeFactor is supposed to
  be assigned at the site level and should not vary from plot-to-plot. The variation
  in the data likely reflects a data entry error. We encourage data users to let NEON
#
  staff know about issues they discover like this one:
  https://www.neonscience.org/about/contact-us
  Correct volumeFactor data for HARV, STEI sites based on fCheck results
tempCDW <- tempCDW %>%
  dplyr::mutate(volumeFactor = as.numeric(volumeFactor)) %>%
  dplyr::mutate(
   volumeFactor = case_when(
      siteID=="HARV" & (volumeFactor!=5 | is.na(volumeFactor)) ~ 5,
      siteID=="STEI" & (volumeFactor!=5 | is.na(volumeFactor)) ~ 5,
     TRUE ~ volumeFactor
   )
  )
### Tally QC checks: Verify that logs in the dataset should be tallied based on
### protocol criteria
## Log Distance: Check for values that exceed maximum transect length as defined in
## protocol, then flag
tempCDW <- tempCDW %>%
  dplyr::mutate(
   logDistQF = case_when(
      is.na(logDistance) ~ 0,
     logDistance > transectLength ~ 1,
      TRUE ~ 0
    )
  )
## Check whether logDistance is <= limitingDistance, as expected, based on
## equivalentLogDiameter, and flag
# Use formula to calculate limiting distance from Affleck 2010, Appendix A:
  limitingDist = ((pi^2)*(roundDiameter^2))/(8*transectNum*fValue), where
# roundDiameter = round diameter of the log (cm); i.e., the equivalentLogDiameter field
  transectNum = the number of transects at a survey point (3 for NEON plots),
  fValue = site-level F value, taken from the volumeFactor field in NEON data
tempCDW <- tempCDW %>%
  dplyr::mutate(
   limitingDist = round(((pi^2)*(equivalentLogDiameter^2))/(8*3*volumeFactor), digits=2)
 ) %>%
  dplyr::mutate(
   limDistQF = case_when(
      is.na(limitingDist) ~ 0,
     logDistance > limitingDist ~ 1,
      TRUE ~ 0
   )
```

```
## Filter out flagged records for logs that should not have been tallied
filterCDW <- tempCDW %>%
  dplyr::filter(
    limDistQF==0,
    logDistQF==0
)
```

## Small Mammal Box Trapping Data QC: Check for duplicates

- mam\_perplotnight: In this table, duplicate records are defined as those with the same plot and date combination (as captured in an auto-generated nightuid).
- mam\_pertrapnight: In this table, duplicate records are defined as those with the same nightuid, trap coordinate, and tagID or individualCode; note that the standard neonOS::removeDups() function used to remove duplicates cannot account for multiple captures of untagged individuals in a single trap.

```
### First, check mam_perplotnight table by nightuid using standard removeDups function
mam_plotNight_nodups <- neonOS::removeDups(</pre>
  data = mam perplotnight,
 variables = variables_10072,
  table = "mam_perplotnight"
# Running this check generated the message: "Primary key fields contain NA values
# and/or empty strings. Results may be unreliable; check input data carefully." -
# followed by an error message.
## To troubleshoot, first query the table to find the problematic records
problems <- which(is.na(mam_perplotnight$nightuid))</pre>
   Next, check the mam_pertrapnight table to see if there are corresponding records
  that might have a nightuid
temp <- mam_pertrapnight %>%
  dplyr::filter(
   plotID %in% mam_perplotnight$plotID[problems] &
      collectDate %in% mam perplotnight$collectDate[problems]
  )
# Since the remaining fields were populated and there are captures for each, these
# records appear to represent a valid night of trapping. So, we will assign custom
# temporary nightuids in both tables here. As an aside, we encourage users to let NEON
# staff know about issues they discover like this one:
# (https://www.neonscience.org/about/contact-us).
    Create custom nightuids for duplicates identified above that are missing primary
    key value
mam_perplotnight_adj <- mam_perplotnight %>%
  dplyr::mutate(nightuid = paste(plotID, '_', collectDate, sep = ''))
mam_pertrapnight_adj <- mam_pertrapnight %>%
 dplyr::mutate(nightuid = paste(plotID, '_', collectDate, sep = ''))
```

```
# Re-run the duplicate removal function
mam_plotNight_nodups <- neonOS::removeDups(</pre>
 data = mam_perplotnight_adj,
 variables = variables 10072,
 table = "mam_perplotnight"
# No duplicate key values found
### Second, check the mam_pertrapnight table by nightuid and trapcoordinate using the
### standard neonOS::removeDups() function.
   Note that RELEASE 2022 contains ~142K records for theSites downloaded, so this
   operation can take some time. In rare cases, multiple animals can be captured in a
  single trap on the same night and not all receive unique tagIDs or individualCodes.
# This means that the duplicate function is not effective for these records; here, we
# remove these records prior to running the standard check, and we then add these
  records back to the data frame afterwards.
mam_trapNight_multipleCaps <- mam_pertrapnight_adj %>%
  dplyr::filter(
    trapStatus == "4 - more than 1 capture in one trap" &
      is.na(tagID) & is.na(individualCode)
mam_trapNight_remainingRecords <- mam_pertrapnight_adj %>%
  dplyr::filter(!(uid %in% mam_trapNight_multipleCaps$uid))
mam_trapNight_nodups <- neonOS::removeDups(</pre>
  data = mam_trapNight_remainingRecords,
 variables = variables_10072,
 table = "mam_pertrapnight"
)
# Output: Two unresolveable duplicates flagged with duplicateRecordQF=2, which
# indicates records that are identical based on primary keys but contain different
# values in other fields; see ?neonOS::removeDups documentation for additional
   information on duplicate record flagging.
## Follow-up trouble-shooting to deal with unresolveable duplicates
   Check for unexpected NAs in primary keys
which(is.na(mam_pertrapnight_adj$nightuid)) #--> yields 0
which(is.na(mam_pertrapnight_adj$trapCoordinate)) #--> yields 0
# Two lines above mean that trapID and individualCode are the source of the NAs, which
# is expected based on the protocol.
# Next, check that the unresolved duplicates will not impact the intended analyses
dupCheck <- mam_trapNight_nodups %>%
  dplyr::filter(
    duplicateRecordQF == 2 & trapStatus != "4 - more than 1 capture in one trap"
# No records in the set, so these unresolved duplicates will not impact the intended
# analyses.
```

```
## Add multiple capture records back to dataset
mam_trapNight_nodups <- mam_trapNight_nodups %>%
   dplyr::mutate(collectDate = as.Date(collectDate)) %>%
   dplyr::bind_rows(mam_trapNight_multipleCaps)
```

# Analysis of Spatially Integrated Data Products at Plot and Site Scales

In this section, we generate Coarse Downed Wood volume estimates and Small Mammal abundance estimates at both plot and site scales in order to demonstrate the spatial integration of NEON TOS data products. Because CDW volume is generated every 5-years and does not change rapidly, barring fires or other site management activities, we compare with Small Mammal data collected the same year as the CDW data  $\pm$ 1 year. For example, for HARV CDW volume data collected in 2019, we compare to Small Mammal data collected from 2018-2020.

Note: NEON publishes site management data via the 'Site management and event reporting' product (DP1.10111.001; https://data.neonscience.org/data-products/DP1.10111.001).

#### Create Coarse Downed Wood Volume Datasets

Using the LIDS method (Affleck 2008, 2010), CDW volume is calculated at the plot scale as:

```
CDW volume density = F * n (m^3/hectare);
```

where F is the site-specific volume factor, and n is the number of qualifying logs tallied across all three transects originating from the centroid of the plot. Importantly, at the plot scale, CDW volume density is colocated with Small Mammal abundance data at a maximum of n=6 plots per site (for the Sites included).

At the site scale, mean CDW volume (m³/hectare) is simply the mean of the plot scale values; the mean site-scale parameter is robust because of the spatially-balanced, stratified-random nature of the TOS Spatial Design. When comparing CDW volume and Small Mammal abundance at the site scale, CDW data from all Distributed plots are used for the calculation.

```
### Calculate CDW volume for each plotID in the cleaned dataset
   First, focus on plots with targetTaxaPresent=="Y" --> i.e., at least one qualifying
    log was tallied within the plot, and remove sampling Impractical != "OK"
plotCdwVol <- filterCDW %>%
  dplyr::filter(is.na(samplingImpractical) | samplingImpractical=="OK") %%
  dplyr::filter(targetTaxaPresent="Y") %>% # Remove records for transects with no logs
  dplyr::group_by(domainID, siteID, volumeFactor, eventID, plotID) %>%
  dplyr::summarise(
   logCount = n()
  ) %>%
  dplyr::mutate(cdwVol = volumeFactor * logCount)
   Second, identify plots where all three transects do not have logs and bind to log
    count output from above
noLogs <- filterCDW %>%
  dplyr::filter(targetTaxaPresent=="N") %>%
  dplyr::group_by(domainID, siteID, volumeFactor, eventID, plotID) %%
  dplyr::summarise(
   transectCount = n() # Identify plots with all 3 transects with ttP=="N"
  dplyr::filter(transectCount==3) %>%
```

```
dplyr::mutate(
    logCount = 0,
   cdwVol = 0
  ) %>%
  dplyr::select(-transectCount)
plotCdwVol <- plotCdwVol %>%
  dplyr::bind rows(noLogs) %>%
  dplyr::arrange(domainID, siteID, eventID, plotID)
  Finally, add back mammalGrid collocation data
temp <- tempCDW %>%
  dplyr::select(plotID, subtype) %>%
  dplyr::distinct()
plotCdwVol <- plotCdwVol %>%
  dplyr::left_join(temp, by = "plotID")
## Display table of plot-level CDW volume estimates
knitr::kable(
 x = plotCdwVol %>%
   dplyr::ungroup() %>%
   dplyr::rename(
      "Log Count" = logCount,
      "CDW Vol (m3/ha)" = cdwVol,
      "Plot subtype" = subtype
   ) %>%
   dplyr::select(-eventID) %>%
   dplyr::slice_head(n = 5),
  row.names = FALSE,
  caption = "Example subset of CDW volume density estimates by plotID for selected sites.
  The 'Plot subtype' column indicates whether a given CDW plot is colocated with a Small
  Mammal sampling grid."
)
```

Table 1: Example subset of CDW volume density estimates by plotID for selected sites. The 'Plot subtype' column indicates whether a given CDW plot is colocated with a Small Mammal sampling grid.

domainID	siteID	volumeFactor	plotID	Log Count	CDW Vol (m3/ha)	Plot subtype
D01	HARV	5	HARV_001	5	25	mammalGrid
D01	HARV	5	$HARV\_002$	3	15	NA
D01	HARV	5	$HARV\_004$	5	25	NA
D01	HARV	5	$HARV\_005$	3	15	NA
D01	HARV	5	$HARV\_006$	6	30	${\rm mammalGrid}$

```
### Calculate CDW Volume for each siteID in the cleaned dataset and display
# Calculate site-level mean CDW volume
siteCdwVol <- plotCdwVol %>%
   dplyr::group_by(domainID, siteID, eventID) %>%
```

```
dplyr::summarise(
   meanCdwVol = round(mean(cdwVol, na.rm = TRUE), digits = 1),
   sdCdwVol = round(sd(cdwVol, na.rm = TRUE), digits = 1),
   nCdwVol = n()
  )
   Display table of site-level CDW volume estimates
knitr::kable(
  x = siteCdwVol %>%
   dplyr::ungroup() %>%
   dplyr::rename(
      "Mean CDW Vol (m3/ha)" = meanCdwVol,
      "StdDev CDW Vol" = sdCdwVol,
      "Plot Number" = nCdwVol
   ),
  row.names = FALSE,
  caption = "CDW volume density estimates by siteID for selected sites and eventIDs."
)
```

Table 2: CDW volume density estimates by siteID for selected sites and eventIDs.

$\overline{\text{domainID}}$	siteID	eventID	Mean CDW Vol (m3/ha)	StdDev CDW Vol	Plot Number
D01	HARV	CDW.2019.HARV	23.8	14.9	20
D03	JERC	CDW.2018.JERC	6.6	8.2	19
D05	STEI	CDW.2016.STEI	27.8	23.1	20
D06	UKFS	CDW.2018.UKFS	20.0	24.2	20
D10	RMNP	CDW.2020.RMNP	37.9	29.4	19
D16	ABBY	CDW.2019.ABBY	197.2	118.5	20

## Create Small Mammal Density datasets

Here, we employ the minimum number known alive (MNKA) approach to estimate total small mammal abundance - e.g., as defined in Slade & Blair (2000). This approach assumes that a marked individual is present at all sampling points between its first and last capture dates, even if it was not actually captured in those interim trapping sessions. To make the dataset meet this assumption, we first add those implicit records to the dataset to make them explicit.

Note that it is necessary to create Small Mammal eventIDs for legacy data collected prior to 2021. For the Small Mammal product, eventIDs correspond to discrete sampling bouts that represent a group of consecutive nights of trapping scheduled around a particular new moon.

```
### Create eventID grouping variable using mam_plotNight data and add to mam_trapNight
### data
# Assign the same eventID to each record with an endDate no more than 10 days later
# than the previously created record.
mam_plotNight_nodups <- mam_plotNight_nodups %>%
    dplyr::mutate(
        year = as.integer(format(collectDate, "%Y")),
        .before = eventID
    ) %>%
    dplyr::group_by(siteID, year) %>%
    dplyr::mutate(
```

```
priorEndDate = data.table::shift(endDate, fill = endDate[1]),
    diffDays = difftime(endDate, priorEndDate, units = "days"),
   boutIncrement = ifelse(diffDays >= 10, 1, 0),
   boutNum = cumsum(boutIncrement),
    .after = endDate
  dplyr::group_by(siteID, year, boutNum) %>%
  dplyr::mutate(
   weekBoutBegan = lubridate::isoweek(min(endDate)),
   eventID = case when(
     is.na(eventID) ~ paste(siteID, year, weekBoutBegan, sep = "."),
     TRUE ~ eventID
   ),
    .after = boutNum
  ) %>%
  dplyr::ungroup() %>%
  dplyr::select(-priorEndDate, -diffDays, -boutIncrement, -boutNum)
# Above, shift() defaults to type = "lag", which returns the previous row value for the
# specified column
   Add nightuid and eventID from mam_plotNight_nodups to mam_trapNight_nodups
mam_trapNight_nodups <- mam_trapNight_nodups %>%
  dplyr::left_join(
   mam_plotNight_nodups %>%
     dplyr::select(nightuid, eventID),
   by = "nightuid"
### Create and finalize Small Mammal capture dataset for selected sites
## First, subset trapping data to only the capture records, i.e., the records that
## describe a captured small mammal, including only those taxa for which the trapping
## protocol is designed. Subsetting can be done with a simple filtering based on
## 'trapStatus' and 'taxonProtocolCategory'; however, we first check to ensure all
## captures have the correct 'trapStatus' - i.e., check if tagIDs exist but trap status
## does not include "capture".
problemRecords <- mam_trapNight_nodups %>%
 dplyr::filter(
    !is.na(tagID),
    !grepl("capture", trapStatus)
  )
  If nrow(problemRecords) > 0, update the corresponding trapStatus fields to
   "5 - capture" or, in the rare case where there are multiple captures in one trap,
  "4 - more than 1 capture in one trap" - this rare case does not occur in the current
  dataset, so it is not addressed here.
mam_trapNight_nodups <- mam_trapNight_nodups %>%
  dplyr::mutate(trapStatus = case_when(
   uid %in% problemRecords$uid ~ "5 - capture",
   TRUE ~ trapStatus
  ))
```

```
## Second, create list of target taxa from taxon list
targetTaxa <- mam.list$data %>%
  dplyr::filter(taxonProtocolCategory == "target") %>%
  dplyr::select(taxonID)
## Third, simplify dataset to targetTaxa and core fields needed for analysis
captures <- mam trapNight nodups %>%
  dplyr::filter(grepl("capture", trapStatus) & taxonID %in% targetTaxa$taxonID) %>%
  dplyr::select(uid, nightuid, plotID, collectDate, tagID)
## Fourth, following the 'Minimum Number Known Alive' (MNKA) approach (Slade & Blair
## 2000), generate a column of all of the unique tagIDs included in the dataset
uTags <- captures %>%
  dplyr::select(tagID) %>%
  dplyr::filter(!is.na(tagID)) %>%
  dplyr::distinct()
  Create empty data frame to populate with 'for()' loop output
capsNew <- dplyr::slice(captures, 0)</pre>
  For each tagged individual, add a record for each night of trapping done in the plots
   in which it was captured between the first and last dates of capture.
for (i in uTags$tagID) {
  temp <- captures %>% filter(tagID == i)
  firstCap <- as.Date(min(temp$collectDate), "YYYY-MM-DD", tz = "UTC")</pre>
  lastCap <- as.Date(max(temp$collectDate), "YYYY-MM-DD", tz = "UTC")</pre>
  possibleDates <- seq(as.Date(firstCap), as.Date(lastCap), by="days")</pre>
  plots <- unique(temp$plotID)</pre>
  potentialNights <- mam_plotNight_nodups %>%
    dplyr::filter(as.character(collectDate) %in% as.character(possibleDates) &
             plotID %in% plots) %>%
   dplyr::select(nightuid,plotID, collectDate) %>%
   dplyr::mutate(tagID = i)
  temp2 <- dplyr::left join(potentialNights, temp)</pre>
  capsNew <- dplyr::bind_rows(capsNew, temp2)</pre>
## To avoid time-consuming processing, save a copy of `for()` loop output
# saveRDS(capsNew, file = "mam_final_dataset.RDS")
# # Read in processed capsNew dataset
# capsNew <- readRDS(file = "mam_final_dataset.RDS")</pre>
   Add untagged individuals back to the dataset
capsNew <- captures %>%
  dplyr::filter(is.na(tagID)) %>%
  dplyr::bind_rows(capsNew)
  Add eventID to enable summarizing by trapping bouts (includes up to 3 nights of
```

```
# trapping for some plots in each bout, 1 night for other plots).
capsNew <- capsNew %>%
    dplyr::left_join(
    mam_plotNight_nodups %>%
        dplyr::select(eventID, nightuid),
    by = "nightuid"
    ) %>%
    dplyr::relocate(eventID, .after = collectDate) %>%
    dplyr::arrange(eventID, plotID, collectDate)
```

Next, we define a function to calculate MNKA for each plot within each bout. The function requires the capsNew data set generated above as an input, as well as a list of plots of interest (plotsOI), in the event that the capsNew data contain records from more plots than are needed for the MNKA estimation.

```
### Define function to calculate MNKA for each plot within bout; 'plotsOI' is a required
### input vector of plotIDs for which MNKA per plot per bout is desired.
mnka per plot per bout <- function(capture data, plots0I) {</pre>
  caps <- capture data %>%
   dplyr::filter(plotID %in% plotsOI)
  ids_by_plot_bout <- capture_data %>%
    dplyr::group_by(eventID, plotID) %>%
    dplyr::distinct(tagID)
  mnka_by_plot_bout <- ids_by_plot_bout %>%
    dplyr::group_by(eventID, plotID) %>%
   dplyr::count() %>%
   dplyr::mutate(
      siteID = stringr::str extract(eventID, "^[A-Z]{4}"),
      year = as.numeric(stringr::str_extract(eventID, "20[0-9]{2}")),
      .before = eventID
   ) %>%
   dplyr::ungroup() %>%
   as.data.frame()
  return(mnka_by_plot_bout)
}
# Output is MNKA (count/hectare), based on 1 ha size of trapping grid
```

We now use the MNKA small mammal density function to calculate MNKA at the plot scale and site scale in order to compare to CDW volume density data calculated above. Because CDW volume does not change rapidly, and small mammal abundance may change substantially both within and among years, here we generate the maximum small mammal abundance for a given plot for a three-year window centered on the year that logs were tallied for CDW.

```
### Calculate MNKA per plot per bout for all data in processed capsNew dataset and
### select 3-year window of data appropriate for each site in order to match up with
### CDW volume data.

plotBoutMNKA <- mnka_per_plot_per_bout(
    capture_data = capsNew,
    plotsOI = unique(capsNew$plotID)
)</pre>
```

```
# Create table of MAM data within 3-year windows by site based on year of CDW
# volume data
mamSiteYears <- plotCdwVol %>%
  dplyr::ungroup() %>%
  dplyr::distinct(domainID, siteID, eventID) %>%
  dplyr::mutate(
   cdwYear = as.numeric(stringr::str_extract(eventID, "20[0-9]{2}")),
   minMamYear = cdwYear - 1,
   maxMamYear = cdwYear + 1
  )
  Calculate plot-specific MNKA across bouts: Join plotBoutMNKA and mamSiteYears, filter
# to records within 3-year window centered on CDW measurement year, and determine
   maximum small mammal abundance for each plot, as well as other per plot metrics.
plotMNKA <- plotBoutMNKA %>%
  dplyr::left_join(
   mamSiteYears %>%
     dplyr::select(
        siteID,
       minMamYear,
       maxMamYear
     ),
   by = "siteID"
  ) %>%
  dplyr::filter(
   year >= minMamYear,
   year <= maxMamYear</pre>
 ) %>%
  dplyr::group_by(siteID, plotID) %>%
  dplyr::summarise(
   plotMeanAbundance = round(mean(n), digits = 1),
   plotMedAbundance = median(n),
   plotMinAbundance = min(n),
   plotMaxAbundance = max(n),
   plotBoutCount = n()
  ) %>%
  dplyr::ungroup()
## Display table of plot-level MNKA parameters
knitr::kable(
 x = plotMNKA %>%
   dplyr::ungroup() %>%
   dplyr::rename(
      "Mean Abundance (count/ha)" = plotMeanAbundance,
     "Median Abundance (count/ha)" = plotMedAbundance,
      "Min Abundance (count/ha)" = plotMinAbundance,
     "Max Abundance (count/ha)" = plotMaxAbundance,
      "# Bouts Sampled" = plotBoutCount
   ) %>%
   dplyr::slice_head(n = 5),
  row.names = FALSE,
  caption = "Example subset of Small Mammal abundance metrics (count/ha) by plotID for
```

```
selected sites for a 3-year window centered on the year that CDW volume was measured."
```

Table 3: Example subset of Small Mammal abundance metrics (count/ha) by plotID for selected sites for a 3-year window centered on the year that CDW volume was measured.

siteID plotID	Mean Abundance (count/ha)	Median Abundance (count/ha)	Min Abundance (count/ha)	Max Abundance (count/ha)	# Bouts Sampled
ABBY ABBY_002	9.2	9.0	2	18	10
ABBY ABBY_004	32.0	31.5	9	58	10
ABBY ABBY_007	7.1	6.0	3	11	9
ABBY ABBY_010	8.5	7.5	2	26	10
ABBY ABBY_014	4.2	5.0	1	10	9

```
### Calculate MNKA metrics per site based on plot-level data
  siteMeanAbundance = mean abundance across plots within a site
  siteMedAbundance = median abundance across plots within a site
  siteMinAbundance = average minimum abundance across plots within a site
    siteMaxAbundance = average maximum abundance across plots within a site
siteMNKA <- plotMNKA %>%
  dplyr::group_by(siteID) %>%
  dplyr::summarise(
   siteMeanAbundance = round(mean(plotMeanAbundance), digits = 1),
   siteMedAbundance = round(median(plotMedAbundance), digits = 1),
   siteMinAbundance = round(mean(plotMinAbundance), digits = 1),
   siteMaxAbundance = round(mean(plotMaxAbundance), digits = 1)
  ) %>%
  dplyr::ungroup()
## Display table of site-level MNKA parameters
knitr::kable(
  x = siteMNKA %>%
   dplyr::ungroup() %>%
   dplyr::rename(
      "Mean Abundance (count/ha)" = siteMeanAbundance,
      "Median Abundance (count/ha)" = siteMedAbundance,
      "Average Min Abundance (count/ha)" = siteMinAbundance,
     "Average Max Abundance (count/ha)" = siteMaxAbundance
   ),
  row.names = FALSE,
  caption = "Small Mammal abundance metrics (count/ha) by siteID for selected sites
  for a 3-year window centered on the year that CDW volume was measured."
```

Table 4: Small Mammal abundance metrics (count/ha) by siteID for selected sites for a 3-year window centered on the year that CDW volume was measured.

siteID	Mean Abundance (count/ha)	Median Abundance (count/ha)	Average Min Abundance (count/ha)	Average Max Abundance (count/ha)
ABBY	13.5	8.2	4.2	26.3
HARV	10.4	8.5	1.0	28.7
JERC	17.8	14.5	4.7	41.3
RMNP	14.1	8.0	1.8	37.0
STEI	20.7	15.8	3.5	52.8
UKFS	8.5	5.8	1.5	22.7

#### Create unified CDW and MAM datasets

To create plot-level and site-level data sets suitable for plotting and correlation analyses, we join the CDW volume and Small Mammal abundance data at both plot and site scales.

```
### Join plot-level CDW and MAM datasets
plotDF <- plotCdwVol %>%
    dplyr::left_join(
        plotMNKA %>%
        dplyr:select(-plotBoutCount),
        by = c("siteID", "plotID")
) %>%
    dplyr::filter(
        !is.na(plotMaxAbundance)
)

### Join site-level CDW and MAM datasets
siteDF <- siteCdwVol %>%
    dplyr::left_join(
        siteMNKA,
        by = "siteID"
)
```

#### Create graphs of CDW and MAM data at plot and site spatial scales

We first examine the potential relationship between CDW volume and Small Mammal abundance using data from all spatially colocated plots from all of the six forested sites we selected (HARV, JERC, STEI, UKFS, RMNP, and ABBY).

```
### Create graph of CDW volume vs. MAM maximum abundance across all plots and sites
plotsGraph <- ggplot2::ggplot(
   data = plotDF,
   mapping = aes(
        x = cdwVol,
        y = plotMaxAbundance
)
) +
   ggplot2::geom_point(
   alpha = 0.5,</pre>
```

```
size = 3
) +
ggplot2::geom_smooth() +
ggplot2::labs(
   x = "CDW Volume (m3/ha)",
   y = "Max Small Mammal Abundance (count/ha)"
)
plotsGraph
```

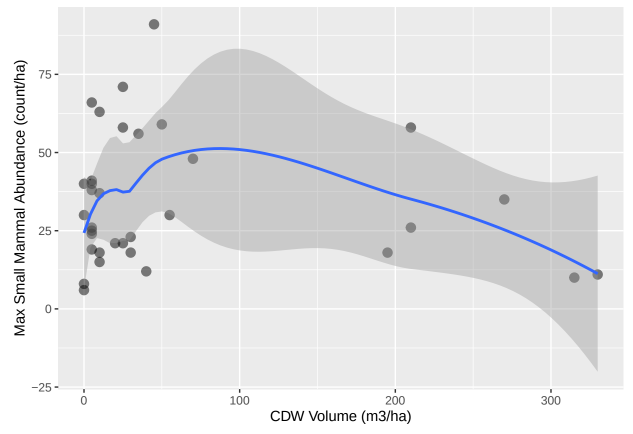


Figure 1. Correlation between CDW Volume and Max Small Mammal Abundance at the plot scale across forested sites.

- Across a selection of close-canopy forested NEON sites, measurements of CDW volume (m³/ha) and Small Mammal abundance (count/ha) reveal no clear correlation at the colocated plot-scale.
- The Pearson's correlation coefficient is: -0.16.

Next, we check for correlations between CDW volume and Small Mammal abundance within sites using colocated plot-scale data.

```
### Create graph of CDW volume vs. MAM maximum abundance across plots within site
sitePlotsGraph <- ggplot2::ggplot(
   data = plotDF,
   mapping = aes(
        x = cdwVol,
        y = plotMaxAbundance
   )
) +</pre>
```

```
ggplot2::geom_point(
    alpha = 0.5,
    size = 3
) +
ggplot2::geom_smooth(method = lm) +
ggplot2::facet_wrap(
    facets = ~siteID,
    ncol = 3,
    scales = "free"
) +
ggplot2::labs(
    x = "CDW Volume (m3/ha)",
    y = "Max Small Mammal Abundance (count/ha)"
)
sitePlotsGraph
```

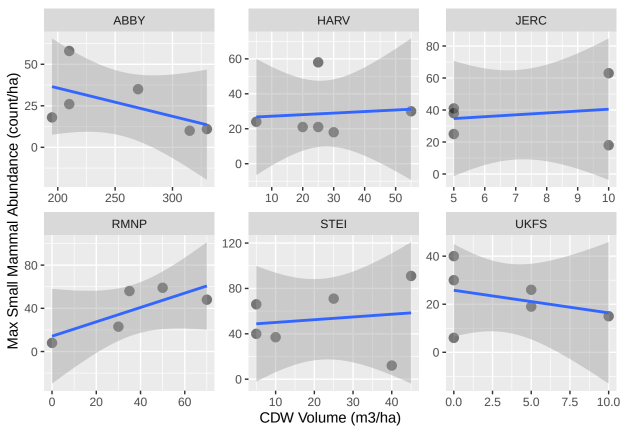


Figure 2. Correlation between CDW Volume and Max Small Mammal Abundance by plot within site.

• Within a selection of close-canopy forested NEON sites, CDW volume ( $\rm m^3/ha$ ) and Small Mammal abundance (count/ha) show positive (RMNP,  $\rm r_{Pearson}=0.77$ ), negative (ABBY,  $\rm r_{Pearson}=-0.55$ ), and weak correlations (HARV, JERC, STEI, UKFS).

Finally, we check for correlations between mean CDW volume and average maximum Small Mammal abundance at the site scale.

```
### Create graph of CDW volume vs. MAM maximum abundance across sites
sitesGraph <- ggplot2::ggplot(</pre>
  data = siteDF,
  mapping = aes(
    x = meanCdwVol,
    y = siteMaxAbundance,
) +
  ggplot2::geom_point(
    alpha = 0.5,
    size = 3
  ggplot2::geom_smooth(method = lm) +
  ggplot2::labs(
    x = "CDW Volume (m3/ha)",
    y = "Average Max Small Mammal Abundance (count/ha)"
  )
sitesGraph
```

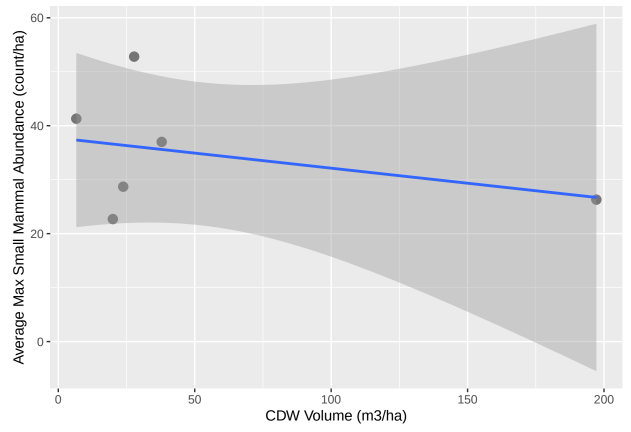


Figure 3. Correlation between CDW Volume and average  $\max$  Small Mammal Abundance across sites.

• Across a selection of closed-canopy forested NEON sites, CDW volume ( $\rm m^3/ha$ ) and Small Mammal abundance (count/ha) show a weak, negative correlation at the site scale ( $\rm r_{Pearson} = -0.36$ ).

## Conclusions

- Coarse Downed Wood volume (m<sup>3</sup>/ha) and Small Mammal abundance (count/ha) can each be calculated at the plot-level, and spatially colocated plots are identified by a shared plotID.
  - It is possible to identify shared plotID values programmatically using the geoNEON::getLocByName() function (lines 165-170), or by downloading the CDW and Small Mammal datasets and joining via the plotID (lines 883-903).
  - Note that more plotID values may be identified as colocated via the geoNEON function than via the table joining method due to the fact that getLocByName() identifieds all plotIDs that have ever been used for a given protocol, and not just those locations that are currently in use.
- Plot-scale spatial collocation of TOS data products allows data users to investigate patterns within NEON sites.
- All NEON TOS data products are colocated at the site-scale. Site-level estimates may be useful for investigating patterns at very large spatial scales.

#### References

Affleck, D.L.R. (2008) A line intersect distance sampling strategy for downed wood inventory. Canadian Journal of Forest Research, 38, 2262-2273.

Affleck, D.L.R. (2010) On the efficiency of line intersect distance sampling. Canadian Journal of Forest Research, 40, 1086-1094.

Slade, N.A. and Blair, S.M. (2000) An empirical test of using counts of individuals captured as indices of population size. *Journal of Mammalogy*, 81, 1035–1045.

# Reproducibility

Code was developed and tested using R v4.1.2, and key packages with versions as shown below. Worked example code may generate errors if key package versions are earlier than those indicated. Source code is available via GitHub: https://github.com/NEONScience/mee-tos-2022

Table 5: Key package versions used to develop and test worked example code.

Package	Version	Depends	Built
data.table devtools	1.14.2 2.4.3	R (>= 3.1.0)	4.1.0 4.1.0
DT devtools	0.21	R (>= 3.0.2), usethis (>= 2.0.1) NA	4.1.0 $4.1.2$
geoNEON	1.0.0.9100	R (>= 3.5.0)	4.1.2
neonOS	0.1.0	R (>= 3.4.0)	4.1.2
neonUtilities tidyverse	2.1.3 1.3.1	R (>= 3.4.0) R (>= 3.3)	4.1.0 4.1.0