# Package 'Aphrodite'

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Type Package
<b>Title</b> Automated PHenotype Routine for Observational Definition Identification Training and Evaluation (APHRODITE) - Phenotype building tool using Fuzzy labels
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<b>Description</b> Aphrodite uses noisy class labels to create silver standard training corpora to construct phenotype models in conjunction with expert knowledge codified in existing ontologies and a comprehensive representation of the patient clinical record to learn phenotype models.
License Apache License 2.0   file LICENSE
<b>Depends</b> R (>= 3.1.0), data.table
Imports DatabaseConnector,  SqlRender, plyr, caret, pROC, devtools, knitr, testthat
Suggests ggplot2
VignetteBuilder knitr
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buildFeatureVector This function builds a feature vector using raw patient data

## **Description**

This function builds a feature vector using raw patient data. Returns a patient feature vector (divided by feature sets).

## Usage

buildFeatureVector(flags, casesS, controlsS)

## **Arguments**

flags	The R dataframe that contains all feature/model flags specified in settings.R.
casesS	Dataframe containing the raw patient data.
controlsS	(OPTIONAL) Dataframe containing the raw patient data.

#### **Details**

This function flattens the patient feature data (per feature set) into a feature vector that will be used as input for caret. This function can optionally flat two sources of patient data (cases and controls)

## Value

An object containing the flattened feature vectors for all given feature sets. Of form: list(observations =  $FV_o$ , visits =  $FV_v$ , labs =  $FV_l$  ab, drugexposures =  $FV_d$ e)

buildKeywordList 3

## **Examples**

## Description

Given any given concept\_id or string of text this function generates keyword and ignore lists based on the expansion of concepts (looking at their synonyms).

## Usage

```
buildKeywordList(connection, aphroditeConceptName, schema, dbms)
```

## **Arguments**

```
connection The connection to the database server.

aphroditeConceptName
The string of text / concept name to use.

schema The database schema being used.

dbms The target DBMS for SQL to be rendered in.
```

sion of concepts.

#### Details

Takes the aphroditeConceptName looks for synonyms and builds a list of related concepts using the vocabulary hierarchies

## Value

A list with two elements: a list of positive keywords found (keywordlist\_ALL), and a list of ignore keywords (ignorelist\_ALL)

```
## Not run:
wordLists <- buildKeywordList(conn, aphrodite_concept_name, cdmSchema, dbms)
## End(Not run)</pre>
```

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buildModel	This function builds a model for the specified feature vector using cases and controls for a certain outcomeName

## Description

This function builds a model for the specified feature vector using cases and controls for a certain outcomeName. Returns a caret trained model.

## Usage

```
buildModel(flags, pp_total, outcomeNameS, saveFolder)
```

## Arguments

The R dataframe that contains all feature/model flags specified in settings.R.

outcomeNameS String description of the outcome for which the model is being built

saveFolder folder in which summary file output will be saved

featureVector

Flattened feature vector returned by combineFeatureVectors function, with labeled cases and controls. Assumed to have one column named "Class\_labels" and one named "pid"

#### **Details**

This function builds a model for the specified outcomeName. The model is specified in the flags dataframe (currently only supports LASSO).

#### Value

An transferable caret Model object

```
## Not run:
model_predictors <- buildModel(flag, fv_all, predictorsNames, outcomeName, saveFolder)
## End(Not run)</pre>
```

combineFeatureVectors 5

```
combineFeatureVectors
```

This function combines all of the desired feature types into one single feature vector

## **Description**

This function combines all of the desired feature types into one single feature vector. This feature vector is ready to be used for training

## Usage

```
combineFeatureVectors(flags, cases_pids, controls_pids, featureVector,
  outcomeNameS)
```

#### **Arguments**

#### **Details**

This function builds a feature vector by concatenating all of the available datasets. If binary features are specified in the settings, this conversion is made. The cases\_pids and control\_pids are patient\_id's used for the labeling of the testing and training sets.

#### Value

fv\_all - The combined feature vector (n patients x n features). The columns are: pid column, predictorNames, outcomeName

```
## Not run:
    fv_full_data <- combineFeatureVectors(flag, cases, controls, fv_all, outcomeName)
## End(Not run)</pre>
```

6 conceptDecoder

conceptDecoder This function returns the concept terms corresponding to an input set of concept IDs.	eptDecode			ot terms correspond	ling to an input se	t
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## Description

This function returns the concept terms corresponding to an input set of concept IDs.

## Usage

```
conceptDecoder(connection, schema, dbms, model, numFeats, breaker = ":",
   typeInd = 1, idInd = 2)
```

## **Arguments**

connection	The connection to the database server.
schema	The database schema being used
dbms	The target DBMS for SQL to be rendered in.
model	The model object; will be used to extract top-ranking features
numFeats	The number of features you'd like returned
breaker=":"	Which sort of breaker is used in feature names (e.g. for "obs:12345" it would be ":")
typeInd1=1	Indice after string split defining which feature class (e.g. [1] for "obs:12345" defines obs)
idInd=1	Indice after string split defining which concept_id (e.g. [2] for "obs:12345 defines "12345")

## **Details**

This function returns the concept terms corresponding to an input set of concept IDs. Use case: to investigate highly-ranked features from classification model

## Value

A list of concept terms and concept ids, corresponding to the IDs of interest

```
## Not run:
high_ranking_concepts <- conceptDecoder(connection, schema, dbms, model, 20)
## End(Not run)</pre>
```

convertFeat VecPortion 7

```
convertFeatVecPortion
```

This function builds a feature vector for a specific subset of features

## Description

This function builds a feature matrix for a specific subset of features, e.g. labs/visits/observations/drug exposures. Returns a feature matrix with all features from all patients included.

#### Usage

```
convertFeatVecPortion(featuresType, key, labIndic)
```

## **Arguments**

featuresType	A set of patient data in the form of a list of data frames. Each data frame contains a pid to label the patient, the names of the features that the patient had present, and the frequency counts of these features in his/her record
key	String descriptor of type of feature (e.g. "obs:" or "visit:"). This will be used to label the feature
labIndic=0	Whether this is for a lab feature. If so, must be converted from factor to numeric. Default is 0=no conversion required; 1=conversion required.

## **Details**

This function takes a list of patient data frames as input. Each patient's data frame contains the features that this patient has present in his/her record. This function flattens this information into the combined feature matrix, with all features (of a certain type - e.g. labs or visits) from all patients included. Clearly, many patients will not have data for many features; their feature counts for any feature that was not present in their record will be set as 0.

#### Value

An data frame of (pts) x (features of input type)

```
## Not run:
FV_converted<-convertFeatVecPortion(featuresType, 'obs:')
#OR
FV_converted<-convertFeatVecPortion(featuresType, 'labs:', labIndic=1)
## End(Not run)</pre>
```

8 executeSQL

## **Description**

This function renders, translates and executes one single SQL statement that produces a result

## Usage

```
executeSQL(connection, schema, query, targetDBMS)
```

## **Arguments**

connection The connection to the database server.

schema The database schema being used.

query The SQL statement to retrieve the data.

targetDBMS The target DBMS for SQL to be rendered in.

#### Details

Renders, translates, and executes a single SQL statement that is expeting to produce and result.

## Value

An object containing the data.

f\_score\_calc 9

f_score_calc This fi	nction creates a summary metric for model training
----------------------	--

## Description

This function creates a new summary metric for model training, specific for unbalanced classes. Inputs as specified in caret.

## Usage

```
f_score_calc(data, lev = levels(data$obs), model = NULL)
```

#### **Arguments**

data	A dataframe of the held-out example cases, with columns for 'obs', 'pred', 'T',
	'F'. 'T' and 'F' have the probabilities of each of these classes
lev	Outcome factor levels for model
model	Character string of model used

#### **Details**

This function returns the F-score for model training optimization. Beta is currently set at 2 - TODO: should make this edit-able in future version.

## Value

f\_score

#### **Examples**

```
## Not run:
  f_score <- f_score_calc(data, lev, model)
## End(Not run)</pre>
```

getAnchors

This function allows Anchor recommendation

## **Description**

This function allows Anchor recommendation based after your initial set of keyword and ignore lists have been provided. This will help improve model by suggesting related features that were not considered initially.

```
getAnchors(connection, dbms, schema, casesList, controlsList, ignores,
    studyName, outcomeName, flag, numAnchors)
```

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#### **Arguments**

connection The connection to the database server.

dbms The target DBMS for SQL to be rendered in.

schema The database schema being used.

casesList The list of case patients (already filtered by keywords or gold standard).

controlsList The list of control patients.

ignores The list of concept\_id's ignored when building the cohort.

studyName The study name(will be used for file naming).

outcomeName (will be use for modeling).

flag The flags variable containg the study configuration - we use this one here to have

flexibility of having two different sets of settings for the same experimental run.

numAnchors The total number of anchors to be returned (top N features).

#### **Details**

This function takes the lists of exclude keywords and fetches all patient data for the patients on the cases and controls list. It then builds a model to identify the top performing features and returns a list of them as anchors. This new keyword list can be feed to the set of Anchors specific functions to use any anchor as a selection criteria for patients

#### Value

A list of anchors containing rank, conceptID, domaiID

## **Examples**

```
## Not run:
numAnchors<-50
anchor_list <- getAnchors(conn, dbms, cdmSchema, cases, controls, as.character(ignoreList
## End(Not run)</pre>
```

getdPatientCohort This function builds a patient cohort (and controls) based on a concept list

#### **Description**

This function will build a patient cohort with its respective controls using an inclusion concept\_id list as well as an exclussion concept\_id list. The user specifies the number of both cases and controls for his cohort.

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#### **Arguments**

#### **Details**

This function takes the lists of include and exclude concept\_ids and finds all patients that satisfy this characteristics from the Observation and Condition\_occurrence tables in CDM V5.

#### Value

A list of dataframes containing both cases and control patient\_id's.

## **Examples**

getNormalizationTerm

This function returns the normalizing factor, based upon the input settings

## **Description**

This function returns the normalizing factor, based upon the input settings

#### Usage

```
getNormalizationTerm(dates, flags, defaultTime = 1)
```

#### **Arguments**

dates The dates of all visits recorded in the record

flags The R dataframe that contains all feature/model flags specified in settings.R. -

specifies which sort of normalization to perform

defaultTime Value by which to normalize patients who only have a single visit, so cannot

say what the follow-up time is  $(=0 \rightarrow \text{undefined})$ . Set default time as 1; as if spreading single observation over an entire year or 1 month (depending on

settings)

#### **Details**

Depending upon the input settings, will return the normalizing term for the feature values. Some normalization settings depend upon the specific feature type; these are addressed within the individual feature types. This is a helper function for getPatientData

#### Value

The value by which to divide term counts

#### **Examples**

```
## Not run:
   timeDiff <- getNormalizationTerm(dates, flags)
## End(Not run)</pre>
```

```
getPatientCohort_w_Anchors
```

This function builds a patient cohort (and controls) based on Anchors and lists

## Description

This function will build a patient cohort with its respective controls using an inclusion concept\_id list as well as an exclussion concept\_id list. The user specifies the number of both cases and controls for his cohort.

```
getPatientCohort_w_Anchors(connection, dbms, includeConceptlist,
  excludeConceptlist, schema, cohortSize, controlSize, flags)
```

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#### **Arguments**

The connection to the database server. connection The target DBMS for SQL to be rendered in. dhms includeConceptlist The list of concept\_id's used to build the cohort. excludeConceptlist The list of concept\_id's used as exclusion criteria for the cohort. schema The database schema being used. cohortSize The number of desired patients to appear in the cohort. controlSize The number of desired patients to be in the control group. flags The flags variable containg the study configuration - we use this one here to have flexibility of having two different sets of settings for the same experimental run.

#### **Details**

This function takes the lists of include and exclude concept\_ids and finds all patients that satisfy this characteristics from the Observation and Condition\_occurrence tables in CDM V5.

#### Value

A list of dataframes containing both cases and control patient\_id's.

## **Examples**

getPatientData

This function fetches all the patient data (generic)

## Description

This function fetches all the patient data (generic). Returns raw patient data.

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#### Usage

```
getPatientData(connection, dbms, patient_ids, keywords, flags, schema,
  removeDomains = c(""))
```

#### **Arguments**

connection The connection to the database server.

dbms The target DBMS for SQL to be rendered in.

patient\_ids The list of case patient id's to extract data from - NOT a data.frame.

keywords The list of concept\_id's that are NOT wanted to be used as features

flags The R dataframe that contains all feature/model flags specified in settings.R.

schema The database schema being used.

removeDomains="

List of domains to not include as features, if any are specified in settings file

## **Details**

Based on the groups of feature sets determined in the flags variable, this function will fetch patient data. The function returns all patient information

#### Value

An object containing the raw feature sets for the patient data.

## **Examples**

```
## Not run:
dataFcontrols <- getPatientData(conn, dbms, controls, flag , cdmSchema)
## End(Not run)</pre>
```

```
getPatientDataCases
```

This function fetches all the patient data (non-generic) designed to work when building a model

#### **Description**

This function fetches all the patient data (non-generic) designed to work when building a model. Returns raw patient data.

```
getPatientDataCases(connection, dbms, patient_ids, keywords, ignores, flags,
    schema, removeDomains = "")
```

#### **Arguments**

connection The connection to the database server.

dbms The target DBMS for SQL to be rendered in.

patient\_ids The list of case patient id's to extract data from.

keywords The list of concept\_id's used to build the cohort.

ignores The list of concept\_id's ignored when building the cohort.

flags The R dataframe that contains all feature/model flags specified in settings.R.

schema The database schema being used.

removeDomains="

List of domains to not include as features, if any are specified in settings file

#### **Details**

Based on the groups of feature sets determined in the flags variable, this function will fetch patient data. The function determines the first mention of the keywords and selects that date to start the data extraction of the remaining patient information

#### Value

An object containing the raw feature sets for the patient data.

## **Examples**

```
getPatientDataFromStartDate
```

This function fetches all the patient data (generic) - from a given start date

#### **Description**

This function fetches all the patient data (generic). Returns raw patient data.

```
getPatientDataFromStartDate(connection, dbms, patient_ids, patIndexDate,
   keywords, flags, schema, removeDomains = c(""))
```

#### **Arguments**

connection The connection to the database server.

dbms The target DBMS for SQL to be rendered in.

patient\_ids The list of case patient id's to extract data from - NOT a data.frame.

patIndexDate The start index date for all patients

keywords The list of concept\_id's that are NOT wanted to be used as features

flags The R dataframe that contains all feature/model flags specified in settings.R.

schema The database schema being used.

removeDomains="

List of domains to not include as features, if any are specified in settings file

#### **Details**

Based on the groups of feature sets determined in the flags variable, this function will fetch patient data. The function returns all patient information

## Value

An object containing the raw feature sets for the patient data.

## **Examples**

```
## Not run:
   patientData <- getPatientDataFromStartDate(conn, dbms, patient_ids, start_dates, ignoreF
## End(Not run)</pre>
```

```
getPatientDataStartEnd
```

This function fetches all the patient data (generic) - from a given start date and with a given end date

## Description

This function fetches all the patient data (generic). Returns raw patient data.

```
getPatientDataStartEnd(connection, dbms, patient_ids, startDate, endDate, flags,
    schema, removeDomains = c(""))
```

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#### **Arguments**

connection The connection to the database server.

dbms The target DBMS for SQL to be rendered in.

patient\_ids The list of case patient id's to extract data from - NOT a data.frame.

startDate The start index date for all patients
endDate The end date to fetch data from patients

flags The R dataframe that contains all feature/model flags specified in settings.R.

schema The database schema being used.

removeDomains="

List of domains to not include as features, if any are specified in settings file

#### **Details**

Based on the groups of feature sets determined in the flags variable, this function will fetch patient data within the specified time range the function returns all patient information

#### Value

An object containing the raw feature sets for the patient data.

## **Examples**

```
## Not run:
   patientData <- getPatientDataStartEnd(conn, dbms, patient_ids, start_dates, end_dates, f
## End(Not run)</pre>
```

manipulateSqlPull This function performs the manipulation of the sql extract data; should be generic for any of the feature types

#### **Description**

This function performs the manipulation of the sql extract data; should be generic for any of the feature types

#### Usage

```
manipulateSqlPull(tmp_fv, flags, timeDiff)
```

## **Arguments**

tmp\_fv Pull from sql query. Should have a column for date and concept\_id flags Flags set in settings - specifies which normalization is needed timeDiff Value to use for normalization

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#### **Details**

This is just a helper function that reduces the repeats of code for the manipulation of the sql extract data, so that it is put in the desired format for compiling all patient features together. This function: gets the counts of codes on a given visit (so multiple codes/terms/drugs/etc are not all counted); normalizes based on the normalization setting; returns a data frame with counts of codes

#### Value

An object containing the re-formatted patient data: ptID x (num concept IDs) - filled with counts, deduplicated by visit

## **Examples**

```
## Not run:
test1 <- manipulateSqlPull(tmp_fv, flags, timeDiff)
## End(Not run)</pre>
```

plotFeatWeightings This function plots the feature importance weightings

## Description

This function plots the feature importance weightings

## Usage

```
plotFeatWeightings(plotSaveFile, weightingsDF)
```

## **Arguments**

```
plotSaveFile The name of the file to save
weightingsDF Data frame of the weightings with their labels
```

#### **Details**

This function returned predicted classes for the input patient list. Use case: evaluate trained model on a set of gold-standard patients.

## Value

(none)

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```
## Not run:
plotFeatWeightings(plotSaveFile, weightingsDF)
## End(Not run)
```

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