Sarah Griffiths - Middle layer user guide

Follow these instructions to use the Business Logic Layer - All code is written in python.

Front End instructions

- 1. You must have all the files saved in 1 directory this should already be set up.
- 2. Import the module BusinessLogicMainFile.py in to your code
- 3. Use following functions as required

Functions To return required lists of records.

getAllGenes() - returns A-Z geneList

getAllProteins() - returns A-Z proteinList

getAllAccessions() - returns A-Z accessionList

getAllChromosomeLocations() – returns ordered list of chromosomeLocationList

4. Call this function to retrieve all associated record data.

getAllEntryData(keyword)

keyword – Any accession/gene/protein/NCBI ID/Chromosomal location

This will return the following python variables

| Variable name (literal) | Explanation of what variable contains | Variable type |
|-------------------------|---|------------------|
| | | |
| accession_number | Returns accession no from data access layer | string |
| NCBI_identifier | NCBI ID | string |
| Chromosome_location | Chromosome location | string |
| Protein_product_name | Protein product name | string |
| parsedSequence | Entire nucleic acid sequence | String |
| codingRegion | Coding region of nucleic acid sequence | String |
| mrnaSequence | Mrna sequence – (t's replaced with u's) | String |
| splitSequence | Mrna sequence split in to codons | List |
| translatedAndAligned | Sequence translated in to AA and aligned with Nucleic acid sequence | 2Aligned Strings |
| justAminoAcids | Just AA sequence | String |
| codonFrequency | Codon frequencies | Dictionary |
| RestrictionEnzymes | Dictionary of associated RE and their respective sequences | Dictionary |

```
Example –
getAllEntryData(U49845)
Returns -
```

parsedSequence = "actgtgggggtcacgtcgta..."

```
codingRegion = "gtcacgtgta..."

mrnaSequence = "gugcacgugaaaguggucgugugugggugucguggggcaaaa...."

splitSequence = gug,cac,gug,aaa,gug,guc,gug,ugu,ggg,ugu,cgu,ggg,gca,aaa...."

translatedAndAligned = 'gugcacgugaaaguggucguguggggugucgugggggcaaaa..', 'T--B--A--G--T--N--R..'

JustAminoAcids = 'TBAGTNR...'

codonFrequencies = codons = {"uuu": 20% , "uuc": 30%, "uua": 20%, "uug": 30% ...}

RestrictionEnzymes = 'EcoRl','BamHl','BsuMl...'

You can then use these variables however you choose
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5. Call this function to find cut sites

```
restrictionEnzymeCutSites(bases) -
```

bases - input the sequence you are looking for eg. tcgaa

This will return a dictionary of the format

(start cut point, end cut point: 'within coding region/not within coding region')

Example

restrictionEnzymeCutSites('tcgaa')

Returns

{(127, 340 : "not in coding region"), (500, 1500: "in coding region"), (1600, 2100: "not in coding region"

Back end instructions

- 1. Import module main_file.py
- 2. To calculate total codon frequency use function

Totalcodonfrequency() – This will return a dictionary containing the 64 codons and their respective frequencies. This can then be stored in the DB and called upon again by the middle layer.