

# Package ‘SVG’

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**Type** Package

**Title** Spatially Variable Genes Detection Methods for Spatial Transcriptomics

**Version** 1.0.0

**Description** A unified framework for detecting spatially variable genes (SVGs) in spatial transcriptomics data. This package integrates multiple state-of-the-art SVG detection methods including 'MERINGUE' (Moran's I based spatial autocorrelation), 'Giotto' binSpect (binary spatial enrichment test), 'SPARK-X' (non-parametric kernel-based test), and 'nnSVG' (nearest-neighbor Gaussian processes). Each method is implemented with optimized performance through vectorization, parallelization, and 'C++' acceleration where applicable. Methods are described in Miller et al. (2021) <doi:10.1101/gr.271288.120>, Dries et al. (2021) <doi:10.1186/s13059-021-02286-2>, Zhu et al. (2021) <doi:10.1186/s13059-021-02404-0>, and Weber et al. (2023) <doi:10.1038/s41467-023-39748-z>.

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ACAT_combine	<i>ACAT: Aggregated Cauchy Association Test</i>
--------------	---

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## Description

Combines multiple p-values using the Aggregated Cauchy Association Test (ACAT). This method is robust and maintains correct type I error even with correlated p-values.

## Usage

```
ACAT_combine(pvals, weights = NULL)
```

## Arguments

pvals	Numeric vector of p-values to combine.
weights	Numeric vector of weights. If NULL (default), equal weights are used.

**Details**

ACAT transforms p-values using the Cauchy distribution and combines them:

$$T = \sum_i w_i \tan(\pi(0.5 - p_i))$$

The combined p-value is then computed from the Cauchy distribution.

This method has several advantages:

- Valid even when p-values are correlated
- Computationally simple
- Handles edge cases (p = 0 or 1) gracefully

**Value**

A single combined p-value.

**References**

Liu, Y. et al. (2019) ACAT: A Fast and Powerful P Value Combination Method for Rare-Variant Analysis in Sequencing Studies. *The American Journal of Human Genetics*.

**Examples**

```
# Combine independent p-values
pvals <- c(0.01, 0.05, 0.3)
combined_p <- ACAT_combine(pvals)
print(combined_p)
```

---

binarize\_expression    *Binarize Gene Expression*

---

**Description**

Converts continuous gene expression values to binary (0/1) using various methods. Used by the binSpect method.

**Usage**

```
binarize_expression(  
  expr_matrix,  
  method = c("kmeans", "rank", "median", "mean"),  
  rank_percent = 30,  
  n_threads = 1L,  
  verbose = FALSE  
)
```

**Arguments**

expr_matrix	Numeric matrix of gene expression. Rows are genes, columns are spots/cells.
method	Character string specifying binarization method. <ul style="list-style-type: none"> <li>• "kmeans" (default): Use k-means clustering (k=2) to separate high and low expression</li> <li>• "rank": Binarize based on expression rank percentile</li> <li>• "median": Values above median are set to 1</li> <li>• "mean": Values above mean are set to 1</li> </ul>
rank_percent	Numeric. For method = "rank", the percentile threshold (0-100). Values in the top rank_percent percent are set to 1. Default is 30.
n_threads	Integer. Number of threads for parallel computation. Default is 1.
verbose	Logical. Whether to print progress. Default is FALSE.

**Details**

**K-means method:** For each gene, k-means clustering with k=2 is applied. The cluster with higher mean expression is labeled as 1, the other as 0.

**Rank method:** For each gene, spots are ranked by expression. The top rank\_percent percent are labeled as 1.

**Value**

Binary matrix with same dimensions as input.

**Examples**

```
# Create example expression matrix
expr <- matrix(rpois(1000, lambda = 10), nrow = 10, ncol = 100)
rownames(expr) <- paste0("gene_", 1:10)

# Binarize using k-means
bin_kmeans <- binarize_expression(expr, method = "kmeans")

# Binarize using rank (top 20%)
bin_rank <- binarize_expression(expr, method = "rank", rank_percent = 20)
```

---

buildSpatialNetwork    *Build Spatial Neighborhood Network*

---

**Description**

Constructs a spatial neighborhood network from spatial coordinates using either Delaunay triangulation or K-nearest neighbors (KNN) approach.

**Usage**

```
buildSpatialNetwork(
  coords,
  method = c("delaunay", "knn"),
  k = 10L,
  filter_dist = NA,
  binary = TRUE,
  verbose = FALSE
)
```

**Arguments**

coords	Numeric matrix of spatial coordinates. Rows are spatial locations, columns are coordinate dimensions (typically x, y).
method	Character string specifying the network construction method. <ul style="list-style-type: none"> <li>"delaunay": Delaunay triangulation (default). Creates a network where neighbors are determined by triangulation. Works well for relatively uniform spatial distributions.</li> <li>"knn": K-nearest neighbors. Each spot is connected to its k nearest neighbors based on Euclidean distance.</li> </ul>
k	Integer. Number of nearest neighbors for KNN method. Default is 10. Ignored when method = "delaunay".
filter_dist	Numeric or NA. Maximum distance threshold for neighbors. Pairs with distance > filter_dist are not considered neighbors. Default is NA (no filtering).
binary	Logical. If TRUE (default), return binary adjacency matrix (0/1). If FALSE, return distance-weighted adjacency matrix.
verbose	Logical. Whether to print progress messages. Default is FALSE.

**Details**

**Delaunay Triangulation:** Creates a network based on Delaunay triangulation, which maximizes the minimum angle of all triangles. This is a natural way to define neighbors in 2D/3D space. Requires the `geometry` package.

**K-Nearest Neighbors:** Connects each point to its k nearest neighbors based on Euclidean distance. More robust to irregular spatial distributions but requires choosing k. Requires the `RANN` package.

**Value**

A square numeric matrix representing the spatial adjacency/weight matrix. Row and column names correspond to the spatial locations (from rownames of `coords`).

- If `binary = TRUE`: Values are 1 (neighbors) or 0 (non-neighbors)
- If `binary = FALSE`: Values are Euclidean distances (0 for non-neighbors)

**See Also**

[getSpatialNeighbors\\_Delaunay](#), [getSpatialNeighbors\\_KNN](#)

## Examples

```
# Generate example coordinates
set.seed(42)
coords <- cbind(x = runif(100), y = runif(100))
rownames(coords) <- paste0("spot_", 1:100)

# Build network using Delaunay (requires geometry package)
if (requireNamespace("geometry", quietly = TRUE)) {
  W_delaunay <- buildSpatialNetwork(coords, method = "delaunay")
}

# Build network using KNN (requires RANN package)
if (requireNamespace("RANN", quietly = TRUE)) {
  W_knn <- buildSpatialNetwork(coords, method = "knn", k = 6)
}
```

---

CalSVG

*Unified Interface for SVG Detection*

---

## Description

A unified interface to run different spatially variable gene (SVG) detection methods. This function provides a consistent API for all supported methods.

## Usage

```
CalSVG(
  expr_matrix,
  spatial_coords,
  method = c("meringue", "seurat", "binspect", "sparkx", "nnsvg", "markvario"),
  n_threads = 1L,
  verbose = TRUE,
  ...
)
```

## Arguments

<code>expr_matrix</code>	Numeric matrix of gene expression values. Rows are genes, columns are spatial locations (spots/cells). Should be normalized (e.g., log-transformed counts).
<code>spatial_coords</code>	Numeric matrix of spatial coordinates. Rows are spatial locations, columns are x and y (and optionally z) coordinates. Row names should match column names of <code>expr_matrix</code> .
<code>method</code>	Character string specifying the SVG detection method. One of: "meringue", "seurat", "binspect", "sparkx", "nnsvg", "markvario".

n_threads	Integer. Number of threads for parallel computation. Default is 1. Set to higher values for faster computation on multi-core systems.
verbose	Logical. Whether to print progress messages. Default is TRUE.
...	Additional arguments passed to the specific method function.

### Details

This function serves as a wrapper around the individual method functions:

- method = "meringue": Calls [CalSVG\\_MERINGUE](#)
- method = "seurat": Calls [CalSVG\\_Seurat](#)
- method = "binspect": Calls [CalSVG\\_binSpect](#)
- method = "sparkx": Calls [CalSVG\\_SPARKX](#)
- method = "nnsvg": Calls [CalSVG\\_nnSVG](#)
- method = "markvario": Calls [CalSVG\\_MarkVario](#)

For method-specific parameters, please refer to the documentation of individual method functions.

### Value

A data.frame containing SVG detection results. The exact columns depend on the method used, but typically include:

- gene: Gene identifiers
- pval or p.value: Raw p-values
- padj or p.adj: Adjusted p-values (multiple testing corrected)
- Method-specific statistics (e.g., Moran's I, LR statistic, odds ratio)

### See Also

[CalSVG\\_MERINGUE](#), [CalSVG\\_binSpect](#), [CalSVG\\_SPARKX](#), [CalSVG\\_nnSVG](#)

### Examples

```
# Simulate example data
set.seed(42)
n_genes <- 20
n_spots <- 100
expr_matrix <- matrix(rpois(n_genes * n_spots, lambda = 10),
                     nrow = n_genes, ncol = n_spots)
rownames(expr_matrix) <- paste0("gene_", seq_len(n_genes))
colnames(expr_matrix) <- paste0("spot_", seq_len(n_spots))

spatial_coords <- cbind(x = runif(n_spots, 0, 100),
                      y = runif(n_spots, 0, 100))
rownames(spatial_coords) <- colnames(expr_matrix)

# Run SPARK-X method (no external dependencies)
results <- CalSVG(expr_matrix, spatial_coords, method = "sparkx",
```

```

kernel_option = "single", verbose = FALSE)
head(results)

```

---

CalSVG\_binSpect

*binSpect: Binary Spatial Enrichment Test for SVG Detection*

---

## Description

Detect spatially variable genes using the binSpect approach from Giotto. This method binarizes gene expression and tests for spatial enrichment of high-expressing cells using Fisher's exact test.

Identifies spatially variable genes by: 1. Binarizing gene expression (high/low) 2. Building a spatial neighborhood network 3. Testing whether high-expressing cells tend to be neighbors of other high-expressing cells more than expected by chance

## Usage

```

CalSVG_binSpect(
  expr_matrix,
  spatial_coords,
  bin_method = c("kmeans", "rank"),
  rank_percent = 30,
  network_method = c("deLaunay", "knn"),
  k = 10L,
  do_fisher_test = TRUE,
  adjust_method = "fdr",
  n_threads = 1L,
  verbose = TRUE
)

```

## Arguments

- |                |   |
|----------------|---|
| expr_matrix    | Numeric matrix of gene expression values. <ul style="list-style-type: none"> <li>• Rows: genes</li> <li>• Columns: spatial locations (spots/cells)</li> <li>• Values: normalized expression (e.g., log counts or normalized counts)</li> </ul>              |
| spatial_coords | Numeric matrix of spatial coordinates. <ul style="list-style-type: none"> <li>• Rows: spatial locations (must match columns of expr_matrix)</li> <li>• Columns: x, y (and optionally z) coordinates</li> </ul>  |
| bin_method     | Character string specifying binarization method. <ul style="list-style-type: none"> <li>• "kmeans" (default): K-means clustering with k=2. Automatically separates high and low expression groups. Robust to different expression distributions.</li> </ul> |

	<ul style="list-style-type: none"> <li>• "rank": Top percentage by expression rank. More consistent across genes with different distributions. Controlled by rank_percent parameter.</li> </ul>
rank_percent	Numeric (0-100). For bin_method = "rank", the percentage of cells to classify as "high expressing". Default is 30 (top 30) <ul style="list-style-type: none"> <li>• Lower values (10-20)</li> <li>• Higher values (40-50)</li> </ul>
network_method	Character string specifying spatial network construction. <ul style="list-style-type: none"> <li>• "deLaunay" (default): Delaunay triangulation</li> <li>• "knn": K-nearest neighbors</li> </ul>
k	Integer. Number of neighbors for KNN network. Default is 10.
do_fisher_test	Logical. Whether to perform Fisher's exact test. Default is TRUE. <ul style="list-style-type: none"> <li>• TRUE: Returns p-values from Fisher's exact test</li> <li>• FALSE: Returns only odds ratios (faster)</li> </ul>
adjust_method	Character string for p-value adjustment. Default is "fdr" (Benjamini-Hochberg). See p.adjust() for options.
n_threads	Integer. Number of parallel threads. Default is 1.
verbose	Logical. Print progress messages. Default is TRUE.

## Details

### Method Overview:

binSpect constructs a 2x2 contingency table for each gene based on:

- Cell A expression: High (1) or Low (0)
- Cell B expression: High (1) or Low (0)

For all pairs of neighboring cells (edges in the spatial network):

	Cell B Low	Cell B High
Cell A Low	n_00	n_01
Cell A High	n_10	n_11

**Statistical Test:** Fisher's exact test is used to test whether n\_11 (both neighbors high) is greater than expected under independence.

### Odds Ratio Interpretation:

- OR = 1: No spatial pattern
- OR > 1: High-expressing cells cluster together (positive spatial pattern)
- OR < 1: High-expressing cells avoid each other (negative pattern)

### Advantages:

- Fast computation (no covariance matrix inversion)
- Robust to outliers through binarization



---

CalSVG\_MarkVario      *Detect SVGs using Mark Variogram Method*

---

## Description

Identifies spatially variable genes using the mark variogram approach, as implemented in Seurat's `FindSpatiallyVariableFeatures` function with `selection.method = "markvariogram"`.

## Usage

```
CalSVG_MarkVario(  
  expr_matrix,  
  spatial_coords,  
  r_metric = 5,  
  normalize = TRUE,  
  n_threads = 1L,  
  verbose = TRUE  
)
```

## Arguments

<code>expr_matrix</code>	Numeric matrix of gene expression values.
<code>spatial_coords</code>	Numeric matrix of spatial coordinates.
<code>r_metric</code>	Numeric. Distance at which to evaluate the variogram. Default is 5. Larger values capture broader spatial patterns.
<code>normalize</code>	Logical. Whether to normalize the variogram. Default is TRUE.
<code>n_threads</code>	Integer. Number of parallel threads. Default is 1.
<code>verbose</code>	Logical. Print progress messages. Default is TRUE.

## Details

### Method Overview:

The mark variogram measures how the correlation between gene expression values changes with distance. It is computed using the `spatstat` package's `markvario` function.

### Interpretation:

- Lower variogram values indicate stronger spatial autocorrelation
- Values near 1 indicate random spatial distribution
- Values < 1 indicate positive spatial autocorrelation (clustering)

**Note:** Requires the `spatstat` package suite to be installed: `spatstat.geom` and `spatstat.explore`.

**Value**

A data.frame with SVG detection results. Columns:

- gene: Gene identifier
- r.metric.X: Variogram value at distance r\_metric
- rank: Rank by variogram value (ascending, lower = more spatially variable)

**References**

Baddeley, A. et al. (2015) Spatial Point Patterns: Methodology and Applications with R. Chapman and Hall/CRC.

**See Also**

[CaSVG\\_Seurat](#)

**Examples**

```
# Load example data
data(example_svg_data)
expr <- example_svg_data$logcounts[1:5, ]
coords <- example_svg_data$spatial_coords

# Requires spatstat packages
if (requireNamespace("spatstat.geom", quietly = TRUE) &&
    requireNamespace("spatstat.explore", quietly = TRUE)) {
  results <- CaSVG_MarkVario(expr, coords, verbose = FALSE)
  head(results)
}
```

---

CaSVG\_MERINGUE

*MERINGUE: Moran's I based Spatially Variable Gene Detection*

---

**Description**

Detect spatially variable genes using the MERINGUE approach based on Moran's I spatial autocorrelation statistic.

Identifies spatially variable genes by computing Moran's I spatial autocorrelation statistic for each gene. Genes with significant positive spatial autocorrelation (similar expression values clustering together) are identified as SVGs.

**Usage**

```
CalSVG_MERINGUE(
  expr_matrix,
  spatial_coords,
  network_method = c("delauanay", "knn"),
  k = 10L,
  filter_dist = NA,
  alternative = c("greater", "less", "two.sided"),
  adjust_method = "BH",
  min_pct_cells = 0.05,
  n_threads = 1L,
  use_cpp = TRUE,
  verbose = TRUE
)
```

**Arguments**

- |                |   |
|----------------|---|
| expr_matrix    | <p>Numeric matrix of gene expression values.</p> <ul style="list-style-type: none"> <li>• Rows: genes</li> <li>• Columns: spatial locations (spots/cells)</li> <li>• Values: normalized expression (e.g., log-transformed counts)</li> </ul> <p>Row names should be gene identifiers; column names should match row names of spatial_coords.</p>  |
| spatial_coords | <p>Numeric matrix of spatial coordinates.</p> <ul style="list-style-type: none"> <li>• Rows: spatial locations (must match columns of expr_matrix)</li> <li>• Columns: coordinate dimensions (x, y, and optionally z)</li> </ul>  |
| network_method | <p>Character string specifying how to construct the spatial neighborhood network.</p> <ul style="list-style-type: none"> <li>• "delauanay" (default): Delaunay triangulation. Creates natural neighbors based on geometric triangulation. Good for relatively uniform spatial distributions.</li> <li>• "knn": K-nearest neighbors. Each spot connected to its k nearest neighbors. More robust for irregular distributions.</li> </ul> |
| k              | <p>Integer. Number of neighbors for KNN method. Default is 10. Ignored when network_method = "delauanay".</p> <ul style="list-style-type: none"> <li>• Smaller k (e.g., 5-6): More local patterns, faster computation</li> <li>• Larger k (e.g., 15-20): Broader patterns, smoother results</li> </ul>  |
| filter_dist    | <p>Numeric or NA. Maximum Euclidean distance for neighbors. Pairs with distance &gt; filter_dist are not considered neighbors. Default is NA (no filtering). Useful for:</p> <ul style="list-style-type: none"> <li>• Removing long-range spurious connections</li> <li>• Focusing on local spatial patterns</li> </ul>   |
| alternative    | <p>Character string specifying the alternative hypothesis for the Moran's I test.</p> <ul style="list-style-type: none"> <li>• "greater" (default): Test for positive autocorrelation (clustering of similar values). Most appropriate for SVG detection.</li> </ul>  |

	<ul style="list-style-type: none"> <li>• "less": Test for negative autocorrelation (dissimilar values as neighbors).</li> <li>• "two.sided": Test for any autocorrelation.</li> </ul>
adjust_method	Character string specifying p-value adjustment method for multiple testing correction. Passed to <code>p.adjust()</code> . Options include: "BH" (default, Benjamini-Hochberg), "bonferroni", "holm", "hochberg", "hommel", "BY", "fdr", "none".
min_pct_cells	Numeric (0-1). Minimum fraction of cells that must contribute to the spatial pattern for a gene to be retained as SVG. Default is 0.05 (5 to filter genes driven by only a few outlier cells. Set to 0 to disable this filter.
n_threads	Integer. Number of threads for parallel computation. Default is 1. <ul style="list-style-type: none"> <li>• For large datasets: Set to number of available cores</li> <li>• Uses R's <code>parallel::mclapply</code> (not available on Windows)</li> </ul>
use_cpp	Logical. Whether to use C++ implementation for faster computation. Default is TRUE. Falls back to R if C++ fails.
verbose	Logical. Whether to print progress messages. Default is TRUE.

## Details

### Method Overview:

MERINGUE uses Moran's I, a classic measure of spatial autocorrelation:

$$I = \frac{n}{W} \frac{\sum_i \sum_j w_{ij} (x_i - \bar{x})(x_j - \bar{x})}{\sum_i (x_i - \bar{x})^2}$$

where:

- n = number of spatial locations
- W = sum of all spatial weights
- w<sub>ij</sub> = spatial weight between locations i and j
- x<sub>i</sub> = expression value at location i

### Interpretation:

- I > 0: Positive autocorrelation (similar values cluster)
- I = 0: Random spatial distribution
- I < 0: Negative autocorrelation (checkerboard pattern)

**Statistical Testing:** P-values are computed using normal approximation based on analytical formulas for the expected value and variance of Moran's I under the null hypothesis of complete spatial randomness.

### Computational Considerations:

- Time complexity: O(n<sup>2</sup>) for network construction, O(n\*m) for testing (n = spots, m = genes)
- Memory: O(n<sup>2</sup>) for storing spatial weights matrix
- For n > 10,000 spots, consider using KNN with small k

**Value**

A data.frame with SVG detection results, sorted by significance. Columns:

- gene: Gene identifier
- observed: Observed Moran's I statistic. Range: [-1, 1]. Positive values indicate clustering, negative indicate dispersion.
- expected: Expected Moran's I under null (approximately  $-1/(n-1)$ )
- sd: Standard deviation under null hypothesis
- z\_score: Standardized test statistic (observed - expected) / sd
- p.value: Raw p-value from normal approximation
- p.adj: Adjusted p-value (multiple testing corrected)

**References**

- Miller, B.F. et al. (2021) Characterizing spatial gene expression heterogeneity in spatially resolved single-cell transcriptomic data with nonuniform cellular densities. *Genome Research*.
- Moran, P.A.P. (1950) Notes on Continuous Stochastic Phenomena. *Biometrika*.
- Cliff, A.D. and Ord, J.K. (1981) *Spatial Processes: Models & Applications*. Pion.

**See Also**

[CalSVG](#) for unified interface, [buildSpatialNetwork](#) for network construction, [moranI\\_test](#) for individual gene testing

**Examples**

```
# Load example data
data(example_svg_data)
expr <- example_svg_data$logcounts[1:20, ] # Use subset for speed
coords <- example_svg_data$spatial_coords

# Basic usage (requires RANN package for KNN)
if (requireNamespace("RANN", quietly = TRUE)) {
  results <- CalSVG_MERINGUE(expr, coords,
                             network_method = "knn", k = 10,
                             verbose = FALSE)

  head(results)

  # Get significant SVGs
  sig_genes <- results$gene[results$p.adj < 0.05]
}
```

CaSVG\_nnSVG

*nnSVG: Nearest-Neighbor Gaussian Process SVG Detection***Description**

Detect spatially variable genes using nnSVG, a method based on nearest-neighbor Gaussian processes for scalable spatial modeling.

nnSVG uses nearest-neighbor Gaussian processes (NNGP) to model spatial correlation structure in gene expression. It performs likelihood ratio tests comparing spatial vs. non-spatial models to identify SVGs.

**Usage**

```
CaSVG_nnSVG(
  expr_matrix,
  spatial_coords,
  X = NULL,
  n_neighbors = 10L,
  order = c("AMMD", "Sum_coords"),
  cov_model = c("exponential", "gaussian", "spherical", "matern"),
  adjust_method = "BH",
  n_threads = 1L,
  verbose = FALSE
)
```

**Arguments**

<code>expr_matrix</code>	Numeric matrix of gene expression values. <ul style="list-style-type: none"> <li>• Rows: genes</li> <li>• Columns: spatial locations (spots/cells)</li> <li>• Values: log-normalized counts (e.g., from <code>scrani::logNormCounts</code>)</li> </ul>
<code>spatial_coords</code>	Numeric matrix of spatial coordinates. <ul style="list-style-type: none"> <li>• Rows: spatial locations (must match columns of <code>expr_matrix</code>)</li> <li>• Columns: x, y coordinates</li> </ul>
<code>X</code>	Optional numeric matrix of covariates to regress out. <ul style="list-style-type: none"> <li>• Rows: spatial locations (same order as <code>spatial_coords</code>)</li> <li>• Columns: covariates (e.g., batch, cell type indicators)</li> </ul> Default is NULL (intercept-only model).
<code>n_neighbors</code>	Integer. Number of nearest neighbors for NNGP model. Default is 10. <ul style="list-style-type: none"> <li>• 5-10: Faster, captures local patterns</li> <li>• 15-20: Better likelihood estimates, slower</li> </ul> Values > 15 rarely improve results but increase computation time.
<code>order</code>	Character string specifying coordinate ordering scheme.

	<ul style="list-style-type: none"> <li>• "AMMD" (default): Approximate Maximum Minimum Distance. Better for most datasets. Requires <math>\geq 65</math> spots.</li> <li>• "Sum_coords": Order by sum of coordinates. Use for very small datasets (<math>&lt; 65</math> spots).</li> </ul>
cov_model	Character string specifying the covariance function. Default is "exponential". <ul style="list-style-type: none"> <li>• "exponential": Most commonly used, computationally stable</li> <li>• "gaussian": Smoother patterns, requires stabilization</li> <li>• "spherical": Finite range correlation</li> <li>• "matern": Flexible smoothness (includes additional nu parameter)</li> </ul>
adjust_method	Character string for p-value adjustment. Default is "BH" (Benjamini-Hochberg).
n_threads	Integer. Number of parallel threads. Default is 1. Set to number of available cores for faster computation.
verbose	Logical. Print progress messages. Default is FALSE.

## Details

### Method Overview:

nnSVG models gene expression as a Gaussian process:

$$y = X\beta + \omega + \epsilon$$

where:

- $y$  = expression vector
- $X$  = covariate matrix,  $\beta$  = coefficients
- $\omega \sim \text{GP}(0, \sigma^2 * C(\phi))$  = spatial random effect
- $\epsilon \sim N(0, \tau^2)$  = non-spatial noise
- $C(\phi)$  = covariance function with range  $\phi$

**Nearest-Neighbor Approximation:** Full GP has  $O(n^3)$  complexity. NNGP approximates using only  $k$  nearest neighbors, reducing complexity to  $O(n * k^3) = O(n)$ .

**Statistical Test:** Likelihood ratio test comparing:

- $H_0$  (null):  $y = X\beta + \epsilon$  (no spatial effect)
- $H_1$  (alternative):  $y = X\beta + \omega + \epsilon$  (with spatial effect)

LR statistic follows chi-squared with  $df = 2$  (testing  $\sigma^2$  and  $\phi$ ).

**Effect Size:** Proportion of spatial variance ( $\text{prop}_{sv}$ ) measures effect size:

- $\text{prop}_{sv}$  near 1: Strong spatial pattern
- $\text{prop}_{sv}$  near 0: Little spatial structure

### Computational Notes:

- Requires BRISC package for NNGP fitting
- $O(n)$  complexity per gene with NNGP approximation
- Parallelization over genes provides good speedup
- Memory:  $O(n * k)$  per gene

**Value**

A data.frame with SVG detection results. Columns:

- gene: Gene identifier
- sigma.sq: Spatial variance estimate ( $\sigma^2$ )
- tau.sq: Nonspatial variance estimate ( $\tau^2$ , nugget)
- phi: Range parameter estimate (controls spatial correlation decay)
- prop\_sv: Proportion of spatial variance =  $\sigma.sq / (\sigma.sq + \tau.sq)$
- loglik: Log-likelihood of spatial model
- loglik\_lm: Log-likelihood of non-spatial model (linear model)
- LR\_stat: Likelihood ratio test statistic =  $-2 * (\loglik\_lm - \loglik)$
- rank: Rank by LR statistic (1 = highest)
- p.value: P-value from chi-squared distribution (df = 2)
- p.adj: Adjusted p-value
- runtime: Computation time per gene (seconds)

**References**

Weber, L.M. et al. (2023) nnSVG for the scalable identification of spatially variable genes using nearest-neighbor Gaussian processes. *Nature Communications*.

Datta, A. et al. (2016) Hierarchical Nearest-Neighbor Gaussian Process Models for Large Geostatistical Datasets. *JASA*.

**See Also**

[CalSVG](#), BRISC package documentation

**Examples**

```
# Load example data
data(example_svg_data)
expr <- example_svg_data$logcounts[1:10, ] # Small subset
coords <- example_svg_data$spatial_coords

# Basic usage (requires BRISC package)
if (requireNamespace("BRISC", quietly = TRUE)) {
  results <- CalSVG_nnSVG(expr, coords, verbose = FALSE)
  head(results)
}
```

---

 CalSVG\_Seurat

*Seurat-style SVG Detection Methods*


---

## Description

Detect spatially variable genes using methods implemented in Seurat, including Moran's I with inverse distance weights and Mark Variogram.

Identifies spatially variable genes using Moran's I statistic with inverse distance squared weighting, as implemented in Seurat's `FindSpatiallyVariableFeatures` function.

## Usage

```
CalSVG_Seurat(
  expr_matrix,
  spatial_coords,
  weight_scheme = c("inverse_squared", "inverse", "gaussian"),
  bandwidth = NULL,
  adjust_method = "BH",
  n_threads = 1L,
  verbose = TRUE
)
```

## Arguments

<code>expr_matrix</code>	Numeric matrix of gene expression values. <ul style="list-style-type: none"> <li>• Rows: genes</li> <li>• Columns: spatial locations (spots/cells)</li> <li>• Values: scaled/normalized expression (Seurat typically uses <code>scale.data</code>)</li> </ul>
<code>spatial_coords</code>	Numeric matrix of spatial coordinates. <ul style="list-style-type: none"> <li>• Rows: spatial locations (must match columns of <code>expr_matrix</code>)</li> <li>• Columns: x, y coordinates</li> </ul>
<code>weight_scheme</code>	Character string specifying the distance-based weighting. <ul style="list-style-type: none"> <li>• "inverse_squared" (default): <math>w_{ij} = 1 / d_{ij}^2</math> (Seurat default, emphasizes local neighbors)</li> <li>• "inverse": <math>w_{ij} = 1 / d_{ij}</math> (less emphasis on close neighbors)</li> <li>• "gaussian": <math>w_{ij} = \exp(-d_{ij}^2 / (2 * bandwidth^2))</math> (controlled by bandwidth parameter)</li> </ul>
<code>bandwidth</code>	Numeric. Bandwidth for Gaussian weighting. Default is NULL (auto-computed as median pairwise distance). Only used when <code>weight_scheme = "gaussian"</code> .
<code>adjust_method</code>	Character string for p-value adjustment. Default is "BH" (Benjamini-Hochberg).
<code>n_threads</code>	Integer. Number of parallel threads. Default is 1.
<code>verbose</code>	Logical. Print progress messages. Default is TRUE.

## Details

### Method Overview:

This function replicates Seurat's `FindSpatiallyVariableFeatures` with `selection.method = "moransi"`. The key difference from other Moran's I implementations is the weighting scheme:

$$w_{ij} = \frac{1}{d_{ij}^2}$$

where `d_ij` is the Euclidean distance between locations `i` and `j`.

### Interpretation:

- Uses continuous distance-based weights (not binary network)
- Emphasizes local spatial relationships
- Higher weights for closer neighbors

### Comparison with MERINGUE:

- MERINGUE: Binary adjacency (neighbors = 1, others = 0)
- Seurat: Continuous weights ( $1/\text{distance}^2$ )
- Seurat method is more sensitive to local patterns

## Value

A data.frame with SVG detection results. Columns:

- `gene`: Gene identifier
- `observed`: Observed Moran's I statistic
- `expected`: Expected Moran's I under null
- `sd`: Standard deviation under null
- `p.value`: Raw p-value
- `p.adj`: Adjusted p-value
- `rank`: Rank by p-value (ascending)

## References

- Hao, Y. et al. (2021) Integrated analysis of multimodal single-cell data. *Cell*.  
Stuart, T. et al. (2019) Comprehensive Integration of Single-Cell Data. *Cell*.

## See Also

[CalSVG](#), [CalSVG\\_MERINGUE](#)

**Examples**

```
# Load example data
data(example_svg_data)
expr <- example_svg_data$logcounts[1:20, ]
coords <- example_svg_data$spatial_coords

# Basic usage
results <- CalSVG_Seurat(expr, coords, verbose = FALSE)
head(results)
```

---

CalSVG\_SPARKX

*SPARK-X: Non-parametric Kernel-based SVG Detection*


---

**Description**

Detect spatially variable genes using SPARK-X, a non-parametric method that tests for spatial expression patterns using multiple kernels.

SPARK-X is a scalable non-parametric method for identifying spatially variable genes. It uses variance component score tests with multiple spatial kernels (projection, Gaussian, and cosine) to detect various types of spatial expression patterns.

**Usage**

```
CalSVG_SPARKX(
  expr_matrix,
  spatial_coords,
  kernel_option = c("mixture", "single"),
  adjust_method = "BY",
  n_threads = 1L,
  verbose = TRUE
)
```

**Arguments**

- `expr_matrix` Numeric matrix of gene expression values.
- Rows: genes
  - Columns: spatial locations (spots/cells)
  - Values: raw counts or normalized counts (NOT log-transformed)
- Note: SPARK-X works best with count data, not log-transformed data.
- `spatial_coords` Numeric matrix of spatial coordinates.
- Rows: spatial locations (must match columns of `expr_matrix`)
  - Columns: x, y coordinates

kernel_option	Character string specifying which kernels to use. <ul style="list-style-type: none"> <li>"mixture" (default): Test with all 11 kernels: 1 projection + 5 Gaussian + 5 cosine. Most comprehensive but slower. Recommended for detecting diverse spatial patterns.</li> <li>"single": Test with projection kernel only. Faster but may miss some pattern types.</li> </ul>
adjust_method	Character string for p-value adjustment. Default is "BY" (Benjamini-Yekutieli), which is more conservative and appropriate when tests may be correlated. Other options: "BH", "bonferroni", "holm", "none".
n_threads	Integer. Number of parallel threads. Default is 1. Higher values significantly speed up computation for large datasets.
verbose	Logical. Print progress messages. Default is TRUE.

## Details

### Method Overview:

SPARK-X uses a variance component score test framework:

$$T_g = \frac{n \cdot y_g^T K y_g}{\|y_g\|^2}$$

where:

- $y_g$  = expression vector for gene  $g$
- $K$  = spatial kernel matrix (derived from coordinates)
- $n$  = number of spatial locations

### Kernel Types:

- Projection kernel: Linear kernel based on scaled coordinates. Detects gradients and linear spatial trends.
- Gaussian kernels: Multiple bandwidth Gaussian RBF kernels. Detect localized hotspots of different sizes.
- Cosine kernels: Multiple frequency periodic kernels. Detect periodic/oscillating spatial patterns.

### P-value Computation:

- Individual kernel p-values: Davies' method for quadratic forms
- Combined p-value: ACAT (Aggregated Cauchy Association Test)

### Advantages:

- Non-parametric: No distributional assumptions
- Scalable:  $O(n)$  complexity, handles millions of cells
- Multiple kernels: Detects diverse pattern types
- Robust: ACAT combination handles correlated tests



---

data_simulation	<i>Simulate Spatial Transcriptomics Data with Known SVGs</i>
-----------------	--

---

### Description

Functions to generate simulated spatial transcriptomics data with known spatially variable genes (ground truth). Useful for benchmarking and testing.

### Value

See individual function documentation for return values.

---

example_svg_data	<i>Example Spatial Transcriptomics Data</i>
------------------	---

---

### Description

A pre-generated example dataset for testing SVG detection methods. Contains 500 spots and 200 genes, with 50 known SVGs.

### Format

A list with components:

**counts** Integer matrix (200 genes  $\times$  500 spots) of raw counts

**logcounts** Numeric matrix of  $\log_2(\text{counts} + 1)$

**spatial\_coords** Numeric matrix (500 spots  $\times$  2) of x, y coordinates

**gene\_info** Data.frame with columns: gene, is\_svg, pattern\_type

### Value

A list containing the example dataset (see Format section).

### Source

Simulated using [simulate\\_spatial\\_data](#)

**Examples**

```

data(example_svg_data)
str(example_svg_data)

# Run SVG detection (requires RANN package)
if (requireNamespace("RANN", quietly = TRUE)) {
  results <- CalSVG_MERINGUE(
    example_svg_data$counts,
    example_svg_data$spatial_coords,
    verbose = FALSE
  )

  # Check accuracy
  truth <- example_svg_data$gene_info$is_svg
  detected <- results$p.adj < 0.05
  print(table(truth, detected))
}

```

---

getSpatialNeighbors\_Delaunay

*Build Spatial Network via Delaunay Triangulation*


---

**Description**

Constructs a spatial adjacency matrix using Delaunay triangulation. Two points are considered neighbors if they share an edge in the triangulation.

**Usage**

```

getSpatialNeighbors_Delaunay(
  coords,
  filter_dist = NA,
  binary = TRUE,
  verbose = FALSE
)

```

**Arguments**

coords	Numeric matrix of spatial coordinates. Rows are spatial locations, columns are x, y (and optionally z) coordinates.
filter_dist	Numeric or NA. Maximum distance threshold for neighbors. Default is NA (no filtering).
binary	Logical. If TRUE (default), return binary adjacency matrix.
verbose	Logical. Whether to print progress messages. Default is FALSE.

**Details**

The function uses Delaunay triangulation from the geometry package. For 2D coordinates, this creates triangles. For 3D, it creates tetrahedra.

Duplicate coordinates are slightly jittered to avoid computational issues.

**Value**

Square numeric matrix of spatial adjacency weights.

**Examples**

```
set.seed(42)
coords <- cbind(x = runif(50), y = runif(50))
rownames(coords) <- paste0("spot_", 1:50)

if (requireNamespace("geometry", quietly = TRUE)) {
  W <- getSpatialNeighbors_Delaunay(coords)
}
```

---

getSpatialNeighbors\_KNN

*Build Spatial Network via K-Nearest Neighbors*

---

**Description**

Constructs a spatial adjacency matrix using K-nearest neighbors. Each point is connected to its k nearest neighbors based on Euclidean distance.

**Usage**

```
getSpatialNeighbors_KNN(
  coords,
  k = 10L,
  mutual = FALSE,
  binary = TRUE,
  verbose = FALSE
)
```

**Arguments**

coords	Numeric matrix of spatial coordinates.
k	Integer. Number of nearest neighbors. Default is 10.
mutual	Logical. If TRUE, only mutual nearest neighbors are connected (both A->B and B->A must exist). Default is FALSE.

binary	Logical. If TRUE (default), return binary adjacency matrix. If FALSE, return distance-weighted matrix.
verbose	Logical. Whether to print progress messages. Default is FALSE.

### Details

Uses the RANN package for efficient nearest neighbor search with KD-trees. The resulting network may be asymmetric (A is neighbor of B doesn't mean B is neighbor of A) unless `mutual = TRUE`.

### Value

Square numeric matrix of spatial adjacency weights.

### Examples

```
set.seed(42)
coords <- cbind(x = runif(50), y = runif(50))
rownames(coords) <- paste0("spot_", 1:50)

if (requireNamespace("RANN", quietly = TRUE)) {
  W <- getSpatialNeighbors_KNN(coords, k = 6)
}
```

---

morani

*Calculate Moran's I Statistic*

---

### Description

Computes Moran's I spatial autocorrelation statistic for a numeric vector given a spatial weights matrix.

### Usage

```
morani(x, W, standardize = TRUE)
```

### Arguments

x	Numeric vector of values (e.g., gene expression).
W	Square numeric matrix of spatial weights. Must have the same dimension as <code>length(x)</code> .
standardize	Logical. If TRUE (default), row-standardize the weights matrix.

## Details

Moran's I is defined as:

$$I = \frac{n}{W} \frac{\sum_i \sum_j w_{ij} (x_i - \bar{x})(x_j - \bar{x})}{\sum_i (x_i - \bar{x})^2}$$

where n is the number of observations, W is the sum of all weights, and w<sub>ij</sub> is the weight between locations i and j.

Under the null hypothesis of no spatial autocorrelation:

- Expected value: E[I] = -1/(n-1)
- Variance is computed using the analytical formula from Cliff and Ord (1981)

## Value

A list containing:

- observed: The observed Moran's I statistic
- expected: Expected value under null hypothesis of no spatial autocorrelation (typically -1/(n-1))
- sd: Standard deviation under null hypothesis

## References

Cliff, A.D. and Ord, J.K. (1981) Spatial Processes: Models & Applications. Pion.

## Examples

```
# Create example data
set.seed(42)
x <- rnorm(100)
coords <- cbind(runif(100), runif(100))

# Calculate Moran's I (requires RANN package)
if (requireNamespace("RANN", quietly = TRUE)) {
  W <- buildSpatialNetwork(coords, method = "knn", k = 6)
  result <- morani(x, W)
  print(result)
}
```

---

morani_test	<i>Moran's I Test for Spatial Autocorrelation</i>
-------------	---

---

### Description

Performs a statistical test for spatial autocorrelation using Moran's I. Returns the test statistic, expected value, standard deviation, and p-value.

### Usage

```
morani_test(
  x,
  W,
  alternative = c("greater", "less", "two.sided"),
  standardize = TRUE
)
```

### Arguments

x	Numeric vector of values.
W	Square numeric matrix of spatial weights.
alternative	Character string specifying the alternative hypothesis. One of "greater" (default), "less", or "two.sided". <ul style="list-style-type: none"> <li>• "greater": Test for positive spatial autocorrelation (similar values cluster together)</li> <li>• "less": Test for negative spatial autocorrelation (dissimilar values are neighbors)</li> <li>• "two.sided": Test for any spatial autocorrelation</li> </ul>
standardize	Logical. If TRUE (default), row-standardize weights.

### Value

A named numeric vector with components:

- observed: Observed Moran's I
- expected: Expected Moran's I under null
- sd: Standard deviation under null
- p.value: P-value from normal approximation

### Examples

```
set.seed(42)
x <- rnorm(100)
coords <- cbind(runif(100), runif(100))
```

```
# Test for spatial autocorrelation (requires RANN package)
if (requireNamespace("RANN", quietly = TRUE)) {
  W <- buildSpatialNetwork(coords, method = "knn", k = 6)
  result <- moranI_test(x, W)
  print(result)
}
```

---

simulate\_spatial\_data *Simulate Spatial Transcriptomics Data*

---

### Description

Generates a simulated spatial transcriptomics dataset with a mixture of spatially variable genes (SVGs) and non-spatially variable genes. Uses scientifically accurate count distributions (Negative Binomial).

### Usage

```
simulate_spatial_data(
  n_spots = 500,
  n_genes = 200,
  n_svg = 50,
  grid_type = c("hexagonal", "square", "random"),
  pattern_types = c("gradient", "hotspot", "periodic", "cluster"),
  mean_counts = 50,
  dispersion = 5
)
```

### Arguments

n_spots	Integer. Number of spatial locations. Default is 500.
n_genes	Integer. Total number of genes. Default is 200.
n_svg	Integer. Number of spatially variable genes. Default is 50.
grid_type	Character. Type of spatial layout. <ul style="list-style-type: none"> <li>• "hexagonal" (default): Visium-like hexagonal grid</li> <li>• "square": Square grid</li> <li>• "random": Random spatial distribution</li> </ul>
pattern_types	Character vector. Types of spatial patterns for SVGs. Any combination of: <ul style="list-style-type: none"> <li>• "gradient": Linear spatial gradient</li> <li>• "hotspot": Localized expression hotspots</li> <li>• "periodic": Periodic/oscillating patterns</li> <li>• "cluster": Clustered expression</li> </ul> Default is all four types.

mean_counts	Numeric. Mean expression level for baseline. Default is 50.
dispersion	Numeric. Dispersion parameter for Negative Binomial. Smaller values = more overdispersion. Default is 5.

## Details

### Spatial Patterns:

- **Gradient:** Expression increases linearly along x-axis
- **Hotspot:** High expression in circular regions
- **Periodic:** Sine wave pattern along x-axis
- **Cluster:** Expression in spatially defined clusters

**Count Distribution:** Counts are drawn from Negative Binomial distribution:

$$X \sim NB(\mu, \phi)$$

where mu is the mean (modulated by spatial pattern) and phi is dispersion.

## Value

A list containing:

- counts: Matrix of gene counts (genes × spots)
- spatial\_coords: Matrix of spatial coordinates (spots × 2)
- gene\_info: Data.frame with gene metadata including is\_svg (TRUE/FALSE) and pattern\_type
- logcounts: Log-normalized counts (log2(counts + 1))

## Examples

```
# Set seed for reproducibility before calling
set.seed(42)
sim_data <- simulate_spatial_data(n_spots = 200, n_genes = 50, n_svg = 10)
str(sim_data, max.level = 1)

# Use with SVG detection (requires RANN)
if (requireNamespace("RANN", quietly = TRUE)) {
  results <- CalSVG_MERINGUE(sim_data$counts, sim_data$spatial_coords,
                             network_method = "knn", k = 10, verbose = FALSE)
}
```

---

utils_spatial	<i>Spatial Network Utilities</i>
---------------	----------------------------------

---

**Description**

Utility functions for building and manipulating spatial neighborhood networks. These functions are used by SVG detection methods to define spatial relationships between spots/cells.

**Value**

See individual function documentation for return values.

---

utils_stats	<i>Statistical Utilities for SVG Detection</i>
-------------	--

---

**Description**

Statistical utility functions used by SVG detection methods, including Moran's I calculation, p-value computation, and expression binarization.

**Value**

See individual function documentation for return values.

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