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## Replicating the Results of this Project

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First, we would recommend, using the environmen.yml.

```
If you have a cuda enabled gpu there is a huge advantage to installing  
torch with cuda.  
this is massively preffered as the experiment suite takes ~50 hours on RTX  
3090, with cuda.
```

To create the environment open the terminal on a Unix-like system;

```
conda env create -f environment.yml
```

Then run,

```
python run_experiments.py
```

## Logging into Weights and Biases

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To run the scripts, you will need to make a Weights and Biases account. On wandb.ai.

After creating the environment and account, you can log into Weights and Biases with,

```
wandb login
```

## Guide on how to run experiments for DMT

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## Architecture for Experiments

The architecture of the experiments is dependent on classes. Any experiment performed in this project are subclasses of a parent class BaseExperiments.

The BaseExperiment class contain all the hyper-parameters needed for models within Dynamic Mutual Training (DMT) and for the Baseline models. These are set to some default values that needs to be overridden for experiments to reflect changes in the experiment conducted.

After all experiment subclasses have been created, they are all registered in a global class Experiments that keeps a registry on all the experiments that will be conducted (i.e., all the subclasses). If the user then wants to run all the experiments, the Experiment class have a functionality for this that will be explained below. However, without proper GPU and memory this will be tedious. Even in the case for running a single experiment for a subclass, the user will rely on having significant computing power (robust GPU).

## How the experiments are created

First, we have the Experiment class

```
class Experiments:
    def register:
        # keeps a experiment registry from Base Experiments

    def run_all():
        # runs all registered experiments
```

Then, we have the BaseExperiment class

```
class BaseExperiment(ABC):
    # initialize and holds all hyper-parameters used for
    # running a DMT/Baseline model

    # the properties of each experiment
    @property
    def description():
        # add model description

    @property
    def model_folder():
        # model folder to save experiment(s)

    # this is the function to override for running
    # constructed experiments
    @abstractmethod
    def run():
        # run some of the below run types

    # the type of runs to do for an experiment
    def _train_baseline_only(**baseline_parameters)
```

```

# trains the baseline model and saves it

def _base_run(**dm_t_parameters):
    # trains and saves dmt model

def _plabel_run(**plabel_parameters):
    # trains model on a varying proportion of
    # labeled data with pseudo labels in training after
    # pre-training

```

Now, for the base experiment, we have three run cases. These are all different with respect to what your experiment is performing, and they can all be used in combination with each other. The functionality of the separate run functions are:

- **\_plabel\_run**: Pseudo-label run method for all experiments. This method is called for pseudo label experiments for subclasses. This method trains the PLabel model and saves the best model (See PLabel class for details). If a baseline model is provided, it is also trained and saved.
- **\_base\_run**: Base run method for all experiments. This method is called by the subclass run method. This method trains the DMT model and saves the best model.
- **\_train\_baseline\_only**: Training the baseline model only. Used for comparison with DMT.

Now, to construct an experiment, what you do is:

```

class MyExperiment(BaseExperiments):
    # create some hyperparameters
    # in here that you want to test

    @property
    def model_folder():
        # set a return string with model folder

    @property
    def description():
        # set a return description of the experiment
    def run():
        # Implement the experiment performed here
        # inside use the run function from the three cases
        # of run experiment that you want to use

```

## How to run experiments

To perform DMT / Baseline experiments in the shell, first navigate:

```
path/to/dir/src/experiments
```

Then, if you want to want to conduct your own experiment you can first import Baseline as followed:

```
python from /path/to/dir/src.experiments import BaseExperiments
```

Then follow the steps above for creating an experiment, and then do:

```
my_experiment = MyExperiment()  
my_experiment.run()
```

## Our Experiments

We have the following list of conducted experiments in this project:

- **TrainBaselines**
- **VaryDifferenceMaximization**
- **VaryDMTEpochs**
- **VaryLabelProportion**
- **PLabelVaryLabelProportion**
- **PlabelDefault**

To conduct any of them follow the steps from the "**How to run experiments**". If you want to conduct all of them at once, do:

```
/path/to/dir/src/experiments python run_experiments.py
```

## Project structure

Plotting scripts to output figures used in the reports.

\*\_scripts.py

run\_experiments.py: Runs all experiments

eval\_data/

    |\_\_dmt\_loss.npy: The IoU of the DMT with varying label proportion on the test set. Sorted by label proportion.

    |\_\_dmt\_loss\_dms.npy: The IoU of DMT with varying difference maximised smapling on the test set. Sorted by dms proportion.

    |\_\_dmt\_loss\_epoch.npy: The IoU of DMT with varying epochs on the test set. Sorted by epoch.

    |\_\_plabel\_loss.npy: The IoU of PLabel with the default hyperparameters repeated 5 times.

    |\_\_plabel\_loss\_label.npy: The IoU of PLabel with varying label proportion on the test set. Sorted by label proportion.

    |\_\_baseline.npy: The IoU of the baseline with the default

```
hyperparameters
    repeated 5 times, Sorted varying label proportion.

final_figs/
    Figures folder.

src/
|__experiments/
    |__experiment.py
        This is the main script for interaction with the code.
        It defines a base class that implements default
        hyperparameters and a wrapper for models,
        and how they are run and evaluated.

|__models/
    The implementation of torch.nn.Modules,
    and their training and evaluation methods.

        |__UNet.py
        |__DMT.py
        |__PLabel.py
|__pet_3/
    |__data.py
        The main interface for data is defined here,
        defining torch datasets, with other convenience methods.

        |__download_utils.py
|__utils/
    Utility functions/classes. Only ref if deeper understanding is
    required.
    |__datasets.py
    |__evaluation.py
    |__loading.py
    |__mixin.py
    |__training.py
|__plotting/
    |__temporaty_plotting_utils.py
```