Calculation of Rule Set 2 score

Requirements:

Software: Python-2.7

Packages:

- 1. scikit-learn 0.16.1 (Other versions will produce runtime errors)
- 2. pickle
- 3. pandas
- 4. numpy
- 5. scipy

Inputs:

- 1. --seq: 30mer nucleotide sequence, which should be of the form NNNN20merNGGNNN
- 2. --aa-cut: This is an optional argument. Specify the position at which the sgRNA cuts the amino acid sequence. Default value: -1
- 3. --per-peptide: This is an optional argument. Specify the percentage of protein cut by the sgRNA. Default value: -1

The optional arguments are helpful for prediction but if this information is unknown, a separate model will be used for prediction where these features are turned off.

Output:

The Rule Set 2 score for the input sequence will be printed on the terminal.

Running the code:

To run this code, make the "analysis" folder your working directory.

- To run full model prediction, type on the terminal: python rs2_score_calculator.py --seq <30mer sequence> --aa-cut <amino acid cut position> --per-peptide <Percent peptide cut by sgRNA>
- To use only sequence information, type on the terminal: python rs2_score_calculator.py --seq <30mer sequence>

Examples:

For full model prediction:

python rs2_score_calculator.py --seq TGGAGGCTGCTTTACCCGCTGTGGGGGCGC --aa-cut 254 --per-peptide 87

Output: Rule set 2 score: 0.5309

For prediction using only sequence information:

python rs2 score calculator.py --seq TGGAGGCTGCTTTACCCGCTGTGGGGGCGC

Output: Rule set 2 score: 0.5656

OR

python rs2_score_calculator.py --seq TGGAGGCTGCTTTACCCGCTGTGGGGGCGC

--aa-cut -1 --per-peptide -1 Output: Rule set 2 score: 0.5656