CSE8803: Big Data Analytics in Healthcare Homework 1

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Deadline: January 31, 2016

- Discussion is encouraged, but each student must write his/her own answers and explicitly mention any collaborators.
- Each student is expected to respect and follow the GT Honor Code.
- Please type the submission with LaTeX or Microsoft Word. We will not accept hand written submissions.
- Please **do not** change the filenames and function definitions in the skeleton code provided, as this will cause the test scripts to fail and subsequently no points will be awarded. Built-in modules of python and the following libraries pandas, numpy, scipy, scikit-learn can be used.
- Update the README.txt file in the folder to include the version of python you have used.

Overview

Preparing the data, computing basic statistics and constructing simple models are essential steps for various data analysis work. In this homework, you will use clinical data as raw input to perform **Mortality Prediction**. For this homework, **Python** programming will be required. See the attached skeleton code as a start-point for the programming questions.

It is your responsibility to make sure that all code and other deliverables are in the correct format and that your submission compiles and runs. We will not manually check your code. Thus non-runnable code will directly lead to 0 score.

1 CITI Certification [10 points]

Georgia Tech policy requires that all personnel involved with human subjects research must pass a training course before doing so. This requirement encompasses all types of interactions with human subjects, including the analysis of data.

Follow the steps below to complete the certification and attach your certification.

- 1. Go to https://www.citiprogram.org/
- 2. Login via Single Sign On (SSO), which will allow to login using your Georgia Tech username and password
- 3. Select Georgia Institute of Technology as the authentication provider
- 4. Once logged in, under Georgia Institute of Technology courses, click on "Add Course or Update Learner Groups"
- 5. Now you will have three main courses to select. You will check the box for "Human Subjects Research"
- 6. Click next, then you will select the radio button "NO, I have NOT completed the basic course"
- 7. Now, you will see three learner groups. You are required to complete Group 1 and Group 2. Let us start with Group 1 (select Group 1) and click next
- 8. Good Clinical Practice is not required so select "N/A", then click next
- 9. Health Information Privacy and Security (HIPS) is required, click "CITI Health Information Privacy and Security (HIPS) for Biomedical Research Investigators"
- 10. Select "RCR for engineering"
- 11. Now under Georgia Tech courses you will have "Group 1 Biomedical research Investigators and Key Personnel" listed as incomplete. You will have to go through every tutorial in that course and complete a quiz for each.
- 12. Once you have completed and passed Group 1, repeat the steps above to complete Group 2 (Social / Behavioral Research Investigators and Key Personnel)

Solution. Sample certification displayed in Figure 1.

About Raw Data

You need to download the homework1 zip from T-Square, after unzipping browse to the homework1/data/train. There are three CSV files which will be the input data in this assignment.

The data provided in *events.csv* are event sequences. Each line of this file consists of a tuple with the format (patient_id, event_id, event_id, event_id, it is in the sequence).

For example,

```
1053, DIAG319049, Acute respiratory failure, 2924-10-08, 1.0
1053, DIAG197320, Acute renal failure syndrome, 2924-10-08, 1.0
1053, DRUG19122121, Insulin, 2924-10-08, 1.0
1053, DRUG19122121, Insulin, 2924-10-11, 1.0
1053, LAB3026361, Erythrocytes in Blood, 2924-10-08, 3.000
```

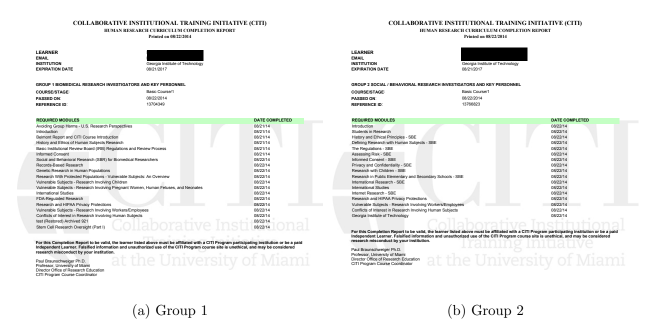


Figure 1: Sample CITI Certification

```
1053, LAB3026361, Erythrocytes in Blood, 2924-10-08, 3.690
1053, LAB3026361, Erythrocytes in Blood, 2924-10-09, 3.240
1053, LAB3026361, Erythrocytes in Blood, 2924-10-10, 3.470
```

- patient_id: Identifies the patients in order to differentiate them from others. For example, the patient in the example above has patient id 1053.
- event_id: Encodes all the clinical events that a patient has had. For example, DRUG19122121 means that a drug with RxNorm code as 19122121 was prescribed to the patient. DIAG319049 means the patient was diagnosed of disease with SNOMED code of 319049 and LAB3026361 means that the laboratory test with a LOINC code of 3026361 was performed on the patient.
- event_description: Shows the description of the event. For example, DIAG319049 is the code for Acute respiratory failure and DRUG19122121 is the code for Insulin.
- **timestamp**: Indicates the date at which the event happened. Here the timestamp is not a real date but a shifted date for protecting privacy of patients.
- value: Contains the value associated to an event. See Table 1 for the detailed description.

The data provided in *mortality_events.csv* contains the patient ids of only the deceased people. They are in the form of a tuple with the format (patient_id, timestamp, label). For example,

event type	sample event_id	value meaning	example
diagnostic code	DIAG319049	diagnosed with a certain disease, value always be 1.0	1.0
drug consumption	DRUG19122121	prescribed a certain medication, value will	1.0
laboratory test	LAB3026361	always be 1.0 test conducted on a patient and its value	3.690

Table 1: Event sequence value explanation

```
37,3265-12-31,1
40,3202-11-11,1
```

The timestamp indicates the death date of a deceased person and a label of 1 indicates death. Patients that are not mentioned in this file are considered alive.

The event_feature_map.csv is a map from an event_id (SNOMED, LOINC and RxNorm) to an integer index. This file contains (int_id, event_id) pairs for all event ids.

Running the tests

Test cases are provided for every module in this assignment and operate on a subset of the data. To run a test, execute the following commands from the base folder i.e. homework1. If any of the test cases fail, an error will be shown. For example to test the statistics computed, the following command should be executed:

nosetests tests/test_statistics.py

A single test can also be run using this syntax:

nosetests tests/< filename >:< test_method >

2 Descriptive Statistics [10 points]

Before starting analytic modeling, it is a good practice to get descriptive statistics of the input raw data. In this homework, you need to write code that computes various metrics on data described previously. A skeleton code is also provided for you as a starting point.

The definition of terms used in the result table are described below:

- Event count: Number of events recorded for a given patient. Note that every line in the input file is an event.
- Encounter count: Count of unique dates on which a given patient visited the ICU.
- **Record length**: Duration (in number of days) between first event and last event for a given patient.
- a. Complete *code/event_statistics.py* for computing statistics required by question Please be aware that **you are not allowed to change the filename, and any existing function declarations.** Only numpy, scipy, scikit-learn and other built-in modules of python will be available for you to use. The use of *pandas* library is suggested.
- **b.** Use *events.csv* and *mortality_events.csv* provided in **train_data** as input and fill Table 2 with actual values.

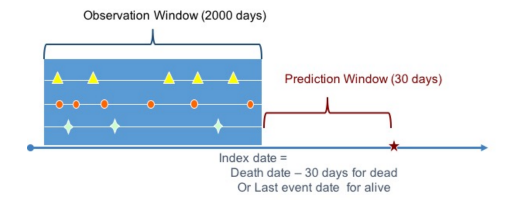
Metric	Deceased patients	Alive patients	Function to complete
Event Count			event_count_metrics
1. Average Event Count			
2. Max Event Count			
3. Min Event Count			
Encounter Count			encounter_count_metrics
1. Average Encounter Count			
2. Max Encounter Count			
3. Min Encounter Count			
Record Length			record_length_metrics
1. Average Record Length			
2. Max Record Length			
3. Min Record Length			

Table 2: Descriptive statistics for alive and dead patients

Deliverable: code/event_statistics.py [10 points]

3 Feature construction [30 points]

It is a common practice to convert raw data into a standard data format before running real machine learning models. In this section, you will work on code/etl.py file and implement the necessary python functions in this script. You will work with events.csv, $mortality_events.csv$ and $event_feature_map.csv$ files provided in **train** folder. The use of pandas library in this question is recommended. Listed below are a few concepts you need to know before beginning feature construction (for details please refer to lectures).



- Observation Window: The time interval you will use to identify relevant events. Only events present in this window should be included while constructing feature vectors. The size of observation window is 2000 days.
- Prediction Window: A fixed time interval that is to be used to make the prediction. Events in this interval should not be included in constructing feature vectors. The size of prediction window is 30 days.
- Index date: The day on which mortality is to be predicted. Index date is evaluated as follows:
 - For deceased patients: Index date is 30 days prior to the death date (timestamp field) in *mortality_events.csv*.
 - For alive patients: Index date is the last event date in *events.csv* for each alive patient.

Step - a. Compute the index date [7.5 points]

Use the definition provided above to compute the index date for all patients. Complete the method $calculate_index_date$ provided in code/etl.py.

Deliverable: code/etl.py, etl_index_dates.csv

Step - b. Filter events [5 points]

Consider an observation window (2000 days) and prediction window (30 days). Remove the events that occur outside the observation window. Complete the method $filter_events$ provided in code/etl.py.

Deliverable: code/etl.py, etl_filtered_events.csv

c. Aggregate events [10 points]

To create features suitable for machine learning, we will need to aggregate the events for each patient as follows:

- sum values for diagnostics and medication events (i.e. event_id starting with DRUG and DIAG) to get their counts.
- average values for lab events (i.e. event_id starting with LAB).

Each event type will become a feature and we will directly use event_id as feature name. For example, given below raw event sequence for a patient,

```
1053, DIAG319049, Acute respiratory failure, 2924-10-08, 1.0 1053, DIAG197320, Acute renal failure syndrome, 2924-10-08, 1.0 1053, DRUG19122121, Insulin, 2924-10-08, 1.0 1053, DRUG19122121, Insulin, 2924-10-11, 1.0 1053, LAB3026361, Erythrocytes in Blood, 2924-10-08, 3.000 1053, LAB3026361, Erythrocytes in Blood, 2924-10-08, 3.690 1053, LAB3026361, Erythrocytes in Blood, 2924-10-09, 3.240 1053, LAB3026361, Erythrocytes in Blood, 2924-10-10, 3.470
```

We can get feature value pairs(event_id, value) for this patient with ID 1053 as

```
(DIAG319049, 1.0)
(DIAG197320, 1.0)
(DRUG19122121, 2.0)
(LAB3026361, 3.35)
```

You will notice there are certain events with no entries in the values column. Handle these missing values by removing all events with null values while constructing the features. Next, replace each <code>event_id</code> with the <code>feature_id</code> provided in <code>event_feature_map.csv</code>

```
(708, 1.0)
(306, 1.0)
(2475, 2.0)
(3030, 3.35)
```

Further, in machine learning algorithm like logistic regression, it is important to normalize different features into the same scale using an approach like min-max normalization (hint: $min(x_i)$ maps to 0 and $max(x_i)$ 1 for feature x_i). Complete the method $aggregate_events$ provided in code/etl.py.

Deliverable: code/etl.py and etl_aggregated_events.csv

d. Save in SVMLight format[7.5 points] If the dimensionality of a feature vector is large but the feature vector is sparse (i.e. it has

only a few nonzero elements), sparse representation should be employed. In this problem you will use the provided data for each patient to construct a feature vector and represent the feature vector in SVMLight format.

```
<!ine> .=. <target> <feature>:<value> <feature>:<value> #<info>
<target> .=. +1 | -1 | 0 | <float>
<feature> .=. <integer> | qid
<value> .=. <float>
<info> .=. <string>
```

For example, the feature vector in SVMLight format will look like:

```
1 2:0.5 3:0.12 10:0.9 2000:0.3
0 4:1.0 78:0.6 1009:0.2
1 33:0.1 34:0.98 1000:0.8 3300:0.2
1 34:0.1 389:0.32
```

where, 1 or 0 will indicate whether the patient is alive or dead i.e. the label and it will be followed by a series of feature-value pairs sorted by the feature index (idx) value.

Deliverable: code/etl.py, featured_symlight.train and features.train

4 Predictive Modelling [45 points]

4.1 Part A [15 points]

In the previous section, you constructed features for each patient in the training set. Now use these features and the corresponding labels to train your classifier. In this question, you will use 3 classifiers: Logistic Regression, SVM and Decision tree classifiers. The skeleton code is provided in *code/models.py*. Once you train your predictive models, you will use it to predict mortality for a few patients, whose features and corresponding labels is provided to you in a separate validation set *data/features_symlight.validate*. To evaluate your predictive models, you will calculate and report the metrics given in Table 3.

Model	Accuracy	AUC	Precision	Recall	F-Score
Logistic Regression SVM Decision Tree					

Table 3: Model performance

Deliverable: code/models.py [15 points]

4.2 Part B: Cross-validation [10 points]

In order to fully utilize available data and get more reliable results, machine learning practitioners use cross-validation. For details, refer to the lecture videos. In this problem, you need to implement two cross-validation strategies and use Logistic Regression as your classifier.

You need to implement

- K-Fold Cross-validation Strategy: divides all the samples in k groups of samples. Each time $\frac{1}{k}$ samples will be used as testing data and other samples as training.
- Randomized Cross-validation Strategy: Iteratively random shuffle the whole dataset and use top specific percentage of data as training set and the rest as validation set.

NOTE: You would use features that you constructed in Section 3 as the entire dataset for this problem.

a. Implement two strategies based on skeleton in code/cross.py. Use k=5 for k-fold CV. Use test data percentage of 0.2 and number of iterations as 5 for Randomized CV.

Report the average Accuracy and AUCs in Table 4

CV strategy	Accuracy	AUC
K-Fold		
Randomized		

Table 4: Cross Validation

Deliverable: code/cross.py [10 points]

4.3 Part C [20 points]

In this part, you will refine your predictive model and set of features to obtain better performance. You are advised to try out different things to improve the performance of your predictive model. One may try out new features, or use feature selection techniques to reduce the feature set, or tune the parameters of the predictive model or try ensemble techniques. However, one **must not** change the observation window and prediction window.

You should use the data available in data/train to construct features and train your model. It is advisable to use the cross validation AUC as a metric to refine your model. Your final model will be evaluated on a (blind) test set for which the labels are unknown to you. The events in the observation window corresponding to the test patients are available in data/test/events.csv. If you are constructing the same features as in Section 3 for the test set, you may use the feature map in $data/test/event_feature_map.csv$.

a. Implement your model in *code/my_model.py*. You are free to use your own features and predictive model. Please ensure that your predicted labels are either 0 or 1. Report

your features of the test patients in *deliverables/test_features.txt*. Submit your predictions in the form of a csv file (patient_id,predicted label) in *deliverables/my_predictions.csv*

b. Write a short paragraph on your best model(based on cross validation AUC) and the models that you tried. What was the rationale behind your approach? Did you improve the model's performance?

Deliverable 1: deliverables/test_features.txt

(Refer to the skeleton code for the required format)

Deliverable 2: code/my_model.py

Deliverable 3: deliverables/my_predictions.csv

(This will be evaluated with the true labels of the test patients. The top 5% of the students with the highest AUC will be awarded 5 bonus points)

5 Submission[5 points]

The folder structure of your submission should be as below. You can use the *tree* command to dispay and verify the folder structure is as seen below. All other unrelated files will be discarded during testing.

```
<your gtid>-<your gt account>-hw1
|-- homework1_answer.pdf
|-- group1.pdf
|-- group2.pdf
|-- hips_certification.pdf
|-- README.txt
|-- code
   |-- event_statistics.py
   |-- etl.py
   |-- models.py
   |-- cross.py
   |-- utils.py
   \-- my_model.py
|-- deliverables
   |-- etl_index_dates.csv
   |-- etl_filtered_events.csv
   |-- etl_aggregated_events.csv
   |-- features_svmlight.train
   |-- features.train
   |-- test_features.txt
   \-- my_predictions.csv
```

Create a tar archive of the folder above with the following command and submit the tar file.

```
tar -czvf <your gtid>-<your gt account>-hw1.tar.gz \
<your gtid>-<your gt account>-hw1
```

Example submission: 901234567-gburdell3-hw1.tar.gz