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# 1 Overview

MyGene.Info provides simple-to-use REST web services to query/retrieve gene annotation data. It's designed with simplicity and performance emphasized. *mygene* is an easy-to-use R wrapper to access MyGene.Info services.

# 2 Gene Annotation Service

# 2.1 getGene

• Use getGene, the wrapper for GET query of "/gene/<geneid>" service, to return the gene object for the given geneid.

```
> gene <- getGene("1017", fields="all")</pre>
> length(gene)
[1] 36
> gene$name
[1] "cyclin-dependent kinase 2"
> gene$taxid
[1] 9606
> gene$uniprot
$`Swiss-Prot`
[1] "P24941"
$TrEMBL
[1] "B4DDL9" "E7ESI2" "G3V317" "G3V5T9"
> gene$refseq
$genomic
[1] "AC_000144"
                   "NC_000012"
                                                   "NG_034014"
                                   "NC_018923"
                                                                  "NT_029419"
[6] "NW_001838059" "NW_004929384"
$rna
[1] "NM_001290230" "NM_001798"
                                   "NM_052827"
$protein
[1] "NP_001277159" "NP_001789"
                                   "NP_439892"
```

# 2.2 getGenes

• Use getGenes, the wrapper for POST query of "/gene" service, to return the list of gene objects for the given character vector of geneids.

```
> getGenes(c("1017","1018","ENSG00000148795"))
chr "symbol,name,taxid,entrezgene"
DataFrame with 3 rows and 6 columns
```

				<character></character>	<character></character>	<integer></integer>	<integer></integer>
1			cyclin-depend	dent kinase 2	CDK2	9606	1017
2			cyclin-depend	dent kinase 3	CDK3	9606	1018
3	cytochrome P450,	family 17,	subfamily A, ]	polypeptide 1	CYP17A1	9606	1586
	query	_id					
	<character></character>	<character></character>					
1	1017	1017					
2	1018	1018					
3	ENSG00000148795	1586					

# 3 Gene Query Service

#### 3.1 query

• Use query, a wrapper for GET query of "/query?q=<query>" service, to return the query result.

```
> query(q="cdk2", size=5)
```

```
$hits
  entrezgene
                                                         symbol
                                                                   _id taxid
                                         name
                                                 _score
1
                    cyclin-dependent kinase 2 389.77063
                                                           CDK2
                                                                  1017 9606
        1017
2
       12566
                    cyclin-dependent kinase 2 348.08942
                                                           Cdk2 12566 10090
3
                    cyclin dependent kinase 2 276.61142
      362817
                                                           Cdk2 362817 10116
4
       52004
                    CDK2-associated protein 2 20.83584 Cdk2ap2 52004 10090
5
      78832 CDK2 associated, cullin domain 1 18.44532 Cacul1
                                                                 78832 10090
```

```
$max_score
[1] 389.7706

$took
[1] 7

$total
[1] 26
> query(q="NM_013993")
$hits
```

entrezgene name \_score symbol \_id taxid 1 780 discoidin domain receptor tyrosine kinase 1 0.524516 DDR1 780 9606

\$max\_score
[1] 0.524516

\$took

[1] 17

\$total

[1] 1

# 3.2 queryMany

• Use queryMany, a wrapper for POST query of "/query" service, to return the batch query result.

```
> queryMany(c('1053_at', '117_at', '121_at', '1255_g_at', '1294_at'),
+ scopes="reporter", species="human")
```

#### Finished

DataFrame with 5 rows and 6 columns

rezgene	taxid	symbol	name	
nteger>	<integer></integer>	<character></character>	<character></character>	
5982	9606	RFC2	actor C (activator 1) 2, 40kDa	1 replication
3310	9606	HSPA6	hock 70kDa protein 6 (HSP70B')	2 heat
7849	9606	PAX8	paired box 8	3
2978	9606	GUCA1A	cyclase activator 1A (retina)	4 guanylat
7318	9606	UBA7	e modifier activating enzyme 7	5 ubiquitin-li
			_id	query
			character>	<character></character>
			5982	1 1053_at
			3310	2 117_at
			7849	3 121_at
			2978	4 1255_g_at
			7318	5 1294_at
	9606	GUCA1A	cyclase activator 1A (retina) e modifier activating enzyme 7 _id character> 5982 3310 7849 2978	4 guanylat 5 ubiquitin-li query <character> 1 1053_at 2 117_at 3 121_at 4 1255_g_at</character>

# 4 makeTxDbFromMyGene

TxDb is a container for storing transcript annotations. makeTxDbFromMyGene allows the user to make a TxDb object in the Genomic Features package from a mygene "exons" query using a default mygene object.

```
> xli <- c('DDX26B',
+ 'CCDC83',
+ 'MAST3',
+ 'RPL11',
+ 'ZDHHC2O',
+ 'LUC7L3',
+ 'SNORD49A',
+ 'CTSH',
+ 'ACOT8')</pre>
```

```
> txdb <- makeTxDbFromMyGene(xli,
+ scopes="symbol", species="human")
> transcripts(txdb)
```

GRanges object with 15 ranges and 2 metadata columns:

	seqnames		ranges	$\operatorname{strand}$	- 1	tx_id	<pre>tx_name</pre>
	<rle></rle>		Ranges	<rle></rle>	- 1	<integer></integer>	<character></character>
[1]	1	[24018268,	24022915]	+	- 1	5	NM_001199802
[2]	1	[24018268,	24022915]	+	- 1	6	NM_000975
[3]	11	[85566143,	85631063]	+	- 1	2	NM_001286159
[4]	11	[85566143,	85631063]	+	- 1	3	NM_173556
[5]	13	[21946709,	22033508]	_	1	10	NM_153251
[11]	17	[ 48796925,	48830072]	+		11	NM_016424
[12]	17	[ 48796925,	48830072]	+	- 1	12	NM_006107
[13]	19	[ 18208602,	18262499]	+	- 1	4	NM_015016
[14]	20	[ 44470359,	44486048]	_	- 1	15	NM_005469
[15]	Х	[134654554,	134716460]	+		1	NM_182540

seqinfo: 8 sequences from an unspecified genome; no seqlengths

makeTxDbFromMyGene invokes either the query or queryMany method and passes the response to construct a TxDb object. See ?TxDb for methods to utilize and access transcript annotations.

# 5 Tutorial, ID mapping

ID mapping is a very common, often not fun, task for every bioinformatician. Supposedly you have a list of gene symbols or reporter ids from an upstream analysis, and then your next analysis requires to use gene ids (e.g. Entrez gene ids or Ensembl gene ids). So you want to convert that list of gene symbols or reporter ids to corresponding gene ids.

Here we want to show you how to do ID mapping quickly and easily.

# 5.1 Mapping gene symbols to Entrez gene ids

Suppose xli is a list of gene symbols you want to convert to entrez gene ids:

```
> xli <- c('DDX26B',
+ 'CCDC83',
+ 'MAST3',
+ 'FLOT1',
+ 'RPL11',
+ 'ZDHHC20',
+ 'LUC7L3',</pre>
```

```
+ 'SNORD49A',
+ 'CTSH',
+ 'ACOT8')
```

You can then call queryMany method, telling it your input is symbol, and you want entrezgene (Entrez gene ids) back.

> queryMany(xli, scopes="symbol", fields="entrezgene", species="human")

#### Finished

DataFrame with 10 rows and 3 columns

	query	entrezgene	_id
	<character></character>	<integer></integer>	<character></character>
1	DDX26B	203522	203522
2	CCDC83	220047	220047
3	MAST3	23031	23031
4	FLOT1	10211	10211
5	RPL11	6135	6135
6	ZDHHC20	253832	253832
7	LUC7L3	51747	51747
8	SNORD49A	26800	26800
9	CTSH	1512	1512
10	ACOT8	10005	10005

# 5.2 Mapping gene symbols to Ensembl gene ids

Now if you want Ensembl gene ids back:

> out <- queryMany(xli, scopes="symbol", fields="ensembl.gene", species="human")
Finished</pre>

> out

DataFrame with 10 rows and 3 columns

	ensembl.gene	_id	query
	<characterlist></characterlist>	<character></character>	<character></character>
1	ENSG00000165359	203522	DDX26B
2	ENSG00000150676	220047	CCDC83
3	ENSG00000099308	23031	MAST3
4	ENSG00000137312, ENSG00000206379, ENSG00000206480,	10211	FLOT1
5	ENSG00000142676	6135	RPL11
6	ENSG00000180776	253832	ZDHHC20
7	ENSG00000108848	51747	LUC7L3
8	ENSG00000277370, ENSG00000175061	26800	SNORD49A
9	ENSG00000103811	1512	CTSH
10	ENSG00000101473	10005	ACOT8

```
> out$ensembl.gene[[4]]
```

```
[1] "ENSG00000137312" "ENSG00000206379" "ENSG00000206480" "ENSG00000223654"
```

```
[5] "ENSG00000224740" "ENSG00000230143" "ENSG00000232280" "ENSG00000236271"
```

### 5.3 When an input has no matching gene

In case that an input id has no matching gene, you will be notified from the output. The returned list for this query term contains notfound value as True.

```
> xli <- c('DDX26B',
+ 'CCDC83',
+ 'MAST3',
+ 'FLOT1',
+ 'RPL11',
+ 'Gm10494')
> queryMany(xli, scopes="symbol", fields="entrezgene", species="human")
```

#### Finished

Pass returnall=TRUE to return lists of duplicate or missing query terms. DataFrame with 6 rows and 4 columns

	query	entrezgene	_id	notfound
	<character></character>	<integer></integer>	<character></character>	<logical></logical>
1	DDX26B	203522	203522	NA
2	CCDC83	220047	220047	NA
3	MAST3	23031	23031	NA
4	FLOT1	10211	10211	NA
5	RPL11	6135	6135	NA
6	Gm10494	NA	NA	TRUE

# 5.4 When input ids are not just symbols

```
> xli <- c('DDX26B',
+ 'CCDC83',
+ 'MAST3',
+ 'FLOT1',
+ 'RPL11',
+ 'Gm10494',
+ '1007_s_at',
+ 'AK125780')
>
```

Above id list contains symbols, reporters and accession numbers, and supposedly we want to get back both Entrez gene ids and uniprot ids. Parameters scopes, fields, species are all flexible enough to support multiple values, either a list or a comma-separated string:

```
> out <- queryMany(xli, scopes=c("symbol", "reporter", "accession"),
+ fields=c("entrezgene", "uniprot"), species="human")</pre>
```

Finished

Pass returnall=TRUE to return lists of duplicate or missing query terms.

> out

DataFrame with 9 rows and 6 columns

	query	${\tt entrezgene}$	<pre>uniprot.Swiss-Prot</pre>	${\tt uniprot.TrEMBL}$	_id	notfound
	<character></character>	<integer></integer>	<character></character>	<list></list>	<character></character>	<logical></logical>
1	DDX26B	203522	Q5JSJ4	#######	203522	NA
2	CCDC83	220047	Q8IWF9	#######	220047	NA
3	MAST3	23031	060307	#######	23031	NA
4	FLOT1	10211	075955	#######	10211	NA
5	RPL11	6135	P62913	#######	6135	NA
6	Gm10494	NA	NA	#######	NA	TRUE
7	1007_s_at	780	Q08345	#######	780	NA
8	1007_s_at	100616237	NA	#######	100616237	NA
9	AK125780	2978	P43080	#######	2978	NA

> out\$`uniprot.Swiss-Prot`[[5]]

[1] "P62913"

# 5.5 When an input id has multiple matching genes

From the previous result, you may have noticed that query term 1007\_s\_at matches two genes. In that case, you will be notified from the output, and the returned result will include both matching genes.

By passing returnall=TRUE, you will get both duplicate or missing query terms

- > queryMany(xli, scopes=c("symbol", "reporter", "accession"),
- + fields=c("entrezgene", "uniprot"), species='human', returnall=TRUE)

Finished

\$response

DataFrame with 9 rows and 6 columns

	query	${\tt entrezgene}$	${\tt uniprot.Swiss-Prot}$	${\tt uniprot.TrEMBL}$	_id	${\tt notfound}$
	<character></character>	<integer></integer>	<character></character>	<list></list>	<character></character>	<logical></logical>
1	DDX26B	203522	Q5JSJ4	#######	203522	NA
2	CCDC83	220047	Q8IWF9	#######	220047	NA
3	MAST3	23031	060307	#######	23031	NA
4	FLOT1	10211	075955	#######	10211	NA
5	RPL11	6135	P62913	#######	6135	NA
6	Gm10494	NA	NA	#######	NA	TRUE
7	1007_s_at	780	Q08345	#######	780	NA
8	1007_s_at	100616237	NA	#######	100616237	NA

9	AK125780	2978	P43080	#######	2978	NA
	plicates 1007_s_at					
1	2					

The returned result above contains out for mapping output, missing for missing query terms (a list), and dup for query terms with multiple matches (including the number of matches).

# 5.6 Can I convert a very large list of ids?

Yes, you can. If you pass an id list (i.e., xli above) larger than 1000 ids, we will do the id mapping in-batch with 1000 ids at a time, and then concatenate the results all together for you. So, from the user-end, it's exactly the same as passing a shorter list. You don't need to worry about saturating our backend servers. Large lists, however, may take a while longer to query, so please wait patiently.

### 6 References

\$missing

[1] "Gm10494"

Wu C, MacLeod I, Su AI (2013) BioGPS and MyGene.info: organizing online, gene-centric information. Nucl. Acids Res. 41(D1): D561-D565. help@mygene.info