# Package 'synapsis'

July 26, 2025

Type Package

**Title** An R package to automate the analysis of double-strand break repair during meiosis

**Version** 1.15.0

Description Synapsis is a Bioconductor software package for automated (unbiased and reproducible) analysis of meiotic immunofluorescence datasets. The primary functions of the software can i) identify cells in meiotic prophase that are labelled by a synaptonemal complex axis or central element protein, ii) isolate individual synaptonemal complexes and measure their physical length, iii) quantify foci and co-localise them with synaptonemal complexes, iv) measure interference between synaptonemal complex-associated foci. The software has applications that extend to multiple species and to the analysis of other proteins that label meiotic prophase chromosomes. The software converts meiotic immunofluorescence images into R data frames that are compatible with machine learning methods. Given a set of microscopy images of meiotic spread slides, synapsis crops images around individual single cells, counts colocalising foci on strands on a per cell basis, and measures the distance between foci on any given strand.

biocViews Software, SingleCell

Depends R (>= 4.1)

Imports EBImage, stats, utils, graphics

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**Encoding** UTF-8 **RoxygenNote** 7.1.1

VignetteBuilder knitr

**Suggests** knitr, rmarkdown, testthat (>= 3.0.0), ggplot2, tidyverse, BiocStyle

Config/testthat/edition 3

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annotate_foci_counting
 annotate_foci_counting
```

#### **Description**

Contains all plotting routines for count foci annotation

```
annotate_foci_counting(
  img_file,
  cell_count,
  img_orig,
  img_orig_foci,
  artificial_amp_factor,
  strands,
  coincident_foci,
  foci_label,
  alone_foci,
  percent_px,
  foci_per_cell
```

#### **Arguments**

```
img_file
                  cell's file name
cell_count
                  unique cell counter
                  original strand crop
img_orig
img_orig_foci
                  cropped foci channel
artificial_amp_factor
                  amplification factor
                  black white mask of strand channel
strands
coincident_foci
                  mask of overlap between strand and foci channel
foci_label
                  black and white mask of foci channel
                  estimated number of foci that are NOT on a strand.
alone_foci
                  percentage of foci mask that coincides with strand channel small number indi-
percent_px
                  cates potentially problematic image.
foci_per_cell
                 number of foci counted per cell
```

## Value

displays key steps from raw image to coincident foci count

## **Description**

Contains all plotting routines for count foci annotation

```
annotate_foci_counting_adjusted(
  img_file,
  cell_count,
  img_orig,
  img_orig_foci,
  artificial_amp_factor,
  strands,
  coincident_foci,
  foci_label,
  alone_foci,
  percent_px,
  foci_per_cell
```

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## **Arguments**

```
img_file
                  cell's file name
cell_count
                  unique cell counter
img_orig
                  original strand crop
                  cropped foci channel
img_orig_foci
artificial_amp_factor
                  amplification factor
                  black white mask of strand channel
strands
coincident_foci
                  mask of overlap between strand and foci channel
foci_label
                  black and white mask of foci channel
alone_foci
                  estimated number of foci that are NOT on a strand.
                  percentage of foci mask that coincides with strand channel small number indi-
percent_px
                  cates potentially problematic image.
foci_per_cell
                  number of foci counted per cell
```

#### Value

displays key steps from raw image to coincident foci count

```
append_data_frame append_data_frame
```

## **Description**

applies new row to data frame

```
append_data_frame(
 WT_str,
 KO_str,
 WT_out,
 KO_out,
  img_file,
  foci_areas,
 df_cells,
 cell_count,
  stage,
  foci_per_cell,
  image_mat,
 percent_px,
 alone_foci,
 discrepant_category,
 C1
)
```

auto\_crop\_fast 5

#### **Arguments**

WT\_str string in filename corresponding to wildtype genotype. Defaults to ++. KO str string in filename corresponding to knockout genotype. Defaults to -. WT\_out string in output csv in genotype column, for knockout. Defaults to +/+. K0\_out string in output csv in genotype column, for knockout. Defaults to -/-.

cell's file name img\_file

foci areas pixel area of each foci df\_cells current data frame cell\_count unique cell counter

meiosis stage of interest. Currently count\_foci determines this with thresholdstage

> ing/object properties in the synaptonemal complex channel by previosly calling the get\_pachytene function. Note that if using this option, the count\_foci function requires that the input directory contains a folder called "pachytene" with

the crops in it.

foci\_per\_cell foci count for cell

image\_mat matrix with all pixel values above zero

percent\_px percentage of foci mask that coincides with strand channel small number indi-

cates potentially problematic image.

alone\_foci estimated number of foci that are NOT on a strand.

discrepant\_category

estimated number of foci that are NOT on a strand.

C1 criteria

#### Value

data frame with new row

auto\_crop\_fast auto\_crop\_fast

# **Description**

crop an image around each viable cell candidate.

```
auto_crop_fast(
 img_path,
 max_cell_area = 20000,
 min_cell_area = 7000,
 mean_pix = 0.08,
 annotation = "off",
 blob_factor = 15,
 bg_blob_factor = 10,
 offset = 0.2,
  final_blob_amp = 10,
```

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```
test_amount = 0,
brush_size_blob = 51,
sigma_blob = 15,
channel3_string = "DAPI",
channel2_string = "SYCP3",
channel1_string = "MLH3",
file_ext = "jpeg",
third_channel = "off",
cell_aspect_ratio = 2,
strand_amp = 2,
path_out = img_path,
resize_1 = 720,
crowded_cells = "FALSE",
watershed_radius = 50,
watershed_tol = 0.2,
cropping_factor = 1.3
```

## **Arguments**

path containing image data to analyse img\_path Maximum pixel area of a cell candidate max\_cell\_area Minimum pixel area of a cell candidate min\_cell\_area Mean pixel intensity of cell crop (in SYCP3 channel) for normalisation mean\_pix annotation Choice to output pipeline choices (recommended to knit) blob\_factor Contrast factor to multiply original image by before smoothing/smudging Contrast factor to multiply original image by to take background. Used prior to bg\_blob\_factor thresholding. offset Pixel value offset from bg\_blob\_factor. Used in thresholding to make blob mask. Contrast factor to multiply smoothed/smudged image. Used in thresholding to final\_blob\_amp make blob mask. Optional number of first N images you want to run function on. For troutest\_amount bleshooting/testing/variable calibration purposes. brush\_size\_blob Brush size for smudging the synaptonemal complex channel to make blobs Sigma in Gaussian brush for smudging the synaptonemal complex channel to sigma\_blob make blobs channel3\_string Optional. String appended to the files showing the channel illuminating cell structures. Defaults to DAPI, if third channel == "on".

channel2\_string

String appended to the files showing the channel illuminating synaptonemal complexes. Defaults to SYCP3

channel1\_string

String appended to the files showing the channel illuminating foci. Defaults to

file\_ext file extension of your images e.g. tif jpeg or png.

Optional, defaults to "off". Set to "on" if you would also like crops of the third third\_channel channel.

count\_foci 7

cell\_aspect\_ratio

Maximum aspect ratio of blob to be defined as a cell

strand\_amp multiplication of strand channel for get\_blobs function.

path\_out user specified output path. Defaults to img\_path

resize\_l length for resized image

crowded\_cells TRUE or FALSE, defaults to FALSE. Set to TRUE if you have many cells in a

frame that almost touch

watershed\_radius

Radius (ext variable) in watershed method used in strand channel. Defaults to 1

(small)

watershed\_tol Intensity tolerance for watershed method. Defaults to 0.05.

cropping\_factor

size of cropping window square, as factor of characteristic blob radius. Defaults

to 1. May need to increase if using watershed.

#### Details

This function takes all images in a directory, and crops around individual cells according to the antibody that stains synaptonemal complexes e.g. SYCP3. First, it increases the brightness and smudges the image with a Gaussian brush, and creates a mask using thresholding (get\_blobs). Then it deletes cell candidates in the mask deemed too large, too small, or too long (keep\_cells). Using the computeFeatures functions from EBImage to locate centre and radius, the cropping area is determined and the original image cropped. These images are saved in either a user specified directory, or a crops folder at the location of the image files.

#### Value

cropped synaptonemal complex and foci channels around single cells, regardless of stage

#### Author(s)

Lucy McNeill

#### **Examples**

```
demo_path = paste0(system.file("extdata",package = "synapsis"))
auto_crop_fast(demo_path, annotation = "on", max_cell_area = 30000,
min_cell_area = 7000, file_ext = "tif",crowded_cells = TRUE)
```

count\_foci

count\_foci

## **Description**

Calculates coincident foci in synaptonemal complex and foci channel, per cell

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#### Usage

```
count_foci(
  img_path,
  stage = "none",
  offset_px = 0.2,
  offset_factor = 2,
  brush_size = 3,
  brush_sigma = 3,
  foci_norm = 0.01,
  annotation = "off",
  channel2_string = "SYCP3",
  channel1_string = "MLH3",
  file_ext = "jpeg",
  KO_str = "--",
  WT_str = "++"
 KO_out = "-/-",
  WT_out = "+/+",
  watershed_stop = "off",
  watershed_radius = 1,
  watershed_tol = 0.05,
  crowded_foci = TRUE,
  artificial_amp_factor = 1,
  strand_amp = 2,
  min_foci = -1,
  disc_size = 51,
  modify_problematic = "off",
  disc_size_foci = 5,
 C1 = 0.02,
  C2 = 0.46,
  C_weigh_foci_number = TRUE
```

# **Arguments**

img_path	path containing crops to analyse	
stage	meiosis stage of interest. Currently count_foci determines this with thresholding/ object properties in the synaptonemal complex channel by previosly calling the get_pachytene function. Note that if using this option, the count_foci function requires that the input directory contains a folder called "pachytene" with the crops in it.	
offset_px	Pixel value offset used in thresholding of synaptonemal complex channel	
offset_factor	Pixel value offset used in thresholding of foci channel	
brush_size	size of brush to smooth the foci channel. Should be small to avoid erasing foci.	
brush_sigma	sigma for Gaussian smooth of foci channel. Should be small to avoid erasing foci.	
foci_norm	Mean intensity to normalise all foci channels to.	
annotation	Choice to output pipeline choices (recommended to knit)	
channel2_string		
	String appended to the files showing the channel illuminating synaptonemal	

complexes. Defaults to SYCP3

count\_foci 9

channel1\_string

String appended to the files showing the channel illuminating foci. Defaults to

MLH3

file\_ext file extension of your images e.g. tiff jpeg or png.

KO\_str string in filename corresponding to knockout genotype. Defaults to -.

WT\_str string in filename corresponding to wildtype genotype. Defaults to ++.

KO\_out string in output csv in genotype column, for knockout. Defaults to -/-.

WT\_out string in output csv in genotype column, for knockout. Defaults to +/+.

watershed\_stop Stop default watershed method with "on"

watershed\_radius

Radius (ext variable) in watershed method used in foci channel. Defaults to 1

(small)

watershed\_tol Intensity tolerance for watershed method. Defaults to 0.05.

 $\label{eq:crowded_foci} \textbf{TRUE} \ \text{or} \ \textbf{FALSE}, \ \textbf{defaults} \ \textbf{to} \ \textbf{FALSE}. \ \textbf{Set} \ \textbf{to} \ \textbf{TRUE} \ \textbf{if} \ \textbf{you} \ \textbf{have} \ \textbf{foci} > 100 \ \textbf{or} \ \textbf{so}.$ 

artificial\_amp\_factor

Amplification of foci channel, for annotation only.

strand\_amp multiplication of strand channel to make masks

min\_foci minimum pixel area for a foci. Depends on your dpi etc. Defaults to 4

disc\_size size of disc for local background calculation in synaptonemal complex channel

modify\_problematic

option for synapsis to try and "save" images which have likely been counted incorrectly due to a number of reasons. Default settings are optimized for mouse

pachytene. Defaults to "off"

disc\_size\_foci size of disc for local background calculation in foci channel

C1 Default crispness criteria = sd(foci\_area)/(mean(foci\_area)+1)

C2 Alternative crisp criteria.

C\_weigh\_foci\_number

choose crispness criteria- defaults to TRUE to use C1 (weighing with number).

Otherwise set to FALSE to use C2

#### **Details**

In this function, masks for the synaptonemal complex (SC) and foci channel are created from the saved crops of single/individual cells. These masks are computed using (optional) input parameters related to meiosis stage/ how well spread chromosomes are (for the former) and related to smoothing, thresholding and how "crowded" foci are for the latter. Finally, these two masks are multiplied, and the number of objects found with EBImage's computeFeatures are the colocalizing foci.

The file, cell number, foci count etc. are output as a data frame.

#### Value

data frame with foci count per cell

#### Author(s)

Lucy McNeill

# **Examples**

# Description

Creates mask for every individual cell candidate in mask

## Usage

```
crop_single_object_fast(
  retained,
  OOI_final,
  counter_final,
  img_orig,
  img_orig_foci,
  img\_orig\_DAPI = "blank",
  file_sc,
  file_foci,
  file_DAPI = "blank",
  cell_count,
  mean_pix,
  annotation,
  file_base,
  img_path,
  r_max,
  cx,
  су,
  channel3_string,
  channel2_string,
  channel1_string,
  file_ext,
  third_channel,
  path_out,
  img_orig_highres,
  resize_l,
  crowded_cells,
  cropping_factor
)
```

# Arguments

retained Mask of cell candidates which meet size criteria. After smoothing/smudging and thresholding.

OOI\_final Objects of interest count. Total number of cell candidates in retained.

counter\_final Counter for single cell we are focussing on. Remove all other cells where

counter\_single not equal to counter\_final.

img\_orig description img\_orig\_foci description img\_orig\_DAPI description

file\_sc filename of synaptonemal complex channel image

file\_foci filename of foci channel image file\_DAPI filename of DAPI channel image

cell\_count counter for successful crops around cells

mean\_pix Mean pixel intensity of cell crop (in SYCP3 channel) for normalisation

annotation Choice to output pipeline choices (recommended to knit)

file\_base filename base common to all three channels i.e. without -MLH3.jpeg etc.

img\_path path containing image data to analyse r\_max maximum radius of blob for cropping

cx centre of blob x
cy centre of blob y

channel3\_string

Optional. String appended to the files showing the channel illuminating cell structures. Defaults to DAPI, if third channel == "on".

channel2\_string

String appended to the files showing the channel illuminating synaptonemal complexes. Defaults to SYCP3

channel1\_string

String appended to the files showing the channel illuminating foci. Defaults to MLH3

file\_ext file extension of your images e.g. tif jpeg or png.

third\_channel Optional, defaults to "off". Set to "on" if you would also like crops of the third

channel.

path\_out user specified output path. Defaults to img\_path

img\_orig\_highres

the original strand image with original resolution

resize\_l length of square to resize original image to.

crowded\_cells TRUE or FALSE, defaults to FALSE. Set to TRUE if you have many cells in a

frame that almost touch

cropping\_factor

size of cropping window square, as factor of characteristic blob radius. Defaults to 1. May need to increase if using watershed.

#### Value

Crops around all candidates in both channels

12 get\_blobs

|--|

# Description

Makes mask of all objects bright enough to be classified as a cell cadidate

# Usage

```
get_blobs(
  img_orig,
  blob_factor,
  bg_blob_factor,
  offset,
  final_blob_amp,
  brush_size_blob,
  sigma_blob,
  watershed_tol,
  watershed_radius,
  crowded_cells,
  annotation
)
```

# Arguments

	img_orig	Original image
	blob_factor	Contrast factor to multiply original image by before smoothing/smudging
	bg_blob_factor	Contrast factor to multiply original image by to take background. Used prior to thresholding.
	offset	Pixel value offset from bg_blob_factor. Used in thresholding to make blob mask.
	final_blob_amp	Contrast factor to multiply smoothed/smudged image. Used in thresholding to make blob mask.
brush_size_blob		
		Brush size for smudging the synaptonemal complex channel to make blobs
	sigma_blob	Sigma in Gaussian brush for smudging the synaptonemal complex channel to make blobs $% \left( 1\right) =\left( 1\right) \left( 1$
	watershed_tol	Intensity tolerance for watershed method. Defaults to 0.05.
watershed_radius		
		Radius (ext variable) in watershed method used in strand channel. Defaults to 1 (small) $$
	crowded_cells	TRUE or FALSE, defaults to FALSE. Set to TRUE if you have many cells in a frame that almost touch
	annotation	Choice to output pipeline choices (recommended to knit) have many cells in a frame that almost touch

# Value

Mask with cell candidates

get\_C1 13

get\_C1 get\_C1

# **Description**

calculates the statistic to compare to crisp\_criteria, which determines whether the foci count will be reliable

# Usage

```
get_C1(foci_areas, foci_per_cell, C_weigh_foci_number)
```

## **Arguments**

```
foci_areas pixel area of each foci

foci_per_cell foci count for cell

C_weigh_foci_number

choose crispness criteria- defaults to TRUE to use C1 (weighing with number).

Otherwise set to FALSE to use C2
```

## Value

statistic to comapre to crisp\_criteria

# **Description**

calculates the statistic to compare to crisp\_criteria, which determines whether the foci count will be reliable

```
get_coincident_foci(
  offset_px,
  offset_factor,
  brush_size,
  brush_sigma,
  annotation,
  watershed_stop,
  watershed_radius,
  watershed_tol,
  crowded_foci,
  artificial_amp_factor,
  strand_amp,
  disc_size,
  disc_size_foci,
  img_file,
```

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```
cell_count,
  img_orig,
  img_orig_foci,
  stage,
 WT_str,
 KO_str,
 WT_out,
 KO_out,
 C1_search,
 discrepant_category,
 C1,
 C2,
 df_cells,
 C_weigh_foci_number
)
```

## **Arguments**

Pixel value offset used in thresholding of synaptonemal complex channel offset\_px

offset\_factor Pixel value offset used in thresholding of foci channel

brush\_size size of brush to smooth the foci channel. Should be small to avoid erasing foci. brush\_sigma sigma for Gaussian smooth of foci channel. Should be small to avoid erasing

foci.

annotation Choice to output pipeline choices (recommended to knit)

watershed\_stop Stop default watershed method with "on"

watershed\_radius

Radius (ext variable) in watershed method used in foci channel. Defaults to 1

(small)

Intensity tolerance for watershed method. Defaults to 0.05. watershed\_tol

TRUE or FALSE, defaults to FALSE. Set to TRUE if you have foci > 100 or so. crowded\_foci

artificial\_amp\_factor

Amplification of foci channel, for annotation only.

multiplication of strand channel to make masks strand\_amp

size of disc for local background calculation in synaptonemal complex channel disc\_size

disc\_size\_foci size of disc for local background calculation in foci channel

cell's file name img\_file cell\_count unique cell counter img\_orig original strand crop cropped foci channel img\_orig\_foci

stage meiosis stage of interest. Currently count foci determines this with threshold-

> ing/object properties in the synaptonemal complex channel by previouly calling the get\_pachytene function. Note that if using this option, the count\_foci function requires that the input directory contains a folder called "pachytene" with

the crops in it.

string in filename corresponding to wildtype genotype. Defaults to ++. WT\_str K0\_str string in filename corresponding to knockout genotype. Defaults to -. WT\_out string in output csv in genotype column, for knockout. Defaults to +/+. get\_foci\_per\_cell 15

KO\_out string in output csv in genotype column, for knockout. Defaults to -/-.

C1\_search TRUE or FALSE whether the image is still being modified until it meets the

crispness criteria

discrepant\_category

estimated number of foci that are NOT on a strand.

C1 Default crispness criteria = sd(foci\_area)/(mean(foci\_area)+1)

C2 Alternative crisp criteria.

df\_cells current data frame

C\_weigh\_foci\_number

choose crispness criteria- defaults to TRUE to use  $\operatorname{C1}$  (weighing with number).

Otherwise set to FALSE to use C2

#### Value

data frame with new row with most recent foci per cell appended

```
get_foci_per_cell get_foci_per_cell
```

#### **Description**

creates mask for coincident foci

# Usage

```
get_foci_per_cell(
   img_file,
   offset_px,
   stage,
   strands,
   watershed_stop,
   foci_label,
   annotation,
   cell_count,
   img_orig,
   img_orig_foci,
   artificial_amp_factor,
   coincident_foci
)
```

## **Arguments**

img\_file cell's file name

offset\_px Pixel value offset used in thresholding of synaptonemal complex channel

stage

meiosis stage of interest. Currently count\_foci determines this with thresholding/object properties in the synaptonemal complex channel by previosly calling the get\_pachytene function. Note that if using this option, the count\_foci function requires that the input directory contains a folder called "pachytene" with the crops in it.

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```
black white mask of strand channel
strands
watershed_stop Stop default watershed method with "on"
foci_label
                 black and white mask of foci channel
                 Choice to output pipeline choices (recommended to knit)
annotation
cell_count
                 unique cell counter
img_orig
                 original strand crop
img_orig_foci
                 cropped foci channel
artificial_amp_factor
                 amplification factor
coincident_foci
                 mask of coincident foci
```

#### Value

number of foci per cell

```
get_overlap_mask
```

#### **Description**

creates mask for coincident foci

## Usage

```
get_overlap_mask(
    strands,
    foci_label,
    watershed_stop,
    img_orig_foci,
    watershed_radius,
    watershed_tol
)
```

# Arguments

```
strands black white mask of strand channel

foci_label black and white mask of foci channel

watershed_stop Stop default watershed method with "on"

img_orig_foci cropped foci channel

watershed_radius

Radius (ext variable) in watershed method used in foci channel. Defaults to 1

(small)

watershed_tol Intensity tolerance for watershed method. Defaults to 0.05.
```

#### Value

mask with coincident foci on strands

get\_pachytene 17

get\_pachytene

# Description

Identifies crops in pachytene

# Usage

```
get_pachytene(
  img_path,
  species_num = 20,
  offset = 0.2,
  ecc\_thresh = 0.85,
  area\_thresh = 0.06,
  annotation = "off",
  channel2_string = "SYCP3",
  channel1_string = "MLH3",
  file_ext = "jpeg",
KO_str = "--",
  WT_str = "++",
  KO_out = "-/-",
  WT_out = "+/+",
  path_out = img_path,
  artificial_amp_factor = 3,
  strand_amp = 2,
  resize_1 = 120
)
```

# **Arguments**

path containing crops analyse		
number of chromosomes in the species		
Pixel value offset used in the rholding for the synaptonemal complex (SYCP3) channel $$		
The minimum average eccentricity of all objects in mask determined by computefeatures, for a cell to be pachytene.		
The minimum ratio of pixels included in mask to total, for a cell to be classified as pachytene.		
Choice to output pipeline choices (recommended to knit)		
channel2_string		
String appended to the files showing the channel illuminating synaptonemal complexes. Defaults to ${\sf SYCP3}$		
channel1_string		
String appended to the files showing the channel illuminating foci. Defaults to MLH3		
file extension of your images e.g. tiff jpeg or png.		
string in filename corresponding to knockout genotype. Defaults to		

18 keep\_cells

```
wT_str string in filename corresponding to wildtype genotype. Defaults to ++.

KO_out string in output csv in genotype column, for knockout. Defaults to -/-.

WT_out string in output csv in genotype column, for knockout. Defaults to +/+.

path_out user specified output path. Defaults to img_path

artificial_amp_factor

Amplification of foci channel, for RGB output files. Deaults to 3.

strand_amp multiplication of strand channel.

resize_1 length of resized square cell image.
```

#### **Details**

This function takes the crops make by auto\_crop fast, and determines the number of synaptonemal complex candidates by considering the local background and using EBImage functions. In general, very bright objects which contrast highly with the background will be classified as the same object. Dim objects will likely be classified as many different objects. If the number of objects is too high compared to the species number (species\_num) then the cell is determined to not be in pachytene. Note that this function has been optimized for mouse cells which can be very well spread / separated.

#### Value

Pairs of foci and synaptonemal channel crops for pachytene

#### Author(s)

Lucy McNeill

#### **Examples**

```
demo_path = paste0(system.file("extdata",package = "synapsis"))
SYCP3_stats <- get_pachytene(demo_path,ecc_thresh = 0.8, area_thresh = 0.04, annotation = "on")</pre>
```

keep\_cells keep\_cells

#### **Description**

Deletes objects in mask which are too small, large, oblong i.e. unlikely to be a cell

```
keep_cells(
  candidate,
  max_cell_area,
  min_cell_area,
  cell_aspect_ratio,
  crowded_cells,
  annotation
)
```

make\_foci\_mask 19

#### **Arguments**

candidate Mask of individual cell candidates

max\_cell\_area Maximum pixel area of a cell candidate

min\_cell\_area Minimum pixel area of a cell candidate

cell\_aspect\_ratio

Maximum aspect ratio of blob to be defined as a cell

crowded\_cells TRUE or FALSE, defaults to FALSE. Set to TRUE if you

annotation Choice to output pipeline choices (recommended to knit) have many cells in a

frame that almost touch

#### Value

Mask of cell candidates which meet size criteria

#### **Description**

creates foci mask for foci channel crop

## Usage

```
make_foci_mask(
   offset_factor,
   bg,
   crowded_foci,
   img_orig_foci,
   brush_size,
   brush_sigma,
   disc_size_foci
)
```

## **Arguments**

offset\_factor Pixel value offset used in thresholding of foci channel

bg background value- currently just mean pixel value of whole image

crowded\_foci TRUE or FALSE, defaults to FALSE. Set to TRUE if you have foci > 100 or so.

img\_orig\_foci cropped foci channel

brush\_size size of brush to smooth the foci channel. Should be small to avoid erasing foci.

brush\_sigma for Gaussian smooth of foci channel. Should be small to avoid erasing foci.

disc\_size\_foci size of disc for local background calculation in foci channel

# Value

foci mask

20 remove\_XY

make\_strand\_mask

make\_strand\_mask

## **Description**

creates strand mask for strand channel crop

# Usage

```
make_strand_mask(
  offset_px,
  stage,
  img_orig,
  disc_size,
  brush_size,
  brush_sigma
)
```

# Arguments

offset\_px Pixel value offset used in thresholding of synaptonemal complex channel meiosis stage of interest. Currently count foci determines this with thresholdstage ing/object properties in the synaptonemal complex channel by previosly calling the get\_pachytene function. Note that if using this option, the count\_foci function requires that the input directory contains a folder called "pachytene" with the crops in it. img\_orig original strand crop size of disc for local background calculation in synaptonemal complex channel disc\_size brush\_size size of brush to smooth the foci channel. Should be small to avoid erasing foci. brush\_sigma sigma for Gaussian smooth of foci channel. Should be small to avoid erasing foci.

# Value

strand mask

remove\_XY

remove\_XY

#### **Description**

applies new row to data frame

```
remove_XY(foci_label, foci_candidates, foci_areas)
```

remove\_XY 21

# Arguments

foci\_label black and white mask of foci channel

foci\_candidates

computeFeatures data frame of foci channel

foci\_areas the areas of the foci objects

# Value

mask with XY blob removed

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