# Hands on with Graph Neural Networks



## Installing Pytorch Geometric and RDKit

- Pytorch Geometric => Build Graph Neural Network
- RDKit => Handle Molecule Data

Note: Currently I didn't find a way to set a specific python version in colab. However, when installing rdkit, only specific python versions are supported... Hence it cannot be ensured that the notebook runs properly. For example Python 3.7.10 is not supported.

```
# Enforce pytorch version 1.6.0
import torch
# if torch.__version__ != '1.6.0':
# !pip uninstall torch -y
# !pip uninstall torchvision -y
# !pip install torch=1.6.0
# !pip install torchvision==0.7.0

# # Check pytorch version and make sure you use a GPU Kernel
# !python -c "import torch; print(torch.__version__)"
# !python -c "import torch; print(torch.version.cuda)"
# !python --version
# !nvidia-smi
```

Make sure you clicked "RESTART RUNTIME" above (if torch version was different)!

```
#@title
# # Install rdkit
# import sys
# import os
# import requests
# import subprocess
# import shutil
# from logging import getLogger, StreamHandler, INFO

# logger = getLogger(__name__)
# logger.addHandler(StreamHandler())
```

```
# logger.setLevel(INFO)
# def install(
          chunk size=4096,
          file name="Miniconda3-latest-Linux-x86 64.sh",
          url base="https://repo.continuum.io/miniconda/"
#
          conda path=os.path.expanduser(os.path.join("~",
"miniconda")),
          rdkit version=None,
          add_python path=True,
#
#
          force=False):
      """install rdkit from miniconda
#
#
#
      import rdkit installer
#
      rdkit installer.install()
#
      0.00
#
#
      python path = os.path.join(
#
          conda path,
#
          "lib",
#
          "python{0}.{1}".format(*sys.version_info),
          "site-packages",
#
#
      if add_python_path and python_path not in sys.path:
#
          logger.info("add {} to PYTHONPATH".format(python path))
#
#
          sys.path.append(python path)
#
      if os.path.isdir(os.path.join(python path, "rdkit")):
          logger.info("rdkit is already installed")
#
#
          if not force:
#
              return
#
          logger.info("force re-install")
#
      url = url_base + file_name
      python_version = "{0}.{1}.{2}".format(*sys.version_info)
#
      logger.info("python version: {}".format(python version))
#
      if os.path.isdir(conda_path):
#
          logger.warning("remove current miniconda")
#
#
          shutil.rmtree(conda_path)
#
      elif os.path.isfile(conda path):
          logger.warning("remove {}".format(conda path))
#
#
          os.remove(conda path)
      logger.info('fetching installer from {}'.format(url))
#
```

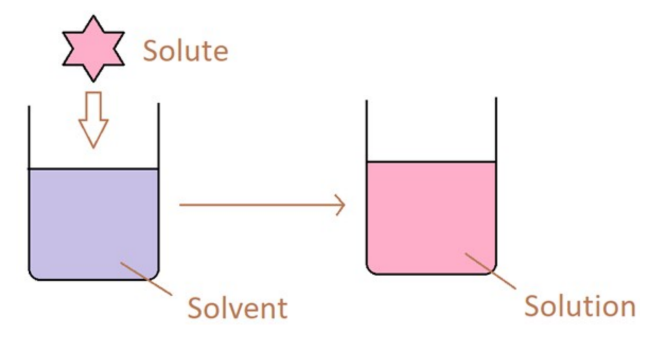
```
res = requests.get(url, stream=True)
#
      res.raise for status()
#
      with open(file_name, 'wb') as f:
          for chunk in res.iter content(chunk size):
#
#
              f.write(chunk)
#
      logger.info('done')
      logger.info('installing miniconda to {}'.format(conda path))
      subprocess.check_call(["bash", file_name, "-b", "-p",
conda path])
      logger.info('done')
      logger.info("installing rdkit")
      subprocess.check_call([
#
          os.path.join(conda path, "bin", "conda"),
#
#
          "install",
#
          "--yes",
          "-c", "rdkit",
          "python==3.7.3",
          "rdkit" if rdkit version is None else
"rdkit=={}".format(rdkit_version)])
      logger.info("done")
      import rdkit
      logger.info("rdkit-{} installation
finished!".format(rdkit.__version__))
# if name == " main ":
      install()
# If something breaks in the notebook it is probably related to a
mismatch between the Python version, CUDA or torch
import torch
# pytorch_version = f"torch-{torch.__version__}.html"
# !pip install --no-index torch-scatter -f https://pytorch-
geometric.com/whl/$pytorch version
# !pip install --no-index torch-sparse -f https://pytorch-
geometric.com/whl/$pytorch version
# !pip install --no-index torch-cluster -f https://pytorch-
geometric.com/whl/$pytorch version
# !pip install --no-index torch-spline-conv -f https://pytorch-
geometric.com/whl/$pytorch version
# !pip install torch-geometric
```

### Background info on the Dataset

In the following we will use a dataset provided in the dataset collection of PyTorch Geometric (Here you find all datasets). The Dataset comes from the MoleculeNet collection, which can be found here.

"ESOL is a small dataset consisting of water solubility data for 1128 compounds. The dataset has been used to train models that estimate solubility directly from chemical structures (as encoded in SMILES strings). Note that these structures don't include 3D coordinates, since solubility is a property of a molecule and not of its particular conformers."

#### >>> Machine Learning task: How are different molecules dissolving in water?



**Source:** https://www.differencebetween.com/difference-between-solubility-product/

### SMILES representation and important sidenodes

**Source:** https://medium.com/@sunitachoudhary103/generating-molecules-using-a-char-rnn-in-pytorch-16885fd9394b

- Using the plain SMILES string as input is not suitable
- This will not consider the molecule structure but rather the grammar of the SMILES string

- The SMILES string can be different for a molecule, depending on the notation (a unique molecule can have multiple SMILES strings)
- Chemical graphs however, are invariant to permutations --> Graph Neural Networks

### Looking into the Dataset

```
import rdkit
from torch_geometric.datasets import MoleculeNet

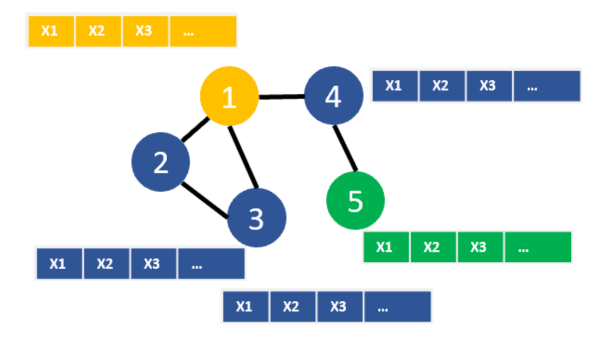
# Load the ESOL dataset
data = MoleculeNet(root=".", name="ESOL")
data

ESOL(1128)
```

Note: There seems to be a change in the Dataset class and somehow the target dim now equals 734 instead of one. You can simply ignore it at this point. :)

```
# Investigating the dataset
print("Dataset type: ", type(data))
print( Dataset type. , type(data))
print("Dataset features: ", data.num_features)
print("Dataset target: ", data.num_classes)
print("Dataset length: ", data.len)
print("Dataset sample: ", data[0])
print("Sample nodes: ", data[0].num_nodes)
print("Sample edges: ", data[0].num_edges)
# edge index = graph connections
# smiles = molecule with its atoms
\# x = node \ features (32 \ nodes \ have each 9 \ features)
\# y = labels (dimension)
Dataset type: <class
'torch geometric.datasets.molecule net.MoleculeNet'>
Dataset features: 9
Dataset target: 734
Dataset length: <bound method InMemoryDataset.len of ESOL(1128)>
Dataset sample: Data(x=[32, 9], edge_index=[2, 68], edge_attr=[68,
3], smiles='0CC30C(0CC20C(0C(C#N)c1ccccc1)C(0)C(0)C20)C(0)C(0)C30',
y=[1, 1]
Sample nodes:
                     32
Sample edges:
                     68
# Investigating the features
# Shape: [num nodes, num node features]
data[0].x
tensor([[8, 0, 2, 5, 1, 0, 4, 0, 0],
           [6, 0, 4, 5, 2, 0, 4, 0, 0],
           [6, 0, 4, 5, 1, 0, 4, 0, 1],
```

```
[8, 0, 2, 5, 0, 0, 4, 0, 1],
[6, 0, 4, 5, 1, 0, 4, 0, 1],
[8, 0, 2, 5, 0, 0, 4, 0, 0],
[6, 0, 4, 5, 2, 0, 4, 0, 0],
[6, 0, 4, 5, 1, 0, 4, 0, 1],
[8, 0, 2, 5, 0, 0, 4, 0, 1],
[6, 0, 4, 5, 1, 0, 4, 0, 1],
[8, 0, 2, 5, 0, 0, 4, 0, 0],
[6, 0, 4, 5, 1, 0, 4, 0, 0],
[6, 0, 2, 5, 0, 0, 2, 0, 0],
[7, 0, 1, 5, 0, 0, 2, 0, 0],
[6, 0, 3, 5, 0, 0, 3, 1, 1],
[6, 0, 3, 5, 1, 0, 3, 1, 1],
[6, 0, 3, 5, 1, 0, 3, 1, 1],
[6, 0, 3, 5, 1, 0, 3, 1, 1],
[6, 0, 3, 5, 1, 0, 3, 1, 1],
[6, 0, 3, 5, 1, 0, 3, 1, 1],
[6, 0, 4, 5, 1, 0, 4, 0, 1],
[8, 0, 2, 5, 1, 0, 4, 0, 0],
[6, 0, 4, 5, 1, 0, 4, 0, 1],
[8, 0, 2, 5, 1, 0, 4, 0, 0],
[6, 0, 4, 5, 1, 0, 4, 0, 1],
[8, 0, 2, 5, 1, 0, 4, 0, 0],
[6, 0, 4, 5, 1, 0, 4, 0, 1],
[8, 0, 2, 5, 1, 0, 4, 0, 0],
[6, 0, 4, 5, 1, 0, 4, 0, 1],
[8, 0, 2, 5, 1, 0, 4, 0, 0],
[6, 0, 4, 5, 1, 0, 4, 0, 1],
[8, 0, 2, 5, 1, 0, 4, 0, 0]])
```



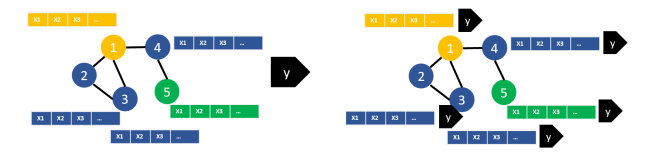
```
# Investigating the edges in sparse COO format
# Shape [2, num_edges]
data[0].edge_index.t()
tensor([[ 0,
              1],
        [ 1,
              0],
          1,
              2],
          2,
               1],
              3],
          2,
        [ 2, 30],
          3,
              2],
          3,
              4],
          4,
               3],
          4,
              5],
          4, 26],
          5,
              4],
          5,
               6],
          6,
               5],
          6,
              7],
        [ 7,
              6],
          7,
              8],
        [7,24],
        [ 8,
              7],
              9],
        [ 8,
        [ 9, 8],
        [ 9, 10],
```

```
[ 9, 20],
          [10, 9],
          [10, 11],
          [11, 10],
          [11, 12],
[11, 14],
          [12, 11],
          [12, 13],
          [13, 12],
          [14, 11],
[14, 15],
          [14, 19],
          [15, 14],
          [15, 16],
          [16, 15],
[16, 17],
          [17, 16],
[17, 18],
          [18, 17],
          [18, 19],
          [19, 14],
          [19, 18],
          [20, 9],
          [20, 21],
          [20, 22],
          [21, 20],
          [22, 20],
          [22, 23],
          [22, 24],
          [23, 22],
          [24, 7],
[24, 22],
          [24, 25],
          [25, 24],
          [26, 4],
          [26, 27],
          [26, 28],
          [27, 26],
          [28, 26],
          [28, 29],
          [28, 30],
          [29, 28],
          [30, 2],
[30, 28],
          [30, 31],
          [31, 30]])
data[0].y
tensor([[-0.7700]])
```

In the following we will perform predictions based on the graph level. This

- 1. List item
- 2. List item

means we have one y-label for the whole graph, as shown on the left image below. The right image would be node-level-predictions.



# Converting SMILES to RDKit molecules - Visualizing molecules

Next we want to have our SMILES molecules as graphs...

```
data[0]["smiles"]
'OCC30C(OCC20C(OC(C#N)clcccccl)C(0)C(0)C(0)C(0)C(0)C(0)C(0)
from rdkit import Chem
from rdkit.Chem.Draw import IPythonConsole
molecule = Chem.MolFromSmiles(data[0]["smiles"])
molecule
```

```
type(molecule)
rdkit.Chem.rdchem.Mol
```

- We can also obtain the features from this RDKit representation
- It tells us everything we need to know e.g. atom features (type, ...), edges, ...
- In our case however, It's even easier as we have the information explicitly given already in the dataset
- Otherwise we would calculate the node features from those atom properties
- -> For datasets containing SMILES representations this would be the way to go

### Implementing the Graph Neural Network

Building a Graph Neural Network works the same way as building a Convolutional

Neural Network, we simple add some layers.

The GCN simply extends torch.nn.Module. GCNConv expects:

- in\_channels = Size of each input sample.
- out\_channels = Size of each output sample.

We apply three convolutional layers, which means we learn the information about 3 neighbor hops. After that we apply a pooling layer to combine the information of the individual nodes, as we want to perform graph-level prediction.

Always keep in mind that different learning problems (node, edge or graph prediction) require different GNN architectures.

For example for node-level prediction you will often encounter masks. For graph-level predictions on the other hand you need to combine the node embeddings.

```
import torch
from torch.nn import Linear
import torch.nn.functional as F
from torch_geometric.nn import GCNConv, TopKPooling,
global mean pool , GATConv
from torch geometric.nn import global mean pool as gap,
global max pool as gmp
embedding_size = 64
class GCN(torch.nn.Module):
    def __init__(self):
        # Init parent
        super(GCN, self).__init__()
        torch.manual seed(42)
        # GCN layers
        self.initial conv = GATConv(data.num features, embedding size)
        self.conv1 = GCNConv(embedding size, embedding size)
        self.conv2 = GCNConv(embedding size, embedding size)
        self.conv3 = GCNConv(embedding size, embedding size)
```

```
# Output layer
        self.out = Linear(embedding size*2, 1)
    def forward(self, x, edge index, batch index):
        # First Conv layer
        hidden = self.initial conv(x, edge index)
        hidden = F.tanh(hidden)
        # Other Conv layers
        hidden = self.conv1(hidden, edge index)
        hidden = F.tanh(hidden)
        hidden = self.conv2(hidden, edge index)
        hidden = F.tanh(hidden)
        # Global Pooling (stack different aggregations)
        hidden = torch.cat([gmp(hidden, batch index),
                            gap(hidden, batch_index)], dim=1)
        # Apply a final (linear) classifier.
        out = self.out(hidden)
        return out, hidden
model = GCN()
print(model)
print("Number of parameters: ", sum(p.numel() for p in
model.parameters()))
GCN(
  (initial conv): GATConv(9, 64, heads=1)
  (conv1): GCNConv(64, 64)
  (conv2): GCNConv(64, 64)
  (conv3): GCNConv(64, 64)
  (out): Linear(in features=128, out features=1, bias=True)
Number of parameters: 13377
```

- We could also reduce the embeddings, but as we have large molecules we use 64
- The more layers we add, the more information we get about the graph
- For the regression problem we use a Linear layer as final output layer
- We try to use not too many parameters, as we only have ~1k samples

### Training the GNN

```
from torch_geometric.data import DataLoader
import warnings
warnings.filterwarnings("ignore")
# Root mean squared error
```

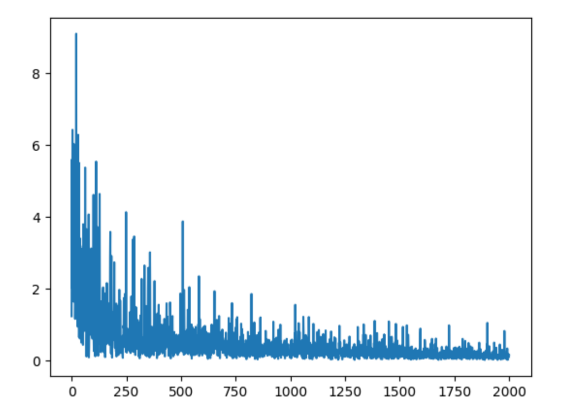
```
loss fn = torch.nn.MSELoss()
optimizer = torch.optim.Adam(model.parameters(), lr=0.0007)
# Use GPU for training
device = torch.device("cuda:0" if torch.cuda.is_available() else
"cpu")
model = model.to(device)
# Wrap data in a data loader
data size = len(data)
NUM GRAPHS PER BATCH = 64
loader = DataLoader(data[:int(data size * 0.8)],
                    batch size=NUM GRAPHS PER BATCH, shuffle=True)
test loader = DataLoader(data[int(data size * 0.8):],
                         batch size=NUM GRAPHS PER BATCH,
shuffle=True)
def train(data):
    # Enumerate over the data
    for batch in loader:
      # Use GPU
      batch.to(device)
      # Reset gradients
      optimizer.zero grad()
      # Passing the node features and the connection info
      pred, embedding = model(batch.x.float(), batch.edge index,
batch.batch)
      # Calculating the loss and gradients
      loss = loss fn(pred, batch.y)
      loss.backward()
      # Update using the gradients
      optimizer.step()
    return loss, embedding
print("Starting training...")
losses = []
for epoch in range(2000):
    loss, h = train(data)
    losses.append(loss)
    if epoch % 100 == 0:
      print(f"Epoch {epoch} | Train Loss {loss}")
Starting training...
Epoch 0 | Train Loss 1.2307624816894531
Epoch 100 | Train Loss 4.608680725097656
Epoch 200 | Train Loss 1.1581676006317139
Epoch 300 | Train Loss 0.31913432478904724
Epoch 400 | Train Loss 0.44638288021087646
Epoch 500 | Train Loss 0.5199289917945862
Epoch 600 | Train Loss 0.9058079719543457
```

```
Epoch 700
          | Train Loss 0.4809274971485138
Epoch 800
           Train Loss 0.05794638767838478
Epoch 900 | Train Loss 0.2919251620769501
Epoch 1000 | Train Loss 0.07834801077842712
Epoch 1100
             Train Loss 0.03218280151486397
Epoch 1200
             Train Loss 0.1406428962945938
Epoch 1300
             Train Loss 0.295724481344223
Epoch 1400
             Train Loss 0.1676671952009201
             Train Loss 0.4162887632846832
Epoch 1500
Epoch 1600
             Train Loss 0.14953072369098663
Epoch 1700
             Train Loss 0.07532656192779541
Epoch 1800 |
             Train Loss 0.09881985932588577
Epoch 1900
             Train Loss 1.0456342697143555
```

### Visualizing the Training loss

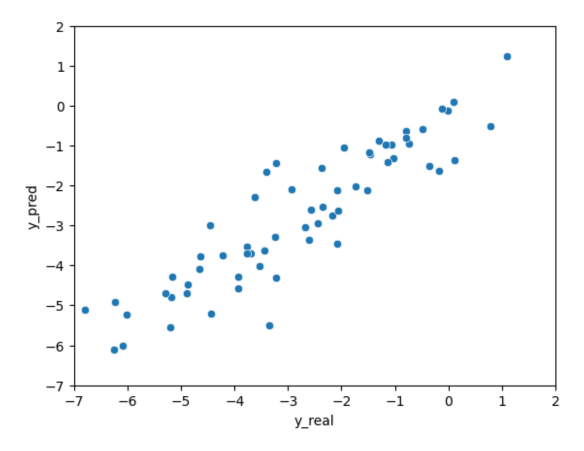
```
# Visualize learning (training loss)
import seaborn as sns
losses_float = [float(loss.cpu().detach().numpy()) for loss in losses]
loss_indices = [i for i,l in enumerate(losses_float)]
plt = sns.lineplot(losses_float)
plt

<Axes: >
```



### Getting a test prediction

```
import pandas as pd
# Analyze the results for one batch
test batch = next(iter(test loader))
with torch.no grad():
    test batch.to(device)
    pred, embed = model(test batch.x.float(), test batch.edge index,
test_batch.batch)
    df = pd.DataFrame()
    df["y real"] = test batch.y.tolist()
    df["y_pred"] = pred.tolist()
df["y_real"] = df["y_real"].apply(lambda row: row[0])
df["y pred"] = df["y pred"].apply(lambda row: row[0])
df
              y_pred
    y_real
     -4.23 -3.743365
1
     -1.06 -0.974107
2
     -1.52 -2.124257
3
    -5.21 -5.555056
4
     -1.30 -0.885036
59
    -3.22 -1.442067
60
   -4.46 -2.989191
61
   -0.80 -0.792187
     -4.66 -4.086609
62
63 -6.25 -6.104234
[64 rows x 2 columns]
plt = sns.scatterplot(data=df, x="y real", y="y pred")
plt.set(xlim=(-7, 2))
plt.set(ylim=(-7, 2))
plt
<Axes: xlabel='y_real', ylabel='y_pred'>
```



## Improving the model / More to play around with

For example you can add:

- Dropouts
- Other (more intelligent) Pooling Layers (all layers here: https://pytorch-geometric.readthedocs.io/en/latest/modules/nn.html#)
- Global Pooling Layers
- Batch Normalization
- More MP layers
- Other hidden state sizes
- Test metrics (test error) and Hyperparameter optimization
- ...

https://github.com/rusty1s/pytorch\_geometric/tree/master/examples