

Package: manureshed (via r-universe)

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Type Package

Title Spatiotemporal Nutrient Balance Analysis Across Agricultural and Municipal Systems

Version 0.1.5

Maintainer Olatunde D. Akanbi <olatunde.akanbi@case.edu>

Description A comprehensive framework for analyzing agricultural nutrient balances across multiple spatial scales (county, 'HUC8', 'HUC2') with integration of wastewater treatment plant ('WWTP') effluent loads for both nitrogen and phosphorus. Supports classification of spatial units as nutrient sources, sinks, or balanced areas based on agricultural surplus and deficit calculations. Includes visualization tools, spatial transition probability analysis, and nutrient flow network mapping. Built-in datasets include agricultural nutrient balance data from the Nutrient Use Geographic Information System ('NuGIS'; The Fertilizer Institute and Plant Nutrition Canada, 1987-2016) <https://nugis.tfi.org/tabular_data/> and U.S. Environmental Protection Agency ('EPA') wastewater discharge data from the 'ECHO' Discharge Monitoring Report ('DMR') Loading Tool (2007-2016) <<https://echo.epa.gov/trends/loading-tool/water-pollution-search>>. Data are downloaded on demand from the Open Science Framework ('OSF') repository to minimize package size while maintaining full functionality. The integrated 'manureshed' framework methodology is described in Akanbi et al. (2025) <[doi:10.1016/j.resconrec.2025.108697](https://doi.org/10.1016/j.resconrec.2025.108697)>. Designed for nutrient management planning, environmental analysis, and circular economy research at watershed/administrative to national scales. This material is based upon financial support by the National Science Foundation, EEC Division of Engineering Education and Centers, NSF Engineering Research Center for Advancing Sustainable and Distributed Fertilizer Production (CASFER), NSF 20-553 Gen-4 Engineering Research Centers award 2133576. We thank Dr. Robert D. Sabo (U.S. Environmental Protection Agency) for his valuable contributions to the

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URL <https://osf.io/g39xa/>, <https://github.com/cwru-sdle/manureshed>,
<https://exelegch.github.io/manureshed-docs/>

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add_centroid_coordinates

Add Centroid Coordinates to Spatial Data

Description

Calculate centroid coordinates for spatial units

Usage

add_centroid_coordinates(spatial_data)

Arguments

spatial_data sf object. Spatial data

Value

Data frame with centroid coordinates added

add_texas_huc8	<i>Add Texas HUC8 Data (Updated for OSF)</i>
----------------	--

Description

Add manually supplied Texas HUC8 data for missing watersheds Uses OSF data loading instead of built-in data

Usage

```
add_texas_huc8(huc8_data, year = 2016, cropland_threshold, verbose = TRUE)
```

Arguments

huc8_data	sf object. Existing HUC8 agricultural data
year	Numeric. Year to extract from Texas data
cropland_threshold	Numeric. Threshold for classification
verbose	Logical. Show progress messages

Value

sf object with Texas data added

agri_classify_complete	<i>Complete Agricultural Classification Pipeline</i>
------------------------	--

Description

Run complete agricultural nutrient classification analysis for both N and P

Usage

```
agri_classify_complete(
  nugis_data,
  scale,
  cropland_threshold = NULL,
  county_data = NULL
)
```

Arguments

nugis_data	Data frame. Raw NuGIS data
scale	Character. Spatial scale: "county", "huc8", or "huc2"
cropland_threshold	Numeric. Optional custom threshold
county_data	Data frame. County data for threshold calculation (if needed)

Value

Data frame with complete agricultural classifications for both nutrients

agri_classify_complete_custom
Complete Agricultural Classification Pipeline with Custom Efficiency Factors

Description

Run complete agricultural nutrient classification analysis for both N and P with user-specified efficiency factors for sensitivity analysis.

Usage

```
agri_classify_complete_custom(
  nugis_data,
  scale,
  cropland_threshold = NULL,
  county_data = NULL,
  n_efficiency = 0.5,
  p_efficiency = 1
)
```

Arguments

nugis_data	Data frame. Raw NuGIS data
scale	Character. Spatial scale: "county", "huc8", or "huc2"
cropland_threshold	Numeric. Optional custom threshold
county_data	Data frame. County data for threshold calculation (if needed)
n_efficiency	Numeric. Nitrogen efficiency factor (default: 0.5)
p_efficiency	Numeric. Phosphorus efficiency factor (default: 1.0)

Value

Data frame with complete agricultural classifications for both nutrients

Examples

```
# Load county data
nugis_data <- load_builtin_nugis("county", 2016)

# Standard analysis
results_standard <- agri_classify_complete_custom(
  nugis_data, "county"
)

# Sensitivity analysis with varied nitrogen efficiency
results_high_n <- agri_classify_complete_custom(
  nugis_data, "county",
  n_efficiency = 0.7
)

# Analysis with both custom efficiencies
results_custom <- agri_classify_complete_custom(
  nugis_data, "county",
  n_efficiency = 0.6,
  p_efficiency = 0.9
)
```

agri_classify_nitrogen

Classify Agricultural Nitrogen Status

Description

Classify spatial units based on nitrogen balance using standard 0.5 efficiency factor

Usage

```
agri_classify_nitrogen(data, cropland_threshold, scale = "huc8")
```

Arguments

data	Data frame with processed agricultural data
cropland_threshold	Numeric. Threshold for excluding small cropland areas
scale	Character. Spatial scale for within-unit classification

Value

Data frame with nitrogen classification added

```
agri_classify_nitrogen_custom
```

Classify Agricultural Nitrogen Status with Custom Efficiency Factor

Description

Classify spatial units based on nitrogen balance with user-specified efficiency factor. This function allows sensitivity analysis by varying the nitrogen efficiency assumption. The default value of 0.5 represents typical losses during nutrient cycling, uptake, and application, but regional conditions may warrant different values.

Usage

```
agri_classify_nitrogen_custom(
  data,
  cropland_threshold,
  scale = "huc8",
  n_efficiency = 0.5
)
```

Arguments

data	Data frame with processed agricultural data
cropland_threshold	Numeric. Threshold for excluding small cropland areas
scale	Character. Spatial scale for within-unit classification
n_efficiency	Numeric. Nitrogen efficiency factor (default: 0.5, range: 0-1)

Value

Data frame with nitrogen classification added

Examples

```
# Load and process data first
nugis_data <- load_built_in_nugis("county", 2016)
processed_data <- agri_process_nugis(nugis_data, "county")
cropland_threshold <- 500 * 2.47105 # 500 ha in acres

# Standard analysis with default 0.5 efficiency
results_default <- agri_classify_nitrogen_custom(
  processed_data, cropland_threshold = cropland_threshold, n_efficiency = 0.5
)

# Sensitivity analysis with higher efficiency (e.g., improved management)
results_high <- agri_classify_nitrogen_custom(
  processed_data, cropland_threshold = cropland_threshold, n_efficiency = 0.7
)
```

```

# Conservative analysis with lower efficiency
results_low <- agri_classify_nitrogen_custom(
  processed_data, cropland_threshold = cropland_threshold, n_efficiency = 0.3
)

# Compare classification changes across efficiency scenarios
table(results_default$N_class)
table(results_high$N_class)
table(results_low$N_class)

```

```
agri_classify_phosphorus
```

Classify Agricultural Phosphorus Status

Description

Classify spatial units based on phosphorus balance (no efficiency factor for P)

Usage

```
agri_classify_phosphorus(data, cropland_threshold, scale = "huc8")
```

Arguments

data	Data frame with processed agricultural data
cropland_threshold	Numeric. Threshold for excluding small cropland areas
scale	Character. Spatial scale for within-unit classification

Value

Data frame with phosphorus classification added

```
agri_classify_phosphorus_custom
```

Classify Agricultural Phosphorus Status with Custom Efficiency Factor

Description

Classify spatial units based on phosphorus balance with user-specified efficiency factor. While standard phosphorus classification uses 100% allows sensitivity analysis by varying the phosphorus efficiency assumption for different management scenarios or application methods.

Usage

```
agri_classify_phosphorus_custom(
  data,
  cropland_threshold,
  scale = "huc8",
  p_efficiency = 1
)
```

Arguments

data	Data frame with processed agricultural data
cropland_threshold	Numeric. Threshold for excluding small cropland areas
scale	Character. Spatial scale for within-unit classification
p_efficiency	Numeric. Phosphorus efficiency factor (default: 1.0, range: 0-1)

Value

Data frame with phosphorus classification added

Examples

```
# Load and process data first
nugis_data <- load_built_in_nugis("county", 2016)
processed_data <- agri_process_nugis(nugis_data, "county")
cropland_threshold <- 500 * 2.47105 # 500 ha in acres

# Standard analysis with default 1.0 efficiency (100%)
results_default <- agri_classify_phosphorus_custom(
  processed_data, cropland_threshold = cropland_threshold, p_efficiency = 1.0
)

# Analysis with reduced efficiency (e.g., accounting for losses)
results_reduced <- agri_classify_phosphorus_custom(
  processed_data, cropland_threshold = cropland_threshold, p_efficiency = 0.8
)

# Conservative analysis with lower efficiency
results_conservative <- agri_classify_phosphorus_custom(
  processed_data, cropland_threshold = cropland_threshold, p_efficiency = 0.6
)

# Compare classification changes across efficiency scenarios
table(results_default$P_class)
table(results_reduced$P_class)
table(results_conservative$P_class)
```

agri_process_nugis *Process NuGIS Data for Manureshed Analysis*

Description

Clean and standardize NuGIS data for agricultural nutrient analysis

Usage

```
agri_process_nugis(nugis_data, scale)
```

Arguments

nugis_data	Data frame. Raw NuGIS data for specified year
scale	Character. Spatial scale: "county", "huc8", or "huc2"

Value

Data frame with standardized columns for analysis

batch_analysis_enhanced
Enhanced Batch Analysis with Full Visualizations

Description

Run batch analysis with comprehensive visualization output for each year

Usage

```
batch_analysis_enhanced(  
  years,  
  scale = "huc8",  
  nutrients = c("nitrogen", "phosphorus"),  
  include_wwtp = TRUE,  
  output_base_dir = tempdir(),  
  create_all_visualizations = TRUE,  
  create_comparative_plots = TRUE,  
  show_progress = TRUE,  
  verbose = TRUE,  
  ...  
)
```

Arguments

years	Numeric vector. Years to analyze
scale	Character. Spatial scale
nutrients	Character vector. Nutrients to analyze
include_wwtp	Logical. Include WWTP analysis
output_base_dir	Character. Base output directory
create_all_visualizations	Logical. Create all maps, networks, and comparisons
create_comparative_plots	Logical. Create year-over-year comparisons
show_progress	Logical. Display progress bar (requires 'progress' package)
verbose	Logical. Show progress
...	Additional arguments

Value

List of results with comprehensive outputs

Examples

```
## Not run:
# This function is computationally intensive
# See vignette("advanced-features") for examples
results <- batch_analysis_enhanced(years = 2015:2016)

## End(Not run)
```

batch_analysis_parallel

Batch Analysis with Parallel Processing

Description

Run batch analysis using multiple cores for faster processing

Usage

```
batch_analysis_parallel(years, n_cores = NULL, ...)
```

Arguments

years	Numeric vector. Years to analyze
n_cores	Integer. Number of cores (default: detectCores() - 1)
...	Arguments passed to run_builtin_analysis

Value

List of results

Examples

```
results <- batch_analysis_parallel(
  years = 2015:2016, # Use valid years only
  n_cores = 2,      # Max 2 cores for CRAN
  scale = "county", # Use county for faster processing
  nutrients = "nitrogen"
)
```

batch_analysis_years *Batch Analysis Across Multiple Years*

Description

Run manureshed analysis across multiple years with consistent parameters

Usage

```
batch_analysis_years(
  years,
  scale = "huc8",
  nutrients = c("nitrogen", "phosphorus"),
  include_wwtp = TRUE,
  output_base_dir = tempdir(),
  create_comparative_plots = TRUE,
  verbose = TRUE,
  ...
)
```

Arguments

years	Numeric vector. Years to analyze
scale	Character. Spatial scale: "county", "huc8", or "huc2"
nutrients	Character vector. Nutrients to analyze
include_wwtp	Logical. Whether to include WWTP (only available for 2007-2016 built-in)
output_base_dir	Character. Base output directory
create_comparative_plots	Logical. Whether to create year-over-year comparisons
verbose	Logical. Whether to print progress
...	Additional arguments passed to run_built_in_analysis

Value

List of results for each year

Examples

```
# Analyze trends with WWTP for subset of supported range
batch_results <- batch_analysis_years(
  years = 2010:2012, # Use smaller range for examples
  scale = "huc8",
  nutrients = "nitrogen",
  include_wwtp = TRUE
)

# Historical analysis without WWTP
historical_results <- batch_analysis_years(
  years = 1990:1992, # Use smaller range
  scale = "county",
  nutrients = c("nitrogen", "phosphorus"),
  include_wwtp = FALSE
)

# Mixed analysis: some years with WWTP, some without
mixed_results <- batch_analysis_years(
  years = c(2005, 2010, 2015), # 2010,2015 will have WWTP
  scale = "huc8",
  nutrients = "nitrogen",
  include_wwtp = TRUE # Will only apply to 2010,2015
)
```

benchmark_analysis *Benchmark Analysis Performance*

Description

Test analysis speed and memory usage

Test analysis speed and memory usage

Usage

```
benchmark_analysis(
  scale = "huc8",
  year = 2016,
  nutrients = "nitrogen",
  n_runs = 3,
  include_wwtp = TRUE
)
```

```
benchmark_analysis(  
  scale = "huc8",  
  year = 2016,  
  nutrients = "nitrogen",  
  n_runs = 3,  
  include_wwtp = TRUE  
)
```

Arguments

scale	Character. Spatial scale
year	Numeric. Year to test
nutrients	Character vector. Nutrients to analyze
n_runs	Integer. Number of benchmark runs (default: 3)
include_wwtp	Logical. Include WWTP processing

Value

List with timing statistics and memory usage

List with timing statistics and memory usage

Examples

```
# Benchmark HUC8 analysis - use smaller scale for faster testing  
benchmark <- benchmark_analysis(  
  scale = "county", # Use county for faster testing  
  year = 2016,  
  nutrients = "nitrogen",  
  n_runs = 2 # Reduce runs for faster testing  
)  
print(benchmark)
```

```
# Benchmark HUC8 analysis - use smaller scale for faster testing  
benchmark <- benchmark_analysis(  
  scale = "county", # Use county for faster testing  
  year = 2016,  
  nutrients = "nitrogen",  
  n_runs = 2 # Reduce runs for faster testing  
)  
print(benchmark)
```

`calculate_cropland_threshold`*Calculate Cropland Threshold for Exclusion*

Description

Calculate cropland threshold for excluding small agricultural areas Uses county 500ha baseline to determine percentile for other scales

Usage

```
calculate_cropland_threshold(  
  county_data,  
  target_data,  
  county_cropland_col,  
  target_cropland_col,  
  baseline_ha = 500  
)
```

Arguments

`county_data` Data frame. County-level NuGIS data with cropland column
`target_data` Data frame. Target scale data (HUC8, HUC2) with cropland column
`county_cropland_col` Character. Name of cropland column in county data
`target_cropland_col` Character. Name of cropland column in target data
`baseline_ha` Numeric. Baseline cropland in hectares for exclusion (default: 500)

Value

Numeric. Threshold value for target scale

`calculate_transition_probabilities`*Calculate Spatial Transition Probabilities*

Description

Calculate transition probabilities between adjacent spatial units

Usage

```
calculate_transition_probabilities(  
  data,  
  class_column,  
  longitude_col = "longitude",  
  latitude_col = "latitude"  
)
```

Arguments

data	Data frame. Data with classification and coordinate columns
class_column	Character. Name of classification column
longitude_col	Character. Name of longitude column (default: "longitude")
latitude_col	Character. Name of latitude column (default: "latitude")

Value

Data frame with transition probabilities as percentages

check_builtin_data	<i>Check Data Availability from OSF</i>
--------------------	---

Description

Check what datasets are available from the OSF repository and which are cached locally.

Usage

```
check_builtin_data(verbose = FALSE)
```

Arguments

verbose	Logical. Show detailed information about cache status
---------	---

Value

List showing available datasets and cache status

`citation_info`*Display Package Citation Information*

Description

Provides citation information for the package and data sources. Prints formatted citation text to the console for the manureshed package, the underlying research methodology paper (Akanbi et al., 2026), and the primary data sources (NuGIS agricultural data and EPA WWTP discharge data). The function is designed for users to easily obtain proper citations for publications and reports.

Usage

```
citation_info()
```

Details

This function takes no arguments. It prints citation information directly to the console using `message()` functions, which can be suppressed with `suppressMessages()` if needed.

Value

No return value, called for side effects. The function prints citation information to the console including:

- Package citation with version and OSF repository
- Research methodology paper citation
- NuGIS data source attribution
- EPA WWTP data source attribution
- Contact information for data sources

Note

This function requires no arguments and can be called simply as `citation_info()`.

See Also

[check_builtin_data](#) for data availability, [health_check](#) for package diagnostics

Examples

```
# Display citation information
citation_info()
```

clean_category_names *Clean Category Names for Display*

Description

Clean Category Names for Display

Clean Category Names for Display

Usage

```
clean_category_names(names)
```

```
clean_category_names(names)
```

Arguments

names Character vector of category names to clean

Value

Character vector of cleaned names

Character vector of cleaned names

clean_text *Clean Text Data*

Description

Remove extra quotes and whitespace from text fields

Remove extra quotes and whitespace from text fields

Usage

```
clean_text(text)
```

```
clean_text(text)
```

Arguments

text Character vector to clean

Value

Character vector of cleaned text

Character vector of cleaned text

clear_data_cache	<i>Clear Data Cache</i>
------------------	-------------------------

Description

Remove cached datasets to free up disk space

Usage

```
clear_data_cache(confirm = TRUE, verbose = TRUE)
```

Arguments

confirm	Logical. Require confirmation before deleting
verbose	Logical. Show what's being deleted

Value

Logical. TRUE if successful

compare_analyses	<i>Compare Two Analysis Results</i>
------------------	-------------------------------------

Description

Compare classifications between two analysis results

Compare classifications between two analysis results

Usage

```
compare_analyses(results1, results2, nutrient = "nitrogen")
```

```
compare_analyses(results1, results2, nutrient = "nitrogen")
```

Arguments

results1	First analysis results
results2	Second analysis results
nutrient	Character. Nutrient to compare ("nitrogen" or "phosphorus")

Value

Data frame with comparison

Data frame with comparison

Examples

```
results_2010 <- run_builtin_analysis(scale = "county", year = 2010)
results_2016 <- run_builtin_analysis(scale = "county", year = 2016)
comparison <- compare_analyses(results_2010, results_2016, "nitrogen")
```

```
results_2010 <- run_builtin_analysis(scale = "county", year = 2010)
results_2016 <- run_builtin_analysis(scale = "county", year = 2016)
comparison <- compare_analyses(results_2010, results_2016, "nitrogen")
```

compare_regions	<i>Compare Multiple Regions</i>
-----------------	---------------------------------

Description

Compare Multiple Regions

Usage

```
compare_regions(  
  regions,  
  scale = "county",  
  year = 2016,  
  nutrients = "nitrogen",  
  include_wwtp = TRUE,  
  verbose = TRUE  
)
```

Arguments

regions	Named list of state vectors, or character vector of built-in region names
scale	Spatial scale
year	Year to analyze
nutrients	Nutrients to analyze
include_wwtp	Include WWTP?
verbose	Show messages?

Value

List with regional comparison results

Examples

```
## Not run:
# Compare built-in regions
comparison <- compare_regions(
  regions = c("corn_belt", "dairy_belt"),
  scale = "county",
  year = 2016,
  nutrients = "nitrogen"
)

# Compare custom regions
comparison <- compare_regions(
  regions = list(
    "Upper Midwest" = c("WI", "MN", "MI"),
    "Lower Midwest" = c("IL", "IN", "OH")
  ),
  scale = "county",
  year = 2016
)

# View comparison
print(comparison$summary)
comparison$plots$bar_chart

## End(Not run)
```

 compare_scenarios

Compare Multiple Analysis Scenarios

Description

Compare results from multiple analysis runs side-by-side with visualizations and summary statistics.

Usage

```
compare_scenarios(
  scenario_list,
  metrics = c("sources", "sinks", "balanced", "excluded"),
  create_plots = TRUE,
  output_dir = NULL
)
```

Arguments

scenario_list Named list of analysis results from `run_built_in_analysis()`

metrics Character vector of metrics to compare. Options: "sources", "sinks", "balanced", "excluded", "total_surplus", "total_deficit"

create_plots Logical. Create comparison plots? (default: TRUE)
 output_dir Character. Directory for saving plots (default: NULL, no save)

Value

List containing:

comparison_data Data frame with metrics for each scenario
 plots List of ggplot objects (if create_plots = TRUE)
 summary Summary statistics

Examples

```
# Create multiple scenarios
base <- run_built_in_analysis(year = 2016, include_wwtp = FALSE,
                             scale = "county", nutrients = "nitrogen")
wwtp <- run_built_in_analysis(year = 2016, include_wwtp = TRUE,
                              scale = "county", nutrients = "nitrogen")

# Compare scenarios
comparison <- compare_scenarios(list(
  "Base (Agricultural Only)" = base,
  "With WWTP" = wwtp
))

# View comparison data
print(comparison$comparison_data)

# View plots
print(comparison$plots$bar_chart)
```

 CONUS_STATES

CONUS States

Description

Vector of Continental United States state abbreviations
 Vector of Continental United States state abbreviations

Usage

CONUS_STATES

CONUS_STATES

Format

An object of class character of length 49.

An object of class character of length 49.

convert_load_units *Convert Load Units to Tons*

Description

Convert loads from various units to US tons

Convert loads from various units to US tons

Usage

```
convert_load_units(load_values, from_unit)
```

```
convert_load_units(load_values, from_unit)
```

Arguments

load_values Numeric vector of load values

from_unit Character. Input unit: "kg", "lbs", "pounds", "tons"

Value

Numeric vector of loads in US tons

Numeric vector of loads in US tons

Examples

```
# Convert from kilograms to tons
kg_loads <- c(1000, 2000, 3000)
tons_loads <- convert_load_units(kg_loads, "kg")

# Convert from pounds to tons
lbs_loads <- c(5000, 10000, 15000)
tons_loads <- convert_load_units(lbs_loads, "lbs")
# Convert from kilograms to tons
kg_loads <- c(1000, 2000, 3000)
tons_loads <- convert_load_units(kg_loads, "kg")

# Convert from pounds to tons
lbs_loads <- c(5000, 10000, 15000)
tons_loads <- convert_load_units(lbs_loads, "lbs")
```

```
create_analysis_report
```

Create Analysis Report

Description

Generate comprehensive HTML or PDF report of analysis results

Usage

```
create_analysis_report(  
  results,  
  output_path,  
  format = "html",  
  title = "Manureshed Analysis Report",  
  include_maps = TRUE  
)
```

Arguments

results	List. Complete analysis results
output_path	Character. Path for output report
format	Character. Report format: "html" or "pdf"
title	Character. Report title
include_maps	Logical. Whether to include maps in report

Value

Character. Path to generated report

Examples

```
# Generate HTML report - use tempdir to avoid check directory pollution  
results <- run_builtin_analysis(scale = "county", year = 2016)  
report_path <- file.path(tempdir(), "analysis_report.html")  
create_analysis_report(results, report_path)
```

`create_classification_summary`*Create Classification Summary Table*

Description

Create summary table of classification counts for both nutrients

Usage

```
create_classification_summary(data, agricultural_col, combined_col)
```

Arguments

<code>data</code>	Data frame. Data with classification columns
<code>agricultural_col</code>	Character. Name of agricultural classification column
<code>combined_col</code>	Character. Name of combined (with WWTP) classification column

Value

Data frame with classification counts and changes

`create_network_plot`*Create Network Plot from Transition Probabilities*

Description

Create network visualization of spatial transition probabilities

Usage

```
create_network_plot(  
  transition_df,  
  nutrient,  
  analysis_type,  
  output_path,  
  highlight_transitions = TRUE  
)
```

Arguments

transition_df	Data frame. Transition probability matrix
nutrient	Character. "nitrogen" or "phosphorus" for coloring
analysis_type	Character. Description of analysis type
output_path	Character. Path to save PNG file
highlight_transitions	Logical. Whether to highlight specific transitions

Value

NULL (saves plot to file)

data_sources	<i>NuGIS and EPA WWTP Data Sources</i>
--------------	--

Description

This package uses two primary data sources:

NuGIS Agricultural Data

The Nutrient Use Geographic Information System (NuGIS) presents cropland nutrient balances for the conterminous United States from 1987-2016.

Source: The Fertilizer Institute (TFI) and Plant Nutrition Canada (PNC)

Website: https://nugis.tfi.org/tabular_data

Contact: nugis@tfi.org

Components:

- County-level crop and livestock data from USDA Census of Agriculture
- Fertilizer use data from AAPFCO
- Geospatial nutrient balance estimates
- Available for counties, HUC8, and HUC2 watersheds

Data Processing: The manureshed package uses cleaned versions of NuGIS data with resolved metadata issues and enhanced spatial integration, as detailed in the manureshed methodology paper (Akanbi et al., 2026).

EPA WWTP Data

Source: U.S. Environmental Protection Agency

System: Discharge Monitoring Report (DMR) Loading Tool via ECHO

Website: <https://echo.epa.gov/trends/loading-tool/water-pollution-search>

Data Years: 2007-2016 (nitrogen and phosphorus loads)

License: Public domain (U.S. Government work)

Data Attribution

When using this package, please cite both the package, methodology paper, and the underlying data sources. Use `citation_info()` to display full citation information.

<code>download_all_data</code>	<i>Download All Datasets from OSF</i>
--------------------------------	---------------------------------------

Description

Convenience function to download all available datasets from OSF

Usage

```
download_all_data(force_download = FALSE, verbose = TRUE)
```

Arguments

`force_download` Logical. Re-download even if files exist in cache
`verbose` Logical. Show progress for each download

Value

Logical. TRUE if all downloads successful

<code>download_osf_data</code>	<i>Download and Cache Data from OSF</i>
--------------------------------	---

Description

Download built-in datasets from OSF repository using Files API

Usage

```
download_osf_data(  
  dataset_name,  
  force_download = FALSE,  
  cache_dir = NULL,  
  verbose = TRUE  
)
```

Arguments

dataset_name	Character. Name of dataset to download
force_download	Logical. Force re-download even if cached version exists
cache_dir	Character. Directory to cache downloaded data. Defaults to tempdir() for CRAN compliance. For persistent caching across sessions, set options(manureshed.cache_dir = tools::R_user_dir("manureshed", "cache")).
verbose	Logical. Show download progress

Value

Path to cached data file

export_for_gis	<i>Export Results for GIS Applications</i>
----------------	--

Description

Export spatial results in GIS-ready formats

Usage

```
export_for_gis(
  results,
  output_dir = file.path(tempdir(), "gis_export"),
  formats = c("shapefile", "geojson")
)
```

Arguments

results	Analysis results object
output_dir	Output directory
formats	Character vector of formats: "shapefile", "geojson", "kml", "gpkg"

Value

List of created files

Examples

```
# Use tempdir to avoid polluting check directory
results <- run_built_in_analysis(scale = "county", year = 2016)
output_dir <- file.path(tempdir(), "gis_outputs")
gis_files <- export_for_gis(results, output_dir)
```

export_for_policy *Export for Policy Briefs*

Description

Create simplified outputs for policy makers

Usage

```
export_for_policy(results, output_dir = file.path(tempdir(), "policy_export"))
```

Arguments

results	Analysis results
output_dir	Output directory

Value

List of created files

export_for_publication
Export for Publication

Description

Create high-resolution outputs suitable for publication

Usage

```
export_for_publication(  
  results,  
  output_dir = file.path(tempdir(), "publication_export"),  
  dpi = 600  
)
```

Arguments

results	Analysis results
output_dir	Output directory
dpi	Resolution (default: 600 for publication quality)

Value

List of created files

filter_by_state	<i>Filter Data by State</i>
-----------------	-----------------------------

Description

Filter spatial data to a specific state

Usage

```
filter_by_state(data, state, scale, boundaries = NULL)
```

Arguments

data	Data frame or sf object. Spatial data with FIPS or HUC codes
state	Character. Two-letter state abbreviation
scale	Character. Spatial scale: "county", "huc8", or "huc2"
boundaries	sf object. Spatial boundaries (optional, for HUC scales)

Value

Filtered data for the specified state

format_huc8	<i>Format HUC8 Codes</i>
-------------	--------------------------

Description

Add leading zeros to 7-digit HUC8 codes to make them 8-digit

Add leading zeros to 7-digit HUC8 codes to make them 8-digit

Usage

```
format_huc8(huc_codes)
```

```
format_huc8(huc_codes)
```

Arguments

huc_codes	Character or numeric vector of HUC codes
-----------	--

Value

Character vector of properly formatted 8-digit HUC codes

Character vector of properly formatted 8-digit HUC codes

```
get_cropland_threshold
```

Get Cropland Threshold by Scale

Description

Get appropriate cropland threshold based on spatial scale

Usage

```
get_cropland_threshold(
  scale,
  county_data = NULL,
  target_data = NULL,
  baseline_ha = 500
)
```

Arguments

scale	Character. Spatial scale: "county", "huc8", or "huc2"
county_data	Data frame. County-level data (required for huc8/huc2)
target_data	Data frame. Target scale data (required for huc8/huc2)
baseline_ha	Numeric. Baseline for county exclusion (default: 500)

Value

Numeric. Threshold value

```
get_nutrient_colors
```

Default Color Schemes for Nutrient Classifications

Description

Default Color Schemes for Nutrient Classifications

Default Color Schemes for Nutrient Classifications

Usage

```
get_nutrient_colors(nutrient = "nitrogen")
```

```
get_nutrient_colors(nutrient = "nitrogen")
```

Arguments

nutrient	Character. Either "nitrogen" or "phosphorus"
----------	--

Value

Named vector of colors for classification categories

Named vector of colors for classification categories

get_region_definitions

Get Built-in Region Definitions

Description

Get Built-in Region Definitions

Usage

`get_region_definitions()`

Value

List of region definitions

get_state_boundaries *Get State Boundaries for Mapping*

Description

Get US state boundaries excluding non-CONUS states

Usage

`get_state_boundaries()`

Value

sf object with state boundaries

get_state_fips	<i>Get State FIPS Code</i>
----------------	----------------------------

Description

Convert state abbreviation to FIPS code

Usage

```
get_state_fips(state_abbr)
```

Arguments

state_abbr Character. Two-letter state abbreviation (e.g., "OH", "TX")

Value

Character. Two-digit state FIPS code

health_check	<i>Check Package Health and Dependencies</i>
--------------	--

Description

Diagnostic function to check package installation and dependencies

Usage

```
health_check(verbose = FALSE)
```

Arguments

verbose Logical. Whether to display detailed information

Value

Logical. TRUE if all checks pass

Examples

```
# Quick health check
health_check()

# Detailed diagnostic information
health_check(verbose = TRUE)
```

integrate_complete *Complete Integration Pipeline (Updated)*

Description

Run complete integration of WWTP and agricultural data for both nutrients

Usage

```
integrate_complete(  
  agri_data,  
  wwtp_nitrogen_aggregated,  
  wwtp_phosphorus_aggregated,  
  cropland_threshold,  
  scale = "huc8",  
  add_texas = FALSE,  
  year = 2016  
)
```

Arguments

agri_data	Data frame or sf object. Agricultural classification data
wwtp_nitrogen_aggregated	Data frame. Aggregated nitrogen WWTP data
wwtp_phosphorus_aggregated	Data frame. Aggregated phosphorus WWTP data
cropland_threshold	Numeric. Threshold for exclusion
scale	Character. Spatial scale
add_texas	Logical. Whether to add Texas HUC8 data (only for HUC8 scale)
year	Numeric. Year for Texas data

Value

List with integrated nitrogen and phosphorus data

```
integrate_wwtp_agricultural
```

Integrate WWTP Data with Agricultural Classifications (Updated)

Description

Combine WWTP loads with agricultural nutrient balance classifications

Usage

```
integrate_wwtp_agricultural(  
  agri_data,  
  wwtp_aggregated,  
  nutrient,  
  cropland_threshold,  
  scale = "huc8"  
)
```

Arguments

agri_data	Data frame. Agricultural classification data
wwtp_aggregated	Data frame. Aggregated WWTP loads by spatial unit
nutrient	Character. "nitrogen" or "phosphorus"
cropland_threshold	Numeric. Threshold for exclusion classification
scale	Character. Spatial scale for within-unit classification

Value

Data frame with combined WWTP + agricultural classifications

```
KG_TO_TONS
```

Conversion Factor: Kilograms to US Tons

Description

Conversion Factor: Kilograms to US Tons

Conversion Factor: Kilograms to US Tons

Usage

```
KG_TO_TONS
```

```
KG_TO_TONS
```

Format

An object of class `numeric` of length 1.

An object of class `numeric` of length 1.

launch_dashboard	<i>Launch Interactive Manureshed Dashboard</i>
------------------	--

Description

Opens an interactive Shiny dashboard for exploring nutrient balance data without writing R code. Users can select analysis parameters, view interactive maps, explore statistics, and download results.

Usage

```
launch_dashboard(port = NULL, launch.browser = TRUE)
```

Arguments

`port` Integer. Port number for the Shiny server (default: random available port)

`launch.browser` Logical. Open browser automatically? (default: TRUE)

Details

This function launches an interactive dashboard with the following features:

- Select analysis parameters (year, scale, nutrients, WWTP inclusion)
- View interactive maps of nutrient classifications
- Explore summary statistics and distributions
- Browse results in interactive data tables
- Download results as CSV files
- Generate and download PDF reports

The dashboard requires several suggested packages. If not installed, run: `install.packages(c("shiny", "shinydashboard", "leaflet", "plotly", "DT"))`

Value

NULL (invisibly). Launches the Shiny application.

Examples

```
## Not run:  
# Launch the dashboard (requires suggested packages)  
launch_dashboard()  
  
# Launch on specific port  
launch_dashboard(port = 3838)  
  
## End(Not run)
```

LBS_TO_KG

Conversion Factor: Pounds to Kilograms

Description

Conversion Factor: Pounds to Kilograms

Conversion Factor: Pounds to Kilograms

Usage

LBS_TO_KG

LBS_TO_KG

Format

An object of class `numeric` of length 1.

An object of class `numeric` of length 1.

LBS_TO_TONS

Conversion Factor: Pounds to US Tons

Description

Conversion Factor: Pounds to US Tons

Conversion Factor: Pounds to US Tons

Usage

LBS_TO_TONS

LBS_TO_TONS

Format

An object of class `numeric` of length 1.

An object of class `numeric` of length 1.

`list_available_years` *List Available Built-in Years*

Description

Show available years for each data type

Show available years for each data type

Usage

```
list_available_years(scale = NULL)
```

```
list_available_years(scale = NULL)
```

Arguments

`scale` Character. Spatial scale (optional)

Value

Data frame with available years by data type

Data frame with available years by data type

`list_regions` *List Available Regions*

Description

List Available Regions

Usage

```
list_regions()
```

 load_builtin_boundaries

Load Built-in Spatial Boundaries from OSF

Description

Load built-in spatial boundary data for specified scale from OSF repository

Usage

```
load_builtin_boundaries(scale, force_download = FALSE, verbose = TRUE)
```

Arguments

scale	Character. Spatial scale: "county", "huc8", or "huc2"
force_download	Logical. Force re-download even if cached
verbose	Logical. Show download progress

Value

sf object with spatial boundaries

load_builtin_nugis

Load Built-in NuGIS Data from OSF

Description

Load built-in NuGIS data from OSF repository for specified year and spatial scale. Data includes all years from 1987 through 2016.

Usage

```
load_builtin_nugis(scale, year = 2016, force_download = FALSE, verbose = TRUE)
```

Arguments

scale	Character. Spatial scale: "county", "huc8", or "huc2"
year	Numeric. Year to filter data (available: 1987-2016)
force_download	Logical. Force re-download even if cached
verbose	Logical. Show download progress

Value

Data frame of NuGIS data for specified scale and year

Examples

```
# Load county data for 2016
county_2016 <- load_builtin_nugis("county", 2016)

# Load HUC8 data for 2010
huc8_2010 <- load_builtin_nugis("huc8", 2010)

# Load county data for 2010, force fresh download
county_2010 <- load_builtin_nugis("county", 2010, force_download = TRUE)
```

load_builtin_wwtp *Load Built-in WWTP Data from OSF*

Description

Load built-in WWTP data for specified year from OSF repository (2007-2016 available)

Usage

```
load_builtin_wwtp(
  nutrient,
  year = 2016,
  force_download = FALSE,
  verbose = TRUE
)
```

Arguments

nutrient	Character. "nitrogen" or "phosphorus"
year	Numeric. Year to filter data (available: 2007-2016)
force_download	Logical. Force re-download even if cached
verbose	Logical. Show download progress

Value

Data frame with cleaned WWTP data for specified year

Examples

```
# Load WWTP nitrogen data for different years (2007-2016 available)
wwtp_n_2016 <- load_builtin_wwtp("nitrogen", 2016)
wwtp_n_2012 <- load_builtin_wwtp("nitrogen", 2012)
wwtp_n_2007 <- load_builtin_wwtp("nitrogen", 2007)

# Load phosphorus data
wwtp_p_2015 <- load_builtin_wwtp("phosphorus", 2015)
```

```

wwtp_p_2010 <- load_builtin_wwtp("phosphorus", 2010)

# Force re-download
wwtp_fresh <- load_builtin_wwtp("nitrogen", 2014, force_download = TRUE)

```

load_user_wwtp	<i>Load User WWTP Data</i>
----------------	----------------------------

Description

Load and standardize user-provided WWTP data with flexible formatting

Load and standardize user-provided WWTP data with flexible formatting

Usage

```

load_user_wwtp(
  file_path,
  nutrient,
  column_mapping = NULL,
  skip_rows = 0,
  header_row = 1,
  load_units = "kg"
)

```

```

load_user_wwtp(
  file_path,
  nutrient,
  column_mapping = NULL,
  skip_rows = 0,
  header_row = 1,
  load_units = "kg"
)

```

Arguments

file_path	Character. Path to WWTP data file
nutrient	Character. "nitrogen" or "phosphorus"
column_mapping	Named list. Custom column mapping (optional)
skip_rows	Numeric. Number of rows to skip (default: 0)
header_row	Numeric. Row containing headers (default: 1)
load_units	Character. Units of pollutant loads: "kg", "lbs", "pounds", "tons"

Value

Data frame with standardized WWTP data

Data frame with standardized WWTP data

Examples

```
## Not run:
# Standard EPA format but will not run because data needs to be supplied as custom
# Load custom WWTP data (for years outside 2007-2016)
wwtp_data <- load_user_wwtp("nitrogen_2020.csv", "nitrogen")

# For years 2007-2016, consider using built-in data:
# wwtp_builtin <- load_builtin_wwtp("nitrogen", 2015)

# Custom format with different units
wwtp_data <- load_user_wwtp("custom_wwtp.csv", "phosphorus",
                           load_units = "lbs", skip_rows = 3)

# Custom column mapping
custom_map <- list(facility_name = "Plant_Name",
                  pollutant_load = "Load_lbs_per_year")
wwtp_data <- load_user_wwtp("custom.csv", "nitrogen", custom_map)

## End(Not run)
## Not run:

# Standard EPA format
wwtp_data <- load_user_wwtp("nitrogen_2020.csv", "nitrogen")

# Custom format with different units
wwtp_data <- load_user_wwtp("custom_wwtp.csv", "phosphorus",
                           load_units = "lbs", skip_rows = 3)

# Custom column mapping
custom_map <- list(facility_name = "Plant_Name",
                  pollutant_load = "Load_lbs_per_year")
wwtp_data <- load_user_wwtp("custom.csv", "nitrogen", custom_map)

## End(Not run)
```

manureshed

manureshed: Manureshed Analysis with WWTP Integration

Description

This package provides comprehensive tools for analyzing agricultural nutrient balances at multiple spatial scales with optional integration of wastewater treatment plant (WWTP) nutrient loads for both nitrogen and phosphorus.

Details

All datasets are downloaded on-demand from OSF repository to minimize package size while maintaining full functionality.

Main Functions

Data Loading and Management::

- `load_builtin_nugis`: Load NuGIS data from OSF
- `load_builtin_boundaries`: Load spatial boundaries from OSF
- `load_builtin_wwtp`: Load WWTP data from OSF (2016)
- `check_builtin_data`: Check available datasets and cache status
- `download_all_data`: Download all datasets at once
- `download_osf_data`: Download specific dataset
- `clear_data_cache`: Clear cached datasets

High-Level Workflows::

- `run_builtin_analysis`: Complete end-to-end analysis workflow
- `quick_analysis`: Analysis with automatic visualizations
- `batch_analysis_years`: Multi-year analysis workflow

Agricultural Classification::

- `agri_process_nugis`: Process and standardize NuGIS data
- `agri_classify_nitrogen`: Classify nitrogen balance
- `agri_classify_phosphorus`: Classify phosphorus balance
- `agri_classify_complete`: Complete agricultural pipeline

WWTP Processing::

- `load_user_wwtp`: Load custom WWTP data with flexible formatting
- `wwtp_clean_data`: Clean and filter WWTP data
- `wwtp_classify_sources`: Classify WWTP facilities by load size
- `convert_load_units`: Handle different load units (kg, lbs, tons)

Data Integration::

- `integrate_wwtp_agricultural`: Combine WWTP and agricultural data
- `integrate_complete`: Complete integration pipeline
- `add_texas_huc8`: Add Texas HUC8 supplemental data

Visualization and Mapping::

- `map_agricultural_classification`: Map nutrient classifications
- `map_wwtp_points`: Map WWTP facility locations
- `map_wwtp_influence`: Map WWTP influence/proportion
- `get_state_boundaries`: Get US state boundaries for mapping

Spatial Analysis::

- `calculate_transition_probabilities`: Spatial transition analysis
- `create_network_plot`: Network visualization of transitions
- `add_centroid_coordinates`: Calculate spatial centroids

Comparison Analysis::

- `create_classification_summary`: Before/after comparison summaries
- `plot_before_after_comparison`: Comparison bar plots
- `plot_impact_ratios`: Impact ratio visualizations
- `plot_absolute_changes`: Absolute change plots

Utility Functions::

- `get_nutrient_colors`: Get color schemes for nutrients
- `clean_category_names`: Clean classification names for display
- `format_huc8`: Format HUC8 codes with leading zeros
- `get_cropland_threshold`: Calculate exclusion thresholds

Spatial Scales

The package supports analysis at three spatial scales:

- **County**: US county boundaries (3,000+ units)
- **HUC8**: 8-digit Hydrologic Unit Code watersheds (2,000+ units)
- **HUC2**: 2-digit Hydrologic Unit Code regions (18 units)

Nutrients Supported

The package supports analysis for both major nutrients with appropriate methodologies:

- **Nitrogen**: Uses 0.5 availability factor for manure nitrogen in calculations
- **Phosphorus**: Direct calculation without availability factor

Users can analyze one nutrient, both nutrients, or different combinations in the same workflow:
`nutrients = c("nitrogen", "phosphorus")`

Classification System

Spatial units are classified into five categories based on nutrient balance:

- **Source**: Net nutrient surplus available for export
- **Sink Deficit**: Total deficit requiring nutrient imports
- **Sink Fertilizer**: Fertilizer surplus available for manure import
- **Within Watershed/County**: Balanced for internal nutrient transfers
- **Excluded**: Insufficient cropland for meaningful analysis (<500 ha equivalent)

Data Sources

The package provides access to comprehensive built-in datasets:

NuGIS Data::

- County-level data (1987 - 2016)
- HUC8 watershed data (1987 - 2016)
- HUC2 regional data (1987 - 2016)

- All nutrient balance components (manure, fertilizer, removal, fixation)

Spatial Boundaries::

- US county boundaries (CONUS)
- HUC8 watershed boundaries
- HUC2 regional boundaries
- All in Albers Equal Area Conic projection (EPSG:5070)

WWTP Data::

- Nitrogen discharge data (2007 - 2016)
- Phosphorus discharge data (2007 - 2016)
- Pre-processed and classification-ready
- Includes facility metadata and spatial coordinates

Supplemental Data::

- Texas HUC8 data (automatically integrated for HUC8 analyses)
- Texas spatial boundaries

OSF Data Repository

All datasets are hosted on OSF and downloaded on-demand:

- **Repository:** <https://osf.io/g39xa/>
- **Automatic caching:** Data downloaded once and reused
- **Flexible loading:** Load only the data you need
- **Version control:** Permanent DOI for reproducibility
- **Size efficiency:** Package <1MB, full datasets ~25MB

WWTP Data Flexibility

The package handles varying EPA WWTP data formats across different years:

- **Unit Conversion:** Automatic conversion between kg, lbs, pounds, and tons
- **Column Mapping:** Flexible mapping to handle EPA naming changes
- **Header Detection:** Support for different header row positions
- **Built-in 2016:** Ready-to-use cleaned data for immediate analysis
- **Custom Integration:** Easy integration of user data for other years

Workflow Examples

```
# Check what data is available
check_builtin_data()

# Download all datasets (optional, ~40MB)
download_all_data()
```

```
# Basic analysis using built-in data - any year 2007-2016
results <- run_builtin_analysis(
  scale = "huc8",
  year = 2012, # Any year 2007-2016 now supported
  nutrients = c("nitrogen", "phosphorus"),
  include_wwtp = TRUE
)

wwtp_n_2010 <- load_builtin_wwtp("nitrogen", year = 2010)
wwtp_p_2015 <- load_builtin_wwtp("phosphorus", year = 2015)

# Quick analysis with automatic visualizations
viz_results <- quick_analysis(
  scale = "county",
  year = 2016,
  nutrients = "nitrogen",
  include_wwtp = TRUE
)

# Historical analysis without WWTP
historical <- run_builtin_analysis(
  scale = "huc8",
  year = 2010,
  nutrients = c("nitrogen", "phosphorus"),
  include_wwtp = FALSE
)

# Load specific datasets manually
county_2016 <- load_builtin_nugis("county", 2016)
boundaries <- load_builtin_boundaries("county")
wwtp_n <- load_builtin_wwtp("nitrogen")
```

Analysis Outputs

The package generates comprehensive outputs for each nutrient analyzed:

- **Spatial Data:** Classification results as sf objects
- **Maps:** Classification, influence, and facility maps
- **Networks:** Transition probability visualizations
- **Comparisons:** Before/after WWTP integration analysis
- **Data Files:** CSV centroids, RDS spatial data
- **Metadata:** Analysis parameters and processing information

Performance and Scalability

- Optimized for CONUS-scale analysis (1000s of spatial units)
- Memory-efficient spatial operations

- On-demand data loading reduces memory footprint
- Progress reporting for long-running analyses
- Automatic garbage collection for memory management

Author(s)

Maintainer: Olatunde D. Akanbi <olatunde.akanbi@case.edu> ([ORCID](#)) [copyright holder]

Authors:

- Vibha Mandayam ([ORCID](#))
- Atharva Gupta ([ORCID](#))
- K. Colton Flynn ([ORCID](#))
- Jeffrey Yarus ([ORCID](#))
- Erika I. Barcelos <erika.barcelos@case.edu> ([ORCID](#)) [copyright holder]
- Roger H. French <roger.french@case.edu> ([ORCID](#)) [copyright holder]

See Also

Useful links:

- <https://osf.io/g39xa/>
- <https://github.com/cwru-sdle/manureshed>
- <https://exelegch.github.io/manureshed-docs/>

MANURESHEDED_CRCS

Standard CRS for Manureshed Analysis

Description

Albers Equal Area Conic projection (EPSG:5070)

Albers Equal Area Conic projection (EPSG:5070)

Usage

MANURESHEDED_CRCS

MANURESHEDED_CRCS

Format

An object of class `numeric` of length 1.

An object of class `numeric` of length 1.

map_agricultural_classification
Create Agricultural Classification Map

Description

Create map showing agricultural nutrient classifications

Usage

```
map_agricultural_classification(data, nutrient, classification_col, title)
```

Arguments

data	sf object. Spatial data with classifications
nutrient	Character. "nitrogen" or "phosphorus"
classification_col	Character. Name of classification column
title	Character. Map title

Value

ggplot object

map_wwtp_columns *Map WWTP Column Names to Standard Format*

Description

Create mapping between EPA WWTP data column names and standardized format. Handles various EPA data formats across different years.

Usage

```
map_wwtp_columns(raw_data, nutrient, custom_mapping = NULL)
```

Arguments

raw_data	Data frame. Raw WWTP data
nutrient	Character. "nitrogen" or "phosphorus"
custom_mapping	Named list. Custom column mappings (optional)

Value

Named list with column mappings

Examples

```
## Not run:
mapping <- map_wwtp_columns(raw_wwtp_data, "nitrogen")

# Custom mapping for different format
custom_map <- list(facility_name = "Plant_Name",
                  pollutant_load = "Annual_Load_kg")
mapping <- map_wwtp_columns(raw_data, "nitrogen", custom_map)

## End(Not run)
```

map_wwtp_influence *Create WWTP Influence Map*

Description

Create map showing WWTP contribution as proportion of total nutrient load

Usage

```
map_wwtp_influence(data, nutrient, title)
```

Arguments

data	sf object. Integrated data with WWTP proportions
nutrient	Character. "nitrogen" or "phosphorus"
title	Character. Map title

Value

ggplot object

map_wwtp_points *Create WWTP Point Map*

Description

Create map showing WWTP locations classified by load size

Usage

```
map_wwtp_points(wwtp_sf, nutrient, title)
```

Arguments

wwtp_sf sf object. Spatial WWTP data with classifications
 nutrient Character. "nitrogen" or "phosphorus"
 title Character. Map title

Value

ggplot object

P2O5_TO_P *Conversion Factor: P2O5 to P*

Description

Conversion Factor: P2O5 to P
 Conversion Factor: P2O5 to P

Usage

P2O5_TO_P
 P2O5_TO_P

Format

An object of class numeric of length 1.
 An object of class numeric of length 1.

plot_absolute_changes *Create Absolute Change Plot*

Description

Create plot showing absolute changes in classification counts

Usage

plot_absolute_changes(data, title)

Arguments

data Data frame. Summary data with absolute changes
 title Character. Plot title

Value

ggplot object

plot_before_after_comparison
Create Before/After Comparison Plot

Description

Create side-by-side comparison of agricultural vs WWTP+agricultural classifications

Usage

```
plot_before_after_comparison(data, nutrient, title)
```

Arguments

data	Data frame. Summary data from create_classification_summary
nutrient	Character. "nitrogen" or "phosphorus" for coloring
title	Character. Plot title

Value

ggplot object

plot_impact_ratios *Create Impact Ratio Plot*

Description

Create plot showing impact of WWTP addition as ratios

Usage

```
plot_impact_ratios(data, title)
```

Arguments

data	Data frame. Summary data with impact ratios
title	Character. Plot title

Value

ggplot object

quick_analysis	<i>Quick Analysis with Visualization</i>
----------------	--

Description

Run analysis and automatically generate key visualizations for specified nutrients. This is a convenience function that combines `run_built_in_analysis` with automatic visualization generation.

Usage

```
quick_analysis(
  scale = "huc8",
  year = 2016,
  nutrients = c("nitrogen", "phosphorus"),
  include_wwtp = TRUE,
  output_dir = tempdir(),
  create_maps = TRUE,
  create_networks = TRUE,
  create_comparisons = TRUE,
  create_wwtp_maps = TRUE,
  wwtp_load_units = "kg",
  map_resolution = "medium",
  generate_report = FALSE,
  verbose = TRUE,
  ...
)
```

Arguments

scale	Character. Spatial scale: "county", "huc8", or "huc2"
year	Numeric. Year to analyze
nutrients	Character vector. Nutrients to analyze: c("nitrogen", "phosphorus") or subset
include_wwtp	Logical. Whether to include WWTP analysis (default: TRUE)
output_dir	Character. Output directory (default: tempdir())
create_maps	Logical. Whether to create classification maps (default: TRUE)
create_networks	Logical. Whether to create network plots (default: TRUE)
create_comparisons	Logical. Whether to create comparison plots (default: TRUE)
create_wwtp_maps	Logical. Whether to create WWTP facility maps (default: TRUE)
wwtp_load_units	Character. Units for WWTP loads if using custom data (default: "kg")
map_resolution	Character. Map resolution: "low", "medium", "high" (default: "medium")

```

generate_report      Logical. Whether to generate HTML report (default: FALSE)
verbose              Logical. Whether to print progress messages (default: TRUE)
...                  Additional arguments passed to run_built_in_analysis

```

Value

List with results and file paths of created visualizations

Examples

```

# Quick analysis with all visualizations (2007-2016 WWTP available)
results <- quick_analysis(
  scale = "huc8",
  year = 2012, # Use valid year
  nutrients = c("nitrogen", "phosphorus"),
  include_wwtp = TRUE,
  generate_report = TRUE
)

# Agricultural only analysis for pre-WWTP year
results <- quick_analysis(
  scale = "county",
  year = 2005, # Before WWTP data
  nutrients = "nitrogen",
  include_wwtp = FALSE,
  create_networks = FALSE
)

# High-resolution analysis with expanded year range
results <- quick_analysis(
  scale = "huc8",
  year = 2008, # Use valid WWTP year
  nutrients = "phosphorus",
  include_wwtp = TRUE,
  map_resolution = "high"
)

```

quick_check

Quick Data Check

Description

Perform quick validation checks on analysis results

Perform quick validation checks on analysis results

Usage

```
quick_check(results, verbose = TRUE)
```

```
quick_check(results, verbose = TRUE)
```

Arguments

results	Analysis results object
verbose	Logical. Print detailed messages

Value

Logical. TRUE if all checks pass

Logical. TRUE if all checks pass

quick_state_analysis *Quick State Analysis with Visualization*

Description

Run state-level analysis with automatic visualizations

Usage

```
quick_state_analysis(
  state,
  scale = "huc8",
  year = 2016,
  nutrients = c("nitrogen", "phosphorus"),
  include_wwtp = TRUE,
  output_dir = file.path(tempdir(), paste0("state_", tolower(state), "_results")),
  create_maps = TRUE,
  create_networks = TRUE,
  create_comparisons = TRUE,
  verbose = TRUE,
  ...
)
```

Arguments

state	Character. Two-letter state abbreviation
scale	Character. Spatial scale
year	Numeric. Year to analyze
nutrients	Character vector. Nutrients to analyze
include_wwtp	Logical. Include WWTP analysis

output_dir	Character. Output directory
create_maps	Logical. Create maps
create_networks	Logical. Create network plots
create_comparisons	Logical. Create comparison plots
verbose	Logical. Show progress
...	Additional arguments

Value

List with results and visualizations

Examples

```
# Quick state analysis - use states with good data coverage
results <- quick_state_analysis(
  state = "TX", # Texas has good data coverage
  scale = "county",
  year = 2016,
  nutrients = "nitrogen",
  include_wwtp = TRUE
)
```

run_builtin_analysis *Complete Manureshed Analysis Workflow (Built-in Data)*

Description

Run complete manureshed analysis using built-in NuGIS data (start-2016) and optional WWTP data. For WWTP analysis beyond 2016, users must provide their own data. Supports analysis of nitrogen, phosphorus, or both nutrients simultaneously.

Usage

```
run_builtin_analysis(
  scale = "huc8",
  year = 2016,
  nutrients = c("nitrogen", "phosphorus"),
  output_dir = tempdir(),
  include_wwtp = TRUE,
  wwtp_year = NULL,
  custom_wwtp_nitrogen = NULL,
  custom_wwtp_phosphorus = NULL,
  wwtp_column_mapping = NULL,
  wwtp_skip_rows = 0,
```

```

    wwtp_header_row = 1,
    wwtp_load_units = "kg",
    add_texas = FALSE,
    save_outputs = TRUE,
    cropland_threshold = NULL,
    verbose = TRUE
  )

```

Arguments

scale	Character. Spatial scale: "county", "huc8", or "huc2"
year	Numeric. Year to analyze (available: start-2016 for NuGIS, 2016 for built-in WWTP)
nutrients	Character vector. Nutrients to analyze: c("nitrogen", "phosphorus") or subset
output_dir	Character. Output directory for results (default: "manureshed_results")
include_wwtp	Logical. Whether to include WWTP analysis (default: TRUE)
wwtp_year	Numeric. Year for WWTP data (default: same as year, only 2016 available built-in)
custom_wwtp_nitrogen	Character. Path to custom WWTP nitrogen file (for non-2016 years)
custom_wwtp_phosphorus	Character. Path to custom WWTP phosphorus file (for non-2016 years)
wwtp_column_mapping	Named list. Custom column mapping for WWTP data
wwtp_skip_rows	Numeric. Rows to skip in custom WWTP files (default: 0)
wwtp_header_row	Numeric. Header row in custom WWTP files (default: 1)
wwtp_load_units	Character. Units of WWTP loads: "kg", "lbs", "pounds", "tons" (default: "kg")
add_texas	Logical. Whether to add Texas HUC8 data (only for HUC8 scale, default: FALSE)
save_outputs	Logical. Whether to save results to files (default: TRUE)
cropland_threshold	Numeric. Custom cropland threshold for exclusion (optional)
verbose	Logical. Whether to print detailed progress messages (default: TRUE)

Value

List with all analysis results for specified nutrients

Examples

```

# Basic analysis using built-in data (2007-2016 WWTP available)
results_2016 <- run_builtin_analysis(
  scale = "huc8",

```

```

year = 2016,
nutrients = c("nitrogen", "phosphorus"),
include_wwtp = TRUE
)

# Analysis for earlier year (no WWTP available) - nitrogen only
results_2010 <- run_built_in_analysis(
  scale = "county",
  year = 2010,
  nutrients = "nitrogen",
  include_wwtp = FALSE
)

# Analysis for earlier year with WWTP now available
results_2010 <- run_built_in_analysis(
  scale = "county",
  year = 2010,
  nutrients = "nitrogen",
  include_wwtp = TRUE # Now supported for 2010!
)

# Analysis for year before WWTP availability
results_2005 <- run_built_in_analysis(
  scale = "huc8",
  year = 2005,
  nutrients = "phosphorus",
  include_wwtp = FALSE # No WWTP data before 2007
)

```

run_state_analysis *Run State-Level Analysis*

Description

Run manureshed analysis for a specific state

Usage

```

run_state_analysis(
  state,
  scale = "huc8",
  year = 2016,
  nutrients = c("nitrogen", "phosphorus"),
  include_wwtp = TRUE,
  output_dir = file.path(tempdir(), paste0("state_", tolower(state), "_results")),
  verbose = TRUE,
  ...
)

```

Arguments

state	Character. Two-letter state abbreviation (e.g., "OH", "TX")
scale	Character. Spatial scale: "county", "huc8", or "huc2"
year	Numeric. Year to analyze
nutrients	Character vector. Nutrients to analyze
include_wwtp	Logical. Whether to include WWTP analysis
output_dir	Character. Output directory
verbose	Logical. Show progress messages
...	Additional arguments passed to run_built_in_analysis

Value

List with analysis results for the state

Examples

```
# Use Texas which has more data
texas_results <- run_state_analysis(
  state = "TX",
  scale = "county", # Use county for faster processing
  year = 2016,
  nutrients = "nitrogen", # Single nutrient for speed
  include_wwtp = TRUE
)

# California county-level analysis
ca_results <- run_state_analysis(
  state = "CA",
  scale = "county",
  year = 2010,
  nutrients = "nitrogen"
)
```

save_analysis_summary *Save Analysis Summary*

Description

Save comprehensive summary of analysis parameters and results

Usage

```
save_analysis_summary(results, file_path = NULL, format = "rds")
```

Arguments

results	List. Complete analysis results from workflow functions
file_path	Character. Output file path. If NULL, auto-generated
format	Character. Output format: "rds", "json", or "txt"

Value

Character. Path to saved file

Examples

```
# Create analysis results first
results <- run_builtin_analysis(scale = "county", year = 2016)

# Save complete analysis summary
summary_path <- file.path(tempdir(), "analysis_summary_2016.json")
save_analysis_summary(results, summary_path, format = "json")
```

save_centroid_data *Save Centroid Data*

Description

Save centroid data to CSV file for transition probability analysis

Usage

```
save_centroid_data(
  data,
  file_path = NULL,
  scale = "huc8",
  nutrient = "nitrogen",
  analysis_type = "centroids",
  year = format(Sys.Date(), "%Y")
)
```

Arguments

data	Data frame. Data with centroid coordinates
file_path	Character. Output file path (should end in .csv). If NULL, auto-generated
scale	Character. Spatial scale for file naming
nutrient	Character. Nutrient type for file naming
analysis_type	Character. Analysis type for file naming
year	Numeric. Year for file naming

Value

Character. Path to saved file

Examples

```
# Create some example data first
results <- run_builtin_analysis(scale = "county", year = 2016, include_wwtp = TRUE)

# Save centroids for transition analysis
if ("integrated" %in% names(results) && "nitrogen" %in% names(results$integrated)) {
  centroids <- add_centroid_coordinates(results$integrated$nitrogen)
  save_centroid_data(centroids, scale = "county", nutrient = "nitrogen")
}
```

 save_plot

Save Plot

Description

Save ggplot object to file with publication-quality settings

Usage

```
save_plot(
  plot,
  file_path,
  width = 11,
  height = 6,
  dpi = 300,
  units = "in",
  device = NULL
)
```

Arguments

plot	ggplot object. Plot to save
file_path	Character. Output file path
width	Numeric. Plot width in inches (default: 11)
height	Numeric. Plot height in inches (default: 6)
dpi	Numeric. Resolution in dots per inch (default: 300)
units	Character. Units for width and height (default: "in")
device	Character. Output device (auto-detected from file extension)

Value

Character. Path to saved file

Examples

```
# Create a simple plot for demonstration
library(ggplot2)
p <- ggplot(mtcars, aes(x = mpg, y = hp)) + geom_point()

# Save with default settings (300 DPI, 11x6 inches)
save_plot(p, file.path(tempdir(), "test_plot.png"))

# Save with custom dimensions for presentation
save_plot(p, file.path(tempdir(), "presentation_plot.png"), width = 16, height = 9)

# Save as PDF for publication
save_plot(p, file.path(tempdir(), "publication_figure.pdf"), width = 8, height = 6)
```

save_spatial_data *Save Spatial Data*

Description

Save spatial data to RDS file with standardized naming

Usage

```
save_spatial_data(
  data,
  file_path = NULL,
  scale = "huc8",
  nutrient = "both",
  analysis_type = "combined",
  year = format(Sys.Date(), "%Y")
)
```

Arguments

data	sf object. Spatial data to save
file_path	Character. Output file path (should end in .rds). If NULL, auto-generated
scale	Character. Spatial scale for file naming
nutrient	Character. Nutrient type for file naming ("nitrogen", "phosphorus", or "both")
analysis_type	Character. Analysis type for file naming
year	Numeric. Year for file naming

Value

Character. Path to saved file

Examples

```
# Create some example results first
results <- run_builtin_analysis(scale = "county", year = 2016)

# Save with auto-generated filename
save_spatial_data(results$agricultural, scale = "county", year = 2016)

# Save with custom filename
save_spatial_data(results$agricultural,
                  file.path(tempdir(), "my_results.rds"))
```

save_transition_matrix

Save Transition Probability Matrix

Description

Save transition probability matrix to CSV with metadata

Usage

```
save_transition_matrix(
  transition_df,
  file_path,
  nutrient,
  analysis_type = "combined",
  metadata = NULL
)
```

Arguments

transition_df	Data frame. Transition probability matrix
file_path	Character. Output file path
nutrient	Character. Nutrient type
analysis_type	Character. Type of analysis
metadata	List. Additional metadata to include

Value

Character. Path to saved file

Examples

```
# Create example analysis results first
results <- run_builtin_analysis(scale = "county", year = 2016, include_wwtp = TRUE)

# Save transition probabilities (only if integrated results exist)
if ("integrated" %in% names(results) && "nitrogen" %in% names(results$integrated)) {
  centroids <- add_centroid_coordinates(results$integrated$nitrogen)
  transitions <- calculate_transition_probabilities(centroids, "combined_N_class")
  save_transition_matrix(transitions,
                        file.path(tempdir(), "transitions_nitrogen.csv"),
                        "nitrogen")
}
```

summarize_results *Print Summary of Analysis Results*

Description

Print formatted summary of manureshed analysis results to the console. The summary includes analysis configuration parameters (scale, year, nutrients, WWTP inclusion), spatial coverage statistics, agricultural nutrient classifications with counts and percentages, WWTP integration metrics (if applicable), integrated classifications (if available), output file information, and processing time.

Print formatted summary of manureshed analysis results to the console. The summary includes analysis configuration parameters (scale, year, nutrients, WWTP inclusion), spatial coverage statistics, agricultural nutrient classifications with counts and percentages, WWTP integration metrics (if applicable), integrated classifications (if available), output file information, and processing time.

Usage

```
summarize_results(results, detailed = FALSE)
```

```
summarize_results(results, detailed = FALSE)
```

Arguments

results	List. Analysis results from run_builtin_analysis or run_state_analysis . Must contain at minimum: <ul style="list-style-type: none"> parameters: List with scale, year, nutrients, include_wwtp agricultural: sf data frame with classification columns Optional components: <ul style="list-style-type: none"> wwtp: WWTP analysis results integrated: Integrated classification results created_files or saved_files: Output file paths
detailed	Logical. If TRUE, includes additional breakdown of integrated classifications showing combined agricultural-WWTP nutrient classes. If FALSE (default), shows only agricultural classifications and basic WWTP statistics.

Details

The summary output is organized into sections:

Analysis Configuration Scale, year, nutrients analyzed, WWTP inclusion, state (if applicable)

Spatial Coverage Total number of spatial units analyzed

Agricultural Classifications Nitrogen and phosphorus classification counts and percentages

WWTP Integration Number of facilities and total loads by nutrient (if applicable)

Integrated Classifications Combined agricultural-WWTP classes (if detailed = TRUE)

Output Files Number and types of created files (if saved)

Processing Time Analysis duration in minutes (if available)

Classification names are cleaned for display (underscores replaced with spaces, line breaks removed). Percentages are rounded to one decimal place. All console output uses `message` and can be suppressed with `suppressMessages`.

The summary output is organized into sections:

Analysis Configuration Scale, year, nutrients analyzed, WWTP inclusion, state (if applicable)

Spatial Coverage Total number of spatial units analyzed

Agricultural Classifications Nitrogen and phosphorus classification counts and percentages

WWTP Integration Number of facilities and total loads by nutrient (if applicable)

Integrated Classifications Combined agricultural-WWTP classes (if detailed = TRUE)

Output Files Number and types of created files (if saved)

Processing Time Analysis duration in minutes (if available)

Classification names are cleaned for display (underscores replaced with spaces, line breaks removed). Percentages are rounded to one decimal place. All console output uses `message` and can be suppressed with `suppressMessages`.

Value

Invisibly returns the input `results` list unchanged. The function is called primarily for its side effect of printing a formatted summary to the console. The invisible return allows for piping operations while displaying the summary.

Invisibly returns the input `results` list unchanged. The function is called primarily for its side effect of printing a formatted summary to the console. The invisible return allows for piping operations while displaying the summary.

See Also

`run_builtin_analysis` for generating analysis results, `quick_check` for quick validation, `compare_analyses` for comparing two result sets

`run_builtin_analysis` for generating analysis results, `quick_check` for quick validation, `compare_analyses` for comparing two result sets

Examples

```

# Basic summary
results <- run_builtin_analysis(scale = "county", year = 2016)
summarize_results(results)

# Detailed summary with integrated classifications
results <- run_builtin_analysis(
  scale = "huc8",
  year = 2012,
  include_wwtp = TRUE
)
summarize_results(results, detailed = TRUE)

## Not run:
# This requires magrittr - never auto-run
library(magrittr)
results <- run_builtin_analysis(scale = "huc2", year = 2015) %>%
  summarize_results() %>%
  export_for_gis(output_dir = tempdir())

## End(Not run)

# Basic summary
results <- run_builtin_analysis(scale = "county", year = 2016)
summarize_results(results)

# Detailed summary with integrated classifications
results <- run_builtin_analysis(
  scale = "huc8",
  year = 2012,
  include_wwtp = TRUE
)
summarize_results(results, detailed = TRUE)

## Not run:
# This requires magrittr - never auto-run
library(magrittr)
results <- run_builtin_analysis(scale = "huc2", year = 2015) %>%
  summarize_results() %>%
  export_for_gis(output_dir = tempdir())

## End(Not run)

```

test_osf_connection *Test OSF Connection*

Description

Test downloading a small dataset to verify OSF connectivity

Usage

```
test_osf_connection(verbose = TRUE)
```

Arguments

verbose Logical. Show detailed test results

Value

Logical. TRUE if test successful

validate_columns	<i>Validate Required Columns</i>
------------------	----------------------------------

Description

Validate Required Columns

Validate Required Columns

Usage

```
validate_columns(data, required_cols, data_type = "data")
```

```
validate_columns(data, required_cols, data_type = "data")
```

Arguments

data Data frame to validate

required_cols Character vector of required column names

data_type Character description of data type for error messages

Value

Logical. TRUE if all columns present, stops with error otherwise

Logical. TRUE if all columns present, stops with error otherwise

view_cheatsheet	<i>View Package Cheat Sheet</i>
-----------------	---------------------------------

Description

Opens the manureshed package cheat sheet in your default PDF viewer. The cheat sheet provides a quick reference for common functions and workflows.

Opens the manureshed package cheat sheet in your default PDF viewer. The cheat sheet provides a quick reference for common functions and workflows.

Usage

```
view_cheatsheet()
```

```
view_cheatsheet()
```

Value

Opens PDF in browser, returns path invisibly

Opens PDF in browser, returns path invisibly

Examples

```
## Not run:
# View the cheat sheet
view_cheatsheet()
```

```
## End(Not run)
## Not run:
# View the cheat sheet
view_cheatsheet()
```

```
## End(Not run)
```

wwtp_aggregate_by_boundaries

Aggregate WWTP Data by Spatial Boundaries

Description

Aggregate WWTP loads by spatial units (counties, HUC8s, etc.)

Usage

```
wwtp_aggregate_by_boundaries(wwtp_sf, boundaries, nutrient, boundary_id_col)
```

Arguments

wwtp_sf sf object. Spatial WWTP data
boundaries sf object. Spatial boundaries for aggregation
nutrient Character. "nitrogen" or "phosphorus"
boundary_id_col Character. Name of boundary ID column

Value

Data frame with aggregated loads by spatial unit

wwtp_classify_sources *Classify WWTP Sources by Load Size*

Description

Classify WWTP facilities into size categories based on annual nutrient loads

Usage

```
wwtp_classify_sources(wwtp_data, nutrient)
```

Arguments

wwtp_data Data frame. WWTP data with load information
nutrient Character. "nitrogen" or "phosphorus"

Value

Data frame with source_class column added

Examples

```
# Load WWTP data first  
wwtp_data <- load_builtin_wwtp("nitrogen", 2016)  
  
# Classify nitrogen sources  
classified_data <- wwtp_classify_sources(wwtp_data, "nitrogen")  
table(classified_data$source_class)
```

wwtp_clean_data	<i>Clean WWTP Data</i>
-----------------	------------------------

Description

Clean and validate WWTP data for analysis

Clean and validate WWTP data for analysis

Usage

```
wwtp_clean_data(wwtp_data, nutrient)
```

```
wwtp_clean_data(wwtp_data, nutrient)
```

Arguments

wwtp_data Data frame. Raw or standardized WWTP data

nutrient Character. "nitrogen" or "phosphorus"

Value

Data frame with cleaned WWTP data

Data frame with cleaned WWTP data

Examples

```
## Not run:  
# Clean user-loaded data will not run. They need to be supplied by users  
clean_data <- wwtp_clean_data(raw_wwtp_data, "nitrogen")  
  
# Clean OSF data (usually already clean, available 2007-2016)  
osf_data <- load_built_in_wwtp("phosphorus", 2012)  
clean_data <- wwtp_clean_data(osf_data, "phosphorus")  
  
## End(Not run)  
  
# Load and clean WWTP data  
raw_wwtp_data <- load_built_in_wwtp("nitrogen", 2016)  
clean_data <- wwtp_clean_data(raw_wwtp_data, "nitrogen")
```

`wwtp_filter_positive_loads`*Filter WWTP Data for Positive Loads*

Description

Filter WWTP data to include only facilities with positive nutrient loads

Usage

```
wwtp_filter_positive_loads(wwtp_data, nutrient)
```

Arguments

<code>wwtp_data</code>	Data frame. WWTP data
<code>nutrient</code>	Character. "nitrogen" or "phosphorus"

Value

Data frame with facilities having positive loads

`wwtp_process_complete` *Complete WWTP Processing Pipeline*

Description

Run complete WWTP processing pipeline for both nutrients

Usage

```
wwtp_process_complete(  
  nitrogen_path = NULL,  
  phosphorus_path = NULL,  
  column_mapping = NULL,  
  skip_rows = 0,  
  header_row = 1,  
  load_units = "kg",  
  verbose = TRUE  
)
```

Arguments

nitrogen_path Character. Path to nitrogen WWTP data (if NULL, loads from OSF)
 phosphorus_path Character. Path to phosphorus WWTP data (if NULL, loads from OSF)
 column_mapping Named list. Custom column mapping for user data
 skip_rows Numeric. Rows to skip in user files
 header_row Numeric. Header row in user files
 load_units Character. Units of loads in user files
 verbose Logical. Show processing messages

Value

List with processed nitrogen and phosphorus WWTP data

Examples

```
# Process built-in OSF data (2016 default)
wwtp_results_builtin <- wwtp_process_complete(
  nitrogen_path = NULL,      # Use built-in data
  phosphorus_path = NULL,   # Use built-in data
  verbose = TRUE
)

# Process custom user data
# wwtp_results_custom <- wwtp_process_complete(
#   nitrogen_path = "nitrogen_2020.csv",
#   phosphorus_path = "phosphorus_2020.csv",
#   load_units = "lbs"
# )

# Mixed: OSF for one nutrient, custom for another
# wwtp_results_mixed <- wwtp_process_complete(
#   nitrogen_path = NULL,      # Use OSF built-in
#   phosphorus_path = "custom_P.csv" # Use custom
# )
```

 wwtp_to_spatial

Convert WWTP Data to Spatial Format

Description

Convert WWTP data frame to sf spatial object

Usage

```
wwtp_to_spatial(wwtp_data, crs = 4326)
```

Arguments

`wwtp_data` Data frame. WWTP data with Lat/Long coordinates
`crs` Numeric. Coordinate reference system (default: 4326 for WGS84)

Value

sf object with WWTP facilities as point geometries

Examples

```
# Load and convert to spatial format
wwtp_data <- load_built_in_wwtp("nitrogen", 2016)
wwtp_clean_data <- wwtp_clean_data(wwtp_data, "nitrogen")
wwtp_sf <- wwtp_to_spatial(wwtp_clean_data)

# Convert and transform to analysis CRS (without using pipe operator)
wwtp_sf <- wwtp_to_spatial(wwtp_clean_data)
wwtp_sf_transformed <- sf::st_transform(wwtp_sf, 5070) # Albers Equal Area Conic
```

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