# Package 'UCSCXenaTools'

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```
Title Download and Explore Datasets from UCSC Xena Data Hubs
Version 1.4.8
Maintainer Shixiang Wang <w_shixiang@163.com>
Description Download and explore datasets from UCSC Xena data hubs, which
     are a collection of UCSC-hosted public databases such as TCGA, ICGC,
     TARGET, GTEx, CCLE, and others. Databases are normalized so they can
     be combined, linked, filtered, explored and downloaded.
License GPL-3
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availTCGA

Get or Check TCGA Available ProjectID, DataType and FileType

### Description

Get or Check TCGA Available ProjectID, DataType and FileType

### Usage

```
availTCGA(which = c("all", "ProjectID", "DataType", "FileType"))
```

### Arguments

which a character of c("All", "ProjectID", "DataType", "FileType")

### Author(s)

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### **Examples**

```
availTCGA("all")
```

cohorts

Get cohorts of XenaHub object

### Description

Get cohorts of XenaHub object

### Usage

```
cohorts(x)
```

### Arguments

Х

a XenaHub object

#### Value

a character vector contains cohorts

### **Examples**

```
xe = XenaGenerate(subset = XenaHostNames == "tcgaHub"); cohorts(xe)
```

datasets

Get datasets of XenaHub object

### Description

Get datasets of XenaHub object

### Usage

```
datasets(x)
```

### Arguments

Х

a XenaHub object

### Value

a character vector contains datasets

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#### **Examples**

```
xe = XenaGenerate(subset = XenaHostNames == "tcgaHub"); datasets(xe)
```

downloadTCGA

Easily Download TCGA Data by Several Options

### **Description**

TCGA is a very useful database and here we provide this function to download TCGA (include TCGA Pancan) datasets in human-friendly way. Users who are not familiar with R operation will benefit from this.

### Usage

```
downloadTCGA(
  project = NULL,
  data_type = NULL,
  file_type = NULL,
  destdir = tempdir(),
  force = FALSE,
   ...
)
```

### **Arguments**

project	default is NULL. Should be one or more of TCGA project id (character vector) provided by Xena. See all available project id, please use availTCGA("ProjectID").
data_type	default is NULL. Should be a character vector specify data type. See all available data types by availTCGA("DataType").
file_type	default is NULL. Should be a character vector specify file type. See all available file types by availTCGA("FileType").
destdir	specify a location to store download data. Default is system temp directory.
force	logical. if TRUE, force to download data no matter whether files exist. Default is FALSE.
	other argument to download.file function

#### **Details**

All availble information about datasets of TCGA can access vis availTCGA() and check with showTCGA().

#### Value

same as XenaDownload() function result.

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#### Author(s)

Shixiang Wang w\_shixiang@163.com

#### See Also

```
XenaQuery(), XenaFilter(), XenaDownload(), XenaPrepare(), availTCGA(), showTCGA()
```

### **Examples**

fetch

Fetch Data from UCSC Xena Hosts

### **Description**

When you want to query just data for several genes/samples from UCSC Xena datasets, a better way is to use these fetch\_ functions instead of downloading a whole dataset. Details about functions please see the following sections.

#### **Usage**

```
fetch(host, dataset)

fetch_dense_values(
  host,
  dataset,
  identifiers = NULL,
  samples = NULL,
  check = TRUE,
   use_probeMap = FALSE,
  time_limit = 30
)

fetch_sparse_values(host, dataset, genes, samples = NULL, time_limit = 30)

fetch_dataset_samples(host, dataset, limit = NULL)

fetch_dataset_identifiers(host, dataset)

has_probeMap(host, dataset, return_url = FALSE)
```

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#### Arguments

host a UCSC Xena host, like "https://toil.xenahubs.net". All available hosts can be

printed by xena\_default\_hosts().

dataset a UCSC Xena dataset, like "tcga\_RSEM\_gene\_tpm". All available datasets can

be printed by running XenaData\$XenaDatasets or obtained from UCSC Xena

datapages.

etc.. If it is NULL, all identifiers in the dataset will be used.

samples ID of samples, like "TCGA-02-0047-01". If it is NULL, all samples in the dataset

will be used. However, it is better to download the whole datasets if you query

many samples and genes.

check if TRUE, check whether specified identifiers and samples exist the dataset

(all failed items will be filtered out). However, if FALSE, the code is much faster.

use\_probeMap if TRUE, will check if the dataset has ProbeMap firstly. When the dataset you

want to query has a identifier-to-gene mapping, identifiers can be gene symbols

even the identifiers of dataset are probes or others.

time\_limit time limit for getting response in seconds.

genes gene names.

limit number of samples, if NULL, return all samples.

return\_url if TRUE, returns the info of probeMap instead of a logical value when the result

exists.

#### **Details**

There are three primary data types: dense matrix (samples by probes (or say identifiers)), sparse (sample, position, variant), and segmented (sample, position, value).

Dense matrices can be genotypic or phenotypic, it is a sample-by-identifiers matrix. Phenotypic matrices have associated field metadata (descriptive names, codes, etc.). Genotypic matricies may have an associated probeMap, which maps probes to genomic locations. If a matrix has hugo probeMap, the probes themselves are gene names. Otherwise, a probeMap is used to map a gene location to a set of probes.

### Value

a matirx or character vector or a list.

#### **Functions**

- fetch\_dense\_values: fetches values from a dense matrix.
- fetch\_sparse\_values: fetches values from a sparse data.frame.
- fetch\_dataset\_samples: fetches samples from a dataset
- fetch\_dataset\_identifiers: fetches identifies from a dataset.
- has\_probeMap: checks if a dataset has ProbeMap.

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#### **Examples**

```
library(UCSCXenaTools)
host <- "https://toil.xenahubs.net"
dataset <- "tcga_RSEM_gene_tpm"
samples <- c("TCGA-02-0047-01", "TCGA-02-0055-01", "TCGA-02-2483-01", "TCGA-02-2485-01")
probes <- c("ENSG00000282740.1", "ENSG0000000005.5", "ENSG000000000419.12")
genes <- c("TP53", "RB1", "PIK3CA")

# Fetch samples
fetch_dataset_samples(host, dataset, 2)
# Fetch identifiers
fetch_dataset_identifiers(host, dataset)
# Fetch expression value by probes
fetch_dense_values(host, dataset, probes, samples, check = FALSE)
# Fetch expression value by gene symbol (if the dataset has probeMap)
has_probeMap(host, dataset)
fetch_dense_values(host, dataset, genes, samples, check = FALSE, use_probeMap = TRUE)</pre>
```

getTCGAdata

Get TCGA Common Data Sets by Project ID and Property

#### Description

This is the most useful function for user to download common TCGA datasets, it is similar to getFirehoseData function in RTCGAToolbox package.

#### Usage

```
getTCGAdata(
  project = NULL,
  clinical = TRUE,
  download = FALSE,
  forceDownload = FALSE,
  destdir = tempdir(),
  mRNASeq = FALSE,
  mRNAArray = FALSE,
  mRNASeqType = "normalized",
  miRNASeq = FALSE,
  exonRNASeq = FALSE,
  RPPAArray = FALSE,
  ReplicateBaseNormalization = FALSE,
  Methylation = FALSE,
  MethylationType = c("27K", "450K"),
  GeneMutation = FALSE,
  SomaticMutation = FALSE,
```

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```
GisticCopyNumber = FALSE,
Gistic2Threshold = TRUE,
CopyNumberSegment = FALSE,
RemoveGermlineCNV = TRUE,
...
)
```

#### **Arguments**

project default is NULL. Should be one or more of TCGA project id (character vector)

provided by Xena. See all available project id, please use availTCGA("ProjectID").

clinical logical. if TRUE, download clinical information. Default is TRUE.

download logical. if TRUE, download data, otherwise return a result list include data infor-

mation. Default is FALSE. You can set this to FALSE if you want to check what you will download or use other function provided by UCSCXenaTools to filter

result datasets you want to download.

forceDownload logical. if TRUE, force to download files no matter if exist. Default is FALSE. destdir specify a location to store download data. Default is system temp directory.

mRNASeq logical. if TRUE, download mRNASeq data. Default is FALSE.

mRNAArray logical. if TRUE, download mRNA microarray data. Default is FALSE.

mRNASeqType character vector. Can be one, two or three in c("normalized", "pancan normalized",

"percentile").

miRNASeq logical. if TRUE, download miRNASeq data. Default is FALSE. exonRNASeq logical. if TRUE, download exon RNASeq data. Default is FALSE.

RPPAArray logical. if TRUE, download RPPA data. Default is FALSE.

ReplicateBaseNormalization

logical. if TRUE, download RPPA data by Replicate Base Normalization (RBN).

Default is FALSE.

Methylation logical. if TRUE, download DNA Methylation data. Default is FALSE.

MethylationType

character vector. Can be one or two in c("27K", "450K").

GeneMutation logical. if TRUE, download gene mutation data. Default is FALSE.

SomaticMutation

logical. if TRUE, download somatic mutation data. Default is FALSE.

GisticCopyNumber

logical. if TRUE, download Gistic2 Copy Number data. Default is FALSE.

Gistic2Threshold

logical. if TRUE, download Threshold Gistic2 data. Default is TRUE.

 ${\tt CopyNumberSegment}$ 

logical. if TRUE, download Copy Number Segment data. Default is FALSE.

RemoveGermlineCNV

logical. if TRUE, download Copy Number Segment data which has removed

germline copy number variation. Default is TRUE.

... other argument to download.file function

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#### **Details**

TCGA Common Data Sets are frequently used for biological analysis. To make easier to achieve these data, this function provide really easy options to choose datasets and behavior. All availble information about datasets of TCGA can access vis availTCGA() and check with showTCGA().

#### Value

if download=TRUE, return data. frame from XenaDownload, otherwise return a list including XenaHub object and datasets information

#### Author(s)

Shixiang Wang w\_shixiang@163.com

#### **Examples**

```
###### get data, but not download

# 1 choose project and data types you wanna download
getTCGAdata(project = "LUAD", mRNASeq = TRUE, mRNAArray = TRUE,
mRNASeqType = "normalized", miRNASeq = TRUE, exonRNASeq = TRUE,
RPPAArray = TRUE, Methylation = TRUE, MethylationType = "450K",
GeneMutation = TRUE, SomaticMutation = TRUE)

# 2 only choose 'LUAD' and its clinical data
getTCGAdata(project = "LUAD")
## Not run:
###### download datasets

# 3 download clinical datasets of LUAD and LUSC
getTCGAdata(project = c("LUAD", "LUSC"), clinical = TRUE, download = TRUE)

# 4 download clinical, RPPA and gene mutation datasets of LUAD and LUSC
# getTCGAdata(project = c("LUAD", "LUSC"), clinical = TRUE, RPPAArray = TRUE, GeneMutation = TRUE)

## End(Not run)
```

hosts

Get hosts of XenaHub object

#### **Description**

Get hosts of XenaHub object

#### Usage

hosts(x)

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### **Arguments**

x a XenaHub object

#### Value

a character vector contains hosts

### **Examples**

```
xe = XenaGenerate(subset = XenaHostNames == "tcgaHub"); hosts(xe)
```

samples

Get Samples of a XenaHub object according to 'by' and 'how' action arguments

### **Description**

One is often interested in identifying samples or features present in each data set, or shared by all data sets, or present in any of several data sets. Identifying these samples, including samples in arbitrarily chosen data sets.

### Usage

```
samples(
    x,
    i = character(),
    by = c("hosts", "cohorts", "datasets"),
    how = c("each", "any", "all")
)
```

### **Arguments**

```
x a XenaHub object
i default is a empty character, it is used to specify the host, cohort or dataset by by option otherwise info will be automatically extracted by code
by a character specify by action
how a character specify how action
```

#### Value

a list include samples

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### **Examples**

```
## Not run:
xe = XenaHub(cohorts = "Cancer Cell Line Encyclopedia (CCLE)")
# samples in each dataset, first host
x = samples(xe, by="datasets", how="each")[[1]]
lengths(x)  # data sets in ccle cohort on first (only) host
## End(Not run)
```

showTCGA

Show TCGA data structure by Project ID or ALL

### **Description**

This can used to check if data type or file type exist in one or more projects by hand.

### Usage

```
showTCGA(project = "all")
```

### Arguments

project

a character vector. Can be "all" or one or more of TCGA Project IDs.

### Value

a data.frame including project data structure information.

### Author(s)

```
Shixiang Wang w_shixiang@163.com
```

#### See Also

```
availTCGA()
```

```
showTCGA("all")
```

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to\_snake

Convert camel case to snake case

### Description

Convert camel case to snake case

### Usage

```
to_snake(name)
```

### Arguments

name

a character vector

#### Value

same length as name but with snake case

### **Examples**

```
to_snake("sparseDataRange")
```

XenaBrowse

View Info of Dataset or Cohort at UCSC Xena Website Using Webbrowser

### Description

This will open dataset/cohort link of UCSC Xena in user's default browser.

### Usage

```
XenaBrowse(x, type = c("dataset", "cohort"), multiple = FALSE)
```

#### **Arguments**

```
x a XenaHub object.
```

type one of "dataset" and "cohort".

multiple if TRUE, browse multiple links instead of throwing error.

```
XenaGenerate(subset = XenaHostNames == "tcgaHub") %>%
  XenaFilter(filterDatasets = "clinical") %>%
  XenaFilter(filterDatasets = "LUAD") -> to_browse
```

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XenaData

Xena Hub Information

### Description

This data.frame is very useful for selecting datasets fastly and independent on APIs of UCSC Xena Hubs.

#### **Format**

A tibble.

#### Source

Generated from UCSC Xena Data Hubs.

### **Examples**

```
data(XenaData)
str(XenaData)
```

XenaDataUpdate

Get or Update Newest Data Information of UCSC Xena Data Hubs

### **Description**

Get or Update Newest Data Information of UCSC Xena Data Hubs

### Usage

```
XenaDataUpdate(saveTolocal = TRUE)
```

### **Arguments**

saveTolocal

logical. Whether save to local R package data directory for permanent use or Not.

#### Value

a data. frame contains all datasets information of Xena.

### Author(s)

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#### **Examples**

```
## Not run:
XenaDataUpdate()
XenaDataUpdate(saveTolocal = TRUE)
## End(Not run)
```

XenaDownload

Download Datasets from UCSC Xena Hubs

### **Description**

Avaliable datasets list: https://xenabrowser.net/datapages/

### Usage

```
XenaDownload(
  xquery,
  destdir = tempdir(),
  download_probeMap = FALSE,
  trans_slash = FALSE,
  force = FALSE,
  max_try = 3L,
  ...
)
```

### Arguments

xquery a tibble object generated by XenaQuery function.

destdir specify a location to store download data. Default is system temp directory.

download\_probeMap

if TRUE, also download ProbeMap data, which used for id mapping.

trans\_slash logical, default is FALSE. If TRUE, transform slash '/' in dataset id to '\_\_'. This

option is for backwards compatibility.

force logical. if TRUE, force to download data no matter whether files exist. Default is

FALSE.

max\_try time limit to try downloading the data.
... other argument to download.file function

#### Value

a tibble

#### Author(s)

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#### **Examples**

```
## Not run:
xe = XenaGenerate(subset = XenaHostNames == "tcgaHub")
hosts(xe)
xe_query = XenaQuery(xe)
xe_download = XenaDownload(xe_query)
## End(Not run)
```

XenaFilter

Filter a XenaHub Object

### Description

One of main functions in **UCSCXenatools**. It is used to filter XenaHub object according to cohorts, datasets. All datasets can be found at <a href="https://xenabrowser.net/datapages/">https://xenabrowser.net/datapages/</a>.

#### Usage

```
XenaFilter(
    x,
    filterCohorts = NULL,
    filterDatasets = NULL,
    ignore.case = TRUE,
    ...
)
```

### Arguments

```
x a XenaHub object

filterCohorts default is NULL. A character used to filter cohorts, regular expression is supported.

filterDatasets default is NULL. A character used to filter datasets, regular expression is supported.

ignore.case if FALSE, the pattern matching is case sensitive and if TRUE, case is ignored during matching.

... other arguments except value passed to base::grep().
```

#### Value

a XenaHub object

### Author(s)

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#### **Examples**

```
# operate TCGA datasets
xe = XenaGenerate(subset = XenaHostNames == "tcgaHub")
xe
# get all names of clinical data
xe2 = XenaFilter(xe, filterDatasets = "clinical")
datasets(xe2)
```

XenaGenerate

Generate and Subset a XenaHub Object from 'XenaData'

### **Description**

Generate and Subset a XenaHub Object from 'XenaData'

### Usage

```
XenaGenerate(XenaData = UCSCXenaTools::XenaData, subset = TRUE)
```

#### **Arguments**

XenaData a data.frame. Default is data(XenaData). The input of this option can only

be data(XenaData) or its subset.

subset logical expression indicating elements or rows to keep.

### Value

a XenaHub object.

#### Author(s)

Shixiang Wang w\_shixiang@163.com

```
# 1 get all datasets
XenaGenerate()
# 2 get TCGA BRCA
XenaGenerate(subset = XenaCohorts == "TCGA Breast Cancer (BRCA)")
# 3 get all datasets containing BRCA
XenaGenerate(subset = grepl("BRCA", XenaCohorts))
```

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XenaHub

Generate a XenaHub Object

#### **Description**

It is used to generate original XenaHub object according to hosts, cohorts, datasets or hostName. If these arguments not specified, all hosts and corresponding datasets will be returned as a XenaHub object. All datasets can be found at https://xenabrowser.net/datapages/.

#### Usage

### **Arguments**

hosts a character vector specify UCSC Xena hosts, all available hosts can be found by

xena\_default\_hosts() function. hostName is a more recommend option.

cohorts default is empty character vector, all cohorts will be returned.

datasets default is empty character vector, all datasets will be returned.

hostName name of host, available options can be accessed by .xena\_hosts This is an

easier option for user than hosts option. Note, this option will overlap hosts.

#### Value

a XenaHub object

#### Author(s)

Shixiang Wang w\_shixiang@163.com

```
## Not run:
#1 query all hosts, cohorts and datasets
xe = XenaHub()
xe
#2 query only TCGA hosts
xe = XenaHub(hostName = "tcgaHub")
xe
hosts(xe) # get hosts
```

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```
cohorts(xe) # get cohorts
datasets(xe) # get datasets
samples(xe) # get samples
## End(Not run)
```

XenaHub-class

Class XenaHub

### Description

a S4 class to represent UCSC Xena Data Hubs

### **Slots**

hosts hosts of data hubs cohorts cohorts of data hubs datasets datasets of data hubs

XenaPrepare

Prepare (Load) Downloaded Datasets to R

### **Description**

Prepare (Load) Downloaded Datasets to R

### Usage

```
XenaPrepare(
  objects,
  objectsName = NULL,
  use_chunk = FALSE,
  chunk_size = 100,
  subset_rows = TRUE,
  select_cols = TRUE,
  callback = NULL,
  comment = "#",
  na = c("", "NA", "[Discrepancy]"),
  ...
)
```

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### Arguments

objects	a object of character vector or data.frame. If objects is data.frame, it should be returned object of XenaDownload function. More easier way is that objects can be character vector specify local files/directory and download urls.
objectsName	specify names for elements of return object, i.e. names of list
use_chunk	default is FALSE. If you want to select subset of original data, please set it to TRUE and specify corresponding arguments: chunk_size, select_direction, select_names, callback.
chunk_size	the number of rows to include in each chunk
subset_rows	logical expression indicating elements or rows to keep: missing values are taken as false. x can be a representation of data frame you wanna do subset operation. Of note, the first colname of most of datasets in Xena will be set to "sample", you can use it to select rows.
select_cols	expression, indicating columns to select from a data frame. 'x' can be a representation of data frame you wanna do subset operation, e.g. select_cols = colnames(x)[1:3] will keep only first to third column.
callback	a function to call on each chunk, default is NULL, this option will overvide operations of subset_rows and select_cols.
comment	a character specify comment rows in files
na	a character vectory specify NA values in files
•••	other arguments transfer to read_tsv function or read_tsv_chunked function (when use_chunk is TRUE) of readr package.

#### Value

a list contains file data, which in way of tibbles

### Author(s)

```
Shixiang Wang w_shixiang@163.com
```

```
## Not run:
xe = XenaGenerate(subset = XenaHostNames == "tcgaHub")
hosts(xe)
xe_query = XenaQuery(xe)

xe_download = XenaDownload(xe_query)
dat = XenaPrepare(xe_download)
## End(Not run)
```

XenaQuery

Query URL of Datasets before Downloading

### Description

Query URL of Datasets before Downloading

### Usage

```
XenaQuery(x)
```

### Arguments

Χ

a XenaHub object

### Value

```
a data. frame contains hosts, datasets and url
```

### Author(s)

Shixiang Wang w\_shixiang@163.com

### **Examples**

```
xe = XenaGenerate(subset = XenaHostNames == "tcgaHub")
hosts(xe)
## Not run:
xe_query = XenaQuery(xe)
## End(Not run)
```

XenaQueryProbeMap

Query ProbeMap URL of Datasets

### Description

If dataset has no ProbeMap, it will be ignored.

### Usage

```
XenaQueryProbeMap(x)
```

### Arguments

Χ

a XenaHub object

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#### Value

```
a data. frame contains hosts, datasets and url
```

#### Author(s)

```
Shixiang Wang w_shixiang@163.com
```

### **Examples**

```
xe = XenaGenerate(subset = XenaHostNames == "tcgaHub")
hosts(xe)
## Not run:
xe_query = XenaQueryProbeMap(xe)
## End(Not run)
```

XenaScan

Scan all rows according to user input by a regular expression

#### **Description**

XenaScan() is a function can be used before XenaGenerate().

### Usage

```
XenaScan(
  XenaData = UCSCXenaTools::XenaData,
  pattern = NULL,
  ignore.case = TRUE
)
```

#### **Arguments**

XenaData a data.frame. Default is data(XenaData). The input of this option can only

be data(XenaData) or its subset.

pattern character string containing a regular expression (or character string for fixed =

TRUE) to be matched in the given character vector. Coerced by as.character to a character string if possible. If a character vector of length 2 or more is supplied, the first element is used with a warning. Missing values are allowed

except for regexpr, gregexpr and regexec.

ignore.case if FALSE, the pattern matching is case sensitive and if TRUE, case is ignored

during matching.

#### Value

```
a data.frame
```

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### **Examples**

```
x1 <- XenaScan(pattern = "Blood")
x2 <- XenaScan(pattern = "LUNG", ignore.case = FALSE)
x1 %>%
   XenaGenerate()
x2 %>%
   XenaGenerate()
```

XenaShiny

Xena Shiny App

### Description

Xena Shiny App

### Usage

XenaShiny()

xena\_default\_hosts

UCSC Xena Default Hosts

### Description

Return Xena default hosts

### Usage

```
xena_default_hosts()
```

### Value

A character vector include current defalut hosts

### Author(s)

Shixiang Wang w\_shixiang@163.com

### See Also

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