## Trial Task Due Date: 12th November 2020

**Mandatory:** Use comment lines, functions and classes where appropriate.

Task 1A: Human Metabolome Database (HMDB) https://hmdb.ca/

1. Download the XML File from the following link (https://hmdb.ca/downloads)

Filename: "All metabolite" released on 2020-09-09

**Caution**: File is too large for the regular browser to process/handle.

Hint: Split the XML nodes by the tag named <metabolite> into smaller XML files

- 2. Review the tags and values under normal and abnormal concentrations of tags
- 3. Calculate the total number of diseases associated with the blood and list all the disease with accession number in CSV file format (important)

**Hint**: <biospecimen>blood</biospecimen> and disease will be tagged as <patient\_information> under the abnormal concentration tag and <subject condition> under normal concentrations tag.

- 4. Filter the XML files with blood as biospecimen on both abnormal and normal concentrations tag

  Caution: Henceforth, we will only work on filtered XML file containing blood as biospecimen on both abnormal and normal concentration tag
- 5. Convert them into given template A

Hint: Select normal concentrations, abnormal concentrations and diseases tag from the original file

6. Standardise the age into a numerical format

**Hint**: Age format YY, MMDD (Y: Year, M: Months, D: Days)

7. Convert the XML files into given template B

**Hint**: The values of the following tags will be the same as given in the template (creation, update, version, completion status and ethnicity tag)

## Task 1B:

**Note**: The following two tasks are independent of the above task. However, they are closely related.

- 1. List all the different units associated with concentration value and develop a system which can convert the values of other units into the desired value (For example, 6 kilograms can be converted into 6,000 grams and vice-versa)
- 2. Develop a search system for a synonym to access its root name. For example, if the synonym for the Iron is "Fe" then whenever "Fe" is called it should return Iron.

## Task 2: Perform web-scraping on metagene.de (<a href="https://www.metagene.de/start.html">https://www.metagene.de/start.html</a>)

- 1. Perform web-scraping on the list of all disease in metagene.de and extract all the values associated on the right side of the panel (Example: disease, symptoms etcetera)
- 2. Convert the extracted values into an XML file.

Hint: Use template C as a reference

- 3. Filter diseases with blood or plasma as a specimen (Check step 4 of Task 1A)
- 4. Propose and standardise the age into the numerical format.
- 5. Analyse, how many of the diseases and metabolite in the metagene overlaps with the HMDB
- 6. Standardise the age in numerical format
- 7. List all the different units associated with concentration value and develop a system which can convert the values of other units into the desired value (For example, 6 kilograms can be converted into 6,000 grams and vice-versa)

## Optional tasks:

- Propose a classification technique to classify the diseases using predictor variables such as age, sex, metabolite concentration etcetera from the above data and provide necessary details or justification.
- Develop an auto/runtime application for Task 1A, which performs all the above task