Package 'animl'

May 13, 2023

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| animl applyPredictions bestGuess buildFileManifest checkFile convertCoordinates cropImageGenerator cropImageTrainGenerator detectObject detectObjectBatch extractBoxes 1 |
| extractBoxesFromFlat |

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Description

Title

Usage

```
animl(imagedir, mdmodel, speciesmodel, classes)
```

applyPredictions 3

Arguments

imagedir description
mdmodel description
speciesmodel description
classes description

Value

none

Examples

```
## Not run:
imagedir <- "examples/test_data/Southwest"
mdmodel <- "/mnt/machinelearning/megaDetector/md_v5b.0.0_saved_model"
modelfile <- "/mnt/machinelearning/Models/Southwest/2022/Southwest_v2.h5"
classes <- "/mnt/machinelearning/Models/Southwest/2022/classes.txt"
animl(imagedir,mdmodel,modelfile,classes)
## End(Not run)</pre>
```

applyPredictions

Apply Classifier Predictions and Merge DataFrames

Description

Apply Classifier Predictions and Merge DataFrames

Usage

```
applyPredictions(animals, pred, classfile, outfile = NULL, counts = FALSE)
```

Arguments

animals Set of animal crops/images

pred Classifier predictions for animal crops/images

classfile .txt file containing common names for species classes

outfile File to which results are saved

counts Returns a table of all predictions, defaults to FALSE

Value

fully merged dataframe with Species predictions and confidence weighted by MD conf

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Examples

```
## Not run:
alldata <- applyPredictions(animals,empty,classfile,pred,counts = FALSE)
## End(Not run)</pre>
```

bestGuess

Select Best Classification From Multiple Frames

Description

Select Best Classification From Multiple Frames

Usage

```
bestGuess(
  manifest,
  sort = "count",
  count = FALSE,
  shrink = FALSE,
  outfile = NULL,
  prompt = TRUE,
  parallel = FALSE,
  workers = 1
)
```

Arguments

| manifest | dataframe of all frames including species classification |
|----------|---|
| sort | method for selecting best prediction, defaults to most frequent |
| count | if true, return column with number of MD crops for that animal (does not work for images) |
| shrink | if true, return a reduced dataframe with one row per image |
| outfile | file path to which the data frame should be saved |
| prompt | if true, prompts the user to confirm overwrite |
| parallel | Toggle for parallel processing, defaults to FALSE |
| workers | number of processors to use if parallel, defaults to 1 |
| | |

Value

dataframe with new prediction in "Species" column

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Examples

```
## Not run:
mdmanifest <- bestGuess(manifest, sort = "conf")
## End(Not run)</pre>
```

buildFileManifest

Extract exif Data and Create File Manifest

Description

Extract exif Data and Create File Manifest

Usage

```
buildFileManifest(imagedir, exif = TRUE, offset = 0, outfile = NULL)
```

Arguments

| imagedir | file path |
|----------|--|
| exif | returns date and time information from exif data, defaults to true |
| offset | add offset to videos, defaults to 0 |
| outfile | file path to which the data frame should be saved |

Value

files dataframe with or without file modify dates

```
## Not run:
files <- extractFiles("C:\\Users\\usr\\Pictures\\")
## End(Not run)</pre>
```

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checkFile

Check for files existence and prompt user if they want to load

Description

Check for files existence and prompt user if they want to load

Usage

```
checkFile(file)
```

Arguments

file

the full path of the file to check

Value

a boolean indicating wether a file was found and the user wants to load or not

Examples

```
## Not run:
    checkFile("path/to/newfile.csv")
## End(Not run)
```

convertCoordinates

Convert bbox from Relative to Absolute Coordinates

Description

Each row is a MD bounding box, there can be multiple bounding boxes per image.

Usage

```
convertCoordinates(results)
```

Arguments

results

list of bounding boxes for each image

Value

A dataframe with one entry for each bounding box

cropImageGenerator 7

Examples

cropImageGenerator

Tensorflow data generator that crops images to bounding box.

Description

Creates an image data generator that crops images based on bounding box coordinates.

Usage

```
cropImageGenerator(
  files,
  boxes,
  resize_height = 456,
  resize_width = 456,
  standardize = FALSE,
  batch = 32
)
```

Arguments

files a vector of file names

boxes a data frame or matrix of bounding box coordinates in the format left, top, width,

height.

resize_height the height the cropped image will be resized to.
resize_width the width the cropped image will be resized to.

standardize standardize the image to the range 0 to 1, TRUE or FALSE.

batch the batch size for the image generator.

Value

A Tensorflow image data generator.

```
## Not run: #' dataset <- cropImageGenerator(images, boxes, standardize = FALSE, batch = batch)
```

cropImageTrainGenerator

Tensorflow data generator for training that crops images to bounding box.

Description

Creates an image data generator that crops images based on bounding box coordinates and returnes an image/label pair.

Usage

```
cropImageTrainGenerator(
  files,
  boxes,
  label,
  classes,
  resize_height = 456,
  resize_width = 456,
  standardize = FALSE,
  augmentation_color = FALSE,
  augmentation_geometry = FALSE,
  shuffle = FALSE,
  cache = FALSE,
  cache_dir = NULL,
  return_iterator = FALSE,
  batch = 32
)
```

a vector of file names

Arguments

files

a data frame or matrix of bounding box coordinates in the format left, top, width, boxes height. label a vector of labels a vector of all classes for the active model classes the height the cropped image will be resized to. resize_height the width the cropped image will be resized to. resize_width standardize standardize the image to the range 0 to 1, TRUE or FALSE. augmentation_color use data augmentation to change the color, TRUE or FALSE. augmentation_geometry use data augmentation to change the geometry of the images, TRUE or FALSE. shuffle return data pairas in random order, TRUE or FALSE.

detectObject 9

cache use caching to reduce reading from disk, TRUE or FALSE.

cache_dir directory used for caching, if none provided chaching will be done in memory.

return_iterator

Should an iterator be returned? If RALSE a tfdataset will be returned.

batch the batch size for the image generator.

Value

A Tensorflow image data generator.

Examples

```
## Not run:
dataset <- cropImageTrainGenerator(images, standardize = FALSE, batch = batch)
## End(Not run)</pre>
```

detectObject

Run MD on a Single Image

Description

Returns the MD bounding boxes, classes, confidence above the min_conf threshold for a single image. #' Requires a an mdsession is already loaded (see loadMDModel()) and the file path of the image in question.

Usage

```
detectObject(mdsession, imagefile, mdversion = 5, min_conf = 0.1)
```

Arguments

mdsession Should be the output from loadMDmodel(model)

imagefile The path for the image in question mdversion MegaDetector version, defaults to 5

min_conf Confidence threshold for returning bounding boxes, defaults to 0.1

Value

a list of MD bounding boxes, classes, and confidence for the image

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Examples

detectObjectBatch

Run MegaDetector on a batch of images

Description

Runs MD on a list of image filepaths. Can resume for a results file and will checkpoint the results after a set number of images

Usage

```
detectObjectBatch(
  mdsession,
  images,
  mdversion = 5,
  min_conf = 0.1,
  batch = 1,
  outfile = NULL,
  checkpoint = 5000
)
```

Arguments

mdsession should be the output from loadMDmodel(model)

images list of image filepaths

mdversion select MegaDetector version, defaults to 5

min_conf Confidence threshold for returning bounding boxes, defaults to 0.1

batch Process images in batches, defaults to 1

outfile File containing previously checkpointed results

checkpoint Bank results after processing a number of images, defaults to 5000

Value

a list of lists of bounding boxes for each image

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Examples

```
## Not run:
images <- read_exif(imagedir,
   tags = c("filename", "directory", "DateTimeOriginal", "FileModifyDate"),
   recursive = TRUE)
colnames(images)[1] <- "FilePath"
mdsession <- loadMDModel(mdmodel)
mdres <- classifyImagesBatchMD(mdsession, images$FilePath,
   outfile = mdoutfile, checkpoint = 2500)
## End(Not run)</pre>
```

 ${\tt extractBoxes}$

Extract bounding boxes and save as new image from a batch of images

Description

Extract bounding boxes and save as new image from a batch of images

Usage

```
extractBoxes(
  images,
  min_conf = 0,
  buffer = 0,
  save = FALSE,
  resize = NA,
  outdir = "",
  quality = 0.8,
  parallel = FALSE,
  nproc = parallel::detectCores()
)
```

Arguments

| images | list of MD output or flat data.frame |
|----------|---|
| min_conf | Confidence threshold (defaults to 0, not in use) |
| buffer | Adds a buffer to the MD bounding box, defaults to 2px |
| save | Toggle to save output cropped, defaults to FALSE |
| resize | Size in pixels to resize cropped images, NA if images are not resized, defaults to NA |
| outdir | Directory in which output cropped images will be saved |
| quality | Compression level of output cropped image, defaults to 0.8 |
| parallel | Toggle to enable parallel processing, defaults to FALSE |
| nproc | Number of workers if parallel = TRUE, defaults to output of detectCores() |

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Details

A variable crop_rel_path in the image list or data.frame can be used to change the path where the crops will be stored.

The final output path will be the outdir plus the crop_rel_path.

Value

a flattened dataframe containing crop information

Examples

```
## Not run:
images <- read_exif(imagedir, tags = c("filename", "directory"), recursive = TRUE)
crops <- extractAllBoxes(images,save=TRUE,out)
## End(Not run)</pre>
```

Description

Extract crops from a single image represented by a processed dataframe

Usage

```
extractBoxesFromFlat(
  image,
  min_conf = 0,
  buffer = 0,
  save = TRUE,
  resize = NA,
  outdir = "",
  quality = 0.8
)
```

Arguments

| ımage | dataframe containing MD output (assumes single row) |
|----------|---|
| min_conf | Confidence threshold (defaults to 0, not in use) |
| buffer | Adds a buffer to the MD bounding box, defaults to 2px |
| save | Toggle to save output cropped, defaults to FALSE |
| resize | Size in pixels to resize cropped images, NA if images are not resized, defaults to NA |
| outdir | Directory in which output cropped images will be saved |
| quality | Compression level of output cropped image, defaults to 0.8 |
| | |

extractBoxesFromMD 13

Details

A variable crop_rel_path in the image list can be used to change the path where the crops will be stored.

The final output path will be the outdir plus the crop_rel_path.

Value

A dataframe containing image and crop paths

Examples

```
## Not run:
crops <- extractBoxesFromFlat(mdresflat[1, ], save = TRUE, out)
## End(Not run)</pre>
```

extractBoxesFromMD

Extract bounding boxes for a single image and save as new images

Description

Requires the unflattened raw MD output

Usage

```
extractBoxesFromMD(
  image,
  min_conf = 0,
  buffer = 0,
  return.crops = FALSE,
  save = FALSE,
  resize = NA,
  outdir = "",
  quality = 0.8
)
```

Arguments

| image | single image, raw MD output format (list) |
|--------------|---|
| min_conf | Confidence threshold (defaults to 0, not in use) |
| buffer | Adds a buffer to the MD bounding box, defaults to 2px |
| return.crops | Toggle to return list of cropped images, defaults to FALSE |
| save | Toggle to save output cropped, defaults to FALSE |
| resize | Size in pixels to resize cropped images, NA if images are not resized, defaults to NA |
| outdir | Directory in which output cropped images will be saved |
| quality | Compression level of output cropped image, defaults to 0.8 |

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Details

A variable crop_rel_path in the image list can be used to change the path where the crops will be stored.

The final output path will be the outdir plus the crop_rel_path.

Value

a flattened data.frame containing crop information

Examples

```
## Not run:
images <- read_exif(imagedir, tags = c("filename","directory"), recursive = TRUE)
crops <- extractBoxesFromMD(images[1, ], return.crops = TRUE, save = TRUE)
## End(Not run)</pre>
```

getAnimals

Return a dataframe of only MD animals

Description

Return a dataframe of only MD animals

Usage

```
getAnimals(manifest)
```

Arguments

manifest all megadetector frames

Value

animal frames classified by MD

```
## Not run:
animals <- getAnimals(imagesall)
## End(Not run)</pre>
```

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getEmpty

Return MD empty, vehicle and human images in a dataframe

Description

Return MD empty, vehicle and human images in a dataframe

Usage

```
getEmpty(manifest)
```

Arguments

manifest

all megadetector frames

Value

list of empty/human/vehicle allframes with md classification

Examples

```
## Not run:
empty <- getEmpty(imagesall)
## End(Not run)</pre>
```

 $image Augmentation {\tt Color}$

Perform image augmentation through random color adjustments on an image/label pair.

Description

Performs image augmentation on a image/label pair for training. Uses random brightness, contrast, saturation, and hue.

Usage

```
imageAugmentationColor(image, label, rng)
```

Arguments

image an image tensor.label a label tensor.

rng a random number generator use to generate a random seed.

16 ImageGenerator

Value

An image and label tensor.

```
imageAugmentationGeometry
```

Perform random geometric transformations on an image.

Description

Returns a keras model that performs random geometric transformations on an image.

Usage

```
imageAugmentationGeometry()
```

Value

A keras model.

ImageGenerator

Tensorflow data generator that resizes images.

Description

Creates an image data generator that resizes images if requested.

Usage

```
ImageGenerator(
   files,
   resize_height = NULL,
   resize_width = NULL,
   standardize = FALSE,
   batch = 1
)
```

Arguments

files a vector of file names

resize_height the height the cropped image will be resized to. If NULL returns original size

images

resize_width the width the cropped image will be resized to. If NULL returns original size

images..

standardize standardize the image to the range 0 to 1, TRUE or FALSE.

batch the batch size for the image generator.

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Value

A Tensorflow image data generator.

Examples

```
## Not run:
dataset <- ImageGenerator(images, standardize = FALSE, batch = batch)
## End(Not run)</pre>
```

ImageGeneratorSize

Tensorflow data generator that resizes images and returns original image size.

Description

Creates an image data generator that resizes images if requested and also returns the original images size needed for MegaDetector.

Usage

```
ImageGeneratorSize(
  files,
  resize_height = NULL,
  resize_width = NULL,
  pad = FALSE,
  standardize = FALSE,
  batch = 1
)
```

Arguments

files a vector of file names

resize_height the height the cropped image will be resized to. If NULL returns original size

images.

resize_width the width the cropped image will be resized to. If NULL returns original size

images..

pad pad the image instead of stretching it, TRUE or FALSE. standardize standardize the image to the range 0 to 1, TRUE or FALSE.

batch the batch size for the image generator.

Value

A Tensorflow image data generator.

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Examples

```
## Not run:
dataset <- ImageGenerator(images, standardize = FALSE, batch = batch)
## End(Not run)</pre>
```

imageLabel Load image and return a tensor with an image and a corresponding

label.

Description

Load image and return a tensor with an image and a corresponding label. Internal function to be called by image generator function.

Usage

```
imageLabel(data, classes, height = 299, width = 299, standardize = FALSE)
```

Arguments

data a list with the first element being an image file path and the second element a

label.

classes list of classes

height the height the cropped image will be resized to.
width the width the cropped image will be resized to.
standardize standardize the image, TRUE or FALSE.

Value

An image and label tensor.

| imageLabelCrop | Load image, crop and return a tensor with an image and a correspond- |
|----------------|--|
| | ing label. |

Description

Load image, crop and return a tensor with an image and a corresponding label. Internal function to be called by image generator function.

Usage

```
imageLabelCrop(data, classes, height = 299, width = 299, standardize = FALSE)
```

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Arguments

data a list with the first element being an image file path, the next four elements being

the bounding box coordinates and the last element a label

classes list of classes

height the height the cropped image will be resized to.
width the width the cropped image will be resized to.
standardize standardize the image, TRUE or FALSE.

Value

An image and label tensor.

imagesFromVideos

Extract frames from video for classification

Description

This function can take

Usage

```
imagesFromVideos(
   files,
   outdir = tempfile(),
   outfile = NULL,
   format = "jpg",
   fps = NULL,
   frames = NULL,
   parallel = FALSE,
   workers = 1,
   checkpoint = 1000
)
```

Arguments

files dataframe of videos outdir directory to save frames to outfile file to which results will be saved format output format for frames, defaults to jpg fps frames per second, otherwise determine mathematically frames number of frames to sample parallel Toggle for parallel processing, defaults to FALSE workers number of processors to use if parallel, defaults to 1 if not parallel, checkpoint ever n files, defaults to 1000 checkpoint

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Value

dataframe of still frames for each video

Examples

```
## Not run:
frames <- imagesFromVideos(videos, outdir = "C:\\Users\\usr\\Videos\\", frames = 5)
## End(Not run)</pre>
```

loadData

Load .csv or .Rdata file

Description

Load .csv or .Rdata file

Usage

```
loadData(file)
```

Arguments

file

the full path of the file to load

Value

data extracted from the file

```
## Not run:
   loadData("path/to/newfile.csv")
## End(Not run)
```

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loadImage

Load an image and return the full size image as an image tensor.

Description

Load an image and return the full size an image tensor. Internal function to be called by image generator function.

Usage

```
loadImage(file, standardize = FALSE)
```

Arguments

file path to a JPEG file

standardize standardize the image, TRUE or FALSE.

Value

An image tensor.

loadImageResize

Load and resize an image and return an image tensor.

Description

Load and resize an image and return an image tensor. Internal function to be called by image generator function.

Usage

```
loadImageResize(
  file,
  height = 299,
  width = 299,
  pad = FALSE,
  standardize = FALSE
)
```

Arguments

file path to a JPEG file

height the height the cropped image will be resized to.
width the width the cropped image will be resized to.

pad logical indicating whether the images should be padded or streched.

standardize standardize the image, TRUE or FALSE.

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Value

An image tensor.

loadImageResizeCrop

Load, resize and crop an image and return an image tensor.

Description

Load a JPEG image and crop it to a bounding box. Internal function to be called by image generator function.

Usage

```
loadImageResizeCrop(data, height = 299, width = 299, standardize = FALSE)
```

Arguments

data a list with the first element being a path to an image file and the next four argu-

ments being the bounding box coordinates.

height the height the cropped image will be resized to.
width the width the cropped image will be resized to.

standardize standardize the image, TRUE or FALSE.

Value

A Tensorflow image data generator.

loadImageResizeSize

Load and resize an image and return an image tensor as well as a tensor with the original image size.

Description

Load and resize an image and return an image tensor as well as a tensor with the original image size. Internal function to be called by image generator function.

Usage

```
loadImageResizeSize(
  file,
  height = 299,
  width = 299,
  pad = FALSE,
  standardize = FALSE
)
```

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Arguments

file path to a JPEG file

height the height the cropped image will be resized to.
width the width the cropped image will be resized to.

pad pad the image instead of stretching it, TRUE or FALSE.

standardize standardize the image, TRUE or FALSE.

Value

An image tensor.

loadMDModel

Load MegaDetector model file from directory or file

Description

Load MegaDetector model file from directory or file

Usage

```
loadMDModel(modelfile)
```

Arguments

modelfile

.pb file or directory obtained from megaDetector

Value

a tfsession containing the MD model

```
## Not run:
mdmodel <- "megadetector_v4.1.pb"
mdsession <- loadMDModel(mdmodel)
## End(Not run)</pre>
```

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parseMD

parse MD results into a simple dataframe

Description

parse MD results into a simple dataframe

Usage

```
parseMD(mdresults, manifest = NULL, outfile = NULL)
```

Arguments

mdresults raw MegaDetector output

manifest dataframe containing all frames outfile file path to save dataframe to

Value

original dataframe including md results

Examples

```
## Not run:
mdresults <- parseMD(mdres)
## End(Not run)</pre>
```

parseMDjson

converte the JSON file produced by the Python version of MegaDetector into the format produced by detectObjectBatch

Description

converte the JSON file produced by the Python version of MegaDetector into the format produced by detectObjectBatch

Usage

```
parseMDjson(json)
```

Arguments

json

json data in a list format

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Value

```
a list of MegaDetector results
```

Examples

```
## Not run:
mdresults <- parseMDjson(json)
## End(Not run)</pre>
```

plotBoxes

Plot bounding boxes on image from md results

Description

Plot bounding boxes on image from md results

Usage

```
plotBoxes(image, label = FALSE, minconf = 0)
```

Arguments

image The mdres for the image

label T/F toggle to plot MD category
minconf minimum confidence to plot box

Value

no return value, produces bounding box in plot panel

```
## Not run:
mdres <- classifyImageMD(mdsession, images$FilePath[30000])
plotBoxes(mdres, minconf = 0.5)
## End(Not run)</pre>
```

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predictSpecies

Classifies Crops Using Specified Models

Description

Classifies Crops Using Specified Models

Usage

```
predictSpecies(
  input,
  model,
  resize = 456,
  standardize = FALSE,
  batch = 1,
  workers = 1
)
```

Arguments

input either dataframe with MD crops or list of filenames

model models with which to classify species

resize resize images before classification, defaults to 299x299px

standardize standardize images, defaults to FALSE

batch number of images processed in each batch (keep small)

workers number of cores

Value

a matrix of likelihoods for each class for each image

process YOLO5 27

| processY0L05 Process Y0L05 outp | out and convert to MD format |
|---------------------------------|------------------------------|
|---------------------------------|------------------------------|

Description

Returns a list with the standard MD output format. Used for batch processing

Usage

```
processYOLO5(n, boxes, classes, scores, selection, batch)
```

Arguments

boxes array of boxes returned by combined_non_max_suppression classes vector of classes returned by combined_non_max_suppression

scores vector of probabilities returned by combined_non_max_suppression

selection vector of number of detected boxes returned by combined_non_max_suppression

batch batch used to detect objects

Value

a list of MD bounding boxes, classes, and confidence for the image

resizePad Resize an image with padding

Description

Resize an image with padding

Usage

```
resizePad(img, size = 256)
```

Arguments

img the image, read by jpeg library

size new size

Value

returns resized jpeg image

Examples

```
## Not run:
crop <- resizePad(cropped_image_path, 256)
## End(Not run)</pre>
```

saveData

Save Data to Given File

Description

Save Data to Given File

Usage

```
saveData(data, outfile, prompt = TRUE)
```

Arguments

data the dataframe to be saved outfile the full path of the saved file

prompt if true, prompts the user to confirm overwrite

Value

none

Examples

```
## Not run:
    saveData(files, "path/to/newfile.csv")
## End(Not run)
```

sequenceClassification

Leverage sequences to classify images

Description

This function applies image classifications at a sequence level by leveraging information from multiple images. A sequence is defined as all images at the same camera/station where the time between consecutive images is <=maxdiff. This can improve classification accuracy, but assumes that only one species is present in each sequence. If you regularly expect multiple species to occur in an image or sequence don't use this function.

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Usage

```
sequenceClassification(
  animals,
  empty = NULL,
  predictions,
  classes,
  emptyclass = "",
  stationcolumn,
  sortcolumns = NULL,
  maxdiff = 60
)
```

Arguments

animals

optional, data frame non-animal images (empty, human and vehicle) that will be merged back with animal imagages

predictions data frame of prediction probabilities from the classifySpecies function

sub-selection of all images that contain MD animals

classes a vector or species corresponding to the columns of 'predictions' emptyclass a string indicating the class that should be considered 'Empty'

stationcolumn a column in the animals and empty data frame that indicates the camera or cam-

era station

sortcolumns optional sort order. The default is 'stationcolumnumn' and DateTime.

maximum difference between images in seconds to be included in a sequence,

defaults to 60

Details

This function retains "Empty" classification even if other images within the sequence are predicted to contain animals. Classification confidence is weighted by MD confidence.

Value

data frame with predictions and confidence values for animals and empty images

30 symlinkMD

setupDirectory

Set Working Directory and Save File Global Variables

Description

Set Working Directory and Save File Global Variables

Usage

```
setupDirectory(workingdir, pkg.env)
```

Arguments

workingdir local directory that contains data to process pkg.env environment to create global variables in

Value

None

Examples

```
## Not run:
setupDirectory(/home/kyra/animl/examples)
## End(Not run)
```

symlinkMD

Create SymLink Directories and Sort Classified Images Based on MD Results

Description

Create SymLink Directories and Sort Classified Images Based on MD Results

Usage

```
symlinkMD(manifest, linkdir, outfile = NULL, copy = FALSE)
```

Arguments

| manifest | DataFrame of classified images |
|----------|------------------------------------|
| linkdir | Destination directory for symlinks |

outfile Results file to save to

copy Toggle to determine copy or hard link, defaults to link

symlinkSpecies 31

Value

manifest with added link columns

Examples

```
## Not run:
symlinkMD(manifest, linkdir)
## End(Not run)
```

symlinkSpecies

Create SymLink Directories and Sort Classified Images

Description

Create SymLink Directories and Sort Classified Images

Usage

```
symlinkSpecies(manifest, linkdir, threshold = 0, outfile = NULL, copy = FALSE)
```

Arguments

manifest DataFrame of classified images
linkdir Destination directory for symlinks

threshold Confidence threshold for determining uncertain predictions, defaults to 0

outfile Results file to save to

copy Toggle to determine copy or hard link, defaults to link

Value

manifest with added link columns

```
## Not run:
manifest <- symlinkSpecies(manifest, linkdir)
## End(Not run)</pre>
```

32 testMD

symUnlink

Remove Symlinks

Description

Remove Symlinks

Usage

```
symUnlink(manifest)
```

Arguments

manifest

DataFrame of classified images

Value

manifest without link column

Examples

```
## Not run:
symlinkMD(manifest, linkdir)
## End(Not run)
```

testMD

Select a Random Image and Run Through MegaDetector

Description

Select a Random Image and Run Through MegaDetector

Usage

```
testMD(input, mdsession, mdversion = 5, minconf = 0)
```

Arguments

input dataframe of all images mdsession MegaDetector mdsession

mdversion megadetector version, defaults to 5

minconf minimum confidence with which to draw boxes, defaults to 0

updateResults 33

Value

Null, plots box on image

Examples

```
## Not run:
testMD(input, mdsession)
## End(Not run)
```

updateResults

Title

Description

Title

Usage

```
updateResults(resultsfile, linkdir)
```

Arguments

resultsfile final results file with predictions, expects a "UniqueName" column linkdir symlink directory that has been validated

Value

dataframe with new "Species" column that contains the verifed species

```
## Not run:
results <- updateResults(resultsfile, linkdir)
## End(Not run)</pre>
```

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