

Package ‘geospatialsuite’

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Title Comprehensive Geospatiotemporal Analysis and Multimodal Integration Toolkit

Version 0.2.0

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Description A comprehensive toolkit for geospatiotemporal analysis featuring 60+ vegetation indices, advanced raster visualization, universal spatial mapping, water quality analysis, CDL crop analysis, spatial interpolation, temporal analysis, and terrain analysis. Designed for agricultural research, environmental monitoring, remote sensing applications, and publication-quality mapping with support for any geographic region and robust error handling. Methods include vegetation indices calculations (Rouse et al. 1974), NDVI and enhanced vegetation indices (Huete et al. 1997) <doi:10.1016/S0034-4257(97)00104-1>, (Akanbi et al. 2024) <doi:10.1007/s41651-023-00164-y>, spatial interpolation techniques (Cressie 1993, ISBN:9780471002556), water quality indices (McFeeters 1996) <doi:10.1080/01431169608948714>, and crop data layer analysis (USDA NASS 2024) <https://www.nass.usda.gov/Research_and_Science/Cropland/>. Funding: 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.

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URL <https://exelegch.github.io/geospatialsuite-docs/>,
<https://github.com/cwru-sdle/geospatialsuite>,
<https://joss.theoj.org/papers/10.21105/joss.09536>

BugReports <https://github.com/cwru-sdle/geospatialsuite/issues>

Depends R (>= 3.5.0)

Imports dplyr, ggplot2, graphics, grDevices, htmlwidgets, leaflet, magrittr, mice, parallel, RColorBrewer, rnaturalearth, sf, stats, stringr, terra, tigris, tools, utils, viridis

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 geospatialsuite-package

geospatialsuite: Comprehensive Geospatiotemporal Analysis and Multimodal Integration Toolkit

Description

geospatialsuite provides universal functions for geospatial analysis and reliable visualization that work with any region for multimodal data. Features include 60+ vegetation indices, efficient terra-based visualization, universal spatial mapping, dynamic crop analysis, water quality assessment, and publication-quality mapping with support for any geographic region and robust error handling.

Details**Key Features::**

Universal Spatial Analysis::

- Universal region support (states, countries, CONUS, custom boundaries)

- Universal spatial join (works with ANY raster-vector combination)
- Multi-dataset integration and temporal analysis
- Spatial interpolation and terrain analysis

Advanced Vegetation Analysis::

- **60+ vegetation indices** including NDVI, EVI, SAVI, ARVI, PRI, SIPI, etc.
- **Specialized crop analysis** with stress detection and yield assessment
- **Auto band detection** from multi-band satellite imagery
- **Quality filtering** and temporal smoothing for time series

Reliable Visualization::

- **Universal mapping** with auto-detection (`quick_map()` function)
- **Terra-based plotting** using reliable `terra::plot()` and `terra::plotRGB()`
- **Interactive maps** with leaflet integration (optional)
- **RGB composites** with stretching algorithms
- **Comparison maps** for before/after analysis

Agricultural Applications::

- **Dynamic CDL crop analysis** (all crop codes and categories)
- **NDVI time series** with classification
- **Crop-specific analysis** (corn, soybeans, wheat, etc.)
- **Water quality assessment** with threshold analysis

Performance & Reliability Features::

- **Standard terra plotting** - no complex dependencies required
- **Robust error handling** throughout all functions
- **Simplified visualization** for maps
- **Smart fallback systems** when optional packages unavailable

Quick Start Examples::

```
# One-line mapping (auto-detects everything!)
quick_map("mydata.shp")

# Auto-geocode data without coordinates
census_data <- data.frame(
  state = c("Ohio", "Pennsylvania", "Michigan"),
  median_income = c(58642, 61744, 59584)
)
spatial_data <- auto_geocode_data(census_data)
quick_map(spatial_data, variable = "median_income")

# Works with HUC codes too (any format: HUC_8, HUC-8, huc8)
watershed_data <- data.frame(
  HUC_8 = c("04100009", "04100012"),
  water_quality = c(72, 65)
)
huc_spatial <- auto_geocode_data(watershed_data)
quick_map(huc_spatial)
```

```
# Calculate multiple vegetation indices
indices <- calculate_multiple_indices(
  red = red_band, nir = nir_band,
  indices = c("NDVI", "EVI", "SAVI", "PRI")
)

# Comprehensive crop analysis
crop_analysis <- analyze_crop_vegetation(
  spectral_data = sentinel_data,
  crop_type = "corn",
  analysis_type = "comprehensive"
)

# Enhanced NDVI calculation
ndvi_enhanced <- calculate_ndvi_enhanced(
  red_data = red_raster,
  nir_data = nir_raster,
  quality_filter = TRUE
)

# Fast, reliable RGB plotting
plot_rgb_raster(satellite_data, r = 4, g = 3, b = 2,
  stretch = "hist", title = "False Color")
```

Recommended Optional Packages::

For enhanced features, consider installing these optional packages:

```
# For interactive mapping
install.packages("leaflet")

# For enhanced colors
install.packages(c("viridis", "RColorBrewer"))

# For advanced remote sensing (optional)
install.packages("RStoolbox")

# For multi-panel plots (optional)
install.packages("patchwork")
```

Core Dependencies Only::

The package works reliably with just the core dependencies:

- terra (raster operations and plotting)
- sf (vector operations)
- ggplot2 (static mapping)
- dplyr (data manipulation)

No complex visualization dependencies required!

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See Also

Useful links:

- <https://exelegch.github.io/geospatialsuite-docs/>
- <https://github.com/cwru-sdle/geospatialsuite>
- <https://joss.theoj.org/papers/10.21105/joss.09536>
- Report bugs at <https://github.com/cwru-sdle/geospatialsuite/issues>

analyze_cdl_crops_dynamic

Analyze CDL crops dynamically

Description

Perform dynamic analysis of CDL crop data including area calculations, classification, and masking for any crop combination. Now accepts file paths directly.

Usage

```
analyze_cdl_crops_dynamic(  
  cdl_data,  
  crop_selection,  
  region_boundary = NULL,  
  analysis_type = "classify",  
  output_folder = NULL,  
  year = NULL  
)
```

Arguments

cdl_data	CDL raster data (file path, directory, or SpatRaster object)
crop_selection	Crop selection (names, codes, or categories)
region_boundary	Region boundary
analysis_type	Type of analysis: "mask", "classify", "area"
output_folder	Output directory
year	Year for analysis (optional)

Details**Usage Tips::***Area Analysis Results::*

```
# Access specific results
result$total_area_ha      # Total area in hectares
result$crop_areas_ha     # Area by crop code
result$crop_areas_ha[["5"]] # Soybean area (code 5)
```

Visualization::

```
# For mask/classify results (SpatRaster):
terra::plot(result)      # Plot the raster

# For area results (list):
print(result$total_area_ha) # Print total area
barplot(unlist(result$crop_areas_ha)) # Simple bar plot
```

Value

Analysis results depend on analysis_type:

- **"mask"**: SpatRaster with binary mask (1 = crop, 0 = other)
- **"classify"**: SpatRaster with binary classification
- **"area"**: List with detailed area analysis:
 - crop_areas_ha: Named list of areas by crop code (hectares)
 - total_area_ha: Total crop area (hectares)
 - pixel_size_ha: Individual pixel size (hectares)
 - crop_codes: CDL codes analyzed
 - crop_name: Crop name(s)
 - region: Region analyzed
 - raster_info: Technical raster information
 - total_pixels: Total number of pixels
 - valid_pixels: Number of valid (non-NA) pixels

Examples

```
## Not run:
# These examples require actual CDL data files
# Analyze soybean area in Iowa - accepts file path directly
soybean_area <- analyze_cdl_crops_dynamic(
  "/path/to/cdl_2023.tif", "soybeans", "Iowa", "area"
)

# Access area results
soybean_area$total_area_ha      # Total hectares
soybean_area$crop_areas_ha[["5"]] # Soybean area (code 5)
soybean_area$total_area_ha * 2.47105 # Convert to acres
```

```

# Create grain classification
grain_classes <- analyze_cdl_crops_dynamic(
  cdl_data, "grains", "CONUS", "classify"
)
terra::plot(grain_classes) # Plot the classification

# Works with directories too
results <- analyze_cdl_crops_dynamic(
  "/path/to/cdl/files/", "corn", "Ohio", "area"
)

## End(Not run)

# Example with mock CDL data (this can run)
# Create sample CDL raster
mock_cdl <- terra::rast(nrows = 10, ncols = 10, xmin = 0, xmax = 10,
  ymin = 0, ymax = 10, crs = "EPSG:4326")
terra::values(mock_cdl) <- sample(c(1, 5, 24), 100, replace = TRUE) # corn, soy, wheat

# Analyze mock data
result <- analyze_cdl_crops_dynamic(mock_cdl, "corn", analysis_type = "mask")
print(class(result)) # Should be SpatRaster

```

analyze_crop_vegetation

Specialized crop vegetation analysis

Description

Perform comprehensive vegetation analysis specifically designed for crop monitoring including growth stage detection, stress identification, and yield prediction support. Handles test scenarios properly with better input validation.

Usage

```

analyze_crop_vegetation(
  spectral_data,
  crop_type = "general",
  growth_stage = "unknown",
  analysis_type = "comprehensive",
  cdl_mask = NULL,
  reference_data = NULL,
  output_folder = NULL,
  verbose = FALSE
)

```

Arguments

spectral_data	Multi-band spectral data (file, directory, or SpatRaster)
crop_type	Crop type for specialized analysis ("corn", "soybeans", "wheat", "general")
growth_stage	Growth stage if known ("early", "mid", "late", "harvest")
analysis_type	Type of analysis: "comprehensive", "stress", "growth", "yield"
cdl_mask	Optional CDL mask for crop-specific analysis
reference_data	Optional reference data for validation
output_folder	Optional output folder for results
verbose	Print detailed progress

Details**Crop-Specific Index Selection::**

- **Corn:** NDVI, EVI, GNDVI, DVI, RVI, PRI
- **Soybeans:** NDVI, EVI, SAVI, GNDVI, PRI
- **Wheat:** NDVI, EVI, SAVI, DVI
- **General:** NDVI, EVI, SAVI, GNDVI, DVI, RVI

Analysis Types::

- **comprehensive:** All analyses (stress, growth, yield)
- **stress:** Focus on stress detection indices
- **growth:** Growth stage analysis
- **yield:** Yield prediction support

Output Structure::

The function returns a list with three main components:

1. vegetation_indices (SpatRaster):

Multi-layer raster with calculated indices (NDVI, EVI, etc.)

*2. analysis_results (List):***stress_analysis** (if requested):

- Percentage of pixels in each stress category
- Categories: healthy (NDVI 0.6-1.0), moderate stress (0.4-0.6), severe stress (0.0-0.4)
- Includes mean, median, std_dev, and thresholds used

growth_analysis (if requested):

- Predicted growth stage based on NDVI patterns
- Stage confidence (0-1 scale)
- Detailed statistics for each index
- Growth stages: emergence, vegetative, reproductive, maturity (crop-specific)

yield_analysis (if requested):

- **Composite Yield Index:** Normalized 0-1 score combining multiple indices
 - 0.0 = Very low yield potential
 - 0.5 = Medium yield potential

- 1.0 = Maximum yield potential
- **Yield Potential Class:** Categorical (Low, Medium, High, Very High)
- **Index Contributions:** How each index contributed to composite score
- Calculation: Each index (NDVI, EVI, GNDVI, DVI, RVI) is normalized to 0-1, then averaged to create composite score

summary_statistics:

- Basic stats (mean, std, min, max, percentiles) for all indices
- Coverage percentage and pixel counts

3. metadata (List):

Processing information: crop_type, indices_used, processing_date, spatial properties

Example Interpretation::

```
result <- analyze_crop_vegetation(data, crop_type = "corn")

# Stress Assessment
stress <- result$analysis_results$stress_analysis$NDVI
cat(sprintf("Healthy: %.1f%%, Stressed: %.1f%%\n",
           stress$healthy_percentage,
           stress$severe_stress_percentage))

# Growth Stage
stage <- result$analysis_results$growth_analysis$predicted_growth_stage
cat(sprintf("Growth stage: %s\n", stage))

# Yield Potential
yield <- result$analysis_results$yield_analysis
cat(sprintf("Yield potential: %s (score: %.2f)\n",
           yield$yield_potential_class,
           yield$composite_yield_index))
```

Important Notes::

- **Composite Yield Index** is a vegetation-based proxy, not a direct yield prediction
- Thresholds are based on literature and may need regional calibration
- Results should be validated with ground truth data
- For detailed output documentation, see package vignette

Analysis Types::

- **comprehensive:** All analyses (stress, growth, yield)
- **stress:** Focus on stress detection indices
- **growth:** Growth stage analysis
- **yield:** Yield prediction support

Value

List with comprehensive vegetation analysis results:

- vegetation_indices: SpatRaster with calculated indices
- analysis_results: Detailed analysis results by type
- metadata: Analysis metadata and parameters

Examples

```
## Not run:
# These examples require actual spectral data
# Comprehensive corn analysis
corn_analysis <- analyze_crop_vegetation(
  spectral_data = sentinel_data,
  crop_type = "corn",
  analysis_type = "comprehensive",
  cdl_mask = corn_mask
)

# Access results
corn_analysis$vegetation_indices      # SpatRaster with indices
corn_analysis$analysis_results$stress_analysis # Stress detection results
corn_analysis$metadata$indices_used    # Which indices were calculated

# Stress detection in soybeans
stress_analysis <- analyze_crop_vegetation(
  spectral_data = landsat_stack,
  crop_type = "soybeans",
  analysis_type = "stress",
  growth_stage = "mid"
)

## End(Not run)

# Example with mock spectral data
# Create mock multi-band raster (simulating satellite data)
red_band <- terra::rast(nrows = 5, ncols = 5, crs = "EPSG:4326")
nir_band <- terra::rast(nrows = 5, ncols = 5, crs = "EPSG:4326")
terra::values(red_band) <- runif(25, 0.1, 0.3) # Typical red values
terra::values(nir_band) <- runif(25, 0.4, 0.8) # Typical NIR values
spectral_stack <- c(red_band, nir_band)
names(spectral_stack) <- c("red", "nir")

# Analyze with mock data
result <- analyze_crop_vegetation(spectral_stack, crop_type = "general")
print(names(result)) # Should show analysis components
```

Description

Analyze temporal changes in raster data including trend analysis, change detection, and seasonal patterns. Works with any time series data.

Usage

```
analyze_temporal_changes(  
  data_list,  
  dates = NULL,  
  region_boundary = NULL,  
  analysis_type = "trend",  
  output_folder = NULL  
)
```

Arguments

data_list	List of raster data for different time periods
dates	Vector of dates corresponding to rasters
region_boundary	Region boundary for analysis
analysis_type	Type of temporal analysis: "trend", "change_detection", "seasonal", "statistics"
output_folder	Output directory for results

Value

Temporal analysis results

Examples

```
## Not run:  
# These examples require external data files not included with the package  
# Analyze NDVI trends over time  
ndvi_trend <- analyze_temporal_changes(  
  data_list = c("ndvi_2020.tif", "ndvi_2021.tif", "ndvi_2022.tif"),  
  dates = c("2020", "2021", "2022"),  
  region_boundary = "Iowa",  
  analysis_type = "trend"  
)  
  
# Detect land cover changes  
land_changes <- analyze_temporal_changes(  
  data_list = land_cover_files,  
  dates = land_cover_dates,  
  analysis_type = "change_detection"  
)  
  
## End(Not run)
```

```
analyze_variable_correlations
```

Analyze correlations between multiple variables

Description

Analyze correlations between multiple raster variables and create correlation matrices and plots.

Usage

```
analyze_variable_correlations(  
  variable_list,  
  output_folder = NULL,  
  region_boundary = NULL,  
  method = "pearson",  
  create_plots = TRUE  
)
```

Arguments

<code>variable_list</code>	Named list of raster variables
<code>output_folder</code>	Output directory for results
<code>region_boundary</code>	Optional region boundary
<code>method</code>	Correlation method
<code>create_plots</code>	Create correlation plots

Value

List with correlation results

Examples

```
## Not run:  
# These examples require directory structures with multiple data files  
# Analyze correlations between multiple variables  
variables <- list(  
  ndvi = "ndvi.tif",  
  nitrogen = "soil_nitrogen.tif",  
  elevation = "dem.tif",  
  precipitation = "precip.tif"  
)  
  
correlation_results <- analyze_variable_correlations(  
  variables,  
  output_folder = "correlations/",  
  region_boundary = "Ohio"
```

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

analyze_water_bodies *Analyze water body characteristics using multiple indices*

Description

Comprehensive water body analysis using multiple water indices to classify and characterize water features.

Usage

```
analyze_water_bodies(  
  green,  
  nir,  
  swir1 = NULL,  
  region_boundary = NULL,  
  water_threshold_ndwi = 0.3,  
  water_threshold_mndwi = 0.5,  
  output_folder = NULL,  
  verbose = FALSE  
)
```

Arguments

green	Green band SpatRaster or file path
nir	NIR band SpatRaster or file path
swir1	SWIR1 band SpatRaster or file path
region_boundary	Optional region boundary for analysis
water_threshold_ndwi	NDWI threshold for water detection (default: 0.3)
water_threshold_mndwi	MNDWI threshold for water detection (default: 0.5)
output_folder	Optional output directory
verbose	Print progress messages

Value

List with water analysis results

Examples

```
## Not run:
# These examples require external data files not included with the package
# Comprehensive water analysis
water_analysis <- analyze_water_bodies(
  green = "green.tif",
  nir = "nir.tif",
  swir1 = "swir1.tif",
  region_boundary = "study_area.shp",
  verbose = TRUE
)

# Access results
water_analysis$water_indices      # All calculated indices
water_analysis$water_mask        # Binary water mask
water_analysis$statistics        # Water body statistics

## End(Not run)
```

analyze_water_quality_comprehensive

Analyze water quality comprehensively with flexible data handling

Description

Complete water quality analysis with flexible data input handling, robust error checking, and comprehensive spatial integration. Supports any water quality dataset format with automatic column detection and standardized processing.

Usage

```
analyze_water_quality_comprehensive(
  water_data,
  variable = NULL,
  region_boundary = NULL,
  river_network = NULL,
  output_folder = tempdir(),
  thresholds = NULL,
  coord_cols = NULL,
  date_column = NULL,
  station_id_col = NULL,
  quality_filters = list(),
  verbose = FALSE
)
```

Arguments

water_data	Water quality data in various formats: <ul style="list-style-type: none"> • File path (CSV, shapefile, GeoJSON) • data.frame with coordinates • sf object • List of datasets for multi-dataset analysis
variable	Variable to analyze (auto-detected if NULL)
region_boundary	Region boundary (optional)
river_network	Optional river network data for context
output_folder	Output directory (default: tempdir())
thresholds	Named list of threshold values for classification (optional)
coord_cols	Coordinate column names (auto-detected if NULL)
date_column	Date/time column name (auto-detected if NULL)
station_id_col	Station ID column name (auto-detected if NULL)
quality_filters	Quality control filters to apply
verbose	Print detailed progress messages

Value

List with comprehensive water quality analysis results:

- water_data: Processed spatial data
- statistics: Summary statistics by variable and category
- spatial_analysis: Spatial pattern analysis
- temporal_analysis: Temporal trends (if date data available)
- threshold_analysis: Threshold exceedance analysis
- output_files: Paths to generated output files
- metadata: Analysis metadata and parameters

Examples

```
## Not run:
# These examples require external data files not included with the package
# Flexible data input - auto-detects columns
results <- analyze_water_quality_comprehensive("water_stations.csv")

# Specify parameters for custom data
results <- analyze_water_quality_comprehensive(
  water_data = my_data,
  variable = "nitrate_concentration",
  region_boundary = "Ohio",
  coord_cols = c("longitude", "latitude"),
```

```

    thresholds = list(
      Normal = c(0, 2),
      Elevated = c(2, 5),
      High = c(5, 10),
      Critical = c(10, Inf)
    )
  )

# Multi-dataset analysis
results <- analyze_water_quality_comprehensive(
  water_data = list(
    surface = "surface_water.csv",
    groundwater = "groundwater.csv"
  ),
  variable = "total_nitrogen"
)

## End(Not run)

```

auto_geocode_data *Auto-geocode data with geographic identifiers*

Description

Automatically detects and geocodes data containing US geographic identifiers (states, counties, FIPS codes, HUC watershed codes, ZIP codes, or city names) without requiring latitude/longitude coordinates.

Usage

```

auto_geocode_data(
  data,
  detect_columns = TRUE,
  entity_column = NULL,
  entity_type = NULL,
  verbose = TRUE
)

```

Arguments

data	Data frame, file path (CSV, shapefile, etc.), or sf object
detect_columns	Auto-detect geographic entity columns (default: TRUE)
entity_column	Explicitly specify the column containing geographic entities (optional)
entity_type	Explicitly specify entity type: "state", "county", "fips", "huc", "zipcode", "city" (optional)
verbose	Print detailed progress messages

Details

Supported geographic entities:

- **States:** Full names or 2-letter abbreviations (e.g., "Ohio", "OH")
- **Counties:** County names, optionally with state
- **FIPS codes:** 5-digit Federal Information Processing Standards codes
- **HUC codes:** Hydrologic Unit Codes (HUC8, HUC10, HUC12)
- **ZIP codes:** 5-digit US postal codes
- **Cities:** City names, works best with state column

Column name variations supported:

- HUC columns: HUC_8, HUC-8, huc8, Huc 8, etc.
- State columns: State, STATE, state_name, StateName, ST, etc.
- All entity types handle spaces, hyphens, underscores, and mixed case

Required packages (installed automatically when needed):

- `tigris`: For US Census boundaries (states, counties, FIPS)
- `nhdplusTools`: For HUC watershed boundaries
- `zipcodeR`: For ZIP code centroids
- `tidygeocoder`: For city name geocoding

Value

sf object with geocoded point or polygon geometries

Examples

```
## Not run:
# Auto-detect and geocode - simplest usage
geodata <- auto_geocode_data("mydata.csv")

# With state names
state_data <- data.frame(
  state = c("California", "Texas", "New York"),
  population = c(39538223, 29145505, 20201249)
)
state_sf <- auto_geocode_data(state_data)

# With FIPS codes
fips_data <- data.frame(
  fips = c("39049", "39035", "39113"), # Ohio counties
  unemployment_rate = c(4.2, 3.8, 5.1)
)
county_sf <- auto_geocode_data(fips_data)

# With HUC codes (handles various formats)
watershed_data <- data.frame(
```

```
HUC_8 = c("04100009", "04100012", "04110002"),
water_quality_index = c(72, 65, 80)
)
huc_sf <- auto_geocode_data(watershed_data)

# Explicit specification
zip_sf <- auto_geocode_data(
  my_data,
  entity_column = "postal_code",
  entity_type = "zipcode"
)

# Then use with your other geospatialsuite functions
quick_map(state_sf, variable = "population")

## End(Not run)
```

calculate_advanced_terrain_metrics

Calculate advanced terrain metrics

Description

Calculate advanced terrain metrics from DEM including curvature, wetness index, and stream power index.

Usage

```
calculate_advanced_terrain_metrics(
  elevation_raster,
  metrics = c("wetness_index", "curvature", "convergence"),
  region_boundary = NULL
)
```

Arguments

elevation_raster	Digital elevation model
metrics	Vector of metrics to calculate
region_boundary	Optional region boundary

Value

List of terrain metric rasters

Examples

```
## Not run:
# These examples require external data files not included with the package
# Calculate advanced terrain metrics
terrain_metrics <- calculate_advanced_terrain_metrics(
  elevation_raster = "dem.tif",
  metrics = c("wetness_index", "curvature", "convergence"),
  region_boundary = "watershed.shp"
)

## End(Not run)
```

```
calculate_multiple_indices
```

Calculate multiple vegetation indices at once

Description

Calculate multiple vegetation indices from the same spectral data in a single operation. Efficient for comparative analysis and comprehensive vegetation assessment. Supports directory input and automatic CRS handling.

Usage

```
calculate_multiple_indices(
  spectral_data = NULL,
  indices = c("NDVI", "EVI", "SAVI"),
  output_stack = TRUE,
  region_boundary = NULL,
  parallel = FALSE,
  verbose = FALSE,
  ...
)
```

Arguments

spectral_data	Multi-band raster, directory path, or individual bands
indices	Vector of index names to calculate
output_stack	Return as single multi-layer raster (TRUE) or list (FALSE)
region_boundary	Optional region boundary for clipping
parallel	Use parallel processing for multiple indices
verbose	Print progress messages
...	Additional arguments passed to calculate_vegetation_index

Value

SpatRaster stack or list of indices

Examples

```
## Not run:
# These examples require satellite imagery files (Landsat/Sentinel data etc.)
# Calculate multiple basic indices from directory
multi_indices <- calculate_multiple_indices(
  spectral_data = "/path/to/sentinel/bands/",
  indices = c("NDVI", "EVI", "SAVI", "MSAVI"),
  auto_detect_bands = TRUE
)

# Comprehensive vegetation analysis from individual files
veg_analysis <- calculate_multiple_indices(
  red = red_band, nir = nir_band, blue = blue_band,
  indices = c("NDVI", "EVI", "ARVI", "GNDVI", "DVI"),
  output_stack = TRUE,
  region_boundary = "Iowa"
)

# Directory with custom band matching
stress_indices <- calculate_multiple_indices(
  spectral_data = "/path/to/bands/",
  indices = c("PRI", "SIPI", "NDRE"),
  band_names = c("red", "green", "nir", "red_edge"),
  output_stack = TRUE
)

## End(Not run)
```

calculate_multiple_water_indices

Calculate multiple water indices at once

Description

Calculate multiple water indices from the same spectral data in a single operation. Efficient for comprehensive water and moisture analysis.

Usage

```
calculate_multiple_water_indices(
  green,
  nir,
  swir1 = NULL,
  indices = c("NDWI", "MNDWI", "NDMI"),
```

```
    output_stack = TRUE,  
    clamp_values = TRUE,  
    mask_invalid = TRUE,  
    verbose = FALSE  
  )
```

Arguments

green	Green band SpatRaster or file path
nir	NIR band SpatRaster or file path
swir1	SWIR1 band SpatRaster or file path
indices	Vector of index names to calculate
output_stack	Return as single multi-layer raster (TRUE) or list (FALSE)
clamp_values	Apply reasonable value clamping
mask_invalid	Mask invalid values
verbose	Print progress messages

Value

SpatRaster stack or list of water indices

Examples

```
## Not run:  
# These examples require external data files not included with the package  
# Calculate multiple water indices  
water_indices <- calculate_multiple_water_indices(  
  green = green_band,  
  nir = nir_band,  
  swir1 = swir1_band,  
  indices = c("NDWI", "MNDWI", "NDMI", "MSI"),  
  output_stack = TRUE,  
  verbose = TRUE  
)  
  
# Access individual indices  
ndwi <- water_indices[["NDWI"]]  
mndwi <- water_indices[["MNDWI"]]  
  
## End(Not run)
```

 calculate_ndvi_enhanced

Calculate NDVI with time series options

Description

NDVI calculation specifically designed for time series analysis with date matching, quality filtering, temporal smoothing, and multi-temporal support. **Use this for time series analysis, use `calculate_vegetation_index()` for single dates.**

Usage

```
calculate_ndvi_enhanced(
  red_data,
  nir_data,
  clamp_range = c(-0.2, 1),
  match_by_date = FALSE,
  quality_filter = FALSE,
  temporal_smoothing = FALSE,
  verbose = FALSE,
  date_patterns = NULL
)
```

Arguments

red_data	Red band data (files, directory, or raster objects)
nir_data	NIR band data (files, directory, or raster objects)
clamp_range	Range to clamp NDVI values (default: c(-0.2, 1))
match_by_date	Logical: match rasters by date using filenames
quality_filter	Apply quality filtering (remove outliers)
temporal_smoothing	Apply temporal smoothing for time series
verbose	Print progress messages
date_patterns	Custom date patterns for matching

Details

When to Use Enhanced vs Basic NDVI::

Use `calculate_ndvi_enhanced()` for::

- **Time series analysis:** Multiple dates, trend analysis
- **Quality control:** Remove outliers, temporal smoothing
- **Date matching:** Automatic pairing of red/NIR by date
- **Multi-temporal studies:** Seasonal analysis, change detection

Use `calculate_vegetation_index(index_type="NDVI")` for::

- **Single date analysis:** One-time calculation
- **Different indices:** Want to calculate EVI, SAVI, etc. too
- **Quick calculations:** Simple, fast NDVI
- **Mixed workflows:** Part of larger vegetation index analysis

Value

SpatRaster with NDVI layers (single or multi-layer for time series)

Examples

```
## Not run:
# These examples require external data files not included with the package
# Time series NDVI with date matching
ndvi_series <- calculate_ndvi_enhanced(
  red_data = "/path/to/red/time_series/",
  nir_data = "/path/to/nir/time_series/",
  match_by_date = TRUE,
  quality_filter = TRUE,
  temporal_smoothing = TRUE
)

# Simple NDVI (single date with quality control)
ndvi_clean <- calculate_ndvi_enhanced(
  red_data = red_raster,
  nir_data = nir_raster,
  quality_filter = TRUE
)

## End(Not run)
```

calculate_spatial_correlation

Calculate spatial correlation between raster layers

Description

Calculate spatial correlation between two raster layers using various methods. Supports pixel-wise correlation and local correlation analysis.

Usage

```
calculate_spatial_correlation(
  raster1,
  raster2,
  method = "pearson",
  local_correlation = FALSE,
  window_size = 3
)
```

Arguments

raster1	First raster layer
raster2	Second raster layer
method	Correlation method: "pearson", "spearman", "kendall"
local_correlation	Calculate local correlation using moving window
window_size	Window size for local correlation (in pixels)

Value

Correlation coefficient or SpatRaster of local correlations

Examples

```
## Not run:  
# These examples require external data files not included with the package  
# Global correlation between NDVI and soil nitrogen  
correlation <- calculate_spatial_correlation(ndvi_raster, nitrogen_raster)  
  
# Local correlation with moving window  
local_corr <- calculate_spatial_correlation(  
  ndvi_raster, nitrogen_raster,  
  local_correlation = TRUE,  
  window_size = 5  
)  
  
## End(Not run)
```

calculate_vegetation_index

Calculate comprehensive vegetation indices

Description

Calculate a wide range of vegetation indices from spectral bands with automatic band detection, comprehensive error handling, and validation. Supports 40+ different vegetation indices for various applications. Accepts directories, file lists, and automatic CRS handling.

Usage

```
calculate_vegetation_index(  
  spectral_data = NULL,  
  red = NULL,  
  nir = NULL,  
  blue = NULL,  
  green = NULL,
```

```

    swir1 = NULL,
    swir2 = NULL,
    red_edge = NULL,
    red_edge2 = NULL,
    coastal = NULL,
    nir2 = NULL,
    index_type = "NDVI",
    auto_detect_bands = FALSE,
    band_names = NULL,
    clamp_range = NULL,
    mask_invalid = TRUE,
    scale_factor = 1,
    auto_crs_fix = TRUE,
    region_boundary = NULL,
    verbose = FALSE
)

```

Arguments

spectral_data	Either individual bands (red, nir, etc.), a multi-band raster, directory path, or list of raster files
red	Red band SpatRaster or file path
nir	NIR band SpatRaster or file path
blue	Optional blue band
green	Optional green band
swir1	Optional SWIR1 band
swir2	Optional SWIR2 band
red_edge	Optional Red Edge band (e.g. Sentinel-2 Band 5, ~705 nm)
red_edge2	Optional second Red Edge band (e.g. Sentinel-2 Band 6, ~740 nm). Required for S2REP index only.
coastal	Optional Coastal/Aerosol band
nir2	Optional second NIR band
index_type	Vegetation index to calculate (see list_vegetation_indices())
auto_detect_bands	Automatically detect bands from multi-band raster
band_names	Custom band names for multi-band input
clamp_range	Range to clamp output values (optional)
mask_invalid	Mask invalid/extreme values
scale_factor	Scaling factor if needed (default: 1)
auto_crs_fix	Automatically fix CRS mismatches between bands
region_boundary	Optional boundary to crop and mask the result. Can be an sf object, file path to shapefile, or region name (if get_region_boundary() is available)
verbose	Print progress messages

Details**Input Format Support::***Single Calculation::*

Individual band files

ndvi <- calculate_vegetation_index(red = "red.tif", nir = "nir.tif", index_type = "NDVI")

Multi-band raster

evi <- calculate_vegetation_index(spectral_data = "landsat.tif", index_type = "EVI",
auto_detect_bands = TRUE)*Directory/Multiple Files::*

Directory with band files

savi <- calculate_vegetation_index(spectral_data = "/path/to/bands/",
band_names = c("red", "nir"), index_type = "SAVI")

File list

arvi <- calculate_vegetation_index(spectral_data = c("red.tif", "nir.tif", "blue.tif"),
index_type = "ARVI")**Enhanced vs Basic NDVI::***Basic calculate_vegetation_index()::*

- Single time point calculation
- 40+ different indices
- Directory/file support
- Automatic CRS fixing
- **Use for:** Single-date analysis, comparing different indices

calculate_ndvi_enhanced()::

- Time series support
- Quality filtering
- Temporal smoothing
- Date matching between red/NIR
- **Use for:** Multi-temporal analysis, time series trends

#' ## Band Naming Conventions:

The function supports case-insensitive band detection:

- **Generic names:** "red"/"RED"/"Red", "nir"/"NIR", "blue"/"BLUE", "green"/"GREEN"
- **Landsat 8/9:** B1-B7 (e.g., B4=Red, B5=NIR)
- **Sentinel-2:** B01-B12 (e.g., B04=Red, B08=NIR, B05=RedEdge)
- **MODIS:** band1-band7

*Satellite-Specific Examples::***Landsat 8/9:**

Bands automatically detected

ndvi <- calculate_vegetation_index(
 spectral_data = "LC08_stack.tif", # Has bands named B1-B7
 index_type = "NDVI",
 auto_detect_bands = TRUE
)

Sentinel-2:

```
# Red Edge indices need Sentinel-2
ndre <- calculate_vegetation_index(
  spectral_data = sentinel_data, # Has B01-B12
  index_type = "NDRE",
  auto_detect_bands = TRUE
)
```

Custom band names:

```
# Rename your bands first
names(my_raster) <- c("red", "nir", "blue", "green")
```

```
# Or specify explicitly
ndvi <- calculate_vegetation_index(
  red = my_raster[[1]],
  nir = my_raster[[4]],
  index_type = "NDVI"
)
```

For complete band naming documentation, see: vignette("vegetation-indices", package = "geospatialsuite")

Value

SpatRaster of vegetation index

Examples

```
## Not run:
# These examples require satellite imagery files (Landsat/Sentinel data etc.)
# Basic NDVI calculation
ndvi <- calculate_vegetation_index(red = red_band, nir = nir_band, index_type = "NDVI")

# Multi-band raster with auto-detection
evi <- calculate_vegetation_index(spectral_data = landsat_stack,
                                index_type = "EVI", auto_detect_bands = TRUE)

# Directory with automatic band detection
savi <- calculate_vegetation_index(spectral_data = "/path/to/sentinel/bands/",
                                  index_type = "SAVI", auto_detect_bands = TRUE)

# S2REP requires two red-edge bands (Sentinel-2 B5 and B6)
s2rep <- calculate_vegetation_index(spectral_data = sentinel_data,
                                   index_type = "S2REP", auto_detect_bands = TRUE)

# Custom band names for multi-band data
ndvi <- calculate_vegetation_index(spectral_data = sentinel_data,
                                   band_names = c("B4", "B3", "B2", "B8"),
                                   index_type = "NDVI")

# With region boundary (auto CRS conversion and crop/mask)
ndvi_ohio <- calculate_vegetation_index(red = red_band, nir = nir_band,
```

```

index_type = "NDVI",
region_boundary = boundary_sf,
verbose = TRUE)

## End(Not run)

```

calculate_water_index *Calculate water indices including both NDWI variants*

Description

Calculate various water indices including NDWI (McFeeters 1996), MNDWI (Xu 2006), and NDMI (Gao 1996) for water body detection and moisture content. Updated formulas based on latest research and satellite missions (2024).

Usage

```

calculate_water_index(
  green,
  nir,
  swir1 = NULL,
  index_type = "NDWI",
  clamp_range = NULL,
  mask_invalid = TRUE,
  verbose = FALSE
)

```

Arguments

green	Green band SpatRaster or file path
nir	NIR band SpatRaster or file path
swir1	SWIR1 band SpatRaster or file path (for MNDWI, NDMI)
index_type	Index type: "NDWI", "MNDWI", "NDMI", "MSI", "NDII", "WI", "SRWI", "LSWI"
clamp_range	Optional range to clamp output values
mask_invalid	Mask invalid/extreme values
verbose	Print progress messages

Details

Available water indices with their specific applications:

Primary Water Detection Indices::

- **NDWI** (McFeeters 1996): $(\text{Green} - \text{NIR}) / (\text{Green} + \text{NIR})$ - **Use:** Open water body detection, flood mapping - **Range:** Values from -1 to 1, water bodies typically > 0.3 - **Pros:** Simple, effective for clear water - **Cons:** Sensitive to built-up areas, can overestimate water

- **MNDWI** (Xu 2006): $(\text{Green} - \text{SWIR1}) / (\text{Green} + \text{SWIR1})$ - **Use:** Enhanced water detection, urban water bodies - **Range:** Values from -1 to 1, water bodies typically > 0.5 - **Pros:** Better separation of water from built-up areas - **Cons:** Requires SWIR band, less effective with turbid water

Vegetation Moisture Indices::

- **NDMI** (Gao 1996): $(\text{NIR} - \text{SWIR1}) / (\text{NIR} + \text{SWIR1})$ - **Use:** Vegetation water content, drought monitoring - **Range:** Values from -1 to 1, higher values = more water content - **Application:** Agriculture, forest fire risk assessment
- **MSI:** $\text{SWIR1} / \text{NIR}$ - Moisture Stress Index - **Use:** Plant water stress detection - **Range:** $[\theta, 5+]$, lower values = higher moisture
- **NDII:** $(\text{NIR} - \text{SWIR1}) / (\text{NIR} + \text{SWIR1})$ - Same as NDMI - **Use:** Alternative name for NDMI, vegetation moisture

Specialized Water Indices::

- **WI:** $\text{NIR} / \text{SWIR1}$ - Water Index (simple ratio)
- **SRWI:** $\text{NIR} / \text{SWIR1}$ - Simple Ratio Water Index
- **LSWI:** $(\text{NIR} - \text{SWIR1}) / (\text{NIR} + \text{SWIR1})$ - Land Surface Water Index

Band Requirements by Satellite::

- **Landsat 8/9:** Green=Band 3, NIR=Band 5, SWIR1=Band 6
- **Sentinel-2:** Green=Band 3, NIR=Band 8, SWIR1=Band 11
- **MODIS:** Green=Band 4, NIR=Band 2, SWIR1=Band 6

Value

SpatRaster of water index

Examples

```
## Not run:
# These examples require external data files not included with the package
# Original NDWI for water body detection
ndwi <- calculate_water_index(green_band, nir_band, index_type = "NDWI")

# Modified NDWI for enhanced water detection (requires SWIR1)
mndwi <- calculate_water_index(green_band, nir_band, swir1_band, index_type = "MNDWI")

# NDMI for vegetation moisture monitoring
ndmi <- calculate_water_index(green_band, nir_band, swir1_band, index_type = "NDMI")

# With quality control
water_index <- calculate_water_index(
  green = "green.tif",
  nir = "nir.tif",
  swir1 = "swir1.tif",
  index_type = "MNDWI",
  clamp_range = c(-1, 1),
  mask_invalid = TRUE,
```

```
    verbose = TRUE
  )

  ## End(Not run)
```

```
compare_interpolation_methods
  Compare interpolation methods
```

Description

Compare multiple interpolation methods using cross-validation and return performance metrics for method selection.

Usage

```
compare_interpolation_methods(
  spatial_data,
  target_variable,
  methods = c("NN", "simple", "spline"),
  cv_folds = 5,
  verbose = TRUE
)
```

Arguments

spatial_data	Spatial data for interpolation
target_variable	Variable to interpolate
methods	Vector of methods to compare
cv_folds	Number of cross-validation folds
verbose	Print comparison results

Value

Data frame with method comparison results

Examples

```
## Not run:
# These examples require external data files not included with the package
# Compare interpolation methods
method_comparison <- compare_interpolation_methods(
  soil_data,
  target_variable = "nitrogen",
  methods = c("NN", "simple", "spline"),
```

```

    cv_folds = 10
  )

  # View results
  print(method_comparison)
  # Best method
  best_method <- method_comparison$method[which.min(method_comparison$rmse)]

  ## End(Not run)

```

create_comparison_map *Create comparison map (before/after, side-by-side)*

Description

Create comparison maps showing before/after analysis or side-by-side comparisons using reliable terra plotting.

Usage

```

create_comparison_map(
  data1,
  data2,
  comparison_type = "side_by_side",
  titles = c("Dataset 1", "Dataset 2"),
  region_boundary = NULL,
  color_scheme = "viridis",
  output_file = NULL,
  verbose = FALSE
)

```

Arguments

data1	First dataset (before, reference)
data2	Second dataset (after, comparison)
comparison_type	Type: "side_by_side", "difference"
titles	Titles for each dataset
region_boundary	Optional region boundary
color_scheme	Color scheme for datasets
output_file	Optional output file path
verbose	Print progress messages

Value

NULL (plots directly to device) or file path if saved

Examples

```
## Not run:
# These examples require external data files not included with the package
# Before/after NDVI comparison
create_comparison_map("ndvi_2020.tif", "ndvi_2023.tif",
                     comparison_type = "side_by_side",
                     titles = c("2020", "2023"),
                     output_file = "ndvi_comparison.png")

## End(Not run)
```

create_crop_mask	<i>Create crop mask from CDL data</i>
------------------	---------------------------------------

Description

Create binary or classified crop mask from USDA CDL data for specified crops. Fixed to handle terra operations properly.

Usage

```
create_crop_mask(
  cdl_data,
  crop_codes,
  region_boundary = NULL,
  mask_type = "binary"
)
```

Arguments

cdl_data	CDL raster data (file path or SpatRaster)
crop_codes	Vector of CDL codes or crop names
region_boundary	Optional region boundary for clipping
mask_type	Type of mask: "binary" (1/0) or "preserve" (keep original codes)

Value

SpatRaster with crop mask

Examples

```
## Not run:
# These examples require actual CDL data files
# Create corn mask
corn_mask <- create_crop_mask("cdl_2023.tif", "corn", "Iowa")

# Create grain crops mask
grain_mask <- create_crop_mask(cdl_raster, "grains", mask_type = "preserve")

## End(Not run)

# Example with mock data (this can run)
mock_cdl <- terra::rast(nrows = 5, ncols = 5, crs = "EPSG:4326")
terra::values(mock_cdl) <- c(1, 1, 5, 5, 24, 1, 5, 5, 24, 24,
                             1, 1, 5, 24, 24, 5, 5, 24, 24, 1,
                             1, 5, 5, 24, 1) # corn, soy, wheat

# Create corn mask from mock data
corn_mask <- create_crop_mask(mock_cdl, "corn")
print(terra::values(corn_mask)) # Should show 1s and 0s
```

```
create_interactive_map
```

Create interactive map using leaflet (if available)

Description

Create interactive maps with leaflet integration when available. Falls back gracefully when leaflet is not installed.

Usage

```
create_interactive_map(
  spatial_data,
  fill_variable = NULL,
  popup_vars = NULL,
  basemap = "terrain",
  color_scheme = "viridis",
  title = "Interactive Map",
  verbose = FALSE
)
```

Arguments

`spatial_data` Spatial data to map (sf object)

fill_variable	Variable for coloring/filling
popup_vars	Variables to show in popups
basemap	Basemap type: "terrain", "satellite", "osm", "light"
color_scheme	Color scheme for continuous variables
title	Map title
verbose	Print progress messages

Value

leaflet map object or NULL if leaflet unavailable

Examples

```
## Not run:
# These examples demonstrate workflows with user's own spatial data
# Simple interactive point map
map <- create_interactive_map(study_sites, fill_variable = "ndvi_mean")

# Polygon map with custom basemap
map <- create_interactive_map(counties, fill_variable = "population",
                             basemap = "satellite")

## End(Not run)
```

create_raster_mosaic *Create raster mosaic with intelligent file selection*

Description

Create mosaics from multiple raster files with various methods and intelligent file selection based on region boundaries.

Usage

```
create_raster_mosaic(
  input_data,
  method = "merge",
  region_boundary = NULL,
  output_file = NULL,
  parallel = FALSE
)
```

Arguments

input_data	Character vector of file paths, directory path, or list of rasters
method	Mosaicing method: "merge", "mosaic", "mean", "max", "min"
region_boundary	Optional region boundary for clipping
output_file	Optional output file path
parallel	Use parallel processing

Value

SpatRaster object

Examples

```
## Not run:
# These examples require external data files not included with the package
# Basic mosaic
mosaic <- create_raster_mosaic("/path/to/rasters", method = "merge")

# Mosaic for specific region
ohio_mosaic <- create_raster_mosaic("/aster/files", "merge", "Ohio")

# Mean composite
mean_mosaic <- create_raster_mosaic(raster_list, method = "mean")

## End(Not run)
```

create_spatial_map *Create universal spatial map with reliable terra plotting*

Description

Universal mapping function that works with any spatial data type using reliable terra and base R plotting. No complex dependencies required. Falls back gracefully when optional packages are unavailable.

Usage

```
create_spatial_map(
  spatial_data,
  fill_variable = NULL,
  coord_cols = c("lon", "lat"),
  region_boundary = NULL,
  map_type = "auto",
  color_scheme = "viridis",
  interactive = FALSE,
```

```

    title = NULL,
    point_size = 3,
    output_file = NULL,
    verbose = FALSE
  )

```

Arguments

spatial_data	sf object, data.frame with coordinates, file path, or SpatRaster
fill_variable	Variable to use for fill/color (for vector data)
coord_cols	Coordinate column names if data.frame provided
region_boundary	Optional region boundary
map_type	Type of map: "points", "polygons", "raster", "auto"
color_scheme	Color scheme: "viridis", "plasma", "ndvi", "terrain", "categorical"
interactive	Create interactive map using leaflet (if available)
title	Map title
point_size	Size of points (for point data)
output_file	Optional output file path
verbose	Print progress messages

Value

ggplot2 object, leaflet map, or file path (depending on options)

Examples

```

## Not run:
# These examples require external data files not included with the package
# Simple point map
create_spatial_map(study_sites, fill_variable = "ndvi_mean")

# Raster map with region boundary
create_spatial_map(ndvi_raster, region_boundary = "Ohio",
  color_scheme = "ndvi")

# Interactive map (if leaflet available)
create_spatial_map(counties, fill_variable = "population",
  interactive = TRUE)

## End(Not run)

```

download_sample_data *Download or access sample datasets for geospatialsuite*

Description

Access built-in sample datasets for learning and testing geospatialsuite functionality. All datasets are in .rds format (CRAN-preferred) and included with the package (~50-80KB total).

Usage

```
download_sample_data(
  dataset = "all",
  dest_dir = tempdir(),
  overwrite = FALSE,
  verbose = TRUE
)
```

Arguments

dataset	Name of dataset to download. See <code>list_sample_datasets()</code> for available options. Use "all" to list all datasets.
dest_dir	Directory to save downloaded data (default: <code>tempdir()</code>)
overwrite	Overwrite existing files? (default: <code>FALSE</code>)
verbose	Print download progress? (default: <code>TRUE</code>)

Details

Built-in Sample Data (No Download Required):

The package includes small sample datasets in `inst/extdata/` using .rds format:

- **sample_red.rds**: Red band SpatRaster (10x10 pixels, Ohio region)
- **sample_nir.rds**: NIR band SpatRaster (10x10 pixels)
- **sample_blue.rds**: Blue band SpatRaster (10x10 pixels)
- **sample_green.rds**: Green band SpatRaster (10x10 pixels)
- **sample_swir1.rds**: SWIR1 band SpatRaster (10x10 pixels)
- **sample_multiband.rds**: Multi-band SpatRaster (5 layers stacked)
- **sample_points.rds**: Sample field locations (20 points, sf object)
- **sample_boundary.rds**: Sample study area polygon (sf object)
- **sample_coordinates.csv**: Sample coordinates with attributes

All .rds files contain native R objects (SpatRaster or sf) that load instantly with `readRDS()` or the `load_sample_data()` helper function.

Value

Character vector of file paths to sample data

Examples

```
# List available datasets
datasets <- list_sample_datasets()
print(datasets)

# Get path to built-in sample data
red_file <- get_sample_data("sample_red.rds")
nir_file <- get_sample_data("sample_nir.rds")

# Load and use sample data
red <- load_sample_data("sample_red.rds")
nir <- load_sample_data("sample_nir.rds")
ndvi <- calculate_vegetation_index(red = red, nir = nir, index_type = "NDVI")
# Plot (users can run this interactively)
# plot(ndvi, main = "NDVI", col = terrain.colors(100))
```

```
extract_dates_universal
```

Extract dates from filenames using various patterns

Description

Universal function to extract dates from filenames or provide custom labels. Enhanced with more flexible regex patterns that work with any filename prefix.

Usage

```
extract_dates_universal(input_data, date_patterns = NULL, verbose = FALSE)
```

Arguments

input_data	Character vector (file paths or folder), or list of raster layers
date_patterns	Named list of custom regex patterns for date extraction
verbose	Print progress messages

Value

Character vector of extracted or inferred date labels

Examples

```
## Not run:
# These examples require external data files not included with the package
# Extract dates from filenames
dates <- extract_dates_universal(c("ndvi_2023-05-15.tif", "evi_2023-06-15.tif"))
```

```
# Custom date patterns
custom_patterns <- list("MMDDYYYY" = "\\b[0-9]{2}[0-9]{2}[0-9]{4}\\b")
dates <- extract_dates_universal(files, custom_patterns)

## End(Not run)
```

geocoding_examples *Geocoding Examples and Use Cases*

Description

This file contains documented examples for the auto-geocoding functionality. These examples demonstrate various use cases for the `auto_geocode_data` function.

Examples

```
## Not run:
# =====
# Example 1: State-level analysis
# =====
state_data <- data.frame(
  state = c("California", "TX", "New York", "FL"),
  gdp_billions = c(3598, 2357, 2053, 1389),
  population_millions = c(39.5, 29.1, 20.2, 22.2)
)

# Auto-detect and geocode
state_sf <- auto_geocode_data(state_data, verbose = TRUE)

# Visualize
quick_map(state_sf, variable = "gdp_billions",
           title = "GDP by State (Billions)")

# =====
# Example 2: County-level with FIPS codes
# =====
county_data <- data.frame(
  fips = c("39049", "39035", "39113", "39061"),
  county_name = c("Franklin", "Cuyahoga", "Montgomery", "Hamilton"),
  unemployment = c(4.2, 5.1, 4.8, 3.9)
)

county_sf <- auto_geocode_data(county_data)
quick_map(county_sf, variable = "unemployment")

# =====
# Example 3: Watershed analysis with HUC codes
# =====
# Works with ANY HUC format: HUC_8, HUC-8, huc8, Huc 8, etc.
```

```

watershed_data <- data.frame(
  HUC_8 = c("04100009", "04100012", "04110002"),
  basin_name = c("Great Miami", "Mill Creek-Cincinnati", "Middle Ohio"),
  water_quality_index = c(72, 65, 80),
  nitrogen_mg_l = c(2.3, 3.1, 1.8)
)

huc_sf <- auto_geocode_data(watershed_data, verbose = TRUE)
quick_map(huc_sf, variable = "water_quality_index",
          title = "Water Quality by Watershed")

# =====
# Example 4: ZIP code analysis
# =====
zip_data <- data.frame(
  zip = c("43215", "44113", "45202", "43017"),
  median_home_price = c(285000, 195000, 320000, 410000)
)

zip_sf <- auto_geocode_data(zip_data)
quick_map(zip_sf, variable = "median_home_price")

# =====
# Example 5: Loading from CSV file
# =====
# Assuming you have a CSV with state data
census_sf <- auto_geocode_data("state_census_data.csv", verbose = TRUE)
quick_map(census_sf)

# =====
# Example 6: Preview before geocoding
# =====
# Check what will be detected without actually geocoding
my_data <- data.frame(
  State = c("Ohio", "Michigan"),
  HUC8 = c("04100009", "04100012"),
  value = c(100, 200)
)

preview_geocoding(my_data)

# =====
# Example 7: Explicit column specification
# =====
# When auto-detection isn't working or you want to be explicit
data_with_weird_names <- data.frame(
  my_state_col = c("CA", "TX", "NY"),
  revenue = c(1000000, 800000, 1200000)
)

result <- auto_geocode_data(
  data_with_weird_names,
  entity_column = "my_state_col",

```

```

    entity_type = "state",
    verbose = TRUE
  )

  # =====
  # Example 8: Integration with other geospatialsuite functions
  # =====
  # Geocode, then use with spatial analysis
  state_sf <- auto_geocode_data(state_data)

  # Use with universal_spatial_join
  raster_data <- terra::rast("ndvi_data.tif")
  result <- universal_spatial_join(
    source_data = state_sf,
    target_data = raster_data,
    method = "extract"
  )

  # Create comprehensive analysis
  quick_map(result, variable = "extracted_mean_ndvi")

  ## End(Not run)

```

```
get_comprehensive_cdl_codes
```

Get comprehensive CDL crop codes

Description

Get USDA Cropland Data Layer (CDL) codes for specific crops or crop categories. Supports all major crops and predefined categories.

Usage

```
get_comprehensive_cdl_codes(crop_type = "all")
```

Arguments

crop_type	Crop type or category name. Options include: <ul style="list-style-type: none"> • Individual crops: "corn", "soybeans", "wheat", etc. • Categories: "grains", "oilseeds", "fruits", "vegetables", etc. • "all" for all available codes
-----------	---

Value

Vector of CDL codes

Examples

```
# Get corn code
corn_codes <- get_comprehensive_cdl_codes("corn")
print(corn_codes) # Should be 1

# Get all grain crop codes
grain_codes <- get_comprehensive_cdl_codes("grains")
print(grain_codes) # Should be vector of grain codes

# See available crop types (this will print to console)
get_comprehensive_cdl_codes("help")
```

get_region_boundary *Get region boundary for any specified region*

Description

Universal function to get region boundaries for any geographic area including US states, countries, CONUS, counties, or custom bounding boxes with comprehensive error handling.

Usage

```
get_region_boundary(region_def, verbose = FALSE)
```

Arguments

region_def	Region definition in various formats: <ul style="list-style-type: none">• Character: "Ohio", "Nigeria", "CONUS"• Character with colon: "Ohio:Franklin" (state:county)• Numeric vector: c(xmin, ymin, xmax, ymax) bounding box• sf object: existing spatial object
verbose	Print progress messages

Value

sf object with boundary geometry

Examples

```
# US State with error handling
ohio_boundary <- get_region_boundary("Ohio")

# Custom bounding box with validation
custom_area <- get_region_boundary(c(-84.5, 39.0, -82.0, 41.0))
```

get_sample_data	<i>Get path to sample data file</i>
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Description

Helper function to get the full path to a sample data file included with geospatialsuite. This is useful when you want the file path rather than loading the data directly.

Usage

```
get_sample_data(filename)
```

Arguments

filename Name of sample data file (e.g., "sample_red.rds")

Value

Character string with full path to file

Examples

```
# Get path to sample raster
red_file <- get_sample_data("sample_red.rds")
nir_file <- get_sample_data("sample_nir.rds")

# Load manually with terra::rast() for PackedSpatRaster
red <- terra::rast(readRDS(red_file))
nir <- terra::rast(readRDS(nir_file))

# Or use load_sample_data() directly (recommended)
red <- load_sample_data("sample_red.rds")
nir <- load_sample_data("sample_nir.rds")

# Calculate NDVI
ndvi <- calculate_vegetation_index(red = red, nir = nir, index_type = "NDVI")
# plot(ndvi)
```

```
integrate_terrain_analysis  
  Integrate terrain analysis with vector data
```

Description

Specialized function for terrain analysis integration. Calculates terrain variables from DEM and extracts values to vector data points/polygons.

Usage

```
integrate_terrain_analysis(  
  vector_data,  
  elevation_raster,  
  terrain_vars = c("slope", "aspect", "TRI", "TPI", "flowdir"),  
  custom_terrain_functions = NULL,  
  extraction_method = "simple"  
)
```

Arguments

vector_data	Vector data (points, lines, or polygons)
elevation_raster	Digital elevation model
terrain_vars	Terrain variables to calculate
custom_terrain_functions	Custom terrain analysis functions
extraction_method	Method for extracting terrain values

Value

sf object with terrain attributes

Examples

```
## Not run:  
# These examples require external data files not included with the package  
# Extract terrain variables for study sites  
sites_with_terrain <- integrate_terrain_analysis(  
  vector_data = "study_sites.shp",  
  elevation_raster = "dem.tif",  
  terrain_vars = c("slope", "aspect", "TRI", "TPI")  
)  
  
# Use custom terrain functions  
custom_functions <- list(  

```

```
ruggedness = function(sf_data) {  
  sf_data$slope * sf_data$TRI  
}  
)  
  
terrain_analysis <- integrate_terrain_analysis(  
  vector_data = field_boundaries,  
  elevation_raster = dem_raster,  
  custom_terrain_functions = custom_functions  
)  
  
## End(Not run)
```

list_sample_datasets *List available sample datasets*

Description

Display information about all sample datasets included with geospatialsuite.

Usage

```
list_sample_datasets(detailed = TRUE)
```

Arguments

detailed Return detailed information? (default: TRUE)

Value

Data frame with dataset information

Examples

```
# List all sample datasets  
datasets <- list_sample_datasets()  
print(datasets)  
  
# Quick list  
datasets_simple <- list_sample_datasets(detailed = FALSE)
```

`list_vegetation_indices`*Get comprehensive list of available vegetation indices*

Description

Returns detailed information about all 40+ available vegetation indices including formulas, required bands, applications, and references.

Usage

```
list_vegetation_indices(  
  category = "all",  
  application = "all",  
  detailed = FALSE  
)
```

Arguments

<code>category</code>	Filter by category: "all", "basic", "enhanced", "specialized", "stress"
<code>application</code>	Filter by application: "general", "agriculture", "forestry", "stress", "water"
<code>detailed</code>	Return detailed information including formulas and references

Value

Data frame with vegetation index information

Examples

```
# All available indices  
all_indices <- list_vegetation_indices()  
  
# Only stress detection indices  
stress_indices <- list_vegetation_indices(category = "stress")  
  
# Detailed information with formulas  
detailed_info <- list_vegetation_indices(detailed = TRUE)  
  
# Agricultural applications only  
ag_indices <- list_vegetation_indices(application = "agriculture")
```

`list_water_indices` *Get comprehensive list of available water indices*

Description

Returns detailed information about all available water indices including formulas, required bands, applications, and interpretation guidelines.

Usage

```
list_water_indices(detailed = FALSE, application_filter = "all")
```

Arguments

`detailed` Return detailed information including formulas and applications
`application_filter` Filter by application: "all", "water_detection", "moisture_monitoring", "drought_assessment"

Value

Data frame with water index information

Examples

```
# All available water indices
water_indices <- list_water_indices()

# Detailed information with formulas
detailed_info <- list_water_indices(detailed = TRUE)

# Only water detection indices
water_detection <- list_water_indices(application_filter = "water_detection")
```

`load_raster_data` *Load raster data from various sources*

Description

Universal function to load raster data from files, directories, or raster objects with comprehensive error handling and validation.

Usage

```
load_raster_data(
  input_data,
  pattern = "\\.(tif|tiff)$",
  recursive = FALSE,
  verbose = FALSE
)
```

Arguments

input_data	Character string (path to file or directory), character vector of file paths, or a SpatRaster/Raster* object
pattern	File pattern for directory search (default: tif files)
recursive	Search subdirectories recursively
verbose	Print progress messages

Value

List of terra SpatRaster objects

Examples

```
## Not run:
# These examples require directory structures with multiple data files
# Load from directory with error handling
rasters <- load_raster_data("/path/to/raster/files")

# Load from file list with validation
rasters <- load_raster_data(c("file1.tif", "file2.tif"))

## End(Not run)
```

load_sample_data	<i>Load sample data directly</i>
------------------	----------------------------------

Description

Convenience function to load sample data directly into R objects. Automatically detects file type and uses appropriate loading function. For .rds files containing SpatRaster objects, properly unpacks from PackedSpatRaster format. For .csv files, uses read.csv().

Usage

```
load_sample_data(filename, ...)
```

Arguments

filename Name of sample data file
... Additional arguments passed to loading function

Value

Loaded data object (SpatRaster, sf, or data.frame)

Examples

```
# Load raster data (returns SpatRaster)
red <- load_sample_data("sample_red.rds")
nir <- load_sample_data("sample_nir.rds")
blue <- load_sample_data("sample_blue.rds")

# Load vector data (returns sf object)
points <- load_sample_data("sample_points.rds")
boundary <- load_sample_data("sample_boundary.rds")

# Load CSV (returns data.frame)
coords <- load_sample_data("sample_coordinates.csv")

# Use loaded data
ndvi <- calculate_vegetation_index(red = red, nir = nir, index_type = "NDVI")
evi <- calculate_vegetation_index(red = red, nir = nir, blue = blue, index_type = "EVI")
# plot(ndvi, main = "NDVI")
# plot(evi, main = "EVI")
```

multiscale_operations *Multi-scale spatial operations*

Description

Handle multi-scale operations including up-scaling, down-scaling, and pyramid operations for efficient processing.

Usage

```
multiscale_operations(  
  spatial_data,  
  target_scales = c(1, 2, 4, 8),  
  operation = "mean",  
  pyramid = FALSE  
)
```

Arguments

spatial_data	Input spatial data
target_scales	Vector of scale factors
operation	Operation to perform at each scale
pyramid	Create image pyramid

Value

List of results at different scales

Examples

```
## Not run:
# These examples require external data files not included with the package
# Create multi-scale analysis
scales <- multiscale_operations("data.tif", c(1, 2, 4, 8), "mean")

## End(Not run)
```

plot_raster_fast *Create fast raster plot using terra*

Description

Create efficient raster plots using terra's native plotting capabilities. Fast and reliable without external dependencies.

Usage

```
plot_raster_fast(
  raster_data,
  title = "Raster Plot",
  color_scheme = "viridis",
  region_boundary = NULL,
  breaks = NULL,
  output_file = NULL,
  verbose = FALSE
)
```

Arguments

raster_data	SpatRaster to plot or file path
title	Plot title
color_scheme	Color scheme to apply

region_boundary	Optional boundary to overlay
breaks	Custom breaks for classification
output_file	Optional output file path
verbose	Print progress messages

Value

NULL (plots directly to device) or file path if saved

Examples

```
## Not run:
# These examples demonstrate workflows with user's own spatial data
# Simple raster plot
plot_raster_fast(ndvi_raster, "NDVI Analysis", "ndvi")

# With custom breaks and save to file
plot_raster_fast(elevation, "Elevation", "terrain",
                 breaks = c(0, 500, 1000, 1500, 2000),
                 output_file = "elevation_map.png")

## End(Not run)
```

plot_rgb_raster	<i>Create multi-band raster RGB plot</i>
-----------------	--

Description

Create RGB plots from multi-band rasters using terra's native RGB plotting. Reliable and fast without external dependencies.

Usage

```
plot_rgb_raster(
  raster_data,
  r = 1,
  g = 2,
  b = 3,
  stretch = "lin",
  title = "RGB Composite",
  output_file = NULL,
  verbose = FALSE
)
```

Arguments

raster_data	Multi-band SpatRaster or file path
r	Red band index (default: 1)
g	Green band index (default: 2)
b	Blue band index (default: 3)
stretch	Stretch method: "lin", "hist", "minmax", "perc"
title	Plot title
output_file	Optional output file path
verbose	Print progress messages

Value

NULL (plots directly to device) or file path if saved

Examples

```
## Not run:
# These examples require external data files not included with the package
# True color composite
plot_rgb_raster(satellite_data, r = 3, g = 2, b = 1, title = "True Color")

# False color composite with histogram stretch
plot_rgb_raster(landsat_data, r = 4, g = 3, b = 2, stretch = "hist",
               title = "False Color Composite", output_file = "rgb_composite.png")

## End(Not run)
```

preview_geocoding *Preview geographic entity detection*

Description

Test what geographic entities will be detected in your data without actually performing the geocoding. Useful for debugging and verification.

Usage

```
preview_geocoding(data, show_sample = TRUE, n_sample = 5)
```

Arguments

data	Data frame to analyze
show_sample	Show sample values (default: TRUE)
n_sample	Number of sample values to show (default: 5)

Value

List with detection results

Examples

```
## Not run:  
# Check what will be detected  
my_data <- data.frame(  
  HUC_8 = c("04100009", "04100012"),  
  State = c("Ohio", "PA"),  
  value = c(100, 200)  
)  
  
preview_geocoding(my_data)  
  
## End(Not run)
```

quick_diagnostic

Quick diagnostic check

Description

Quick diagnostic to identify what might be wrong with the package.

Usage

```
quick_diagnostic()
```

Value

List containing diagnostic results with components:

r_version Character string of R version

minimal_works Logical indicating if basic functionality works

function_status Logical indicating function availability status

Called primarily for side effects (printing diagnostic messages).

`quick_map`*Quick map function - one-line mapping with auto-detection*

Description

Ultra-simple function for quick spatial mapping. Auto-detects data type and creates appropriate map.

Usage

```
quick_map(spatial_data, variable = NULL, title = NULL, ...)
```

Arguments

<code>spatial_data</code>	Any spatial data
<code>variable</code>	Variable to visualize (optional, auto-detected)
<code>title</code>	Map title (optional)
<code>...</code>	Additional arguments passed to <code>create_spatial_map</code>

Value

Map object

Examples

```
## Not run:  
# These examples require external data files not included with the package  
quick_map("data.shp")  
quick_map(my_raster)  
quick_map(points_data, interactive = TRUE)  
  
## End(Not run)
```

`raster_to_raster_ops`*Raster to Raster Operations*

Description

Specialized function for mathematical and overlay operations between rasters. Handles alignment, projection, and complex operations with comprehensive error handling and performance optimization.

Usage

```
raster_to_raster_ops(
  raster1,
  raster2,
  operation = "overlay",
  align_method = "resample",
  summary_function = "mean",
  handle_na = "propagate",
  mask_value = NA,
  output_file = NULL,
  verbose = FALSE
)
```

Arguments

raster1	First raster (SpatRaster or file path)
raster2	Second raster (SpatRaster or file path)
operation	Character. Mathematical operation: <ul style="list-style-type: none"> • "add": Add rasters (raster1 + raster2) • "subtract": Subtract rasters (raster1 - raster2) • "multiply": Multiply rasters (raster1 * raster2) • "divide": Divide rasters (raster1 / raster2) • "mask": Mask raster1 with raster2 • "overlay": Combine with summary function • "difference": Absolute difference raster1 - raster2 • "ratio": Ratio raster1 / raster2 (with zero handling)
align_method	Character. How to align mismatched rasters: <ul style="list-style-type: none"> • "resample": Resample raster2 to match raster1 (default) • "crop": Crop both to common extent • "extend": Extend smaller raster to match larger • "project": Reproject raster2 to raster1 CRS
summary_function	Character. Function for overlay operation
handle_na	Character. How to handle NA values: <ul style="list-style-type: none"> • "propagate": NA + value = NA (default) • "ignore": Skip NAs in calculations • "zero": Treat NAs as zero
mask_value	Numeric. Value to use for masking (default: NA)
output_file	Character. Optional output file path
verbose	Logical. Print processing details

Value

SpatRaster with operation results

Examples

```
## Not run:
# These examples require external data files not included with the package
# Mathematical operations
sum_raster <- raster_to_raster_ops("ndvi.tif", "evi.tif", "add")
diff_raster <- raster_to_raster_ops("before.tif", "after.tif", "subtract")

# Masking operations
masked <- raster_to_raster_ops("data.tif", "mask.tif", "mask")

# Complex overlay with alignment
overlay <- raster_to_raster_ops(
  raster1 = "fine_res.tif",
  raster2 = "coarse_res.tif",
  operation = "overlay",
  align_method = "resample",
  summary_function = "mean",
  verbose = TRUE
)

## End(Not run)
```

run_comprehensive_geospatial_workflow

Run comprehensive geospatial workflow -

Description

Execute complete geospatial analysis workflows with simplified visualization. to handle test cases and provide robust error handling without complex dependencies.

Usage

```
run_comprehensive_geospatial_workflow(analysis_config)
```

Arguments

analysis_config

List containing analysis configuration with required fields:

- analysis_type: "ndvi_crop_analysis", "water_quality_analysis", "terrain_analysis", "temporal_analysis", "vegetation_comprehensive", "mosaic_analysis", "interactive_mapping"
- input_data: Input data paths or objects
- region_boundary: Region boundary specification
- output_folder: Output directory (optional)
- visualization_config: Visualization settings (optional)

Value

List containing analysis results, visualizations, summary, and configuration

Examples

```
## Not run:  
# These examples require external data files not included with the package  
# Simple NDVI crop analysis workflow  
config <- list(  
  analysis_type = "ndvi_crop_analysis",  
  input_data = list(red = red_raster, nir = nir_raster),  
  region_boundary = "Ohio",  
  output_folder = "results/"  
)  
results <- run_comprehensive_geospatial_workflow(config)  
  
## End(Not run)
```

run_comprehensive_vegetation_workflow

Run comprehensive vegetation analysis workflow -

Description

Complete vegetation analysis using multiple indices with simplified processing. for reliability and test compatibility.

Usage

```
run_comprehensive_vegetation_workflow(config, output_folder = tempdir())
```

Arguments

config	Analysis configuration
output_folder	Output directory

Value

Comprehensive vegetation analysis results

`run_enhanced_ndvi_crop_workflow`*Run enhanced NDVI crop analysis workflow -*

Description

Enhanced NDVI workflow with quality filtering, temporal analysis, and visualization. to handle test scenarios and provide robust error handling.

Usage

```
run_enhanced_ndvi_crop_workflow(config, output_folder = tempdir())
```

Arguments

<code>config</code>	Analysis configuration
<code>output_folder</code>	Output directory

Value

List with enhanced NDVI results and visualizations

`run_interactive_mapping_workflow`*Run interactive mapping workflow*

Description

Create interactive mapping workflow with multiple data types and layers. Simplified for reliability.

Usage

```
run_interactive_mapping_workflow(config, output_folder = tempdir())
```

Arguments

<code>config</code>	Analysis configuration
<code>output_folder</code>	Output directory

Value

Interactive mapping results

`select_rasters_for_region`*Select rasters for specific region with intelligent filtering*

Description

Intelligently select raster files that overlap with a specified region. For ASTER files (GDEM and WBD), uses filename-based coordinate extraction for fast filtering without loading full rasters.

Usage

```
select_rasters_for_region(  
  input_folder,  
  region_boundary,  
  overlap = TRUE,  
  buffer_size = 0.1  
)
```

Arguments

<code>input_folder</code>	Directory containing raster files
<code>region_boundary</code>	Region boundary or bounding box
<code>overlap</code>	Logical. If TRUE (default), checks for actual overlap with region. If FALSE, only includes tiles whose centroid falls within region. Only applies to ASTER files; non-ASTER files always use overlap detection.
<code>buffer_size</code>	Buffer around region (in degrees)

Value

Character vector of relevant file paths

Examples

```
# Select ASTER files for Michigan using overlap detection  
michigan_files <- select_rasters_for_region("/aster/files", "Michigan")  
  
# Use centroid-based selection (faster, fewer tiles)  
michigan_files_centroid <- select_rasters_for_region("/aster/files", "Michigan",  
  overlap = FALSE)  
  
# Select with custom buffer  
nevada_files <- select_rasters_for_region("/data", "Nevada", buffer_size = 0.2)
```

spatial_interpolation *Legacy spatial interpolation function (for backward compatibility)*

Description

Simplified version of spatial interpolation maintaining backward compatibility. For new projects, use `spatial_interpolation_comprehensive()` instead.

Usage

```
spatial_interpolation(  
  spatial_data,  
  target_variables,  
  method = "NN",  
  power = 2,  
  mice_method = "pmm"  
)
```

Arguments

<code>spatial_data</code>	sf object with some missing values
<code>target_variables</code>	Variables to interpolate
<code>method</code>	Interpolation method: "NN", "simple", "mice"
<code>power</code>	Power parameter for simple method (default: 2)
<code>mice_method</code>	MICE method for multivariate imputation

Value

sf object with interpolated values

Examples

```
## Not run:  
# These examples require external data files not included with the package  
# Simple interpolation (legacy interface)  
interpolated_data <- spatial_interpolation(  
  soil_data,  
  target_variables = c("nitrogen", "carbon"),  
  method = "NN"  
)  
  
## End(Not run)
```

 spatial_interpolation_comprehensive

Perform spatial interpolation for missing data

Description

Perform spatial interpolation using reliable methods to fill missing values in spatial datasets. Supports nearest neighbor, spline interpolation, and multivariate imputation with comprehensive error handling.

Usage

```
spatial_interpolation_comprehensive(
  spatial_data,
  target_variables,
  method = "NN",
  target_grid = NULL,
  region_boundary = NULL,
  power = 2,
  max_distance = Inf,
  min_points = 3,
  max_points = 50,
  cross_validation = FALSE,
  cv_folds = 5,
  handle_outliers = "none",
  outlier_threshold = 3,
  coord_cols = c("lon", "lat"),
  mice_method = "pmm",
  mice_iterations = 10,
  output_format = "sf",
  output_file = NULL,
  verbose = FALSE
)
```

Arguments

<code>spatial_data</code>	Spatial data to interpolate. Can be: <ul style="list-style-type: none"> • sf object with point geometries • data.frame with coordinate columns • File path to spatial data (CSV, SHP, GeoJSON)
<code>target_variables</code>	Character vector of variables to interpolate
<code>method</code>	Interpolation method: <ul style="list-style-type: none"> • "NN": Nearest neighbor (default) • "simple": Simple distance weighting

	<ul style="list-style-type: none"> • "spline": Thin plate spline interpolation • "mice": Multivariate imputation (requires mice package) • "auto": Automatically select best method based on data
target_grid	Target grid for interpolation. Can be: <ul style="list-style-type: none"> • SpatRaster template for raster output • sf object with target locations • NULL for point-to-point interpolation only
region_boundary	Optional region boundary for clipping results
power	Power parameter for simple distance weighting (default: 2)
max_distance	Maximum distance for interpolation (map units)
min_points	Minimum number of points for interpolation
max_points	Maximum number of points to use for each prediction
cross_validation	Perform cross-validation for accuracy assessment
cv_folds	Number of folds for cross-validation (default: 5)
handle_outliers	Method for outlier handling: "none", "remove", "cap"
outlier_threshold	Z-score threshold for outlier detection (default: 3)
coord_cols	Coordinate column names for data.frame input
mice_method	MICE method for multivariate imputation
mice_iterations	Number of MICE iterations (default: 10)
output_format	Output format: "sf", "raster", "both"
output_file	Optional output file path
verbose	Print detailed progress messages

Details

Supported Interpolation Methods::

Distance-Based Methods::

- **NN** (Nearest Neighbor): Assigns nearest known value - Best for: Categorical data or when preserving exact values - Fast and creates Voronoi-like patterns - No assumptions about data distribution
- **Simple** (Simple distance weighting): Basic distance-based averaging - Best for: Quick estimates with minimal computation - Uses inverse distance weighting without external dependencies

Statistical Methods::

- **Spline**: Smooth surface interpolation using thin plate splines - Best for: Smooth, continuous phenomena - Creates smooth surfaces without sharp changes - Good for environmental data with gradual spatial variation

Multivariate Methods::

- **MICE**: Multivariate imputation by chained equations - Best for: Multiple correlated variables with missing values - Handles complex missing data patterns - Preserves relationships between variables - Requires mice package

Input Data Support::

- sf objects with point geometries
- data.frame with coordinate columns
- File paths (CSV, shapefile, GeoJSON)
- Target grids for raster output

Quality Control Features::

- Cross-validation for method comparison
- Outlier detection and handling
- Performance metrics calculation
- Robust error handling

Value

Depending on output_format:

"sf" sf object with interpolated values

"raster" SpatRaster with interpolated surfaces

"both" List containing both sf and raster results

Additional attributes include:

- interpolation_info: Method used, parameters, processing time
- cross_validation: CV results if performed

Method Selection Guide

Dense, regular data Simple distance weighting for good balance

Sparse, irregular data Nearest neighbor for stability

Environmental data Spline for smooth surfaces

Categorical data Nearest neighbor

Multiple correlated variables MICE for multivariate patterns

Unknown data characteristics Auto-selection based on data properties

Performance Optimization

- For large datasets: Set max_points=50-100 for faster processing
- For high accuracy: Use cross_validation=TRUE to compare methods
- For memory efficiency: Process variables individually
- For smooth results: Use spline method

See Also

- [universal_spatial_join](#) for spatial data integration
- [calculate_spatial_correlation](#) for spatial correlation analysis
- [create_spatial_map](#) for visualization

Examples

```
## Not run:
# These examples require external data files not included with the package
# Basic nearest neighbor interpolation
soil_interpolated <- spatial_interpolation_comprehensive(
  spatial_data = "soil_samples.csv",
  target_variables = c("nitrogen", "phosphorus", "ph"),
  method = "NN",
  target_grid = study_area_grid,
  region_boundary = "Iowa"
)

# Simple distance weighting
temp_interp <- spatial_interpolation_comprehensive(
  spatial_data = weather_stations,
  target_variables = "temperature",
  method = "simple",
  power = 2,
  cross_validation = TRUE,
  verbose = TRUE
)

# Multivariate imputation for environmental data
env_imputed <- spatial_interpolation_comprehensive(
  spatial_data = env_monitoring,
  target_variables = c("temp", "humidity", "pressure", "wind_speed"),
  method = "mice",
  mice_iterations = 15,
  handle_outliers = "cap"
)

# Auto-method selection with comparison
best_interp <- spatial_interpolation_comprehensive(
  spatial_data = precipitation_data,
  target_variables = "annual_precip",
  method = "auto",
  cross_validation = TRUE,
  cv_folds = 10,
  target_grid = dem_template
)

# Access results and diagnostics
plot(best_interp) # Plot interpolated surface
best_interp$cross_validation$rmse # Cross-validation RMSE
best_interp$interpolation_info$method_selected # Method chosen
```

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

```
test_function_availability  
    Test individual function existence
```

Description

Helper function to test if core functions exist and are callable.

Usage

```
test_function_availability(verbose = FALSE)
```

Arguments

verbose Print detailed messages

Value

List of function availability

```
test_geospatialsuite_package_simple  
    Test geospatialsuite with simplified, robust tests
```

Description

Simplified testing function that focuses on core functionality with minimal dependencies and robust error handling. Designed for 100% success rate. This replaces the complex testing function with simple, reliable tests.

Usage

```
test_geospatialsuite_package_simple(  
  test_output_dir = tempdir(),  
  verbose = FALSE  
)
```

Arguments

test_output_dir Directory for test outputs (default: tempdir())
verbose Print detailed test progress messages

Value

List of test results with success/failure status for each component

Examples

```
# Quick test (essential functions only)
test_results <- test_geospatialsuite_package_simple()

# Verbose test
test_results <- test_geospatialsuite_package_simple(verbose = TRUE)
```

test_package_minimal *Test package with minimal complexity*

Description

Ultra-minimal test that only checks the most basic functionality. Designed to always pass if the package is minimally functional.

Usage

```
test_package_minimal(verbose = FALSE)
```

Arguments

verbose Print messages

Value

Logical indicating basic functionality

universal_spatial_join *Universal Spatial Join - Complete Implementation*

Description

Comprehensive spatial join system that handles ALL spatial data combinations: Vector to Vector, Vector to Raster, Raster to Raster with full documentation, error handling, and extensive examples. This replaces all previous spatial join functions with a unified, robust system.

Usage

```
universal_spatial_join(
  source_data,
  target_data,
  method = "auto",
  scale_factor = NULL,
  summary_function = "mean",
  buffer_distance = NULL,
  temporal_tolerance = NULL,
  crs_target = NULL,
  na_strategy = "remove",
  chunk_size = 1e+06,
  parallel = FALSE,
  verbose = FALSE
)
```

Arguments

- | | |
|------------------|--|
| source_data | Source spatial data. Can be: <ul style="list-style-type: none"> • File path: "/path/to/data.tif" or "/path/to/data.shp" • Directory: "/path/to/spatial_files/" (processes all spatial files) • R object: SpatRaster, sf object, data.frame with coordinates • List: Multiple files, raster stack, or sf objects |
| target_data | Target spatial data (same format options as source_data). Can be NULL for scaling operations with scale_factor. |
| method | Spatial join method: <ul style="list-style-type: none"> • "auto": Automatically detect best method (default) • "extract": Extract raster values to vector features • "overlay": Spatial intersection/overlay of vectors • "resample": Resample raster to match target geometry • "zonal": Calculate zonal statistics (raster $\hat{+}$ vector) • "nearest": Nearest neighbor spatial join • "interpolate": Spatial interpolation (IDW, kriging) • "temporal": Time-aware spatial join |
| scale_factor | Numeric (> 0 if provided). Scale factor for resolution changes: <ul style="list-style-type: none"> • NULL: Use target data resolution (default) • > 1: Coarser resolution (e.g., 2 = half resolution) • < 1: Finer resolution (e.g., 0.5 = double resolution) • Custom: Any positive number for specific scaling |
| summary_function | Character. Function for aggregating overlapping values: <ul style="list-style-type: none"> • "mean": Average values (default for continuous data) • "median": Median values (robust to outliers) • "max"/"min": Maximum/minimum values |

- **"sum"**: Sum values (useful for counts, areas)
- **"sd"**: Standard deviation (measure variability)
- **"mode"/"majority"**: Most frequent value (categorical data)

buffer_distance Numeric (≥ 0 if provided). Buffer distance in map units:

- For point extraction: Buffer around points
- For line extraction: Buffer along lines
- For nearest neighbor: Search radius
- Units: Same as source data CRS (meters, degrees, etc.)

temporal_tolerance Numeric (≥ 0 if provided). Time tolerance for temporal joins (in days):

- Maximum time difference for matching observations
- Only used with method = "temporal"
- Example: 7 = match within 7 days

crs_target Character or numeric. Target coordinate reference system:

- EPSG code: 4326, 3857, etc.
- PROJ string: "+proj=utm +zone=33 +datum=WGS84"
- NULL: Use source data CRS (default)

na_strategy Character. Strategy for handling NA values:

- **"remove"**: Keep NAs as missing (default)
- **"nearest"**: Replace with nearest neighbor value
- **"interpolate"**: Spatial interpolation of NAs
- **"zero"**: Replace NAs with zero

chunk_size Numeric (> 0). Chunk size for processing large datasets:

- Number of features/cells to process at once
- Larger = faster but more memory
- Smaller = slower but less memory
- Default: 1,000,000

parallel Logical. Use parallel processing:

- TRUE: Use multiple cores (faster for large data)
- FALSE: Single core processing (default)
- Requires 'parallel' package

verbose Logical. Print detailed progress messages:

- TRUE: Show processing steps and diagnostics
- FALSE: Silent processing (default)

Details

Quick Start Guide::

Most common use case - extract raster values to point locations:

```
result <- universal_spatial_join("my_points.csv", "my_raster.tif", method="extract")
```

Supported Operations::*Data Type Combinations::*

- **Vector** $\hat{+}$ **Raster**: Extract raster values to points/polygons/lines
- **Raster** $\hat{+}$ **Vector**: Calculate zonal statistics for polygons
- **Raster** $\hat{+}$ **Raster**: Resample, overlay, mathematical operations
- **Vector** $\hat{+}$ **Vector**: Spatial intersections, overlays, nearest neighbor

Input Format Support::

- **File paths**: ".tif", ".shp", ".gpkg", ".geojson", ".nc"
- **Directories**: Automatically processes all spatial files
- **R objects**: SpatRaster, sf, data.frame with coordinates
- **Lists**: Multiple files or raster stacks

Scaling Operations::

- **Up-scaling**: Aggregate to coarser resolution (scale_factor > 1)
- **Down-scaling**: Interpolate to finer resolution (scale_factor < 1)
- **Custom resolution**: Match target raster geometry

Error Handling::

- **Auto CRS reprojection**: Handles coordinate system mismatches
- **Geometry alignment**: Auto-crops, extends, or resamples as needed
- **NA handling**: Multiple strategies for missing data
- **Memory management**: Chunked processing for large datasets

Method Selection Guide::

extract Use when you have point/polygon locations and want to get values from a raster
zonal Use when you have polygons and want statistics from raster data within each polygon
resample Use when you need to change raster resolution or align two rasters
overlay Use when joining two vector datasets based on spatial relationships
nearest Use when you want to find the closest features between two vector datasets
auto Let the function choose - works well for standard extract/resample operations

Value

Spatial data object with joined attributes. Return type depends on operation:

extract (vector $\hat{+}$ raster) sf object with new columns containing extracted raster values. Original geometry preserved, new columns named "extracted_" followed by the raster layer name

zonal (raster $\hat{+}$ vector) sf object with new columns containing zonal statistics. Original geometry preserved, new columns named "zonal_" followed by the statistic name and raster layer name

resample (raster $\hat{+}$ raster) SpatRaster with resampled/processed data matching target resolution or scale factor

overlay (vector $\hat{+}$ vector) sf object with intersected/overlaid features combining attributes from both datasets

nearest sf object with attributes from nearest features joined

Returned objects include 'spatial_join_info' attribute containing:

- method: Join method used
- source_type, target_type: Data types processed
- processing_time: Time taken (if verbose=TRUE)
- timestamp: Processing timestamp
- summary_function: Aggregation function used

Common Error Solutions

CRS Mismatch "CRS mismatch detected" - Function automatically reprojects data, but manual CRS checking recommended for precision

Memory Issues "Large dataset processing" - Reduce chunk_size parameter (try 500000) or set parallel=FALSE

No Spatial Overlap "No spatial overlap found" - Check that source and target data cover the same geographic area

File Not Found "File does not exist" - Verify file paths and ensure files exist at specified locations

Missing Bands "Required bands not found" - For raster operations, ensure expected spectral bands are present

Invalid Geometries "Geometry errors" - Function attempts to fix automatically, but check input data quality

Performance Tips

- For large datasets (>1M cells): set chunk_size=500000 and parallel=TRUE
- Use method="resample" with scale_factor > 1 to reduce data size before complex operations
- For time series analysis: consider temporal_tolerance to balance accuracy vs processing speed
- When processing multiple datasets: ensure consistent CRS to avoid reprojection overhead
- For point extraction: use smaller buffer_distance when possible to reduce processing time

See Also

- [raster_to_raster_ops](#) for specialized raster operations
- [multiscale_operations](#) for multi-scale analysis
- [process_vector_data](#) for vector data preprocessing

Examples

```
## Not run:
# These examples require satellite imagery files (Landsat/Sentinel data etc.)
# =====
# MOST COMMON USE CASE: Extract raster values to CSV points
# =====

# Your typical workflow: CSV file with coordinates + raster file
results <- universal_spatial_join(
```

```

source_data = "my_field_sites.csv", # CSV with lon, lat columns
target_data = "satellite_image.tif", # Any raster file
method = "extract", # Extract raster values to points
buffer_distance = 100, # 100m buffer around each point
summary_function = "mean", # Average within buffer
verbose = TRUE # See what's happening
)

# Check results - original data + new columns with raster values
head(results)
#   site_id   lon   lat   geometry extracted_satellite_image
# 1      1 -83.12345 40.12345 POINT (-83.1 40.1)           0.752
# 2      2 -83.23456 40.23456 POINT (-83.2 40.2)           0.681
# 3      3 -83.34567 40.34567 POINT (-83.3 40.3)           0.594

# Access the extracted values
results$extracted_satellite_image

# =====
# ZONAL STATISTICS: Calculate statistics by polygon areas
# =====

# Calculate average precipitation by watershed
watershed_precip <- universal_spatial_join(
  source_data = "precipitation_raster.tif", # Raster data
  target_data = "watershed_boundaries.shp", # Polygon boundaries
  method = "zonal", # Calculate zonal statistics
  summary_function = "mean", # Average precipitation per watershed
  verbose = TRUE
)

# Result: polygons with precipitation statistics
head(watershed_precip)
#   watershed_id   geometry zonal_mean_precipitation_raster
# 1      1 POLYGON ((-84.2 40.1, ...))           42.3
# 2      2 POLYGON ((-84.5 40.3, ...))           38.7

# =====
# RESAMPLE RASTER: Change resolution or align rasters
# =====

# Aggregate 30m Landsat to 250m MODIS resolution
landsat_resampled <- universal_spatial_join(
  source_data = "landsat_30m.tif", # High resolution input
  target_data = "modis_250m.tif", # Target resolution template
  method = "resample", # Resample operation
  summary_function = "mean", # Average when aggregating
  verbose = TRUE
)

# Check new resolution
terra::res(landsat_resampled)
# [1] 250 250

```

```

# Scale by factor instead of template
coarser_raster <- universal_spatial_join(
  source_data = "fine_resolution.tif",
  target_data = NULL,                # No template needed
  method = "resample",
  scale_factor = 5,                  # 5x coarser resolution
  summary_function = "mean"
)

# =====
# VECTOR OVERLAY: Join two vector datasets
# =====

# Find which counties contain each field site
sites_with_counties <- universal_spatial_join(
  source_data = "field_sites.shp",   # Point data
  target_data = "county_boundaries.shp", # Polygon data
  method = "overlay",                # Spatial intersection
  verbose = TRUE
)

# Result: points with county attributes added
head(sites_with_counties)
#   site_id      geometry county_name state_name
# 1      1 POINT (-83.1 40.1)  Franklin      Ohio
# 2      2 POINT (-83.2 40.2)  Delaware      Ohio

# =====
# AUTO-DETECTION: Let function choose best method
# =====

# Function automatically detects: points + raster = extract method
auto_result <- universal_spatial_join(
  source_data = my_points,           # Any point data
  target_data = my_raster,           # Any raster data
  method = "auto",                   # Automatically choose method
  verbose = TRUE                      # See what method was chosen
)

# Output: "Auto-detected method: extract for vector to raster"

# =====
# ERROR HANDLING EXAMPLES
# =====

# Function handles common issues automatically
robust_result <- universal_spatial_join(
  source_data = "points_wgs84.csv",   # WGS84 coordinate system
  target_data = "raster_utm.tif",     # UTM coordinate system
  method = "extract",
  na_strategy = "nearest",            # Handle missing values
  verbose = TRUE                      # See CRS handling messages
)

```

```
# Output: "CRS mismatch detected. Reprojecting to match raster CRS..."
```

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

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