SWMPr: An R package for retrieving, organizing, and analyzing environmental data for estuaries

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## Abstract

Standardized monitoring programs have vastly improved the quantity and quality of data that form the basis of environmental decision-making. One example in the United States is the System Wide Monitoring Program (SWMP) that was implemented in 1995 by the federally-funded National Estuarine Research Reserve System (NERRS). This program has provided two decades of continuous monitoring data at over 300 fixed stations in 28 estuaries across the United States. SWMP data have been used in a variety applications with the general objective of describing dynamics of estuarine ecosystems to better inform effective coastal management. However, simple tools for processing and evaluating the large and increasing quantity of data provided by the monitoring network have prevented large-scale comparisons between systems and, in some cases, simple trend analysis of water quality parameters at individual sites. We describe a new open-source software package, SWMPr, developed in program R for use with SWMP environmental data. The package provides several functions that facilitate data retrieval, organization, and analysis of time series data to describe water quality, weather, and nutrient dynamics in the reserve estuaries. Previously unavailable functions for estuaries are also provided to estimate rates of ecosystem metabolism using the open-water method. Tools included with the SWMPr package have facilitated a cross-reserve comparison of trends, including simple evaluation of changes over time and comparisons of patterns in primary productivity. Overall, the package provides an effective approach to link quantitative information with analysis tools that will greatly inform management programs aimed at coastal protection and restoration.

Introduction

The development of low-cost, automated sensors that have the ability to collect data in near real-time has enabled a proliferation of standardized environmental monitoring programs [1,2]. These programs provide access to invaluable sources of data that can be used to address a variety of research and management objectives. Applications from automated remote sensors are numerous with notable examples including prediction of harmful algal blooms and toxicants in aquatic systems [3], development of a hydrometeorological monitoring network to support flash flood warning programs [4], and automated detection of airborne chemical warfare agents [5]. Automated remote monitoring programs offer several advantages over traditional site-specific, field-based

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methods including streamlining of data acquisition, minimizing human error, and reducing the overall cost of the collection process [1]. However, the growing quantity of available information to address relevant questions has contributed to the notion of 'big data' science where analysis limitations are more often defined by compututational requirements and signal identification in the presence of noise rather than the availability of information. Growing concerns over the use of adequate quality assurance and control methods, challenges for synthesis and interpretation, and increased emphasis on exploratory-based analytical techniques have characterized applications of data from automated monitoring programs [6, 7].

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The National Estuarine Research Reserve System (NERRS, http://www.nerrs.noaa.gov/) is a network of 28 estuarine reserves in the United States that was established by the Coastal Zone Management Act of 1972. The reserves represent different biogeographic regions and estuarine types that were chosen to address multiple goals for long-term research, monitoring, education, and stewardship in support of coastal management. As part of this effort, the System Wide Monitoring Program (SWMP) program was implemented in 1995 at over 300 stations at each of the reserves to provide a robust, long-term monitoring system for water quality, weather, and land-use/habitat change. The SWMP network has provided a continous source of data collected at near real-time at each of the reserves with the intent to evaluate natural and anthrogogenic causes of spatiotemporal variation in environmental condition and ecosystem function. These data have been useful for evaluating relevant characteristics at individual reserves (eg., [8,9]) and differences between reserves (e.g., ecosystem metabolism [10, 11], tidal characteristics [12], dissolved oxygen [13]). However, no cross-reserve comparisons have been conducted within the last decade despite the online availability of current SWMP data. NERRS researchers and staff have also expressed a need for quantitative analysis tools to evaluate trends in water quality time series given the quantity of data provided by SWMP [14].

This article describes a software package that was developed to address research needs of the NERRS program using the open-source statistical programming language R [15]. SWMPr (pronounced 'swamper') is an R package that contains functions for retrieving, organizing, and analyzing estuary monitoring data from the System Wide Monitoring Program (SWMP). Functions provided by SWMPr address many of the common issues working with large datasets created from automated sensor networks, such as data pre-processing to remove unwanted information, combining data from different sources, and exploratory analyses to identify key parameters of interest. Additionally, a cross-reserve comparison of current ecosystem metabolism estimates is provided to illustrate a potential application using the functions in this package. The software is provided specifically for use with NERRS data, although many of the applications are relevant addressing common challenges working with large datasets.

Methods

### Installing the package

The SWMPr package was developed for use with the R statistical programming language and a recent version of R (v3.0.0 or greater) should be installed (see <a href="http://cran.r-project.org/">http://cran.r-project.org/</a>). The SWMPr package can be installed from GitHub by executing the following commands at the R terminal. The package is loaded in the current workspace by using the library command.

install.packages('devtools')

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```
library(devtools)
install_github('fawda123/SWMPr')
library(SWMPr)
```

#### SWMP overview and data retrieval

Four core data elements are collected through the SWMP monitoring network: abiotic monitoring data, biotic observations, habitat and land use mapping, and sentinel monitoring. The SWMPr package is developed for the continuous abiotic monitoring network which includes a majority of the data collected by SWMP. Abjoric elements monitored at each reserve include water quality (water temperature, specific conductivity, salinity, dissolved oxygen concentration, dissolved oxygen saturation, depth, pH, turbidity, chlorophyll fluorescence), weather (air temperature, relative humidity, barometric pressure, wind speed, wind direction, photosynthetically active radiation, precipitation), and nutrient data (orthophosphate, ammonium, nitrite, nitrate, nitrite + nitrate, chlorophyll a). Each reserve has no less than four water quality stations and one weather station at fixed locations. Water quality and weather data are collected at 15 minute intervals, whereas nutrient data are collected monthly at each water quality station. All data are made accessible through the Centralized Data Management Office (CDMO) web portal, where multiple quality assurance/quality control (QAQC) measures are used to screen the information. The final data include all observations with relevant QAQC flags indicating the appropriate qualifier (view codes).

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The CDMO web portal was established to support priority areas of SWMP that focus on the continuation and advancement of data and information management. As such, CDMO provides access to over 35 million water quality, weather, and nutrient records that have been authenticated through systematic QAQC procedures. Estuary data must be obtained from CDMO prior to using most of the functions within the SWMPr package. In most cases, the analysis needs will typically define the location, date range, and parameters of interest that need to be obtained from the CDMO. All stations in the SWMP network are identified by a 7 or 8 character name that specifies the reserve, station, and parameter type. For example, 'apaebwq' is the water quality identifier for the East Bay station at the Apalachicola reserve. Similarly, a suffix of 'met' or 'nut' would specify the weather (meteorological) or nutrients station.

SWMP data can be used in R after they are obtained directly from the CDMO through an online query or by using the retrieval functions provided in this package. Prior to any data request, the site, parameter type, and date ranges need to be identified based on the analysis needs. All reserve names, stations, and date ranges for the water quality, weather, or nutrients data can be viewed on the CDMO website. Alternatively, the site\_codes (all sites) or site\_codes\_ind (single site) functions provided by the SWMPr package can be used to view the same information. As noted below, the computer's IP address must be registered by CDMO staff before using the function.

### Structure of the SWMPr package

The SWMPr package was developed by considering a standard workflow that categorizes the functions as one of three steps based on their intended use: retrieving, organizing, and analyzing. Functions for retrieving are used to import the data into R as a swmpr object class. Functions for organizing and analyzing the data provide methods for working with the swmpr object class. An additional class of functions, termed 'miscellaneous', are also included as helpers for the main functions. The following

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Table 1.	Data retrieval	functions	available from	the	SWMPr	package.
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Function	Description				
all_params	Retrieve up to 100 records starting with the most re-				
	cent at a given station, all parameters. Wrapper to exportAllParamsXMLNew function on web services.				
all_params_dtrng	Retrieve records of all parameters within a given date range for a station. Optional argument for a single				
	parameter. Maximum of 1000 records. Wrapper to exportAllParamsDateRangeXMLNew.				
import_local	Import files from a local path. The files must be in a specific format, specifically those returned from the CDMO using the zip downloads option for a reserve.				
${ t single\_param}$	Retrieve up to 100 records for a single parameter starting with the most recent at a given station. Wrapper to exportSingleParamXMLNew function on web services.				
site_codes	Metadata for all stations, wrapper to exportStationCodesXMLNew function on web services.				
site_codes_ind	Metadata for all stations at a single site, wrapper to NERRFilterStationCodesXMLNew function on web services.				

describes a general approach for using each category of functions based on a standard data workflow.

Data retrieval SWMP data must be obtained from the CDMO prior to organizing and analyzing the information. Two approaches can be used. First, the data can be obtained outside of R using the online query forms on the CDMO website (see here). Second, functions from the SWMPr package can be used to import the data directly from the online server (Table 1). In the latter case, the IP address for the computer making the request must be registered with CDMO. This can be done by following instructions here. The site\_codes\_ind functions can be used to view the available metadata after a computer is registered with CDMO.

```
# retrieve metadata for all sites
site_codes()

# retrieve metadata for a single site
site_codes_ind('apa')
```

Due to rate limitations on the CDMO server, the retrieval functions return a limited number of records. The functions are more useful for evaluating short time periods, although these functions could be used iteratively (i.e., with for loops) to obtain longer time series. Data retrieval functions to access the CDMO include all\_params, all\_params\_dtrng, and single\_param. These are functions that call the existing web protocol methods on the CDMO web services. all\_params returns the most recent records of all parameters at a station, all\_params\_dtrng returns all records within a date range for all parameters or a single parameter, and single\_param is identical to all\_params except that a single parameter is requested.

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```
# all parameters for a station, most recent
all_params('hudscwq')

# get all parameters within a date range
all_params_dtrng('hudscwq', c('09/10/2012', '02/8/2013'))

# get single parameter within a date range
all_params_dtrng('hudscwq', c('09/10/2012', '02/8/2013'),
    param = 'do_mgl')

# single parameter for a station, most recent
single_param('hudscwq', 'do_mgl')
```

For larger requests, data are easier to obtain outside of R using the CDMO query system and then importing using the import\_local function. Data can be retrieved from the CDMO several ways. The import\_local function is designed for data from the zip downloads feature in the advanced query section of the CDMO. The function may also work using data from the data export system, but this feature has not been extensively tested. The zip downloads feature is an easy way to obtain data from multiple stations in one request. The downloaded data will be in a compressed folder that includes multiple .csv files by year for a given data type (e.g., apacpwq2002.csv, apacpwq2003.csv, apacpnut2002.csv, etc.). The import\_local function can be used after the folder is decompressed.

Occasionally, duplicate time stamps are present in the raw data. The import\_local function handles duplicate entries differently depending on the data type (water quality, weather, or nutrients). For water quality and nutrient data, duplicate time stamps are simply removed. Note that nutrient data often contain replicate samples with similar but not duplicated time stamps within a few minutes of each other. Replicates with unique time stamps are not removed but can be further processed using rem\_reps. Weather data prior to 2007 may contain duplicate time stamps at frequencies for 60 (hourly) and 144 (daily) averages, in addition to 15 minute frequencies. Duplicate values that correspond to the smallest value in the frequency column (15 minutes) are retained.

```
# import data for apaebmet that you downloaded

# this is an example path with the csv files, change as needed
path <- 'C:/my_path/'

# import, do not include file extension
import_local(path, 'apaebmet')</pre>
```

All data retrieval functions return a swmpr object that includes relevant data and several attributes describing the dataset. The data include a datetimestamp column in the appropriate timezone for a station. Note that the datetimestamp is standard time for each timezone and does not include daylight savings. Additional columns include parameters for a given data type (weather, nutrients, or water quality) and correspondingg QAQC columns if returned from the initial data request. The attributes for a swmpr object include names of the dataset, row.names of the dataset, class (character string indicating swmpr and data.frame) station (7 or 8 characters identifying the station), parameters (character vector of data columns, e.g., 'do\_mgl'), qaqc\_cols (logical T or F if present or not), date\_rng (POSIXct vector of minimum/maximum dates), timezone (text string in country/city format), and

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stamp\_class (class of datetimestamp vector, POSIXct or Date). Attributes of a swmpr object can be viewed as follows.

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```
# import binary data
data(apadbwq)
dat <- apadbwq
# verify that dat is swmpr class
class(dat)
## [1] "swmpr"
                     "data.frame"
# all attributes of dat
names(attributes(dat))
## [1] "names"
                      "row.names"
                                    "class"
                                                   "station"
## [5] "parameters"
                      "qaqc_cols"
                                    "date_rng"
                                                   "timezone"
## [9] "stamp_class"
# a single attribute of dat
attr(dat, 'station')
## [1] "apadbwq"
```

The swmpr object class was created for use with specific methods following the S3 object definition approach [16]. A swmpr object also secondarily inherits methods from the data.frame class, such that common data.frame methods also apply to swmpr objects. Available methods for the swmpr class are described below and can also be viewed:

```
# available methods for sumpr class
methods(class = 'swmpr')
    [1] aggregate.swmpr
                               aggregate_metab.swmpr comb.swmpr
    [4] decomp.swmpr
                               decomp_cj.swmpr
                                                     ecometab.swmpr
##
   [7] hist.swmpr
                              lines.swmpr
                                                     na.approx.swmpr
## [10] plot.swmpr
                               plot_metab.swmpr
                                                     plot_summary.swmpr
## [13] qaqc.swmpr
                               qaqcchk.swmpr
                                                     rem_reps.swmpr
## [16] setstep.swmpr
                               smoother.swmpr
                                                     subset.swmpr
```

Example data as raw, comma-separated files have not been included in the package due to size limitations. However, a sample dataset can be downloaded for use with the examples below. This dataset has an identical format as the data returned from the zip downloads feature of the CDMO. Processed verions of the raw data are included with the package as binary data files (.RData) to decrease processing times with the examples. Information for each binary file can be viewed as follows.

```
# view help files for complementary data
```

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```
# all files are samples from Apalachicola Bay

# cat point station, nutrients
?apacpnut

# cat point station, water quality
?apacpwq

# dry bar station, water quality
?apadbwq

# east bay station, weater
?apaebmet
```

**Data organizing** The retrieval functions import the data into R as a swmpr object for use with the organize and analyze functions. The organize functions are used to clean or prepare the data for analysis, including removal of QAQC flags, subsetting, creating a standardized time series vector, and combining data of different types.

The qaqc function is a simple screen to retain values from the data with specified QAQC flags (described here). Each parameter in the swmpr data typically has a corresponding QAQC column of the same name with the added prefix f. Values in the QAQC column specify a flag from -5 to 5. Generally, only data with the 0 QAQC flag should be used, which is the default option for the QAQC function. Data that do not satisfy QAQC criteria are converted to NA values. Additionally, simple filters are used to remove obviously bad values, e.g., wind speed values less than zero or pH values greater than 12. Erroneous data entered as -99 are also removed. Processed data will have QAQC columns removed, in addition to removal of values in the actual parameter columns that do not meet the criteria.

```
# qaqc screen for a swmpr object, retain only '0'
qaqc(dat)

# retain all data regardless of flag
qaqc(dat, qaqc_keep = NULL)

# retain only '0' and '-1' flags
qaqc(dat, qaqc_keep = c(0, -1))
```

Viewing the number of observations for each parameter that are assigned to a QAQC flag may be useful for deciding how to process the data with ), qaqc. The qaqcchk function can be used to view this information. Consult the online documentation for a description of each QAQC flag.

```
# view the number of observations in each QAQC flag
qaqcchk(dat)
```

Raw nutrient data obtained from the CDMO will usually include replicate samples that were taken within a few minutes of each other. The rem\_reps.swmpr function combines nutrient data that occur on the same day to preserve an approximate monthly time step. The datetimestamp column will always be averaged for replicates, but the actual observations will be combined based on the user-supplied function which defauls

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to the mean. Other suggested functions include the median, min, or max. The entire function call including treatment of NA values should be passed to the FUN argument (see the examples). The function is meant to be used after qaqc processing, although it works with a warning if QAQC columns are present.

```
# get nutrient data
data(apacpnut)
swmp1 <- apacpnut
swmp1 <- qaqc(swmp1)

# remove replicate nutrient data
rem_reps(swmp1)

# use different function to aggregate replicates
func <- function(x) max(x, na.rm = T)
rem_reps(swmp1, FUN = func)</pre>
```

A subset method added to the existing subset function is available for swmpr objects. This function is used to subset the data by date and/or a selected parameter. The date can be a single value or as two dates to select records within the range. The former case requires a binary operator input as a character string passed to the argument, such as > or <. The subset argument for the date(s) must also be a character string of the format YYYY-mm-dd HH:MM for each element (i.e., %Y-%m%-%d %H:%M in POSIX standards). Be aware that an error may be returned using this function if the subset argument is in the correct format but the calendar date does not exist, e.g. 2012-11-31 12:00. Finally, the function can be used to remove rows and columns that do not contain data.

```
# select two parameters from dat
subset(dat, select = c('rh', 'bp'))

# subset records greater than or equal to a date
subset(dat, subset = '2013-01-01 0:00', operator = '>=')

# subset records within a date range
subset(dat, subset = c('2012-07-01 6:00', '2012-08-01 18:15'))

# subset records within a date range, select two parameters
subset(dat, subset = c('2012-07-01 6:00', '2012-08-01 18:15'),
    select = c('atemp', 'totsorad'))

# remove rows/columns that do not contain data
subset(dat, rem_rows = T, rem_cols = T)
```

The setstep function formats a swmpr object to a continuous time series at a given time step. This function is not necessary for most stations but can be useful for combining data or converting an existing time series to a set interval. The first argument of the function, timestep, specifies the desired time step in minutes starting from the nearest hour of the first observation. The second argument, differ, specifies the allowable tolerance in minutes for matching existing observations to user-defined time steps in cases where the two are dissimilar. Values for differ that are greater than one half the value of timestep are not allowed to prevent duplication of existing data. Likewise, the default value for differ is one half the time step. Rows that do not

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match any existing data within the limits of the differ argument are not discarded. Output from the setstep function can be used with subset and to create a time series at a set interval with empty data removed.

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```
# convert time series to two hour invervals
# tolerance of +/- 30 minutes for matching existing data
setstep(dat, timestep = 120, differ = 30)

# convert a nutrient time series to a continuous time series
# then remove empty rows and columns
data(apacpnut)
dat_nut <- apacpnut
dat_nut <- setstep(dat_nut, timestep = 60)
subset(dat_nut, rem_rows = T, rem_cols = T)</pre>
```

The comb function is used to combine multiple swmpr objects into a single object with a continuous time series at a given step. The timestep function is used internally such that timestep and differ are accepted arguments for comb. The function requires one or more swmpr objects as input as separate, undefined arguments. The remaining arguments must be called explicitly since an arbitrary number of objects can be used as input. In general, the function combines data by creating a master time series that is used to iteratively merge all swmpr objects. The time series for merging depends on the value passed to the method argument. Passing union to method will create a time series that is continuous starting from the earliest date and the latest date for all input objects. Passing intersect to method will create a time series that is continuous from the set of dates that are shared between all input objects. Finally, a seven or eight character station name passed to method will merge all input objects based on a continuous time series for the given station. The specified station must be present in the input data. Currently, combining data types from different stations is not possible, excluding weather data which are typically at a single, dedicated station.

```
# get nuts, wq, and met data as separate objects for the same station
# note that most sites usually have one weather station
data(apacpnut)
data(apacpwq)
data(apaebmet)
swmp1 <- apacpnut
swmp2 <- apacpwq
swmp3 <- apaebmet

# combine nuts and wq data by union
comb(swmp1, swmp2, method = 'union')

# combine nuts and wq data by intersect
comb(swmp1, swmp3, method = 'intersect')

# combine nuts, wq, and met data by nuts time series, two hour time step
comb(swmp1, swmp2, swmp3, timestep = 120, method = 'apacpnut')</pre>
```

**Data analysis** The analysis functions range from general purpose tools for time series analysis to more specific functions for working with continuous monitoring data in estuaries. The latter category includes a limited number of functions that were

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developed by myself or others. The general purpose tools are swmpr methods that were developed for existing generic functions in the R base installation or relevant packages. These functions include swmpr methods for aggregate, filter, and approx to deal with missing or noisy data and more general functions for exploratory data analysis, such as plot, summary, and hist methods. Decomposition functions decomp and decomp\_cj) are provided as relatively simple approaches for decomposing time series into additive or multiplicative components. The analysis functions may or may not return a swmpr object depending on whether further processing with swmpr methods is possible from the output.

The aggregate function aggregates parameter data for a swmpr object by set periods of observation. This function is most useful for aggregating noisy data to evaluate trends on longer time scales, or to simply reduce the size of a dataset. Data can be aggregated by years, quarters, months, weeks, days, or hours for a user-defined function, which defaults to the mean. A swmpr object is returned for the aggregated data, although the datetimestamp vector will be converted to a date object if the aggregation period is a day or longer. Days are assigned to the date vector if the aggregation period is a week or longer based on the round method for IDate objects data.table package. This approach was used to facilitate plotting using predefined methods for Date and POSIX objects. Additionally, the method of treating NA values for the aggregation function should be noted since this may greatly affect the quantity of data that are returned (see the example below). Finally, the default argument for na.action is set to na.pass for swmpr objects to preserve the time series of the input data.

```
# combine, qaqc, remove empty columns
dat <- comb(swmp1, swmp2, method = 'union')
dat <- qaqc(dat)
swmpr_in <- subset(dat, rem_cols = T)

# get mean DO by quarters
aggregate(swmpr_in, 'quarters', params = c('do_mgl'))

# get mean DO by quarters, remove NA when calculating means
fun_in <- function(x) mean(x, na.rm = T)
aggregate(swmpr_in, FUN = fun_in, 'quarters', params = c('do_mgl'))</pre>
```

Time series can be smoothed to better characterize a signal independent of noise (Fig. 1). Although there are many approaches to smoothing, a moving window average is intuitive and commonly used. The smoother function can be used to smooth parameters in a swmpr object using a specified window size. This method is a simple wrapper to filter. The window argument specifies the number of observations included in the moving average. The sides argument specifies how the average is calculated for each observation (see the documentation for filter). A value of 1 will filter observations within the window that are previous to the current observation, whereas a value of 2 will filter all observations within the window centered at zero lag from the current observation. As before, the params argument specifies which parameters to smooth. See Fig. 1 for the output from the code.

```
# import data
```

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```
data(apadbwq)
swmp1 <- apadbwq

# qaqc and subset imported data
dat <- qaqc(swmp1)
dat <- subset(dat, subset = c('2012-07-09 00:00', '2012-07-24 00:00'))

# filter
test <- smoother(dat, window = 50, params = 'do_mgl')

# plot to see the difference
plot(do_mgl ~ datetimestamp, data = dat, type = 'l')
lines(test, select = 'do_mgl', col = 'red', lwd = 2)</pre>
```

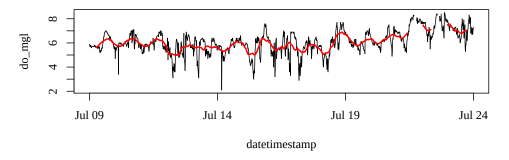


Fig. 1. Raw and smoothed dissolved oxygen data for a two-week period after using the smoother function.

A common issue with any statistical analysis is the treatment of missing values. Missing data can be excluded from the analysis, included but treated as true zeroes, or interpolated based on similar values. In either case, an analyst should have a strong rationale for the chosen method. A common approach used to handle missing data in time series analysis is linear interpolation. A simple curve fitting method is used to create a continuous set of records between observations separated by missing data. A challenge with linear interpolation is an appropriate gap size for fitting missing observations. The ability of the interpolated data to approximate actual trends is a function of the gap size. Interpolation between larger gaps are less likely to resemble patterns of an actual parameter, whereas interpolation between smaller gaps are more likely to resemble actual patterns. An appropriate gap size limit depends on the unique characteristics of specific datasets or parameters. The na.approx function can be used to interpolate gaps in a swmpr object. A required argument for the function is maxgap which defines the maximum gap size for interpolation. See Fig. 2 for the output from the following code.

```
# qet data
```

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```
data(apadbwq)
swmp1 <- apadbwq
# gagc and subset imported data
dat <- gagc(swmp1)</pre>
dat \leftarrow subset(dat, subset = c('2013-01-22 00:00', '2013-01-26 00:00'))
# interpolate, maxgap of 10 records
test <- na.approx(dat, params = 'do_mgl', maxgap = 10)
# interpolate maxgap of 30 records
test2 <- na.approx(dat, params = 'do_mgl', maxgap = 30)</pre>
# plot for comparison
par(mfrow = c(3, 1))
plot(do_mgl ~ datetimestamp, dat, main = 'Raw', type = 'l')
plot(do_mgl ~ datetimestamp, test, col = 'red',
  main = 'Interpolation - maximum gap of 10 records', type = '1')
lines(dat, select = 'do_mgl')
plot(do_mgl ~ datetimestamp, test2, col = 'red',
  main = 'Interpolation - maximum gap of 30 records', type = '1')
lines(dat, select = 'do_mgl')
```

The decomp function is a simple wrapper to decompose that separates a time series into additive or multiplicative components describing a trend, cyclical variation (e.g., daily or seasonal), and the remainder. The additive decomposition assumes that the cyclical component of the time series is stationary (i.e., the variance is constant), whereas a multiplicative decomposition accounts for non-stationarity. By default, a moving average with a symmetric window is used to filter the seasonal component. Alternatively, a vector of filter coefficients in reverse time order can be supplied (see the help documentation for decompose).

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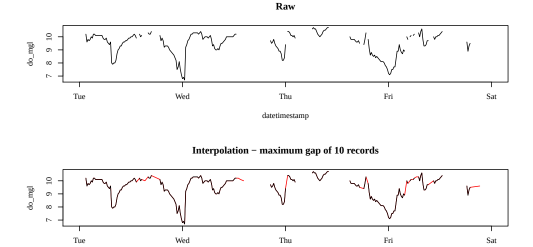
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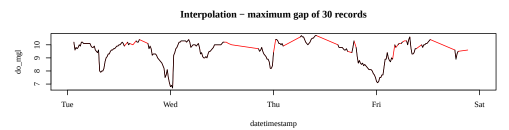
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The decompose function requires a ts object with a specified frequency as input. The decomp function converts the input swmpr vector to a ts object prior to decompose. This requires an explicit input defining the frequency of the parameter in the time series. For example, the frequency of a parameter with diurnal periodicity would be 96 if the time step is 15 minutes (4 \* 24). The frequency of a parameter with seasonal periodicity would be 35040 (4 \* 24 \* 365). For simplicity, character strings of 'daily' or 'seasonal' can be supplied in place of numeric values. A starting value of the time series must be supplied in the latter case. Use of the setstep function is also required to standardize the time step prior to decomposition. Note that the decompose function is a relatively simple approach and alternative methods should be investigated if a more sophisticated decomposition is desired. ?? is an example of the decomp function.

```
# get data
```

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datetimestamp

Fig. 2. Examples illustrating use of the na.approx function to fill gaps of different sizes in a dissolved oxygen time series for a four day period.

```
data(apadbwq)
swmp1 <- apadbwq

# subset for daily decomposition
dat <- subset(swmp1, subset = c('2013-07-01 00:00', '2013-07-31 00:00'))

# decomposition and plot
test <- decomp(dat, param = 'do_mgl', frequency = 'daily')
plot(test)</pre>
```

An alternative approach to time series decomposition is provided by the <code>decomp\_cj</code> function, which is a simple wrapper to the <code>decompTs</code> function in the wq package. Theory describing this method is provided by Cloern and Jassby [17]. The function is similar to <code>decomp.swmpr</code> with a few key differences. The <code>decomp.swmpr</code> function decomposes the time series into a trend, seasonal, and random component, whereas the current function decomposes into the grandmean, annual, seasonal, and events components. For both functions, the random or events components, respectively, can be considered anomalies that do not follow the trends in the remaining categories. The <code>decomp\_cj</code> function provides only a monthly decomposition, which is appropriate for characterizing relatively long-term trends. This approach works best for nutrient data that are typically obtained on a monthly cycle. The function will also work with continuous

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### Decomposition of additive time series

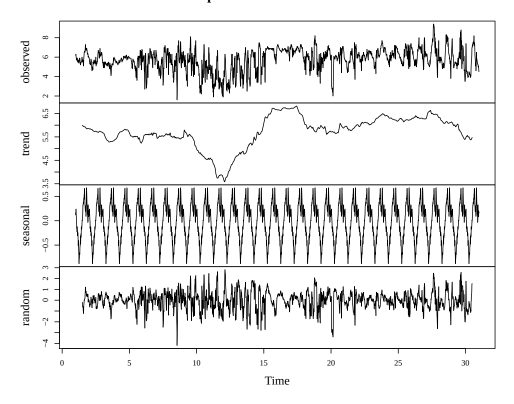


Fig. 3. An additive decomposition of dissolved oxygen into a trend, seasonal, and random component using the decomp function.

water quality or weather data but note that the data must first aggregated on the monthly scale before decomposition. Additional arguments passed to decompTs can be used with decomp\_cj, such as startyr, endyr, and type. Values passed to type are mult (default) or add, referring to multiplicative or additive decomposition. ?? shows the results from the decomp\_cj function applied to a multi-year chlorophyll time series.

```
# get data
data(apacpnut)
dat <- apacpnut
dat <- qaqc(dat, qaqc_keep = NULL)

# decomposition of chl, ggplot
decomp_cj(dat, param = 'chla_n')</pre>
```

Several graphics showing seasonal and annual trends for a given SWMP parameter can be obtained using the plot\_summary function. The plots include monthly distributions, monthly anomalies, and annual anomalies in multiple formats. Anomalies are defined as the difference between the monthly or annual average from the grand mean for the parameter. Monthly anomalies are in relation to the grand mean for the same month across all years. All data are aggregated for quicker plotting. Nutrient data are based on monthly averages, whereas weather and water quality data are based on daily averages. Cumulative precipitation data are based on the daily maximum. The

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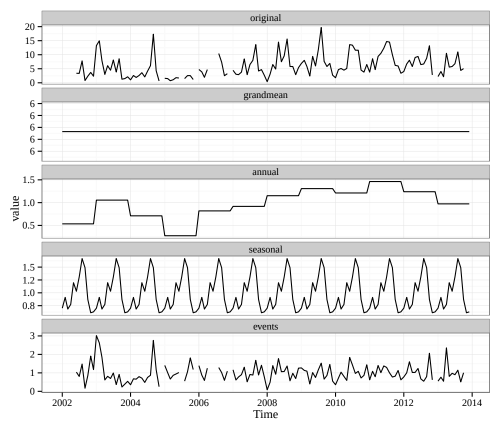


Fig. 4. Additive decomposition of a multi-year chlorophyll time series into the grandmean, annual, seasonal, and events components using the decomp\_cj function.

function returns a graphics object (Grob) of multiple ggplot objects. An interactive Shiny application [18] that uses this function is available (see the Supporting Information).

```
## import data
data(apacpnut)
dat <- qaqc(apacpnut)

## plot
plot_summary(dat, param = 'chla_n')</pre>
```

Estimates of ecosystem metabolism provide a useful measure of overall system productivity. These estimates are commonly used to evaluate whether an ecosystem is a net source or sink of organic material. The open-water method is a common approach to quantify net ecosystem metabolism using a mass balance equation that describes the change in dissolved oxygen over time from the balance between photosynthetic and respiration rates, corrected for air-sea gas diffusion at the surface [19]. The diffusion-corrected DO flux estimates are averaged during day and night for each 24 hour period in the time series, where flux is an hourly rate of DO change. DO flux is averaged during night hours for respiration and averaged during day hours for net production. Respiration rates are assumed constant during day and night such that total daily rates are calculated as hourly respiration multiplied by 24. The metabolic

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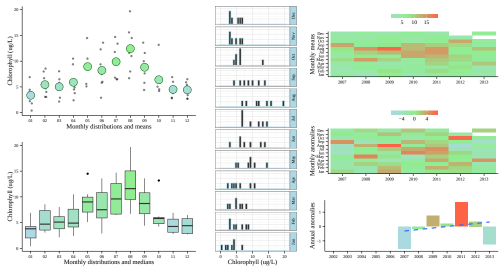


Fig. 5. Summaries of a multi-year chlorophyll time series using the plot\_summary function. Summaries include monthly distributions (means on top left, quantiles on bottom left), monthly histograms (center), monthly means by year (top right), deviation from monthly means (middle right), and annual trends as deviations from the grand mean (bottom right)

day is considered the 24 hour period between sunsets on two adjacent calendar days. Respiration is subtracted from daily net production estimates to yield gross production.

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The ecometab function is used to implement an adaptation of the open-water method [19, 20]. Several assumptions must be met for a valid interpretation of the results. In general, the dissolved oxygen time series is assumed to represent the same water mass over time. Tidal advection may have a significant influence on the time series, which can contribute to a significant amount of noise in metabolic estimates. The extent to which tidal advection influences the dissolved oxygen signal depends on various site-level characteristics and an intimate knowledge of the site may be required. Volumetric rates for gross production and total respiration are also based on total depth of the water column, which is assumed to be mixed. Water column depth is based on mean value for the depth variable across the time series and is floored at 1 meter for very shallow stations. Additionally, the volumetric reaeration coefficient requires an estimate of the anemometer height of the weather station, which is set as 10 meters by default. The metadata should be consulted for exact height. Other assumptions may apply and the user should consult the relevant literature (see the references in the help file). All estimates are in mmol of oxygen but can be converted to grams by changing the default arguments (i.e., 1mmol O2 = 32 mg O2, 1000 mg = 1 g, multiply all estimates by 32/1000).

The following is an example that shows how to use the function from a combined water quality and weather data set. The results can be plotted using plot\_metab (??).

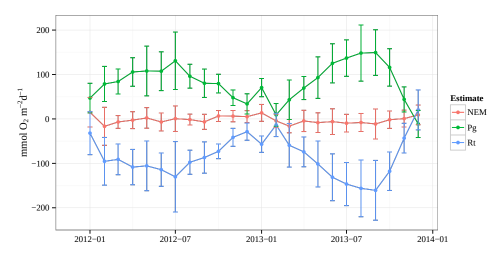
## import water quality and weather data

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```
data(apadbwq)
data(apaebmet)

## qaqc, combine
wq <- qaqc(apadbwq)
met <- qaqc(apaebmet)
dat <- comb(wq, met)

## estimate metabolism
res <- ecometab(dat, trace = FALSE)
plot_metab(res)</pre>
```



**Fig. 6.** Monthly aggregations of ecosystem metabolism estimates (net ecosystem metabolism, gross production, and total respiration) for combined water quality and weather data at Apalachicola Bay, Florida.

Miscellaneous functions Several additional functions are provided that cannot be described by the above categories. These functions are generally used within the main functions but may be useful for more customized evaluation of SWMP data.

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Finally, a reserve map with all stations can be obtained using the map\_reserve function. This function is a simple wrapper to functions in the ggmap package [21]. The current function is limited to Google maps, which allows four map types that can be set with the map\_type argument: terrain (default), satellite, roadmap, or hybrid. The zoom argument may have to be chosen through trial and error depending on the spatial extent of the reserve. See the help documentation for ggmap for more info on zoom.

```
# plot the stations at Jacques Cousteau reserve
map_reserve('jac')
```

Function list

Organize comb.swmpr Combines swmpr objects to a common time series using setstep, such as combining the weather, nutrients, and water quality data for a single station. Only different data types can be combined.

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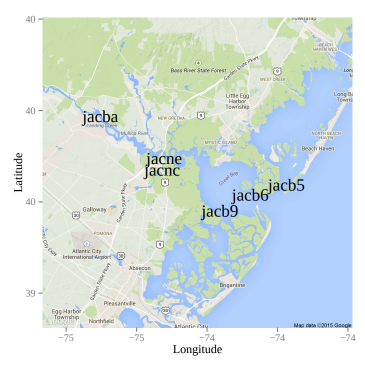


Fig. 7. Locations of all sites at the Jacques Cousteau reserve using the map\_reserve function.

qaqc.swmpr Remove QAQC columns and remove data based on QAQC flag values for a swmpr object. Only applies if QAQC columns are present.

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qaqcchk.swmpr View a summary of the number of observations in a swmpr object that are assigned to different QAQC flags used by CDMO. The output is used to inform further processing but is not used explicitly.

rem\_reps.swmpr Remove replicate nutrient data that occur on the same day. The default is to average replicates.

setstep.swmpr Format data from a swmpr object to a continuous time series at a given timestep. The function is used in comb.swmpr and can also be used with individual stations.

subset.swmpr Subset by dates and/or columns for a swmpr object. This is a method passed to the generic subset function provided in the base package.

Analyze aggregate.swmpr Aggregate swmpr objects for different time periods - years, quarters, months, weeks, days, or hours. Aggregation function is user-supplied but defaults to mean.

aggregate\_metab Aggregate metabolism data from a swmpr object. This is primarly used within plot\_metab but may be useful for simple summaries of raw daily data.

ecometab.swmpr Estimate ecosystem metabolism for a combined water quality and weather dataset using the open-water method.

decomp.swmpr Decompose a swmpr time series into trend, seasonal, and residual components. This is a simple wrapper to decompose. Decomposition of monthly or daily trends is possible.

decomp\_cj.swmpr Decompose a swmpr time series into grandmean, annual, seasonal, and events components. This is a simple wrapper to decompTs in the wq package. Only monthly decomposition is possible.

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hist.swmpr Plot a histogram for a swmpr object.

lines.swmpr Add lines to an existing swmpr plot.

na.approx.swmpr Linearly interpolate missing data (NA values) in a swmpr object. The maximum gap size that is interpolated is defined as a maximum number of records with missing data.

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plot.swmpr Plot a univariate time series for a swmpr object. The parameter name must be specified.

plot\_metab Plot ecosystem metabolism estimates after running ecometab on a swmpr object.

plot\_summary Create summary plots of seasonal/annual trends and anomalies for a water quality or weather parameter.

smoother.swmpr Smooth swmpr objects with a moving window average. Window size and sides can be specified, passed to filter.

**Miscellaneous** calckL Estimate the reaeration coefficient for air-sea gas exchange. This is only used within the ecometab function.

map\_reserve Create a map of all stations in a reserve using the ggmap package.

metab\_day Identify the metabolic day for each approximate 24 period in an hourly time series. This is only used within the ecometab function.

param\_names Returns column names as a list for the parameter type(s) (nutrients, weather, or water quality). Includes QAQC columns with f\_ prefix. Used internally in other functions.

parser Parses html returned from CDMO web services, used internally in retrieval functions.

swmpr Creates object of swmpr class, used internally in retrieval functions.

time\_vec Converts time vectors to POSIX objects with correct time zone for a site/station, used internally in retrieval functions.

Results

Discussion

# **Supporting Information**

## Trends in SWMP parameters

https://beckmw.shinyapps.io/swmp\_comp This widget is an interactive tool to evaluate

trends in SWMP data within and between sites. Trends are described by an increase or decrease in values over time using a simple linear regression of summarized data. The regression for each station can be viewed by clicking on a map location. Trends at each station are plotted as circles that identify the direction and significance of the trend. The trend direction is blue for decreasing and red for increasing. The significance is indicated by radius of the circle and color shading where larger points with darker colors indicate a strong trend.

## Monthly and annual summary of SWMP parameters

https://beckmw.shinyapps.io/swmp\_summary/

This interactive widget provides graphical summaries of water quality, weather, and nutrient station data from SWMP. The drop down menus can be used to select the station, date range, and parameter for plotting. The raw data used for plotting include

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all SWMP records from the earliest date at each station after processing to remove QAQC flags. The data were downloaded from the CDMO web services on November 25th, 2014 and include observations up to that date. Plots are based on daily averages for each parameter. Cumulative precipitation data are based on the daily maximum.

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