Package 'AssessORFData'

December 4, 2025

Type Package

```
Title Data and Files for the AssessORF Package
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Description This package provides access to mapping and results objects generated by the AssessORF
      package, as well as the genome sequences for the strains corresponding to those objects.
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      Pseudomonas_aeruginosa_Data, Staphylococcus_aureus_Data,
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```

AP1

Contents

	AP1	2
	AssessORF_StrainIDs	4
	ATCC11842	4
	ATCC13032	6
	ATCC17978	7
	ATCC700084	9
	BW25113	10
	CCMP1375	12
	CECT5344	13
	CNRZ327	15
	СОН1	16
	D_UW_3_CX	18
	EGD e	19
	GetDataMapObj	21
	GetGeneSources	
	GetResultsObj	22
	GetStrainIDs	23
	H37Rv	23
	HG001	25
	Houston_1	26
	K_12_MG1655	
	LAL14 1	
	MG1363	32
	MGAS5005	34
	NCIB 3610	35
	PAO1	37
	SaveGenomeToPath	38
		39
	Strain10403S	41
	Strain168	
	TCH1516	44
Index		46

AP1

Assessment Objects for Streptococcus pyogenes strain AP1

Description

Objects of class Assessment and either subclass DataMap or subclass Results for $\it Streptococcus pyogenes strain AP1$

Usage

```
data(AP1_PreSaved_DataMapObj)
data(AP1_PreSaved_ResultsObj_GenBank)
data(AP1_PreSaved_ResultsObj_GeneMarkS2)
data(AP1_PreSaved_ResultsObj_Glimmer)
data(AP1_PreSaved_ResultsObj_Prodigal)
```

AP1 3

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Details

'AP1_PreSaved_DataMapObj' is an object of subclass DataMap.

 $'AP1_PreSaved_ResultsObj_GenBank' is an object of subclass \verb|Results| with predicted genes from the GenBank database.$

'AP1_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'AP1_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'AP1_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

• StrainID: AP1

• Species: S. pyogenes

Mapping object

The mapping object, 'AP1_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD006345. The related genomes used to determine evolutionary conservation all came from the family, Streptococcaceae exculding anomalous genomes and non-complete *Streptococcus pneumoniae* genomes. Links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'AP1_PreSaved_ResultsObj_GenBank', 'AP1_PreSaved_ResultsObj_GeneMarkS2', 'AP1_PreSaved_ResultsObj_Glimmer', and 'AP1_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("AP1", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

4 ATCC11842

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD006345 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/ GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/CP007537.1

AssessORF_StrainIDs Vector of strain IDs used in the package

Description

Vector of strain IDs, which describes the strains for which the package has data

Usage

```
data(AssessORF_StrainIDs)
```

Format

Character vector of length 26

ATCC11842

Assessment *Objects for* Lactobacillus delbrueckii *subsp.* bulgaricus *strain ATCC 11842*

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Lactobacillus delbrueckii* subsp. *bulgaricus* strain ATCC 11842

Usage

```
data(ATCC11842_PreSaved_DataMapObj)
data(ATCC11842_PreSaved_ResultsObj_GenBank)
data(ATCC11842_PreSaved_ResultsObj_GeneMarkS2)
data(ATCC11842_PreSaved_ResultsObj_Glimmer)
data(ATCC11842_PreSaved_ResultsObj_Prodigal)
```

Format

ATCC11842 5

Details

'ATCC11842_PreSaved_DataMapObj' is an object of subclass DataMap.

'ATCC11842_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'ATCC11842_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'ATCC11842_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'ATCC11842_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

• StrainID: ATCC11842

• Species: L. delbrueckii bulgaricus

Mapping object

The mapping object, 'ATCC11842_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD006551. The related genomes used to determine evolutionary conservation all came from the genus Lactobacillus, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'ATCC11842_PreSaved_ResultsObj_GenBank', 'ATCC11842_PreSaved_ResultsObj_GeneMarkS' 'ATCC11842_PreSaved_ResultsObj_Glimmer', and 'ATCC11842_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("ATCC11842", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD006551 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/NC_008054.1

6 ATCC13032

ATCC13032 Assessment <i>Objects for</i> Corynebacterium glutamicum <i>strain ATC</i> 13032

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Corynebacterium glutamicum* strain ATCC 13032

Usage

```
data(ATCC13032_PreSaved_DataMapObj)
data(ATCC13032_PreSaved_ResultsObj_GenBank)
data(ATCC13032_PreSaved_ResultsObj_GeneMarkS2)
data(ATCC13032_PreSaved_ResultsObj_Glimmer)
data(ATCC13032_PreSaved_ResultsObj_Prodigal)
```

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Details

'ATCC13032_PreSaved_DataMapObj' is an object of subclass DataMap.

'ATCC13032_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'ATCC13032_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'ATCC13032_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'ATCC13032_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

StrainID: ATCC13032Species: C. glutamicum

Mapping object

The mapping object, 'ATCC13032_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD005812. The related genomes used to determine evolutionary conservation came from the genera Corynebacterium, Dietzia, and Tsukamurella, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

ATCC17978 7

Results objects

The 4 results objects, 'ATCC13032_PreSaved_ResultsObj_GenBank', 'ATCC13032_PreSaved_ResultsObj_GeneMarkS' 'ATCC13032_PreSaved_ResultsObj_Glimmer', and 'ATCC13032_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("ATCC13032", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD005812 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/ GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/BA000036.3

ATCC17978

Assessment Objects for Acinetobacter baumannii strain ATCC 17978

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Acinetobacter baumannii* strain ATCC 17978

Usage

```
data(ATCC17978_PreSaved_DataMapObj)
data(ATCC17978_PreSaved_ResultsObj_GenBank)
data(ATCC17978_PreSaved_ResultsObj_GeneMarkS2)
data(ATCC17978_PreSaved_ResultsObj_Glimmer)
data(ATCC17978_PreSaved_ResultsObj_Prodigal)
```

Format

8 ATCC17978

Details

'ATCC17978 PreSaved DataMapObj' is an object of subclass DataMap.

'ATCC17978_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'ATCC17978_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'ATCC17978_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'ATCC17978_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

StrainID: ATCC17978Species: A. baumannii

Mapping object

The mapping object, 'ATCC17978_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The proteomics data was generated in-house and collected as part of the work for the AssessORF paper. The raw mass spectra data can be found at ProteomeXchange dataset PXD012539. The related genomes used to determine evolutionary conservation all came from the genus Acinetobacter, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'ATCC17978_PreSaved_ResultsObj_GenBank', 'ATCC17978_PreSaved_ResultsObj_GeneMarkS' 'ATCC17978_PreSaved_ResultsObj_Glimmer', and 'ATCC17978_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("ATCC17978", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

ATCC700084 9

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD012539 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/CP000521.1

ATCC700084 Assessment *Objects for* Mycobacterium smegmatis *strain ATCC 700084*

Description

Objects of class Assessment and either subclass DataMap or subclass Results for Mycobacterium smegmatis strain ATCC 700084

Usage

```
data(ATCC700084_PreSaved_DataMapObj)
data(ATCC700084_PreSaved_ResultsObj_GenBank)
data(ATCC700084_PreSaved_ResultsObj_GeneMarkS2)
data(ATCC700084_PreSaved_ResultsObj_Glimmer)
data(ATCC700084_PreSaved_ResultsObj_Prodigal)
```

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Details

'ATCC700084_PreSaved_DataMapObj' is an object of subclass DataMap.

'ATCC700084_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'ATCC700084_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'ATCC700084_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'ATCC700084_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

StrainID: ATCC700084Species: M. smegmatis

Mapping object

The mapping object, 'ATCC700084_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD003500. The related genomes used to determine evolutionary conservation all came from the genus Mycobacterium, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the AssessORF package.

10 BW25113

Results objects

The 4 results objects, 'ATCC700084_PreSaved_ResultsObj_GenBank', 'ATCC700084_PreSaved_ResultsObj_GeneMan' ATCC700084_PreSaved_ResultsObj_Glimmer', and 'ATCC700084_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("ATCC700084", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine. Please note that there is no genome sequence available for strain ATCC 700084 so the reference genome from strain MC2 155 was used instead.

Source

```
Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD003500 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/NC_008596.1
```

BW25113

Assessment Objects for Escherichia coli strain BW25113

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Escherichia coli* strain BW25113

Usage

```
data(BW25113_PreSaved_DataMapObj)
data(BW25113_PreSaved_ResultsObj_GenBank)
data(BW25113_PreSaved_ResultsObj_GeneMarkS2)
data(BW25113_PreSaved_ResultsObj_Glimmer)
data(BW25113_PreSaved_ResultsObj_Prodigal)
```

Format

BW25113

Details

'BW25113_PreSaved_DataMapObj' is an object of subclass DataMap.

'BW25113_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'BW25113_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'BW25113_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'BW25113_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

StrainID: BW25113Species: E. coli

Mapping object

The mapping object, 'BW25113_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD000498. The related genomes used to determine evolutionary conservation all came from the genus Escherichia, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'BW25113_PreSaved_ResultsObj_GenBank', 'BW25113_PreSaved_ResultsObj_GeneMarkS2', 'BW25113_PreSaved_ResultsObj_Glimmer', and 'BW25113_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("BW25113", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD000498 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/NZ_CP009273.1

12 CCMP1375

CCMP1375	Assessment <i>Objects for</i> Prochlorococcus marinus <i>subsp.</i> marinus <i>strain CCMP1375</i>

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Prochlorococcus marinus* subsp. *marinus* strain CCMP1375

Usage

```
data(CCMP1375_PreSaved_DataMapObj)
data(CCMP1375_PreSaved_ResultsObj_GenBank)
data(CCMP1375_PreSaved_ResultsObj_GeneMarkS2)
data(CCMP1375_PreSaved_ResultsObj_Glimmer)
data(CCMP1375_PreSaved_ResultsObj_Prodigal)
```

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Details

'CCMP1375_PreSaved_DataMapObj' is an object of subclass DataMap.

'CCMP1375_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'CCMP1375_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'CCMP1375_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'CCMP1375_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

StrainID: CCMP1375Species: P. marinus

Mapping object

The mapping object, 'CCMP1375_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD005745. The related genomes used to determine evolutionary conservation all came from the order Synechococcales, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

CECT5344 13

Results objects

The 4 results objects, 'CCMP1375_PreSaved_ResultsObj_GenBank', 'CCMP1375_PreSaved_ResultsObj_GeneMarkS2' CCMP1375_PreSaved_ResultsObj_Glimmer', and 'CCMP1375_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("CCMP1375", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

```
Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD005745 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/NC_005042.1
```

CECT5344 Assessment Objects for Pseudomonas pseudoalcaligenes strain $CECT\,5344$

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Pseudomonas* pseudoalcaligenes strain CECT 5344

Usage

```
data(CECT5344_PreSaved_DataMapObj)
data(CECT5344_PreSaved_ResultsObj_GenBank)
data(CECT5344_PreSaved_ResultsObj_GeneMarkS2)
data(CECT5344_PreSaved_ResultsObj_Glimmer)
data(CECT5344_PreSaved_ResultsObj_Prodigal)
```

Format

14 CECT5344

Details

'CECT5344_PreSaved_DataMapObj' is an object of subclass DataMap.

'CECT5344_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'CECT5344_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'CECT5344_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'CECT5344_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

• StrainID: CECT5344

• Species: P. pseudoalcaligenes

Mapping object

The mapping object, 'CECT5344_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD005745. The related genomes used to determine evolutionary conservation all came from the genus Pseudomonas, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'CECT5344_PreSaved_ResultsObj_GenBank', 'CECT5344_PreSaved_ResultsObj_GeneMarkS2' 'CECT5344_PreSaved_ResultsObj_Glimmer', and 'CECT5344_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("CECT5344", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD005745 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/HG916826.1

CNRZ327 15

CNRZ327 Assessment <i>Objects for</i> Lactobacillus delbrueckii <i>subsp.</i> lactis <i>stra LBCNRZ327_V11</i>	ıin
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Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Lactobacillus delbrueckii* subsp. *lactis* strain LBCNRZ327_V11

Usage

```
data(CNRZ327_PreSaved_DataMapObj)
data(CNRZ327_PreSaved_ResultsObj_GenBank)
data(CNRZ327_PreSaved_ResultsObj_GeneMarkS2)
data(CNRZ327_PreSaved_ResultsObj_Glimmer)
data(CNRZ327_PreSaved_ResultsObj_Prodigal)
```

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Details

'CNRZ327_PreSaved_DataMapObj' is an object of subclass DataMap.

'CNRZ327_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'CNRZ327_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'CNRZ327_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'CNRZ327_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

• StrainID: CNRZ327

• Species: L. delbrueckii lactis

Mapping object

The mapping object, 'CNRZ327_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD006551. The related genomes used to determine evolutionary conservation all came from the genus Lactobacillus, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the <code>AssessORF</code> package.

16 COH1

Results objects

The 4 results objects, 'CNRZ327_PreSaved_ResultsObj_GenBank', 'CNRZ327_PreSaved_ResultsObj_GeneMarkS2', 'CNRZ327_PreSaved_ResultsObj_Glimmer', and 'CNRZ327_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("CNRZ327", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

```
Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD006551 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/CCDV01000001.1
```

COH1

Assessment Objects for Streptococcus agalactiae strain COH1

Description

Objects of class Assessment and either subclass DataMap or subclass Results for Streptococcus agalactiae strain COH1

Usage

```
data(COH1_PreSaved_DataMapObj)
data(COH1_PreSaved_ResultsObj_GenBank)
data(COH1_PreSaved_ResultsObj_GeneMarkS2)
data(COH1_PreSaved_ResultsObj_Glimmer)
data(COH1_PreSaved_ResultsObj_Prodigal)
```

Format

COH1 17

Details

'COH1_PreSaved_DataMapObj' is an object of subclass DataMap.

'COH1_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'COH1_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'COH1_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'COH1_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

• StrainID: COH1

• Species: S. agalactiae

Mapping object

The mapping object, 'COH1_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The proteomics data was generated in-house and collected as part of the work for the AssessORF paper. The raw mass spectra data can be found at ProteomeXchange dataset PXD012567. The related genomes used to determine evolutionary conservation all came from the family Streptococcaceae, exculding anomalous genomes and non-complete *Streptococcus pneumoniae* genomes. Links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'COH1_PreSaved_ResultsObj_GenBank', 'COH1_PreSaved_ResultsObj_GeneMarkS2', 'COH1_PreSaved_ResultsObj_Glimmer', and 'COH1_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("COH1", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

 $D_{_}UW_{_}3_{_}CX$

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD012567 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/ GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/HG939456.1

D_UW_3_CX

Assessment Objects for Chlamydia trachomatis strain D/UW-3/CX

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Chlamydia tra- chomatis* strain D/UW-3/CX

Usage

```
data(D_UW_3_CX_PreSaved_DataMapObj)
data(D_UW_3_CX_PreSaved_ResultsObj_GenBank)
data(D_UW_3_CX_PreSaved_ResultsObj_GeneMarkS2)
data(D_UW_3_CX_PreSaved_ResultsObj_Glimmer)
data(D_UW_3_CX_PreSaved_ResultsObj_Prodigal)
```

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Details

'D_UW_3_CX_PreSaved_DataMapObj' is an object of subclass DataMap.

'D_UW_3_CX_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'D_UW_3_CX_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'D_UW_3_CX_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'D_UW_3_CX_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

StrainID: D_UW_3_CXSpecies: C. trachomatis

Mapping object

The mapping object, 'D_UW_3_CX_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD003883. The related genomes used to determine evolutionary conservation all came from the phylum Chlamydiae, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

EGD_e 19

Results objects

The 4 results objects, 'D_UW_3_CX_PreSaved_ResultsObj_GenBank', 'D_UW_3_CX_PreSaved_ResultsObj_GeneMa' 'D_UW_3_CX_PreSaved_ResultsObj_Glimmer', and 'D_UW_3_CX_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("D_UW_3_CX", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

```
Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD003883 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/AE001273.1
```

EGD e

Assessment Objects for Listeria monocytogenes strain EGD-e

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Listeria mono-cytogenes* strain EGD-e

Usage

```
data(EGD_e_PreSaved_DataMapObj)
data(EGD_e_PreSaved_ResultsObj_GenBank)
data(EGD_e_PreSaved_ResultsObj_GeneMarkS2)
data(EGD_e_PreSaved_ResultsObj_Glimmer)
data(EGD_e_PreSaved_ResultsObj_Prodigal)
```

Format

 EGD_e

Details

'EGD_e_PreSaved_DataMapObj' is an object of subclass DataMap.

'EGD_e_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'EGD_e_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'EGD_e_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'EGD_e_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

• StrainID: EGD_e

• Species: L. monocytogenes

Mapping object

The mapping object, 'EGD_e_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD000890. The related genomes used to determine evolutionary conservation all came from the genus Listeria, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'EGD_e_PreSaved_ResultsObj_GenBank', 'EGD_e_PreSaved_ResultsObj_GeneMarkS2', 'EGD_e_PreSaved_ResultsObj_Glimmer', and 'EGD_e_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use $SaveGenomeToPath("EGD_e", <INSERT FILE PATH HERE>)$ to save the genome sequence for the strain to your local machine.

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD000890 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/NC_003210.1

GetDataMapObj 21

GetDataMapObj

Get a Data Map Object

Description

Gets and returns the data map object for a specific strain

Usage

```
GetDataMapObj(strainID)
```

Arguments

strainID

Character string corresponding to the strain identifier.

Details

GetDataMapObj returns an object of class Assessment and subclass DataMap corresponding to the given strain ID. The given strain ID must be a part of the AssessORF set, and there is no partial matching. Otherwise, the function will error.

Value

An object of class ${\tt Assessment}$ and subclass ${\tt DataMap}$

Examples

```
mapObj <- GetDataMapObj("MGAS5005")</pre>
```

GetGeneSources

Get the Gene Sources

Description

Returns the list of gene sources used in making the results objects

Usage

GetGeneSources()

Details

GetGeneSources returns the list of gene sources (programs and databases) used in making the results objects in the AssessORF set.

Value

A character vector where each element corresponds to a single gene source

22 GetResultsObj

Examples

```
geneSourceSet <- GetGeneSources()</pre>
```

GetResultsObj

Get a Results Object

Description

Gets and returns the results object for a specific strain-gene source combo

Usage

```
GetResultsObj(strainID, geneSource = "Prodigal")
```

Arguments

strainID Character string corresponding to the strain identifier.
geneSource Character string corresponding to the gene source.

Details

GetDataMapObj returns an object of class Assessment and subclass Results corresponding to the given strain ID and the given gene source. The given strain ID must be a part of the AssessORF set, and there is no partial matching. Otherwise, the function will error. The given gene source must also be a part of the AssessORF set, but the function ignores case when checking if the given gene source is a part of the set.

Value

An object of class Assessment and subclass Results

Examples

```
resObj1 <- GetResultsObj("MGAS5005", "Prodigal")
resObj2 <- GetResultsObj("MGAS5005", "GenBank")
resObj3 <- GetResultsObj("MGAS5005", "GeneMarkS2")
resObj4 <- GetResultsObj("MGAS5005", "Glimmer")</pre>
```

GetStrainIDs 23

GetStrainIDs

Get the Strain Identifiers

Description

Returns the list of strain identifiers for which the package has data

Usage

```
GetStrainIDs()
```

Details

GetStrainIDs returns the list of strains in the AssessORF set. This function is a shorter alternative to data("AssessORF_StrainIDs").

Value

A character vector where each element corresponds to a single strain identifier

Examples

```
allStrainIDs <- GetStrainIDs()</pre>
```

H37Rv

Assessment Objects for Mycobacterium tuberculosis strain H37Rv

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Mycobacterium tuberculosis* strain H37Rv

Usage

```
data(H37Rv_PreSaved_DataMapObj)
data(H37Rv_PreSaved_ResultsObj_GenBank)
data(H37Rv_PreSaved_ResultsObj_GeneMarkS2)
data(H37Rv_PreSaved_ResultsObj_Glimmer)
data(H37Rv_PreSaved_ResultsObj_Prodigal)
```

Format

24 H37Rv

Details

'H37Rv_PreSaved_DataMapObj' is an object of subclass DataMap.

'H37Rv_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'H37Rv_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'H37Rv_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

 $'H37Rv_PreSaved_ResultsObj_Prodigal' \ is \ an \ object \ of \ subclass \ Results \ with \ predicted \ genes \ from \ the \ program \ Prodigal.$

For all 5 objects, values of two key identifying list items within each object are listed below.

• StrainID: H37Rv

• Species: M. tuberculosis

Mapping object

The mapping object, 'H37Rv_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD006117. The related genomes used to determine evolutionary conservation all came from the genus Mycobacterium, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'H37Rv_PreSaved_ResultsObj_GenBank', 'H37Rv_PreSaved_ResultsObj_GeneMarkS2', 'H37Rv_PreSaved_ResultsObj_Glimmer', and 'H37Rv_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("H37Rv", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD006117 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/AL123456.3

HG001 25

HG001	Assessment Objects for Staphylococcus aureus subsp. aureus strain HG001

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Staphylococcus aureus* subsp. *aureus* strain HG001

Usage

```
data(HG001_PreSaved_DataMapObj)
data(HG001_PreSaved_ResultsObj_GenBank)
data(HG001_PreSaved_ResultsObj_GeneMarkS2)
data(HG001_PreSaved_ResultsObj_Glimmer)
data(HG001_PreSaved_ResultsObj_Prodigal)
```

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Details

'HG001_PreSaved_DataMapObj' is an object of subclass DataMap.

'HG001_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'HG001_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

 $'HG001_PreSaved_ResultsObj_Glimmer'$ is an object of subclass Results with predicted genes from the program Glimmer.

'HG001_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

StrainID: HG001Species: S. aureus

Mapping object

The mapping object, 'HG001_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD000702. The related genomes used to determine evolutionary conservation all came from the genus Staphylococcus, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

26 Houston_1

Results objects

The 4 results objects, 'HG001_PreSaved_ResultsObj_GenBank', 'HG001_PreSaved_ResultsObj_GeneMarkS2', 'HG001_PreSaved_ResultsObj_Glimmer', and 'HG001_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("HG001", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine. Note that this genome comes from accession CP018205.1, which has the same genome sequence as accession NZ_CP018205.1. Unlike NZ_CP018205.1 however, CP018205.1 does not have any associated GenBank genes.

Source

```
Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD000702
NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/
GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/NZ_CP018205.1
```

Houston_1

Assessment Objects for Bartonella henselae strain Houston-1

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Bartonella henselae* strain Houston-1

Usage

```
data(Houston_1_PreSaved_DataMapObj)
data(Houston_1_PreSaved_ResultsObj_GenBank)
data(Houston_1_PreSaved_ResultsObj_GeneMarkS2)
data(Houston_1_PreSaved_ResultsObj_Glimmer)
data(Houston_1_PreSaved_ResultsObj_Prodigal)
```

Format

Houston_1 27

Details

'Houston_1_PreSaved_DataMapObj' is an object of subclass DataMap.

'Houston_1_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'Houston_1_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'Houston_1_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'Houston_1_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

StrainID: Houston_1Species: B. henselae

Mapping object

The mapping object, 'Houston_1_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD000153. The related genomes used to determine evolutionary conservation came from the families Bartonellaceae, Brucellaceae, Phyllobacteriaceae, Rhizobiaceae. Links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'Houston_1_PreSaved_ResultsObj_GenBank', 'Houston_1_PreSaved_ResultsObj_GeneMarkS2', 'Houston_1_PreSaved_ResultsObj_Glimmer', and 'Houston_1_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("Houston_1", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD000153 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/BX897699.1

28 II1403

I11403

Assessment Objects for Lactococcus lactis subsp. lactis strain Il1403

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Lactococcus lactis* subsp. *lactis* strain II1403

Usage

```
data(Il1403_PreSaved_DataMapObj)
data(Il1403_PreSaved_ResultsObj_GenBank)
data(Il1403_PreSaved_ResultsObj_GeneMarkS2)
data(Il1403_PreSaved_ResultsObj_Glimmer)
data(Il1403_PreSaved_ResultsObj_Prodigal)
```

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Details

'II1403_PreSaved_DataMapObj' is an object of subclass DataMap.

'II1403_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'II1403_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'II1403_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'II1403_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

StrainID: Il1403Species: L. lactis

Mapping object

The mapping object, 'II1403_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD000494. The related genomes used to determine evolutionary conservation all came from the family, Streptococcaceae exculding anomalous genomes and non-complete *Streptococcus pneumoniae* genomes. Links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

K_12_MG1655

Results objects

The 4 results objects, 'Il1403_PreSaved_ResultsObj_GenBank', 'Il1403_PreSaved_ResultsObj_GeneMarkS2', 'Il1403_PreSaved_ResultsObj_Glimmer', and 'Il1403_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the AssessORF package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("Il1403", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

```
Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD000494
NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/
GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/AE005176.1
```

K_12_MG1655 Assessment Objects for Escherichia coli strain K-12 substrain MG1655

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Escherichia coli* strain K-12 substrain MG1655

Usage

```
data(K_12_MG1655_PreSaved_DataMapObj)
data(K_12_MG1655_PreSaved_ResultsObj_GenBank)
data(K_12_MG1655_PreSaved_ResultsObj_GeneMarkS2)
data(K_12_MG1655_PreSaved_ResultsObj_Glimmer)
data(K_12_MG1655_PreSaved_ResultsObj_Prodigal)
```

Format

30 K_12_MG1655

Details

'K_12_MG1655_PreSaved_DataMapObj' is an object of subclass DataMap.

 ${\rm `K_12_MG1655_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.}$

'K_12_MG1655_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'K_12_MG1655_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

 ${\rm `K_12_MG1655_PreSaved_ResultsObj_Prodigal' is an object of subclass \ Results \ with \ predicted genes from the program Prodigal.}$

For all 5 objects, values of two key identifying list items within each object are listed below.

• StrainID: K_12_MG1655

• Species: E. coli

Mapping object

The mapping object, 'K_12_MG1655_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD005901. The related genomes used to determine evolutionary conservation all came from the genus Escherichia, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'K_12_MG1655_PreSaved_ResultsObj_GenBank', 'K_12_MG1655_PreSaved_ResultsObj_Gene 'K_12_MG1655_PreSaved_ResultsObj_Glimmer', and 'K_12_MG1655_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("K_12_MG1655", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD005901 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/CP025268.1

LAL14_1 31

LAL14_1

Assessment Objects for Sulfolobus islandicus strain LAL14/1

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Sulfolobus islandicus* strain LAL14/1

Usage

```
data(LAL14_1_PreSaved_DataMapObj)
data(LAL14_1_PreSaved_ResultsObj_GenBank)
data(LAL14_1_PreSaved_ResultsObj_GeneMarkS2)
data(LAL14_1_PreSaved_ResultsObj_Glimmer)
data(LAL14_1_PreSaved_ResultsObj_Prodigal)
```

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Details

'LAL14_1_PreSaved_DataMapObj' is an object of subclass DataMap.

'LAL14_1_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'LAL14_1_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'LAL14_1_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'LAL14_1_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

StrainID: LAL14_1Species: S. islandicus

Mapping object

The mapping object, 'LAL14_1_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD003074. The related genomes used to determine evolutionary conservation all came from the phylum Crenarchaeota, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the <code>AssessORF</code> package.

32 MG1363

Results objects

The 4 results objects, 'LAL14_1_PreSaved_ResultsObj_GenBank', 'LAL14_1_PreSaved_ResultsObj_GeneMarkS2', 'LAL14_1_PreSaved_ResultsObj_Glimmer', and 'LAL14_1_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("LAL14_1", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD003074 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/ GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/CP003928.1

MG1363

Assessment *Objects for* Lactococcus lactis *subsp.* cremoris *strain MG1363*

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Lactococcus lactis* subsp. *cremoris* strain MG1363

Usage

```
data(MG1363_PreSaved_DataMapObj)
data(MG1363_PreSaved_ResultsObj_GenBank)
data(MG1363_PreSaved_ResultsObj_GeneMarkS2)
data(MG1363_PreSaved_ResultsObj_Glimmer)
data(MG1363_PreSaved_ResultsObj_Prodigal)
```

Format

MG1363 33

Details

'MG1363_PreSaved_DataMapObj' is an object of subclass DataMap.

'MG1363_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'MG1363_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'MG1363_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'MG1363_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

StrainID: MG1363Species: L. lactis

Mapping object

The mapping object, 'MG1363_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD011263. The related genomes used to determine evolutionary conservation all came from the family, Streptococcaceae exculding anomalous genomes and non-complete *Streptococcus pneumoniae* genomes. Links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'MG1363_PreSaved_ResultsObj_GenBank', 'MG1363_PreSaved_ResultsObj_GeneMarkS2', 'MG1363_PreSaved_ResultsObj_Glimmer', and 'MG1363_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("MG1363", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

34 MGAS5005

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD011263

NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/

GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/AM406671.1

MGAS5005

Assessment Objects for Streptococcus pyogenes strain MGAS5005

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Streptococcus pyogenes* strain MGAS5005

Usage

```
data(MGAS5005_PreSaved_DataMapObj)
data(MGAS5005_PreSaved_ResultsObj_GenBank)
data(MGAS5005_PreSaved_ResultsObj_GeneMarkS2)
data(MGAS5005_PreSaved_ResultsObj_Glimmer)
data(MGAS5005_PreSaved_ResultsObj_Prodigal)
```

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Details

'MGAS5005_PreSaved_DataMapObj' is an object of subclass DataMap.

'MGAS5005_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'MGAS5005_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'MGAS5005_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'MGAS5005_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

• StrainID: MGAS5005

• Species: S. pyogenes

NCIB_3610 35

Mapping object

The mapping object, 'MGAS5005_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The proteomics data was generated in-house and collected as part of the work for the AssessORF paper. The raw mass spectra data can be found at ProteomeXchange dataset PXD012568. The related genomes used to determine evolutionary conservation all came from the family, Streptococcaeae exculding anomalous genomes and non-complete *Streptococcus pneumoniae* genomes. Links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'MGAS5005_PreSaved_ResultsObj_GenBank', 'MGAS5005_PreSaved_ResultsObj_GeneMarkS' 'MGAS5005_PreSaved_ResultsObj_Glimmer', and 'MGAS5005_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("MGAS5005", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

 $Proteomcis\ data:\ http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD012568$

NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/

GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/CP000017.2

NCIB_3610 Assessment *Objects for* Bacillus subtilis *subsp.* subtilis *strain NCIB* 3610

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Bacillus subtilis* subsp. *subtilis* strain NCIB 3610

36 NCIB_3610

Usage

```
data(NCIB_3610_PreSaved_DataMapObj)
data(NCIB_3610_PreSaved_ResultsObj_GenBank)
data(NCIB_3610_PreSaved_ResultsObj_GeneMarkS2)
data(NCIB_3610_PreSaved_ResultsObj_Glimmer)
data(NCIB_3610_PreSaved_ResultsObj_Prodigal)
```

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Details

'NCIB_3610_PreSaved_DataMapObj' is an object of subclass DataMap.

'NCIB_3610_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'NCIB_3610_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'NCIB_3610_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'NCIB_3610_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

StrainID: NCIB_3610Species: B. subtilis

Mapping object

The mapping object, 'NCIB_3610_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD006444. The related genomes used to determine evolutionary conservation all came from the genus Bacillus, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'NCIB_3610_PreSaved_ResultsObj_GenBank', 'NCIB_3610_PreSaved_ResultsObj_GeneMarkS' 'NCIB_3610_PreSaved_ResultsObj_Glimmer', and 'NCIB_3610_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

PAO1 37

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("NCIB_3610", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

```
Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD006444
NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/
GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/NZ_CM000488.1
```

PA01

Assessment Objects for Pseudomonas aeruginosa strain PAO1

Description

Objects of class Assessment and either subclass DataMap or subclass Results for Pseudomonas aeruginosa strain PAO1

Usage

```
data(PA01_PreSaved_DataMapObj)
data(PA01_PreSaved_ResultsObj_GenBank)
data(PA01_PreSaved_ResultsObj_GeneMarkS2)
data(PA01_PreSaved_ResultsObj_Glimmer)
data(PA01_PreSaved_ResultsObj_Prodigal)
```

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Details

'PAO1_PreSaved_DataMapObj' is an object of subclass DataMap.

'PAO1_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'PAO1_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'PAO1_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'PAO1_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

• StrainID: PAO1

• Species: P. aeruginosa

38 SaveGenomeToPath

Mapping object

The mapping object, 'PAO1_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD004560. The related genomes used to determine evolutionary conservation all came from the genus Pseudomonas, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'PAO1_PreSaved_ResultsObj_GenBank', 'PAO1_PreSaved_ResultsObj_GeneMarkS2', 'PAO1_PreSaved_ResultsObj_Glimmer', and 'PAO1_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("PAO1", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD004560 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/AE004091.2

SaveGenomeToPath

Save a Strain's Genome to a Directory

Description

Saves the genome for a specified strain to a given directory

Usage

SaveGenomeToPath(strainID, filePath)

Arguments

strainID Character string corresponding to the strain identifier.

filePath Character string corresponding to the path to the file path. Must end in '.fasta'.

SL1344 39

Details

SaveGenomeToPath saves the genome for the specified strain ID to the given file path. If the file specified by the path already exists, it will be overwitten (with a warning). The given strain ID must be a part of the AssessORF set, and there is no partial matching. Otherwise, the function will error.

Note: there is no genome for strain ATCC700084, so a reference genome for the species (strain MC2155) is used instead.

Value

Invisibly returns filePath

Examples

```
tmpFile <- paste0(tempfile(), ".fasta")
SaveGenomeToPath("MGAS5005", tmpFile)
unlink(tmpFile)</pre>
```

SL1344

Assessment Objects for Salmonella enterica subsp. enterica serovar Typhimurium strain SL1344

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Salmonella enterica* subsp. *enterica* serovar Typhimurium strain SL1344

Usage

```
data(SL1344_PreSaved_DataMapObj)
data(SL1344_PreSaved_ResultsObj_GenBank)
data(SL1344_PreSaved_ResultsObj_GeneMarkS2)
data(SL1344_PreSaved_ResultsObj_Glimmer)
data(SL1344_PreSaved_ResultsObj_Prodigal)
```

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Details

'SL1344_PreSaved_DataMapObj' is an object of subclass DataMap.

'SL1344_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'SL1344_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'SL1344_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

40 SL1344

'SL1344_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

• StrainID: SL1344

• Species: S. typhimurium

Mapping object

The mapping object, 'SL1344_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD005579. The related genomes used to determine evolutionary conservation all came from the genus Salmonella, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'SL1344_PreSaved_ResultsObj_GenBank', 'SL1344_PreSaved_ResultsObj_GeneMarkS2', 'SL1344_PreSaved_ResultsObj_Glimmer', and 'SL1344_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("SL1344", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD005579

NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/

GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/FQ312003.1

Strain10403S 41

Strain10403S

Assessment Objects for Listeria monocytogenes strain 10403S

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Listeria monocytogenes* strain 10403S

Usage

```
data(Strain10403S_PreSaved_DataMapObj)
data(Strain10403S_PreSaved_ResultsObj_GenBank)
data(Strain10403S_PreSaved_ResultsObj_GeneMarkS2)
data(Strain10403S_PreSaved_ResultsObj_Glimmer)
data(Strain10403S_PreSaved_ResultsObj_Prodigal)
```

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Details

'Strain10403S_PreSaved_DataMapObj' is an object of subclass DataMap.

'Strain10403S_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'Strain10403S_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'Strain10403S_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'Strain10403S_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

• StrainID: Strain10403S

• Species: L. monocytogenes

Mapping object

The mapping object, 'Strain10403S_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD010000. The related genomes used to determine evolutionary conservation all came from the genus Listeria, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

42 Strain168

Results objects

The 4 results objects, 'Strain10403S_PreSaved_ResultsObj_GenBank', 'Strain10403S_PreSaved_ResultsObj_GeneMar 'Strain10403S_PreSaved_ResultsObj_Glimmer', and 'Strain10403S_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("Strain10403S", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

```
Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD010000 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/NC_017544.1
```

Strain168

Assessment Objects for Bacillus subtilis subsp. subtilis strain 168

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Bacillus subtilis* subsp. *subtilis* strain 168

Usage

```
data(Strain168_PreSaved_DataMapObj)
data(Strain168_PreSaved_ResultsObj_GenBank)
data(Strain168_PreSaved_ResultsObj_GeneMarkS2)
data(Strain168_PreSaved_ResultsObj_Glimmer)
data(Strain168_PreSaved_ResultsObj_Prodigal)
```

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Strain168 43

Details

'Strain168_PreSaved_DataMapObj' is an object of subclass DataMap.

'Strain168_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'Strain168_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'Strain168_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'Strain168_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

StrainID: Strain168Species: B. subtilis

Mapping object

The mapping object, 'Strain168_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The raw proteomics mass spectra data came from ProteomeXchange dataset PXD004565. The related genomes used to determine evolutionary conservation all came from the genus Bacillus, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

Results objects

The 4 results objects, 'Strain168_PreSaved_ResultsObj_GenBank', 'Strain168_PreSaved_ResultsObj_GeneMarkS2', 'Strain168_PreSaved_ResultsObj_Glimmer', and 'Strain168_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("Strain168", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

Proteomcis data: http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD004565 NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/ GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/CM000487.1 44 TCH1516

TCH1516 Assessment *Objects for* Staphylococcus aureus *subsp.* aureus *strain USA300_TCH1516*

Description

Objects of class Assessment and either subclass DataMap or subclass Results for *Staphylococcus aureus* subsp. *aureus* strain USA300_TCH1516

Usage

```
data(TCH1516_PreSaved_DataMapObj)
data(TCH1516_PreSaved_ResultsObj_GenBank)
data(TCH1516_PreSaved_ResultsObj_GeneMarkS2)
data(TCH1516_PreSaved_ResultsObj_Glimmer)
data(TCH1516_PreSaved_ResultsObj_Prodigal)
```

Format

All 5 objects have a list structure. For specifics on what the two types of Assessment objects contain, please see the **AssessORF** package.

Details

'TCH1516_PreSaved_DataMapObj' is an object of subclass DataMap.

'TCH1516_PreSaved_ResultsObj_GenBank' is an object of subclass Results with predicted genes from the GenBank database.

'TCH1516_PreSaved_ResultsObj_GeneMarkS2' is an object of subclass Results with predicted genes from the program GeneMarkS-2.

'TCH1516_PreSaved_ResultsObj_Glimmer' is an object of subclass Results with predicted genes from the program Glimmer.

'TCH1516_PreSaved_ResultsObj_Prodigal' is an object of subclass Results with predicted genes from the program Prodigal.

For all 5 objects, values of two key identifying list items within each object are listed below.

StrainID: TCH1516Species: S. aureus

Mapping object

The mapping object, 'TCH1516_PreSaved_DataMapObj', stores the mapping of proteomics evidence and evolutionary conservation evidence to the strain's genome. The proteomics data was generated in-house and collected as part of the work for the AssessORF paper. The raw mass spectra data can be found at ProteomeXchange dataset PXD012538. The related genomes used to determine evolutionary conservation all came from the genus Staphylococcus, and links to their sequences were downloaded from NCBI's Genome Browser. The object was built using the MapAssessmentData function from the **AssessORF** package.

TCH1516 45

Results objects

The 4 results objects, 'TCH1516_PreSaved_ResultsObj_GenBank', 'TCH1516_PreSaved_ResultsObj_GeneMarkS2', 'TCH1516_PreSaved_ResultsObj_Glimmer', and 'TCH1516_PreSaved_ResultsObj_Prodigal', store how much evidence there is supporting or against each gene in a set of predicted genes for the strain's genome. The four objects were built using the AssessGenes function from the **AssessORF** package, a set of genes, and the mapping object described here.

For the GenBank object, the genes were downloaded from the corresponding record in the GenBank database. For the other three objects, GenemarkS-2 (web server), Prodigal (2.6.3), and Glimmer (3.02) were run at default settings in order to generate the set of predicted genes for the corresponding results object.

The GeneLeftPos, GeneRightPos, and the GeneStrand within each of the four results objects provide positional information on the set of predicted genes used to generate that object.

The GeneSource list item within each of the four results objects describes where the predicted genes came from (either "GenBank", "GeneMarkS2", "Glimmer", or "Prodigal" respectively).

Getting the strain's genome

Use SaveGenomeToPath("TCH1516", <INSERT FILE PATH HERE>) to save the genome sequence for the strain to your local machine.

Source

 $Proteomcis\ data:\ http://proteomecentral.proteomexchange.org/cgi/GetDataset?ID=PXD012538$

NCBI's Genome Browser: https://www.ncbi.nlm.nih.gov/genome/browse/#!/prokaryotes/

GenBank record: https://www.ncbi.nlm.nih.gov/nuccore/CP000730.1

Index

* datasets	ATCC17978_PreSaved_ResultsObj_Prodigal
AssessORF_StrainIDs, 4	(ATCC17978), 7
	ATCC700084, 9
AP1, 2	ATCC700084_PreSaved_DataMapObj
AP1_PreSaved_DataMapObj (AP1), 2	(ATCC700084), 9
AP1_PreSaved_ResultsObj_GenBank (AP1), 2	ATCC700084_PreSaved_ResultsObj_GenBank
AP1_PreSaved_ResultsObj_GeneMarkS2	(ATCC700084), 9
(AP1), 2	ATCC700084_PreSaved_ResultsObj_GeneMarkS2
AP1_PreSaved_ResultsObj_Glimmer(AP1), 2	(ATCC700084), 9
AP1_PreSaved_ResultsObj_Prodigal (AP1),	ATCC700084_PreSaved_ResultsObj_Glimmer (ATCC700084), 9
AssessORF_StrainIDs, 4	ATCC700084_PreSaved_ResultsObj_Prodigal
ATCC11842, 4	(ATCC700084), 9
ATCC11842_PreSaved_DataMapObj	
(ATCC11842), 4	BW25113, 10
ATCC11842_PreSaved_ResultsObj_GenBank	BW25113_PreSaved_DataMapObj(BW25113),
(ATCC11842), 4	10
ATCC11842_PreSaved_ResultsObj_GeneMarkS2	BW25113_PreSaved_ResultsObj_GenBank
(ATCC11842), 4	(BW25113), 10
ATCC11842_PreSaved_ResultsObj_Glimmer	BW25113_PreSaved_ResultsObj_GeneMarkS2
(ATCC11842), 4	(BW25113), 10
ATCC11842_PreSaved_ResultsObj_Prodigal	BW25113_PreSaved_ResultsObj_Glimmer
(ATCC11842), 4	(BW25113), 10
ATCC13032, 6	BW25113_PreSaved_ResultsObj_Prodigal
ATCC13032_PreSaved_DataMapObj	(BW25113), 10
(ATCC13032), 6	CCMD127E 12
ATCC13032_PreSaved_ResultsObj_GenBank	CCMP1375, 12 CCMP1375_PreSaved_DataMapObj
(ATCC13032), 6	(CCMP1375), 12
ATCC13032_PreSaved_ResultsObj_GeneMarkS2	CCMP1375_PreSaved_ResultsObj_GenBank
(ATCC13032), 6	(CCMP1375), 12
ATCC13032_PreSaved_ResultsObj_Glimmer	CCMP1375_PreSaved_ResultsObj_GeneMarkS2
(ATCC13032), 6	(CCMP1375), 12
ATCC13032_PreSaved_ResultsObj_Prodigal	CCMP1375_PreSaved_ResultsObj_Glimmer
(ATCC13032), 6	(CCMP1375), 12
ATCC17978, 7	CCMP1375_PreSaved_ResultsObj_Prodigal
ATCC17978_PreSaved_DataMapObj	(CCMP1375), 12
(ATCC17978), 7	CECT5344, 13
ATCC17978_PreSaved_ResultsObj_GenBank	CECT5344_PreSaved_DataMapObj
(ATCC17978), 7	(CECT5344), 13
ATCC17978_PreSaved_ResultsObj_GeneMarkS2	CECT5344_PreSaved_ResultsObj_GenBank
(ATCC17978), 7	(CECT5344), 13
ATCC17978_PreSaved_ResultsObj_Glimmer	CECT5344_PreSaved_ResultsObj_GeneMarkS2
(ATCC17978), 7	(CECT5344), 13

INDEX 47

CECT5344_PreSaved_ResultsObj_Glimmer	H37Rv, 23
(CECT5344), 13	H37Rv_PreSaved_DataMapObj(H37Rv), 23
CECT5344_PreSaved_ResultsObj_Prodigal	H37Rv_PreSaved_ResultsObj_GenBank
(CECT5344), 13	(H37Rv), 23
CNRZ327, 15	H37Rv_PreSaved_ResultsObj_GeneMarkS2
CNRZ327_PreSaved_DataMapObj(CNRZ327),	(H37Rv), 23
15	H37Rv_PreSaved_ResultsObj_Glimmer
CNRZ327_PreSaved_ResultsObj_GenBank	(H37Rv), 23
(CNRZ327), 15	H37Rv_PreSaved_ResultsObj_Prodigal
CNRZ327_PreSaved_ResultsObj_GeneMarkS2	(H37Rv), 23
(CNRZ327), 15	HG001, 25
CNRZ327_PreSaved_ResultsObj_Glimmer	HG001_PreSaved_DataMapObj(HG001), 25
(CNRZ327), 15	HG001_PreSaved_ResultsObj_GenBank
CNRZ327_PreSaved_ResultsObj_Prodigal	(HG001), 25
(CNRZ327), 15	HG001_PreSaved_ResultsObj_GeneMarkS2
COH1, 16	(HG001), 25
COH1_PreSaved_DataMapObj(COH1), 16	HG001_PreSaved_ResultsObj_Glimmer
COH1_PreSaved_ResultsObj_GenBank	(HG001), 25
(COH1), 16	HG001_PreSaved_ResultsObj_Prodigal
COH1_PreSaved_ResultsObj_GeneMarkS2	(HG001), 25
(COH1), 16	Houston_1, 26
COH1_PreSaved_ResultsObj_Glimmer	
(COH1), 16	Houston_1_PreSaved_DataMapObj
COH1_PreSaved_ResultsObj_Prodigal	(Houston_1), 26
(COH1), 16	Houston_1_PreSaved_ResultsObj_GenBank
(6011), 10	(Houston_1), 26
D_UW_3_CX, 18	Houston_1_PreSaved_ResultsObj_GeneMarkS2
D_UW_3_CX_PreSaved_DataMapObj	(Houston_1), 26
(D_UW_3_CX), 18	Houston_1_PreSaved_ResultsObj_Glimmer
D_UW_3_CX_PreSaved_ResultsObj_GenBank	(Houston_1), 26
(D_UW_3_CX), 18	Houston_1_PreSaved_ResultsObj_Prodigal
D_UW_3_CX_PreSaved_ResultsObj_GeneMarkS2	(Houston_1), 26
(D_UW_3_CX), 18	
D_UW_3_CX_PreSaved_ResultsObj_Glimmer	I11403, 28
(D_UW_3_CX), 18	Il1403_PreSaved_DataMapObj(Il1403),28
D_UW_3_CX_PreSaved_ResultsObj_Prodigal	Il1403_PreSaved_ResultsObj_GenBank
(D_UW_3_CX), 18	(I11403), <u>28</u>
(5_5,15,5), 15	Il1403_PreSaved_ResultsObj_GeneMarkS2
EGD_e, 19	(I11403), <u>28</u>
EGD_e_PreSaved_DataMapObj(EGD_e), 19	Il1403_PreSaved_ResultsObj_Glimmer
EGD_e_PreSaved_ResultsObj_GenBank	(Il1403), <u>28</u>
(EGD_e), 19	Il1403_PreSaved_ResultsObj_Prodigal
EGD_e_PreSaved_ResultsObj_GeneMarkS2	(Il1403), <u>28</u>
(EGD_e), 19	
EGD_e_PreSaved_ResultsObj_Glimmer	K_12_MG1655, 29
(EGD_e), 19	K_12_MG1655_PreSaved_DataMapObj
EGD_e_PreSaved_ResultsObj_Prodigal	(K_12_MG1655), 29
(EGD_e), 19	K_12_MG1655_PreSaved_ResultsObj_GenBank
((K_12_MG1655), 29
GetDataMapObj, 21	K_12_MG1655_PreSaved_ResultsObj_GeneMarkS2
GetGeneSources, 21	(K_12_MG1655), 29
GetResultsObj, 22	K_12_MG1655_PreSaved_ResultsObj_Glimmer
GetStrainIDs, 23	(K_12_MG1655), 29

48 INDEX

<pre>K_12_MG1655_PreSaved_ResultsObj_Prodigal</pre>	PAO1_PreSaved_ResultsObj_GeneMarkS2 (PAO1), 37
(K_12_H01033), 27	PAO1_PreSaved_ResultsObj_Glimmer
LAL14_1, 31	(PAO1), 37
LAL14_1_PreSaved_DataMapObj(LAL14_1),	PA01_PreSaved_ResultsObj_Prodigal
31	
	(PA01), 37
LAL14_1_PreSaved_ResultsObj_GenBank	SavoCanomaTaDath 20
(LAL14_1), 31	SaveGenomeToPath, 38 SL1344, 39
LAL14_1_PreSaved_ResultsObj_GeneMarkS2	
(LAL14_1), 31	SL1344_PreSaved_DataMapObj (SL1344), 39
LAL14_1_PreSaved_ResultsObj_Glimmer	SL1344_PreSaved_ResultsObj_GenBank
(LAL14_1), 31	(SL1344), 39
LAL14_1_PreSaved_ResultsObj_Prodigal	SL1344_PreSaved_ResultsObj_GeneMarkS2
(LAL14_1), 31	(SL1344), 39
	SL1344_PreSaved_ResultsObj_Glimmer
MG1363, 32	(SL1344), 39
MG1363_PreSaved_DataMapObj (MG1363), 32	SL1344_PreSaved_ResultsObj_Prodigal
MG1363_PreSaved_ResultsObj_GenBank	(SL1344), 39
(MG1363), 32	Strain10403S, 41
MG1363_PreSaved_ResultsObj_GeneMarkS2	Strain10403S_PreSaved_DataMapObj
(MG1363), 32	(Strain10403S), 41
MG1363_PreSaved_ResultsObj_Glimmer	Strain10403S_PreSaved_ResultsObj_GenBank
(MG1363), 32	(Strain10403S), 41
MG1363_PreSaved_ResultsObj_Prodigal	Strain10403S_PreSaved_ResultsObj_GeneMarkS2
(MG1363), 32	(Strain10403S), 41
MGAS5005, 34	Strain10403S_PreSaved_ResultsObj_Glimmer
MGAS5005_PreSaved_DataMapObj	(Strain10403S), 41
(MGAS5005), 34	Strain10403S_PreSaved_ResultsObj_Prodigal
MGAS5005_PreSaved_ResultsObj_GenBank	(Strain10403S), 41
(MGAS5005), 34	Strain168, 42
MGAS5005_PreSaved_ResultsObj_GeneMarkS2	Strain168_PreSaved_DataMapObj
(MGAS5005), 34	(Strain168), 42
MGAS5005_PreSaved_ResultsObj_Glimmer	Strain168_PreSaved_ResultsObj_GenBank
(MGAS5005), 34	(Strain168), 42
MGAS5005_PreSaved_ResultsObj_Prodigal	Strain168_PreSaved_ResultsObj_GeneMarkS2
	(Strain168), 42
(MGAS5005), 34	Strain168_PreSaved_ResultsObj_Glimmer
NCTP 2610 35	(Strain168), 42
NCIB_3610, 35	Strain168_PreSaved_ResultsObj_Prodigal
NCIB_3610_PreSaved_DataMapObj	(Strain168), 42
(NCIB_3610), 35	(301 a111100), 42
NCIB_3610_PreSaved_ResultsObj_GenBank	TCH1516, 44
(NCIB_3610), 35	TCH1516_PreSaved_DataMapObj (TCH1516),
NCIB_3610_PreSaved_ResultsObj_GeneMarkS2	44
(NCIB_3610), 35	TCH1516_PreSaved_ResultsObj_GenBank
NCIB_3610_PreSaved_ResultsObj_Glimmer	(TCH1516), 44
(NCIB_3610), 35	TCH1516_PreSaved_ResultsObj_GeneMarkS2
NCIB_3610_PreSaved_ResultsObj_Prodigal	(TCH1516), 44
(NCIB_3610), 35	TCH1516_PreSaved_ResultsObj_Glimmer
D.01. 37	(TCH1516), 44
PA01, 37	
PA01_PreSaved_DataMapObj (PA01), 37	TCH1516_PreSaved_ResultsObj_Prodigal
PAO1_PreSaved_ResultsObj_GenBank	(TCH1516), 44

(PAO1), 37