My Project

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Namespace Index

1.1 Namespace List

Here is a list of all documented namespaces with brief descriptions:

chemistry2quant.chem2quant_analysis	5
chemistry2quant.chem2quant_chembl2pandas	6
chemistry2quant.chem2quant_lipophilicity	6
chemistry2quant.chem2quant_mol2vec_ChEMBL	7
chemistry2quant.chem2quant NN	7

2 Namespace Index

Class Index

2.1 Class List

Here are the classes, structs, unions and interfaces with brief descriptions:

chemistry2quant.chem2quant_gen_psi4_input.chem2quant_psi4	9
chemistry2quant.chem2quant_chembl2pandas.chemblConnect	9
chemistry2quant.chem2quant_lipophilicity.rdkit_lipophilicity	1 (
chemistry2quant.chem2quant_analysis.rdkitProcessDf	1
chemistry2quant.chem2quant_analysis.rdkitPsi4DataGenerator	12
chemistry2quant.chem2quant_screening.two_step_screen	12

4 Class Index

Namespace Documentation

3.1 chemistry2quant.chem2quant_analysis Namespace Reference

Classes

- · class rdkitProcessDf
- · class rdkitPsi4DataGenerator

Functions

- def neuron_layer (X, n_neurons, name, activation=None)
 A neural network layer for use later.
- def sdfToMol (sdf)
- def substructure_search (substruct, struct_array)

Variables

- string **directory** = "/home/noh/Desktop/CURRENT_WORK_IN_PROGRESS/Chemiinformatics/RDK IT/rdkit/Docs/Book/data"
- string **sdf_file** = 'bzr.sdf'
- **process** = rdkit_processdf(directory, sdf_file)
- molList = process.returnMol()
- molSmiles = process.MoltoSmiles()
- list mol2VecList = [mol2alt_sentence(x,1) for x in molList]
- int **n_hidden1** = 300
- int **n_hidden2** = 100
- int n hidden3 = 100
- feature_colummns = tf.contrib.learn infer real valued columns from input(X train)
- def hidden1 = neuron_layer(X, n_hidden1, "hidden1", activation = "relu")
- def hidden2 = neuron_layer(hidden1, n_hidden2, "hidden2", activation = "relu")
- def logits = neuron_layer(hidden2, n_outputs, "outputs")
- **init** = tf.global variables initalizer()
- **saver** = tf.train.Saver()

3.1.1 Detailed Description

```
@package docstring
Documentation for this module
More details here
```

3.1.2 Function Documentation

3.1.2.1 sdfToMol()

```
def chemistry2quant.chem2quant_analysis.sdfToMol ( sdf \ ) Returns array of mols from sdf
```

3.2 chemistry2quant.chem2quant_chembl2pandas Namespace Reference

Classes

class chemblConnect

Variables

- **cur** = conn.cursor()
- database = cur.fetchall()

3.2.1 Detailed Description

Python module which returns the chembl database as a pandas table, depending on the specifications you would like in the end.

Make sure that the permissions settings on the table has been granted to the USER.

3.3 chemistry2quant.chem2quant_lipophilicity Namespace Reference

Classes

· class rdkit_lipophilicity

Variables

- **zipFileDir** = zipfile.ZipFile('../zip/lipophilicity.zip')
- Pd_df = pd.read_csv(zipFileDir.open('Lipophilicity.csv'))
- · batch_x
- batch y
- train op
- · feed_dict
- loss
- acc

3.3.1 Detailed Description

3.4 chemistry2quant.chem2quant_mol2vec_ChEMBL Namespace Reference

Functions

- def emolecule_command (sql_command, user, password)
- def chembl24_command (sql_command, user, password)

Variables

- def **table_names** = chembl24_command("SELECT table_name FROM information_schema.tables WHERE table_schema='public'", "sang", "silver!!")
- list data_list = []
- string sql_command = "SELECT * FROM {} LIMIT 100;".format(str(title))

3.4.1 Detailed Description

Module for downloading ChemBL data

3.5 chemistry2quant.chem2quant_NN Namespace Reference

Functions

- def next_batch (num, data, labels)
- def mol2arr (mol)

Variables

```
• int n hidden1 = 300
```

- int **n_hidden2** = 300
- int **n_hidden3** = 300
- float learning rate = 0.01
- int **n_outputs** = 3
- string datadir = 'data'
- list train_mol = [mol for mol in Chem.SDMolSupplier(os.path.join(datadir,'solubility.train.sdf')) if mol != None]
- list **test_mol** = [mol for mol in Chem.SDMolSupplier(os.path.join(datadir,'solubility.test.sdf')) if mol != None]
- **cls_mol** = list(set([mol.GetProp('SOL classification') for mol in train mol]))
- dictionary cls_dic = {}
- train_X = np.array([mol2arr(mol) for mol in train_mol])
- train_y = np.array([cls_dic[mol.GetProp('SOL_classification')] for mol in train_mol])
- test_X = np.array([mol2arr(mol) for mol in test_mol])
- test_y = np.array([cls_dic[mol.GetProp('SOL_classification')] for mol in test_mol])
- **X** = tf.placeholder(tf.float32, shape = (None, np.shape(train_X[0])[0]), name = "X")
- y = tf.placeholder(tf.int64, shape=(None), name="y")
- **he_init** = tf.contrib.layers.variance_scaling_initializer()
- hidden1 = fully_connected(X, n_hidden1, weights_initializer = he_init, scope = "hidden1")
- hidden2 = fully connected(hidden1, n hidden2, weights initializer = he init, scope = "hidden2")
- logits = fully_connected(hidden2, n_outputs, weights_initializer = he_init, scope = "outputs", activation_fn = None)
- xentropy = tf.nn.sparse_softmax_cross_entropy_with_logits(labels = y, logits = logits)
- loss = tf.reduce mean(xentropy, name = "loss")
- correct = tf.nn.in_top_k(logits, y ,1)
- accuracy = tf.reduce_mean(tf.cast(correct, tf.float32))
- **optimizer** = tf.train.GradientDescentOptimizer(learning_rate)
- training_op = optimizer.minimize(loss)
- init = tf.global_variables_initializer()
- **saver** = tf.train.Saver()
- int **n_epochs** = 20
- int **batch_size** = 100
- · X batch
- · y_batch
- feed_dict
- acc_train = accuracy.eval(feed_dict={X: X_batch, y: y_batch})
- acc_test = accuracy.eval(feed_dict={X: test_X, y: test_y})
- save_path = saver.save(sess, "./my_model_final.ckpt")

3.5.1 Detailed Description

Tensorflow implementation of

3.5.2 Function Documentation

3.5.2.1 next_batch()

Class Documentation

4.1 chemistry2quant.chem2quant_gen_psi4_input.chem2quant_psi4 Class Reference

Public Member Functions

· def mol2psi4 (mol)

4.1.1 Detailed Description

Generate coordinate file from smiles for calculation with $\ensuremath{\operatorname{psi4}}$

The documentation for this class was generated from the following file:

- /home/oohnohnoh1/Desktop/GIT/Chemiinformatics_work/Chemistry2quant/src/chemistry2quant/chem2quant
 —gen_psi4_input.py
- 4.2 chemistry2quant.chem2quant_chembl2pandas.chemblConnect Class Reference

Public Member Functions

- def __init__ (self, database, user, host, password)
- def issue_command (self)

Public Attributes

- database
- user
- host
- password
- string

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Static Public Attributes

• **conn** = psycopg2.connect(dbname='emolecules', user='sang', host='localhost', password='Blad1bl@1234', port=5432)

The documentation for this class was generated from the following file:

/home/oohnohnoh1/Desktop/GIT/Chemiinformatics_work/Chemistry2quant/src/chemistry2quant/chem2quant
 —chembl2pandas.py

4.3 chemistry2quant.chem2quant_lipophilicity.rdkit_lipophilicity Class Reference

Public Member Functions

- def __init__ (zipFile, name, featurizer, reload=True, move mean=True)
- def load_lipo (featurizer='ECFP', split='index', reload=True, move_mean=True)
- def process data column ()
- def mols2feat ()
- def load_tensorflow (learning_rate, training_epochs, batch_size)
- def weights ()
- def neural_net (x)
- def fingerprints (mols)
- def categories (fingerprints, index)

Public Attributes

- rdkit_zip
- rdkit_csv
- · lipophilicity_csv
- · initial score
- · initial_smiles
- Mol
- MFingerprints
- np_fps
- X
- Y
- · batch size
- · learning_rate
- · training_epochs
- · weights
- biases

4.3.1 Detailed Description

TODO

4.3.2 Member Function Documentation

4.3.2.1 mols2feat()

```
def chemistry2quant.chem2quant_lipophilicity.rdkit_lipophilicity.mols2feat ( )
Converting fingerprint data into vector data

4.3.2.2 process_data_column()

def chemistry2quant.chem2quant_lipophilicity.rdkit_lipophilicity.process_data_column ( )
```

The documentation for this class was generated from the following file:

/home/oohnohnoh1/Desktop/GIT/Chemiinformatics_work/Chemistry2quant/src/chemistry2quant/chem2quant
 — lipophilicity.py

4.4 chemistry2quant.chem2quant_analysis.rdkitProcessDf Class Reference

Processing data columns in csv to pandas, then to Morgan Fingerprints ($^{\prime\prime}$)

Public Member Functions

• def __init__ (self, directory, sdf_file_name)

The constructor for the class.

def returnMol (self)

Return the data of the molecule in the Mol format (more details and links here: PLACEHOLDEr)

• def MoltoSmiles (self)

Convert Mol to smiles.

• def MACCSfingerprintList (self)

Simple fingerprint list.

• def torsionalfingerprintList (self)

Torsional fingerprint list.

Public Attributes

- · rdkit_directory
- lig_data
- dataMol
- · ms smiles
- MACCSlist
- Pairslist

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4.4.1 Detailed Description

The documentation for this class was generated from the following file:

/home/oohnohnoh1/Desktop/GIT/Chemiinformatics_work/Chemistry2quant/src/chemistry2quant/chem2quant
 analysis.py

4.5 chemistry2quant.chem2quant_analysis.rdkitPsi4DataGenerator Class Reference

Public Member Functions

- def __init__ (molfile)
- def molToPsi4 (self)
- def forEachSimilarity (ref, array)
- def storeMolecule ()

Public Attributes

molfile

4.5.1 Detailed Description

Here, we want to translate the smilestoMol file into a psi4 file and run DFT calculations for each. Based on the code seen in "https://iwatobipen.wordpress.com/2018/08/24/calculate-homo-and-lumo-with-psi4-rdkit

The documentation for this class was generated from the following file:

/home/oohnohnoh1/Desktop/GIT/Chemiinformatics_work/Chemistry2quant/src/chemistry2quant/chem2quant
 —analysis.py

4.6 chemistry2quant.chem2quant_screening.two_step_screen Class Reference

Public Member Functions

- def __init__ (self, smilesList, molecule)
- def first screen ()
- def second_screen ()

Public Attributes

- smilesList
- molecule
- smileMol

4.6.1 Detailed Description

Initial screening of a database with a substructure - The first screen removes 99% of all the options in the tFor example, the emoleucles or the chembl database. The second screen studies the work using a number of algin

4.6.2 Member Function Documentation

4.6.2.1 first_screen()

```
def chemistry2quant.chem2quant_screening.two_step_screen.first_screen ( )
Remove 99% of the structural database which doesn't match the general structure
at all
```

4.6.2.2 second_screen()

```
def chemistry2quant.chem2quant_screening.two_step_screen.second_screen ( )
Building from a simple screening
```

4.6.3 Member Data Documentation

4.6.3.1 smileMol

```
chemistry2quant.chem2quant_screening.two_step_screen.smileMol

Remove 99% of the structural database which doesn't match the general structure
at all
```

The documentation for this class was generated from the following file:

/home/oohnohnoh1/Desktop/GIT/Chemiinformatics_work/Chemistry2quant/src/chemistry2quant/chem2quant
 _screening.py

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