# Package 'globalKinhom'

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Type Package

<b>Title</b> Inhomogeneous K- And Pair Correlation Functions Using Global Estimators			
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<b>Depends</b> R ( $>= 3.5.0$ ), spatstat.explore ( $>= 3.0$ )			
<b>Imports</b> stats, utils, grDevices, spatstat.geom (>= 3.1), spatstat.random (>= 2.1.0), spatstat.univar			
<b>Description</b> Second-order summary statistics K- and pair-correlation functions describe interactions in point pattern data. This package provides computations to estimate those statistics on inhomogeneous point processes, using the methods of in T Shaw, J Møller, R Waagepetersen, 2020 <doi:10.48550 arxiv.2004.00527="">.</doi:10.48550>			
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globalKinhom-package Inhomogeneous K- And Pair Correlation Functions Using Global Estimators

#### **Description**

Second-order summary statistics K- and pair-correlation functions describe interactions in point pattern data. This package provides computations to estimate those statistics on inhomogeneous point processes, using the methods of in T Shaw, J Møller, R Waagepetersen, 2020 <doi:10.48550/arXiv.2004.00527>.

#### **Details**

#### The DESCRIPTION file:

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Description: Second-order summary statistics K- and pair-correlation functions describe interactions in point pattern data.

License: GPL (>= 2)

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Kglobal (cross) K functions with a global intensity

reweighting

expectedPairs Expected pairs in an inhomogeneous poisson

process

globalKinhom-package Inhomogeneous K- And Pair Correlation Functions

Using Global Estimators

pcfglobal (cross) pair correlation functions with a

global intensity reweighting

This package accompanies Shaw et al (2020). It provides "global" estimators for the non-parametric K- and pair correlation functions, which summarize the second order interactions of second-order intensity-reweighted stationary (SOIRS) point processes. These estimators provide an alternative to those proposed by Baddeley et al (2000) for SOIRS point processes, which we refer to as "local" estimators. The local estimators are implemented in the spatstat.explore package as pcfinhom and Kinhom, with pcfcross.inhom and Kcross.inhom for the corresponding cross-pcf and cross-K-function.

Where possible, the interfaces are made to match those used by the spatstat.explore package.

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#### Author(s)

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#### References

T Shaw, J Møller, R Waagepetersen. 2020. "Globally Intensity-Reweighted Estimators for *K*- and pair correlation functions". arXiv:2004.00527 [stat.ME].

A Baddeley, J Møller, R Waagepetersen. 2000. "Non- and Semi-Parametric Estimation of Interaction in Inhomogeneous Point Patterns". Statistica Neerlandica 54, 329-350.

#### See Also

```
spatstat.explore, Kglobal, link{pcfglobal}
```

expectedPairs

Expected pairs in an inhomogeneous poisson process

# Description

Compute the expected number of pairs at a given displacement h in a poisson process with a given intensity function. This corresponds to the integrals  $\gamma$  of Shaw et al. 2020. The various functions correspond to the univariate and bivariate versions of the anisotropic or isotropic versions of  $\gamma$ . The final two options (expectedPairs\_kernloo and expectedPairs\_iso\_kernloo), provide implementations of the leave-out kernel estimates of  $\gamma$ :  $\bar{\gamma}(h)$  and  $\bar{\gamma}^{\rm iso}(r)$ . In those cases, the point pattern X itself is passed to the routine, rather than the (true or estimated) intensities rho etc. The estimators for  $\bar{\gamma}(h)$  are only applicable to univariate processes. See Shaw et al, 2020 for details.

#### Usage

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# Arguments

rho1, rho2, rho	Intensity functions, either of class im or funxy. This may be produced by density.ppp or densityfun.ppp, or provided by a fitted intensity model.
X	Point pattern of class ppp with the points of the pattern for which $\bar{\gamma}$ is to be estimated.
hx, hy	For expectedPairs and expectedCrossPairs (i.e. $\gamma(h)$ ), the displacements $h \in \mathbb{R}^2$ to evaluate $\gamma$ at. These can be in any format supported by xy.coords.
r	For the isotropic versions $\gamma^{\rm iso}(r),$ the separations $r$ at which $\gamma^{\rm iso}$ is to be evaluated.
method	Either mc (the default) or lattice. Compute integral using monte-carlo or on a lattice.
tol	A tolerance for how precise the integral should be. This is compared to a standard error for the mc estimate.
sigma	Smoothing bandwidth for direct kernel-based estimators $\bar{\gamma}$ .
leaveoneout	Use leave-out estimators. This should generally be true except for the purpose of evaluating the bias of the standard estimators. See Shaw et al 2020 for details.
maxeval	Maximum number of evaluations of rho per iteration. Prevents memory-related crashes that can occur.
maxsamp	Maximum number of monte carlo samples per iteration. If this is too large, you may do more work than required to achieve tol.
dx	if method=="lattice", a lattice spacing for the computation. defaults to .01.

# Value

The return value is a numeric vector with length equal to the number of displacements h passed

# Author(s)

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# References

T Shaw, J Møller, R Waagepetersen. 2020. "Globally Intensity-Reweighted Estimators for K- and pair correlation functions". arXiv:2004.00527 [stat.ME].

# See Also

pcfglobal, Kglobal, which use these functions to compute the normalization functions  $\gamma$ .

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Kglobal

(cross) K functions with a global intensity reweighting

# Description

Compute  $K_{global}$ 

# Usage

#### **Arguments**

interpolate

X, Y lambda, lambdaX,	point process of type ppp, on which to evaluate the (cross) $K$ -function , lambdaY intensity function estimates corresponding to X and Y. If omitted, intensity functions will be computed using <code>density.ppp</code> or <code>densityfun.ppp</code> (see discrete.lambda below)	
	extra args passed to density.ppp or densityfun.ppp, if applicable.	
sigma	Bandwidth value to use for kernel-based intensity estimation, intensity functions and exp_prs are not provided by the user.	
r	Values of $r$ to evaluate $K(r)$ at. If omitted, a sensible default is chosen, using the same conventions as Kest and Kinhom.	
rmax	Maximum $r$ to evaluate $K(r)$ at. rmax is used to generate values for r, if omitted. If missing, a sensible default is chosen.	
breaks	For internal use only.	
normtol	A tolerance to use for expectedPairs or expectedCrossPairs when computing monte-carlo estimates of the normalizing factor $\gamma$ . Expressed as a maximum fractional standard error.	
discrete.lambda		
	If TRUE, and intensity function(s) are not supplied, estimate intensities by interpolating the values on a discrete lattice (using interp.im and density.ppp), instead of exactly (using densityfun.ppp).	

at the exact displacements observed in the pattern.

If TRUE, evaluate the expectedCrossPairs on a lattice and interpolate, rather than

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interpolate.fac

If interpolate, the lattice spacing will be sigma/interpolate.fac.

isotropic Set to TRUE to use the isotropic estimators  $\gamma_{\rm iso}$ .

leaveoneout Use the leave-one-out estimator for  $\gamma$ . See Shaw et al, 2020 for details.

exp\_prs A function that returns values for  $\gamma_{iso}(r)$ . If  $\gamma$  is known explicitly, or the same

calculation is being used for several point patterns, it can be much faster to compute it once and provide the function as exp\_prs, since the computation of

 $\gamma$  is usually the slowest part.

interpolate.maxdx

Upper bound on allowable lattice spacing for interpolation.

dump For debugging purposes, include computed values of  $\gamma$  with the output, as attrs.

#### Value

The return value is an object of class fv, just as for Kest and Kinhom. The object contains columns r, theo, and global, corresponding respectively to the argument r, the theoretical values of K(r) for a Poisson process, and  $K_{\rm global}(r)$ .

#### Author(s)

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#### References

T Shaw, J Møller, R Waagepetersen. 2020. "Globally Intensity-Reweighted Estimators for K- and pair correlation functions". arXiv:2004.00527 [stat.ME].

# See Also

```
expectedPairs
```

# **Examples**

```
rho <- funxy(function(x,y) 80*(1+x), owin())
X <- rpoispp(rho)
K <- Kglobal(X)
#plot(K)</pre>
```

pcfglobal

(cross) pair correlation functions with a global intensity reweighting

# **Description**

Compute  $g_{global}$  or  $c_{global}$ 

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#### Usage

```
pcfglobal(X, lambda=NULL, ..., sigma=bw.CvL(X), r=NULL, rmax=NULL,
    kernel="epanechnikov", bw=NULL, stoyan=0.15, normtol=.005, ratio=FALSE,
    discrete.lambda=FALSE, divisor=c("r", "d"),
    leaveoneout=TRUE, interpolate=TRUE, interpolate.fac=10, exp_prs=NULL,
    interpolate.maxdx=diameter(as.owin(X))/100, dump=FALSE)

pcfcross.global(X,Y, lambdaX=NULL, lambdaY=NULL, ...,
    sigma=bw.CvL(X), r=NULL, rmax=NULL, kernel="epanechnikov", bw=NULL,
    stoyan=0.15, normtol=.005, ratio=FALSE, discrete.lambda=FALSE,
    divisor=c("r", "d"), analytical=NULL, interpolate=TRUE,
    interpolate.fac=10, exp_prs=NULL,
    interpolate.maxdx=diameter(as.owin(X))/100, dump=FALSE)
```

# **Arguments**

X, Y point process of type ppp, on which to evaluate the (cross) K-function

lambda, lambdaX, lambdaY

intensity function estimates corresponding to X and Y. If omitted, intensity functions will be computed using density.ppp or densityfun.ppp (see discrete.lambda

below)

... extra args passed to density.ppp or densityfun.ppp, if applicable.

sigma Bandwidth value to use for kernel-based intensity estimation, intensity functions

and exp\_prs are not provided by the user.

Values of r to evaluate K(r) at. If omitted, a sensible default is chosen, using

the same conventions as Kest and Kinhom.

rmax Maximum r to evaluate K(r) at. rmax is used to generate values for r, if omit-

ted. If missing, a sensible default is chosen.

kernel Kernel type for smoothing of pcf.

bw Kernel bandwidth for smoothing of pcf.

stoyan Coefficient for Stoyan's bandwidth selection rule. See pcf.ppp.

normtol A tolerance to use for expectedPairs or expectedCrossPairs when computing

monte-carlo estimates of the normalizing factor  $\gamma$ . Expressed as a maximum

fractional standard error.

ratio If TRUE, assemble numerator and denominator of pcf estimator separately.

divisor Whether to use the evaluation distance ("r") or the distance between points

("d") to normalize the contribution of each point pair.

analytical If TRUE, use Diggle-Jones weights

discrete.lambda

If TRUE, and intensity function(s) are not supplied, estimate intensities by interpolating the values on a discrete lattice (using interp.im and density.ppp),

instead of exactly (using densityfun.ppp).

interpolate If TRUE, evaluate the expected Cross Pairs on a lattice and interpolate, rather than

at the exact displacements observed in the pattern.

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interpolate.fac

If interpolate, the lattice spacing will be sigma/interpolate.fac.

leave-one-out estimator for  $\gamma$ . See Shaw et al 2020 for details.

exp\_prs A function that returns values for  $\gamma_{iso}(r)$ . If  $\gamma$  is known explicitly, or the same

calculation is being used for several point patterns, it can be much faster to compute it once and provide the function as exp\_prs, since the computation of

 $\gamma$  is usually the slowest part.

interpolate.maxdx

Upper bound on allowable lattice spacing for interpolation.

dump For debugging purposes, include computed values of  $\gamma$  with the output, as attrs.

#### Value

The return value is an object of class fv, just as for pcf and pcfinhom. The object contains columns r, theo, and global, corresponding respectively to the argument r, the theoretical values of g(r) for a Poisson process, and  $g_{\rm global}(r)$ .

#### Author(s)

Thomas Shaw <shawtr@umich.edu>

#### References

T Shaw, J Møller, R Waagepetersen. 2020. "Globally Intensity-Reweighted Estimators for K- and pair correlation functions". arXiv:2004.00527 [stat.ME].

# See Also

```
expectedPairs
```

# Examples

```
rho <- funxy(function(x,y) 80*(1+x), owin())
X <- rpoispp(rho)
g <- pcfglobal(X)
#plot(g)</pre>
```

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