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Pre-Training Graph Neural Networks for Data-Efficient Brain Network Analysis

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Abstract

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The human brain is the central hub of the neurobiological system, controlling behavior and cognition in complex ways. Recent advances in neuroscience and neuroimaging analysis have shown a growing interest in the interactions between brain regions of interest (ROIs) and their impact on neural development and disorder diagnosis. As a powerful deep model for analyzing structural data, Graph Neural Networks (GNNs) have been applied for brain network analysis. However, effective training of deep models requires large amounts of labeled data, which is often scarce in brain network datasets due to the complexities of data acquisition and sharing restrictions. To make the most out of available training data, this work examines data- and label-efficient training of GNN model. In particular, the goal is to pre-train GNN to capture intrinsic brain network structures, regardless of clinical outcomes, and is easily adaptable to various downstream tasks. To this end, the proposed framework comprises three key components: (1) a meta-learning based multi-task pre-training platform with dynamic task adaptive reweighing consideration that learns a generalizable model initialization with efficient optimization schedule (2) an unsupervised pre-training objective designed specifically for brain networks, which enables learning from large-scale datasets without task-specific labels; (3) a data-driven atlas mapping pipeline with variance-based ROI alignment mechanism that facilitates knowledge transfer across datasets with different ROI systems. Extensive empirical evaluations using various GNN backbones have demonstrated the robust and superior performance of the proposed framework compared to baseline methods.
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Chapter 1

Introduction

1.1 Background and Motivation

It has long been an enticing pursuit for neuroscience researchers and mental disorder clinicians to understand the functions and structures of human brains, which are known to be related to many complicated diseases, including bipolar disorder (BP), immunodeficiency virus infection (HIV), and Parkinson’s disease (PPMI) [96] which this study will mainly focus on. In the last decade, the development of neuroimaging techniques, such as magnetic resonance imaging (MRI), functional MRI, diffusion tensor imaging (DTI), etc., provides an important source of information that facilitates the diagnosis of various brain diseases. Based on neuroimaging data, one can build brain networks that encode brain anatomical regions as nodes and their connections as edges. This kind of data representation characterizes the complex connections among different regions of interest (ROI). Effective brain network analysis plays a pivotal role in understanding the biological structures and functions of complex neural systems, which potentially helps the early diagnosis of neurological disorders and facilitates neuroscience research [61, 88, 53].
Graph Neural Networks (GNNs) have emerged as a powerful tool for analyzing graph-structured data, delivering impressive results on a wide range of network datasets, including social networks, recommender systems, knowledge graphs, protein and gene networks, and molecules, among others [47, 28, 73, 79, 87, 92, 95, 54, 86]. These models have proven their ability to learn powerful representations and efficiently compute complex graph structures, making them ill-suited for various downstream tasks. In the field of neuroscience, GNN has been applied to brain network analysis, specifically for graph-level classification/regression [92, 87, 21] and important vertex/edge identification [91, 57, 83], towards tasks such as connectome-based disease prediction and multi-level neural pattern discovery. However, deep learning models, including GNNs, require large amounts of labeled data to achieve optimal performance [38, 93, 99]. While neuroimaging datasets are available from national neuroimaging studies such as the ABCD [10], ADNI [32], and PPMI [2], these datasets are still relatively small compared to graph datasets from other domains, such as datasets with 41K to 452K graphs on OGB [39] and datasets with thousands to millions of graphs on NetRepo [71]). The limited amount of data can result in GNNs having difficulty in learning informative knowledge and easily overfitting the data distribution.

Recently, to improve data efficiency, the framework of transfer learning has attracted a lot of attention in many application domains, which allows a model pre-trained on large-scale source datasets to be adapted to smaller target datasets while maintaining robust performance. However, the success of transfer learning depends on the availability of similar supervision labels on the source and target dataset. This is not always feasible in large-scale public studies, particularly in the field of brain network analysis. One other major challenge is the inconsistent ROI parcellation systems in constructing different brain network datasets, which hinders the transferability of pre-trained models across datasets. The process of parcellating raw imaging data into
brain networks is highly complex and usually done ad hoc by domain experts for each study, making it unrealistic to expect every institution to follow the same parcellation system. Although some institutions may release preconstructed brain network datasets [20], the requirement for universal adherence to a single parcellation system is infeasible.

1.2 Overview of Proposed Solutions

To tackle these challenges, this work aims to explore meta-learning techniques and self-supervised pre-training for GNNs. The framework of meta-learning, also known as learning to learn, aims at learning over multiple, seemingly diverse tasks during the pre-training phase, with the goal of deriving a generalized initialization of the model such that it can be adapted to any arbitrary unseen tasks with efficient convergence. One of the advantages of this approach is that the model can be trained by multiple tasks simultaneously. Considering the multimodal nature of the brain network datasets where multiple interrelated types of connections exist among ROIs (e.g., structural and functional connections), we propose to leverage every such modality as one training task and meta-train the models using multiple training tasks. With sufficient amount of meta-training, we have a pre-trained model that simultaneously performs well on all these training tasks, which we believe to be generic and easily transferable to new target tasks.

On the other hand, self-supervised pre-training has been shown to be effective in various domains, such as computer vision [29, 12], natural language processing [18, 69], and graph mining [76]. This work also aims to explore a self-supervised pre-training approach for GNNs on brain networks that is not restricted by task-specific supervision labels. Despite the promising potential, unique challenges still need to be
addressed to achieve effective disease prediction. In particular, this work proposes a novel two-level contrastive learning strategy based on the naturally aligned node systems of brain networks across individuals.

Based on the meta-learning framework and the contrastive self-supervised training strategy, this work further improves the model with brain-network-oriented designs. At first, the datasets used for training and testing usually use different ROI atlas mappings for constructing the brain networks, resulting in different numbers and physical regions of nodes, which hinders the transferability of GNNs. To mitigate this discrepancy, this work proposes to leverage a linear autoencoder model that transforms the original features into low-dimensional representations in a uniform embedding space and aligns them using variance-based projection, which incorporates regularizations that preserve spatial relationships, consider neural modules, and promote sparsity. Secondly, in the meta-training phase, different training tasks may contribute differently to the learning of generic and transferable knowledge which may limit the generalization performance. This work motivates the design consideration by visualizing the relative contribution of the source tasks towards the learning on the target task, where the data-driven observation corroborates with existing clinical research. Based on this findings, this work then proposes an adaptive task reweighing scheme to dynamically adjust the learning rate and weight decay parameters according to the contribution of each meta-training task. Extensive experiments and ablation studies conducted on real-world brain network datasets verify the effectiveness of these proposed strategies.

1.3 Summary of Contribution

In summary, the contribution of this work is four-folded:
• This work is the first to highlight the inherent challenge of limited training samples for learning with brain network data. This work formulates this problem into a data-efficient learning objective with the goal of pre-training the model to a generalizable initialization that can effectively adapt to unseen downstream objectives.

• This work proposes to leverage meta-learning strategies to pre-train a given model on available source tasks. In addition, the pre-training process is powered by a novel two-level contrastive sampling strategy that considers special properties of brain network data.

• This work also addresses unique challenges in multi-dataset and cross-dataset learning on brain networks by proposing brain-network-specific design components featured by a linear autoencoder network with customized regularizations, a dynamic task reweighing mechanism for multi-task pre-training, and a variance-based sorting algorithm to promote ROI alignment after dataset-specific atlas tranformation.

• This work also conducts extensive experiments to benchmark the working effectiveness of the proposed framework against a multitude of state-of-the-art methods adapted to our setting. In addition, the work also investigates the contribution of each constituent parts of the framework through series of ablation studies.
Chapter 2

Related Work

2.1 GNNs for Brain Network Analysis.

In recent years, graph neural networks (GNNs) have attracted broad interest due to their established power for analyzing graph-structured data [80, 87, 48]. Several pioneering deep models have been devised to predict brain diseases by learning the graph structures of brain networks. For example, BrainGNN [52] proposes ROI-aware graph convolutional layers and ROI-selection pooling layers for predicting neurological biomarkers. BrainNetCNN [44] designs a CNN that includes edge-to-edge, edge-to-node, and node-to-graph convolutional filters, leveraging the topological locality of brain connectome structures. BrainNetTF [43] introduces a transformer architecture with an orthonormal clustering readout function that considers ROI similarity within functional modules. Additionally, various studies [15, 42, 100, 14] have shown that, when data is sufficient, GNNs can greatly improve performance in tasks such as disease prediction. However, in reality, the lack of training data is a common issue in neuroscience research, particularly for specific domains and clinical tasks. Despite this, there has been little research into the ability of GNNs to effectively train for brain network analysis when data is limited.
2.2 Meta-Learning for Graph Classification

Recently, meta-learning has drawn significant attention in the machine learning community since it is able to address the problem of limited training data. There are also several attempts of meta-learning for GNN-based graph classification. For example, [11] recognize unseen classes with limited labeled graph samples using meta-training. [7] attempt to develop a general framework that can adapt to three-level tasks — graph classification, node classification, and link prediction with meta-learning, but without considering the unique characteristics of brain networks. [59] use the shared sub-structures between training classes and test classes to design a better meta-learning framework. However, none of the shared sub-structures can be utilized since brain networks are complete graphs. Meta-MGNN [27] proposes a self-supervised learning objective that predicts atom types for molecular datasets. However, there is no precise label for each node for prediction in brain networks.

2.3 Unsupervised Graph Representation Learning and GNN Pre-training.

Unsupervised learning is a widely used technique for training complex models when resources are limited. Recent advancements in contrastive learning [13, 29] have led to various techniques for graphs. For instance, GBT [5] designs a Barlow Twins [94] loss function based on the empirical cross-correlation of node representations learned from two different views of the graph [97]. Similarly, GraphCL [93] involves a comparison of graph-level representations obtained from two different augmentations of the same graph. DGI [82] contrasts graph and node representations learned from the original graph and its corruption.
To obtain strong models for particular downstream tasks, unsupervised training techniques can be used to pre-train a model, which is then fine-tuned on the downstream tasks to reduce the dependence on labeled training data. The approach has proven highly successful in computer vision [9, 24], natural language processing [19, 69, 68], and multi-modality (e.g. text-image pair) learning [49, 90]. There are various strategies for pre-training GNNs as well. GPT-GNN [40] proposes graph-oriented pretext tasks, such as masked attribute and edge reconstruction. L2P-GNN [56] introduces dual adaptation by simultaneously optimizing the encoder on a node-level link prediction objective and a graph-level self-supervision task similar to DGI. Others, such as GMPT [36] adopt an inter-graph message-passing approach to obtain context-aware node embedding and optimize the model concurrently under supervision and self-supervision. To the best of our knowledge, the effectiveness of both contrastive learning and pre-training has not been investigated in the context of the unique properties of brain networks.
Chapter 3

Problem Definition

This work considers the problem of disease prediction with multiple brain network datasets. Formally, given a dataset for one specific disease $\mathcal{D} = \{\mathcal{G}_i\}^N_{i=1}$ containing $N$ subjects, where $\mathcal{G}_i$ represents the $i$th brain network instance. Each brain network object can be considered as an edge weighted graph $\mathcal{G}_i = (\mathcal{V}, \mathcal{E}_i, A_i)$, where $\mathcal{V} = \{v_i\}^M_{i=1}$ is the node set of size $M$ describing the defined region of interests (ROIs), $\mathcal{E}_i = \mathcal{V} \times \mathcal{V}$ is the weighted edge set, and $A_i \in \mathbb{R}^{M \times M}$ is the weighted adjacency matrix representing the connectivity among ROIs. Since each disease can be recorded in multiple datasets and each dataset can have multiple views of brain networks, we define a training task to be the prediction of one disease on a specific view of brain networks (e.g., different types of functional networks and structural networks). In our cross-dataset multitask learning setting, one aims to train a $\Theta$ parameterized model $f(\cdot)$ on a set of source tasks $\mathcal{S} = \{S_k\}_{k=0}^K$ to obtain $\Theta_0$ such that the weights capture generalized domain knowledge of brain structures that are useful and transferable to an unseen target task $\mathcal{T}$, where $\mathcal{S}$ and $\mathcal{T}$ do not necessarily concern the same type of disease. One can then aim to fine-tune $f(\cdot)$ on $\mathcal{T}$ such that the model can efficiently adapt to the target task optimal $\Theta^*$ given that available training samples in $\mathcal{T}$ are much fewer than those in $\mathcal{S}$. 
Figure 3.1: Overview of the proposed framework. The initial features of the source datasets are projected to a fixed dimension through atlas transformation followed by variance-based feature alignment, which facilitates self-supervised GNN pre-training on multiple datasets via the novel two-level contrastive learning objective. The learned model can serve as the parameter initialization and be further fine-tuned on target tasks.
Chapter 4

Data-Efficient Training Strategies

Graph neural networks are powerful in learning representations of graph-structured objects such as brain networks. However, under a direct train- and-test setting, GNN needs a relatively large-sized dataset for proper training. With small-sized datasets like brain networks, GNNs may suffer from overfitting and fail to generalize the learned knowledge, which leads to a deterioriated performance in downstream tasks. In this chapter, the paper studies the problem of data-efficient training using multiple sources of datasets. Specifically, given one large-sized dataset and the other smaller-sized datasets, the goal is to study how to pre-train the model on the larger dataset (i.e., the source dataset) and use the learned knowledge to improve the performance on smaller ones (i.e., the target datasets).

In the following, this work proposes to study two data-efficient training strategies for brain network analysis — single-task transfer learning and multi-task meta-learning, both of which are representative techniques in dealing with the absence of sufficient training data. In addition, the work presents two other baseline techniques, namely, learning without pre-training and multi-task transfer learning (without the meta-learning portion).
4.1 Method 1: Learning Without Pre-training (NPT)

As a baseline investigation, one directly applies and trains a randomly initialized model on a given task. The model is optimized under a given objective function. The discussion of the setup of objective functions for pre-training is deferred to the subsequent chapter when introducing the self-supervised training strategies in Chapter 5. Specifically under this setting, the source dataset (i.e., the larger-sized dataset) is not used. Instead, the model is directly evaluated on the smaller-sized target dataset, which indeed have limited training samples. Since the downstream objective is to perform binary classification on a specific disease (i.e., determine whether infected or not), the binary cross-entropy loss is used throughout. In particular, the loss is given as:

\[
L_{\text{bce}} = -\frac{1}{|D|} \sum_{(G_i, y_i) \sim D} y_i \log \sigma(f_\theta(G_i)) + (1 - y_i) \log(1 - \sigma(f_\theta(G_i))),
\]

where \(y_i\) stands for the ground truth label, and \(\sigma(x) = \frac{1}{1+e^{-x}}\) is the sigmoid activation function on the output logits. The testing performance is reported using \(k\)-fold cross-validation and this work reports an averaged metric along with standard deviation.

4.2 Method 2: Single-task Transfer Learning (STT)

At first, this work follows the pre-training and fine-tuning scheme in transfer learning [64] to distill knowledge from source task to target task in a sample-efficient way. This framework consists of two consecutive phases: pre-training and fine-tuning. Specifically, one first trains the encoder model on the source task and apply the model weights to train another encoder on the target task.

In the first pre-training phase, one trains the model on the source tasks using the objective described in Eq. (5.2). Then, in the fine-tuning phase, the trained weights
\( \Theta_0 \) are used to initialize another encoder model. This model is then fine-tuned on the target task with the same objective function as Eq. (4.1). Since the model has already learned generic knowledge underneath the source task, one uses a smaller learning rate to optimize the model in the fine-tuning phase. This method is summarized in Algorithm 1. Note that although the source dataset may contain multiple structural views, here this study defines only one view as the source task since in the pre-training phase, the model is trained based on one unified objective function and cannot distinguish between multiple tasks, if they are arbitrarily grouped together.

### 4.3 Method 3: Multi-task Transfer Learning (MTT)

Pre-training on a singular source task is vulnerable to the inherent risk of information loss during transfer learning since the knowledge gaps among source and target domains are not readily quantifiable. This motivates the investigation to train a model that is initialized on some shared knowledge in multiple source tasks when they are available, such that the fine-tuning performance is not conditioned upon any particular knowledge inconsistencies from a source and target pair.

As an immediate solution, this work extends STT into a multi-task setting by expanding the pre-training phase into simultaneously co-learning over multiple source objectives. That is, one can regard each modality from the dataset as an individual task and the model now learns over multiple modalities. To this end, this work formulates all tasks into a distribution, where during pre-training, the model is trained on several objectives sampled from the task distribution, hence the name “multi-task” for this method.

Specifically for each pre-training iteration, one can optimize the model parameters
on a merged objective function which takes the sum over the pre-training objective as given in Eq. (5.2) on all source tasks. For an efficient computation, each iteration processes a mini-batch of data sampled from the source dataset. The learned weights are then used in the fine-tuning phase on the target task following the conventional downstream evaluation procedure, identical to NPT.

4.4 Method 4: Multi-task Meta-Learning (MML)

Meta-learning aims at learning a meta model that is capable of generalizing over a variety of source objectives and can quickly adapt to an arbitrary unseen task. Different from MTT, meta-learning aims at finding an optimal model initialization that enables similarly good performance on multiple pre-training tasks rather than directly combining individual models that are good for each pre-training task through averaging the model weights. This means that meta-learning can achieve better generalization, allowing efficient adaptation to unseen objectives through minimizing the risk of over-fitting the model to outperform on certain tasks while under-perform on others, which is a typical underlying concern of MTT.

Based on such intuition, this work follows the widely adopted model-agnostic meta-learning (MAML) [23] method in brain network learning framework. According to [70], MAML is characterized by two iconic features: (1) rapid learning and (2) feature reuse, which also refers to the outer-loop update and inner-loop adaptation. Specifically, the model is first separately trained on each objective using fast weights during the inner loop function, then the meta parameters of the model are updated by evaluating the loss against the adapted fast weights via the outer-loop module. In other words, the model is optimized by updating on the second-order Hessian of the parameters, which leads to quicker convergence since the optimizer incorporates the
additional curvature information of the loss function that helps estimate the optimal step-size along the optimization trajectory [77]. This effectively reduces the number of training iterations required to achieve a generic model. In addition, the feature reuse inner-loop performs task-specific adaptation, which results in the meta-initialization to be an informative approximation to every task. Due to the fact that the meta-trained model does not pertain to any particular task knowledge, such initialization is therefore non-over-fitting and generically applicable to any unseen target tasks.

To be specific about pipeline design, in the first meta-training phase, one randomly draws $n$ training tasks with a support set (used in inner-loop) and a query set (used in outer-loop) each containing $k$ samples from the pool of training datasets. Then, given the encoder model, the fast weights of the parameters is updated using the objective given in Eq. (5.2) for every pre-training ($i.e.,$ source) task. After training the model on all tasks, one updates the meta parameters, $i.e.,$ model initialization in our case. Thereafter, in the meta-test phase, one performs the conventional classification procedure on the target data identical to NPT. This method is summarized in Algorithm 2.
Algorithm 1 Single-task supervised transfer learning (STT)
1: **Input:** pre-train task $S$, fine-tune task $T$, encoder $f(\theta)$
2: **Require:** $\alpha$: learning rate hyperparameter
3: Randomly initialize $\theta$
4: $\triangleright$ Pre-training phase
5: while not done do
6: Evaluate the gradient $\nabla_\theta \mathcal{L}_S f(\theta)$
7: Update parameters with SGD: $\theta \leftarrow \theta - \alpha \nabla_\theta \mathcal{L}_S f(\theta)$
8: end while
9: $\triangleright$ Fine-tuning phase
10: Split $T$ into $T_{train}$ and $T_{eval}$ into $K$ folds
11: for split in $K$ folds do
12: Get split-specific parameters $\hat{\theta}$ ← $\theta$
13: while not done do
14: Evaluate the gradient $\nabla_{\hat{\theta}} \mathcal{L}_{T_{train}} f(\hat{\theta})$
15: Update parameters with SGD $\hat{\theta} \leftarrow \hat{\theta} - \alpha \nabla_{\hat{\theta}} \mathcal{L}_{T_{train}} f(\hat{\theta})$
16: end while
17: Evaluate ACC, AUC from $f_{\hat{\theta}}(T_{eval})$
18: end for

Algorithm 2 Multi-task meta-learning (MML)
1: **Input:** meta-train task pool $S_\tau$, meta-test task $T$, encoder $f(\theta)$
2: **Require:** $\alpha$, $\beta$: learning rate hyperparameters
3: Randomly initialize $\theta$
4: $\triangleright$ Meta-training phase
5: while not done do
6: for each task $\tau_i$ in $S_\tau$ do
7: Sample $k$ datapoints $\mathcal{D}_i$ from $\tau_i$
8: Evaluate the gradient $\nabla_\theta \mathcal{L}_{\mathcal{D}_i} f(\theta)$
9: Compute the adapted parameters $\theta'_{i} \leftarrow \theta - \beta \nabla_\theta \mathcal{L}_{\mathcal{D}_i} f(\theta)$
10: Sample another set of datapoints $\mathcal{D}'_{i}$ from $\tau_i$
11: end for
12: Update parameters $\theta \leftarrow \theta - \alpha \nabla_\theta \sum_{\mathcal{D}'_i, \theta'_i \sim S_\tau} \mathcal{L}_{\mathcal{D}'_i} f(\theta'_i)$
13: end while
14: $\triangleright$ Meta-test phase
15: Perform $k$-fold evaluation on target tasks
Chapter 5

Unsupervised Brain Network Pre-training

Given the high cost of acquiring labeled training data for brain network analysis, the pre-training pipeline of this work adopts to the effective label-free learning strategy of contrastive learning (CL). CL aims to maximize the mutual information (MI) between an anchor point of investigation $X$ from a data distribution $\mathcal{H}$ and its positive samples $X^+$, while minimizing MI with its negative samples $X^-$. The contrastive objective function is formulated as follows:

$$J_{\text{con}} = \arg \min_h \left[ -I(X; X^+) + I(X; X^-) \right]. \quad (5.1)$$

In the context of graph CL, given an anchor node representation $z_\alpha$, a set of positive samples $S^+$, and a set of negative samples $S^-$, the training objective is based on the Jensen-Shannon divergence [34],

$$J_{\text{JSD}}(z_\alpha) = \arg \min \left[ -I(z_\alpha; S^+) + I(z_\alpha; S^-) \right], \quad (5.2)$$
where

\[
I(z_\alpha; S^+) = \frac{1}{|S^+|} \sum_{z_s^+ \in S^+} \text{sp}\left( \frac{z_\alpha^T z_s^+}{\|z_\alpha\| \|z_s^+\|} \right),
\]

(5.3)

\[
I(z_\alpha; S^-) = \frac{1}{|S^-|} \sum_{z_s^- \in S^-} \text{sp}\left( \frac{z_\alpha^T z_s^-}{\|z_\alpha\| \|z_s^-\|} \right),
\]

(5.4)

and \( \text{sp}(\cdot) = \log(1 + e^\cdot) \) is softplus nonlinearity.

The ultimate goal of our framework is to localize effective GNN CL learning [102] for brain networks. Given a dataset \( D \) and an anchor node \( i \) from graph \( G_p \in D \) with the learned representation \( z_{i,p} \), this work proposes to categorize the possible sample selections into three fundamental types (a visualization is shown in Figure 5.1):

- **\( S_1 \):** \( \{z_{j,p} : j \in N_k(i,p)\} \) refers to the node representation set within the the \( k \)-hop neighborhood of the anchor in graph \( G_p \).

- **\( S_2 \):** \( \{z_{j,p} : j \notin N_k(i,p)\} \) refers to the remaining node representation set in graph \( G_p \) that are not in the the \( k \)-hop neighborhood of the anchor.

- **\( S_3 \):** \( \{z_{j,q} : G_q \in D, j \in G_q, q \neq p\} \) refers to the node representation set of nodes in all the other graphs of dataset \( D \).

Notice that this framework leverages the \( k \)-hop substructure around the anchor node to further differentiate \( S_1 \) and \( S_2 \) for contrastive optimization. This design is driven by two considerations: (1) Regarding GNN learning. Given that node representations are learned from the information aggregation of its \( k \)-hop neighborhood, maximizing the MI of an anchor to its \( k \)-hop neighbors naturally enhances lossless message passing of GNN convolutions. (2) Regarding the uniqueness of brain networks. Brain networks can be anatomically segmented into smaller neural
system modules [16], thus capturing subgraph-level knowledge can provide valuable signals for brain-related analysis.

Building on these three fundamental types of samples, one can take advantage of the property of brain networks that ROI identities and orders are fixed across samples to introduce an additional sample type. This encourages the GNN to extract shared substructure knowledge by evaluating the MI of an anchor against its presence in other graphs. Given an anchor representation $z_{i,p}$ of node $i$ from graph $G_p \in \mathcal{D}$, the novel inter-graph sample type is defined as:

- $\mathbf{S}_4$: \{ $z_{j,q} : j \in \mathcal{N}_k(i,q) \cap \mathcal{N}_k(i,p), G_q \in \mathcal{D}, q \neq p$ \}, refers to the node representation set within the $k$-hop neighborhood of node $i$ in all other graphs in $\mathcal{D}$.

Conceptually, $\mathbf{S}_4$ is a special subset of $\mathbf{S}_3$.

It is important to note that for an anchor node $i$, its $k$-hop neighborhood structures might not be identical among different graphs. As a result, one can only consider shared neighborhoods when evaluating the mutual information across multiple graphs. To encourage the learning of unique neighborhood knowledge within a single brain network instance and shared substructure knowledge across the entire dataset, the proposed pipeline configures $\mathbf{S}_1$ and $\mathbf{S}_4$ as positive samples while $\mathbf{S}_2$ and the set $\mathbf{S}_3 - \mathbf{S}_4$
Figure 5.2: The sampling configuration of the proposed framework. $S_1$ and $S_4$ are positive samples, $S_2$ and the set $S_3 - S_4$ are negative samples.

as negative samples, as illustrated in Figure 5.2. Furthermore, the proposed sampling
categorization can also help understand the objective formulations in various state-of-the-art graph CL frameworks [82, 67, 85, 75, 99]. The findings are summarized in Table 5.1. Specifically, “+” denotes positive sampling; “-” denotes negative sampling; and “/” means that the sample type is not considered. It can be observed that DGI and InfoGraph (InfoG) use graph representation pooled from node representations as a special sample, which is essentially equivalent to jointly considering $S_1$ and $S_2$ without explicit differentiation. On the other hand, GCC and EGI, which are more closely related to the proposed framework, leverage neighborhood mutual information maximization on a single graph, but fail to extend this to a multi-graph setting.

<table>
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<tr>
<th></th>
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<th>$S_3$</th>
<th>$S_4$</th>
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<td>/</td>
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Chapter 6

Brain Network Oriented Design Considerations

Unlike conventional graph-structured datasets, brain networks have some unique properties. In this section, this paper first identifies two challenges concerning learning with brain networked data. Accordingly, two design considerations are presented to address these two challenges.

6.1 Data-driven Brain Atlas Mapping

6.1.1 Challenges

For brain network data, ROI templates describe the mapping relationship between nodes and brain atlas. Once the template is chosen, all graphs in a dataset share the same amount of nodes and their physical meanings. In our cross-dataset setting, considering that the source and target datasets are based on different templates, it is difficult to directly transfer the learned knowledge from source to target datasets due to the misalignment of nodes and dimensions in the graphs. Although GNNs are capable of handling input graphs of varied sizes, the model is essentially learning
predictive signals regarding the structures of local subgraphs [50], and thus simply
transferring the model parameters without manipulating data-level correspondence
may lead to a significant loss of information. Note that we may directly convert
different atlas to a unified one through manual mapping. However, finding all such
mappings exhaustively is costly and demands tremendous expert efforts because the
mapping varies across different pairs of atlas and there is often a lack of ground truth.

To address this issue, this study aims to provide a data-driven atlas mapping solution
that is easily accessible and eliminates the strong dependency on network construc-
tion. The data-driven atlas mapping solution, which transforms the original node
features into lower-dimensional representations that preserve the original connectiv-
ity information and align features across datasets, is learned independently on each
dataset prior to GNN pre-training.

6.1.2 Autoencoder with Customized Regularizers

The proposed framework adopts a one-layer linear autoencoder (AE) as the base
structure that transforms source data into a target dimension with fixed representa-
tion in an unsupervised fashion. The AE consists of a linear projection encoder $W$
and a transposed decoder $W^T$, with the goal of learning a low-dimensional projection
that can easily reconstruct the original presentation. The loss function is defined as
minimizing the reconstruction error

$$L_{\text{rec}} = \frac{1}{M} \|X - XWW^T\|_2^2,$$

(6.1)

where $X \in \mathbb{R}^{M \times M}$ is the input and $W \in \mathbb{R}^{M \times D}$ is the learnable projection [33]. To
further enhance the feature compression and to guide the overall AE optimization, this
work proposes to incorporate several regularizers that take into account the unique
characteristics of brain networks:

**Locality-Preserving Regularizer (LR)**

We aim to ensure that the compressed features preserve the spatial relationships of the original brain surface. To achieve this, we incorporate a locality preserving regularizer \cite{30} to the AE objective. The regularizer is formulated as \( \mathcal{L}_{\text{loc}} = (1/M)\|\mathbf{Y} - T\mathbf{Y}\|^2 \), where \( \mathbf{Y} \in \mathbb{R}^{M \times D} \) represents the projected features from the AE and \( T \in \mathbb{R}^{M \times M} \) is a transition matrix constructed from the \( k\)-NN graph of the 3D coordinates of ROIs.

**Modularity-Aware Regularizer (CR)**

Brain networks can be segmented into various neural system modules that characterize functional subsets of ROIs. In graph terminology, they are community structures. The projected feature should also capture information about neural system membership. However, obtaining ground-truth segmentations is a difficult task that requires expert knowledge. To overcome this challenge, we resort to community detection methods on graphs, specifically based on modularity maximization. The regularizer \cite{72} is defined as minimizing

\[
\mathcal{L}_{\text{com}} = -\frac{1}{2D} \sum_{i,j=1}^{M} \left[ A_{ij} - \frac{k_i k_j}{2D} \right] \exp\left(-\|y_i - y_j\|_2^2\right), \quad (6.2)
\]

where \( A \in \mathbb{R}^{M \times M} \) is the graph adjacency matrix, \( k_i \) denotes degree of node \( i \), and \( y_i \) is the AE projected features. Essentially, this optimization minimizes the \( L_2 \) distance between representations of nodes within the same communities, as measured by the modularity score, and maximizes the distance between representations of nodes in different communities.
Sparsity-Oriented Regularizer (SC)

Sparse networks have proven to be effective in learning robust representations from noisy data [41, 74, 60]. In brain connectome analysis, sparsity has also been shown to improve the interpretation of task-specific ROI connections in generation and classification tasks [42]. To this end, we implement the popular KL-divergence smoothing to enforce sparsity in the parameters of the linear projection encoder, $W$. This is formulated as:

$$L_{KL} = \sum_{i=1}^{M} \sum_{j=1}^{D} \left[ \rho \log \left( \frac{\rho}{\hat{\rho}_{ij}} \right) + (1 - \rho) \log \left( \frac{1 - \rho}{1 - \hat{\rho}_{ij}} \right) \right], \quad (6.3)$$

where $\rho$ is a small positive float set as the target sparsity value, and $\hat{\rho}_{ij}$ represents the element-wise activation of the encoder projection matrix $W \in \mathbb{R}^{M \times D}$.

### 6.1.3 Variance-based Dimension Sorting

In addition to transforming dataset-specific features, cross-dataset alignment of feature signals is also crucial for improving model adaptation. The one-layer AE transforms the original feature vectors into weighted combinations of multiple dimensions, creating new feature dimensions which this work names as virtual ROIs. In the context of brain networks, this process helps to group ROIs and their signals. This idea is inspired by the well-studied functional brain modules [66, 3, 31, 6, 98], which provide a higher-level and generic organization of the brain surface, as opposed to fine-grained ROI systems. Since the variations in ROI parcellations are due to differences in clinical conventions, it is reasonable to assume that there exists a shared virtual ROI system underlying different parcellation systems, similar to the discretization of functional brain modules. The community learning and neighborhood preserving regularizers, introduced in Section 6.1.2, allow one to capture these shared virtual ROIs in a data-driven manner. Our ultimate goal is to align the discovered virtual ROIs
across datasets, so that each virtual ROI characterizes the same functional module in
the human brain, regardless of its origin. This cross-dataset alignment of virtual ROIs
ensures that the model can effectively adapt to new datasets and provide meaningful
insights into the different downstream analyses.

The objective of the one-layer linear AE is similar to PCA, as discussed in more
detail in Appendix A.1, with the added benefit of incorporating additional regulariz-
ers. PCA orders dimensions based on decreasing levels of sample variance [35]. The
proposed framework leverages this approach by utilizing the learned parameters of
the AE projection to estimate the variance of each virtual ROI (i.e., projected feature
dimension). The sample variance of each virtual ROI indicates its representativeness
of the original data variations. Given the shared patterns across different parcellation
systems, one can expect that similar virtual ROIs in datasets with different atlas tem-
plates will have similar variance scores, especially in terms of their order. By sorting
the same number of virtual ROIs based on their sample variance in each dataset, the
proposed framework aims to align virtual ROI cross datasets, so that each virtual ROI
represents the same functional unit in the human brain. The procedure is explained
in detail in Algorithm 4 in Appendix A.2.

6.2 Source Task Reweighing

6.2.1 Challenges

Another challenge of cross-dataset brain network analysis is that the previous base
meta-learning pipeline fails to consider relative difficulty of different individual tasks.
It is possible that varying the choice of source task does not lead to uniform im-
provements on the target performance. This means that one can suspect that this is
because some tasks are easier to learn than others, which will converge faster during
the meta training phase. In other words, the base meta-learning pipeline fails to equally capture the latent knowledge of all source datasets, which potentially hinders the ability of generalization.

6.2.2 Dynamic Task Reweighing

The first step is to investigate the data-level task correlations. In particular, this work analyzes the task similarity between the HIV and BP modalities (i.e., target datasets) with respect to the PPMI modalities (i.e., source dataset). A detailed introduction of the datasets including the variety of data modalities and their pre-processing information are deferred to Chapter 7. Inspired by task2vec [1], for each task, one calculates a respective task embedding that stores information regarding its learning difficulty and latent knowledge. In particular, the embedding is derived from the Fisher information estimation of the positive semidefinite upper bound of the Hessian matrix, on which the model is trained on an encoder model using the same objective in Eq. (5.2). This work visualizes the task correlation in cosine similarity among the embeddings on HIV and BP in 6.2.2 respectively. It can be seen that there is an inherent correlation among source (i.e., PPMI) and target (i.e., HIV, BP) datasets which indicates that there exists shared properties and latent information among the three categories of brain network data. This observation can be corroborated with existing clinical research presented in earlier studies [62, 17, 63, 22], where detailed analyses on the coexistence and co-influence among BP, HIV, and PPMI disease are discussed. This validates the working effectiveness of the cross-dataset learning setting since useful and transferable inter-domain knowledge and shared features can be discovered by learning on a source data. In addition, the visualization also shows a non-uniform task correlation, which suggests that the source tasks are prescribed to varying level of learning and adaptation difficulty relative to the given target task. This demonstrates that the optimizer tends to distribute unequal attention within
Figure 6.1: Task correlations among different data modalities from source and target datasets. The Fisher information estimation is first computed to derive from the Hessian matrix by training each task using the same architecture as in Section 4.4. The task embedding is then composed of layer-wise concatenation of the flattened Fisher information matrix.

Following the mechanism proposed by ALFA [4], during the task-specific inner-loop update, the proposed framework implements a trainable hyperparameter generator that guides the rate of convergence for the gradient-descent update. The generator processes the learning state as input, which is consisted of a stacked layer-wise value of model parameter and gradient estimate. The generator then outputs a layer-wise learning rate and weight decay coefficient conditioned on the current learning state. Then, its parameters are updated by the query loss objective as in Eq. (5.2). Different from the original ALFA, where the encoder parameters are frozen from updating at the outer-loop phase, we allow the encoder to be trainable on the query set for quicker adaptation. This variant is summarized (dubbed AR) in Algorithm 3.
Algorithm 3 Multi-task meta-learning with adaptive task reweighing (AR)

1: **Input:** meta-train tasks $S_T$, meta-test task $T$, encoder $f(\theta)$, hyperparameter generator $g(\phi)$
2: **Require:** $\eta$: outer-loop learning rate
3: Randomly initialize $\theta$, $\phi$
4: $\triangleright$ Meta-training phase
5: while not done do
6: for each task $\tau_i$ in $S_T$ do
7: Sample $n$ datapoints $D_i$ from $\tau_i$
8: Evaluate the gradient $\nabla_\theta L_{D_i} f(\theta)$
9: Obtain the task-specific learning state $\rho_i = [\nabla_\theta L_{D_i} f(\theta), \theta]$
10: Generate hyperparameters $\alpha$, $\beta = g(\rho_i)$
11: Compute the adapted parameters $\theta_i' \leftarrow \beta \odot \theta - \alpha \odot \nabla_\theta L_{D_i} f(\theta)$
12: Sample another set of datapoints $D_i'$ from $\tau_i$
13: end for
14: Update parameters $\theta \leftarrow \theta - \eta \nabla_\theta \sum_{D_i', \theta_i' \sim S_T} L_{D_i'} f(\theta_i')$
15: Update parameters $\phi \leftarrow \phi - \eta \nabla_\phi \sum_{D_i', \theta_i' \sim S_T} L_{D_i'} f(\theta_i')$
16: end while
17: Perform $k$-fold evaluation on target tasks
Chapter 7

Dataset and Experimental Configuration

7.1 Dataset Details

The empirical study in this work uses three real-world brain network datasets: 1) the Bipolar Disorder (BP) dataset, 2) the Human Immunodeficiency Virus Infection (HIV) dataset, and 3) the Parkinson’s Progress Markers Initiative (PPMI) dataset. The BP and HIV are private datasets, while the large-scale PPMI dataset\(^1\) is publicly available for authorized users. The study has been approved by an Institutional Review Board (IRB) to ensure the ethical and responsible use of human subjects in research. The IRB reviewed and approved the study protocols and consent forms, ensuring that the rights and welfare of the participants are protected. The study strictly adheres to the Good Clinical Practice guidelines and U.S. 21 CFR Part 50 (Protection of Human Subjects) to ensure the safety and privacy of the participants. All the data used in this work is processed anonymously to protect the privacy of participants, and no personally identifiable information is used or disclosed.

\(^1\)https://www.ppmi-info.org/
7.1.1 Parkinson’s Progression Markers Initiative (PPMI)

This is a restrictively public available dataset\(^2\) to speed breakthroughs and support validation on Parkinson’s Progression research. This dataset contains 718 subjects, where 569 subjects are Parkinson’s Disease (PD) patients and the rest 149 are Healthy Control (HC). The raw imaging signals are pre-processed by Eddy-current and head motion correction using FSL\(^3\) and the brain networks are extracted using the same tool. The EPI-induced susceptibility artifacts correction is handled using Advanced Normalization Tools (ANT)\(^4\). In the meantime, 84 ROIs are parcellated from T1-weighted structural MRI using Freesurfer\(^5\). The brain networks are constructed using three whole brain tractography algorithms namely the Probabilistic Index of Connectivity (PICo), Hough voting (Hough), and FSL. Each resulted network for each subject is \(84 \times 84\). Each brain network is normalized by the maximum value to avoid computation bias for the later feature extraction and evaluation, since matrices derived from different tractography algorithms differ in scales and ranges. The final brain networks were parcellated according to the Desikan-Killiany 84 template.

7.1.2 Bipolar Disorders (BP)

This local dataset is composed of the resting-state fMRI and DTI image data of 52 Bipolar I subjects who are in euthymia and 45 Healthy Controls (HCs) with matched age and gender [8, 58]. The fMRI data was acquired on a 3T Siemens Trio scanner using a T2* echo planar imaging (EPI) gradient-echo pulse sequence with integrated parallel acquisition technique (IPAT) and DTI data were acquired on a Siemens 3T Trio scanner. The brain networks are constructed using the CONN\(^6\) toolbox and are parcellated using the Brodmann 82 template. A normalization and smoothing after

\(^2\)https://www.ppmi-info.org/
\(^3\)https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/
\(^4\)http://stnava.github.io/ANTS/
\(^5\)https://surfer.nmr.mgh.harvard.edu/
\(^6\)http://www.nitrc.org/projects/conn/
first realigning and co-registering were performed on the raw EPI pictures. After that, the signal was regressed to remove the confounding effects of the motion artifact, white matter, and CSF. The 82 defined ROIs, also identified as cortical and subcortical gray matter regions were produced by Freesurfer, and pairwise signal correlations were used to build the brain networks.

7.1.3 Human Immunodeficiency Virus Infection (HIV)

This local dataset involves fMRI and DTI brain networks for 70 subjects, with 35 of them early HIV patients and the other 35 Healthy Controls (HCs). These two groups of subjects do not differ in demographic distributions such as age and biological sex. The preprocessings for fMRI including brain extraction, slice timing correction and realignment are managed with the DPARSF\(^7\) toolbox, while the preprocessings for DTI such as distortion correction are finished with the help of FSL\(^3\) toolbox. Finally, brain networks with 90 regions of interest are parcellated based on the automated anatomical labeling (AAL 90) atlas template [78].

7.2 Experimental Setup

7.2.1 Backbone Selection and Evaluation Metric

The proposed framework employs GCN as the backbone for the GNN [47] encoder. The experiment also benchmarks using GAT [81] and GIN [87], and the results are provided in Appendix B.1. The hyperparameter tuning follows the standard designs in related studies such as in [89, 84, 37]. The downstream evaluation is binary graph classification for disease prediction. To assess the performance, this experiment uses the two widely used metrics in the medical field [51, 14]: accuracy score (ACC) and the area under the receiver operating characteristic curve (AUC).

\(^7\)http://rfmri.org/DPARSF/
7.2.2 GNN Setup

The GCN encoder is composed of 4 graph convolution layers with hidden dimensions of 32, 16, 16, and 8. Similarly, the GAT encoder is built from 4 graph attention layers with hidden dimensions of 32, 16, 16, and 8. Regarding GIN, which is slightly different, the encoder consists of 4 MLP layers with each MLP containing 2 linear layers with a unifying hidden dimension of 8.

7.2.3 Pre-training Pipeline Setup

For two-level node contrastive sampling, we set $k = 2$ as the radius regarding $k$-hop neighborhood sampling for $S_1$ and $S_4$. To enable efficient computation on multi-graph MI evaluation, we resort to mini-batching and we set a default batch size of 32. In addition, we leverage the popular Adam [45] optimizer with the learning rate set to 0.002 as well as the cosine annealing scheduler [55] to facilitate GNN training. In general, a complete pre-training cycle takes 400 epochs with an active deployment of early stopping.

7.2.4 Atlas Mapping Regularizer Setup

Following the discussion in section 6.1.2, the total running loss of the AE projection is given as:

$$\mathcal{L} = \mathcal{L}_{\text{rec}} + \alpha \mathcal{L}_{\text{loc}} + \beta \mathcal{L}_{\text{com}} + \gamma \mathcal{L}_{\text{KL}},$$

(7.1)

in particular, we set $\alpha, \beta = 0.8$ and $\gamma = 0.01$. The one-layer AE encoder transforms the feature signals from all given datasets into a universally projected dimension of 32. For the details of locality-preserving regularizer (i.e., $\mathcal{L}_{\text{loc}}$), the transition matrix $\mathbf{T}$ is built from the 5-nearest-neighbor graph from the 3D coordinates of each atlas templates. For the sparsity-oriented regularizer (i.e., $\mathcal{L}_{\text{KL}}$), the target sparsity value $\rho$ is set to $1e^{-5}$. The overall optimization process, which is similar to model pre-training,
takes a total of 100 epochs with a learning rate of 0.02.

7.2.5 Downstream Evaluation Setup

For each target evaluation, the fine-tuning process features a 5-fold cross-validation, which approximately splits the dataset into 70% training, 10% validation, and 20% testing. To prevent model over-fitting, we implement a $L_2$ penalty with a coefficient of $1e^{-4}$. Overall, the model fine-tuning process, which is nearly identical to the other two training procedures, takes a total of 200 epochs with a learning rate of 0.001 and a cosine annealing scheduler.
Chapter 8

Experiments and Analysis

The effectiveness of the proposed framework is evaluated through extensive experiments on real brain network datasets, with a focus on the following research questions:

- **RQ1**: How does the proposed framework compare with other unsupervised GNN pre-training frameworks adapted to the scenario of brain networks?

- **RQ2**: What is the contribution of each major component in the proposed framework to the overall performance?

- **RQ3**: How does the choice of sampling method affect model convergence during pre-training and performance in downstream adapting?

- **RQ4**: How effective is the variance-based sorting in aligning virtual ROIs among different parcellation systems?

### 8.1 Overall Performance Comparison (RQ1)

A comprehensive comparison of the target performance between the proposed framework and popular unsupervised learning strategies is presented in Table 8.1. To fairly compare the methods, the atlas mapping pre-processing, the multi-task meta-learning
Table 8.1: Disease prediction performance comparison. All results are averaged from 5-fold cross-validation along with standard deviations. The best result is highlighted in bold and runner-up (excluding the w/o AR variant) is underlined.

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<th>Type</th>
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<th>BP-IMRI AUC</th>
<th>BP-DTI ACC</th>
<th>BP-DTI AUC</th>
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<td>60.44 ± 0.44</td>
<td>72.46 ± 2.74</td>
<td>72.94 ± 2.64</td>
<td>61.79 ± 1.84</td>
<td>61.57 ± 2.45</td>
</tr>
<tr>
<td>EGS</td>
<td>GCC</td>
<td>63.45 ± 1.24</td>
<td>62.39 ± 0.30</td>
<td>60.44 ± 0.04</td>
<td>60.29 ± 1.03</td>
<td>70.97 ± 0.31</td>
<td>72.48 ± 1.16</td>
<td>61.27 ± 0.66</td>
<td>61.38 ± 0.72</td>
</tr>
<tr>
<td></td>
<td>EGI</td>
<td>63.35 ± 0.83</td>
<td>63.58 ± 0.82</td>
<td>61.82 ± 1.13</td>
<td>61.57 ± 0.27</td>
<td>73.46 ± 2.42</td>
<td>73.28 ± 3.50</td>
<td>60.89 ± 0.37</td>
<td>62.41 ± 0.50</td>
</tr>
<tr>
<td>Ours</td>
<td>w/o AR</td>
<td>64.15 ± 0.63</td>
<td>64.24 ± 0.31</td>
<td>62.41 ± 0.52</td>
<td>65.84 ± 0.74</td>
<td>74.93 ± 0.14</td>
<td>75.74 ± 0.50</td>
<td>64.39 ± 0.41</td>
<td>64.38 ± 0.55</td>
</tr>
<tr>
<td></td>
<td>Full</td>
<td>68.84 ± 2.26</td>
<td>68.45 ± 2.96</td>
<td>66.57 ± 2.87</td>
<td>68.31 ± 2.39</td>
<td>77.80 ± 2.76</td>
<td>77.22 ± 2.74</td>
<td>67.51 ± 2.67</td>
<td>67.74 ± 2.59</td>
</tr>
</tbody>
</table>

learning backbone, and the task adaptive reweighing algorithm discussed in section 4.4 are applied to all benchmarked methods, with only few exceptions including STT, MTT, and Ours w/o AR (i.e., without task adaptive reweighing). The purpose of this comparison is to effectively highlight the impact of the proposed two-level contrastive pre-training and there will be further analysis on the effect of atlas mapping in subsequent sections. In addition, for a clearer presentation, this experiment groups the selected baselines according to their optimization strategies:

- No pre-training (NPT): the backbone with randomly initialized parameters for target evaluation.
- Transfer learning (TFL): methods that are formed based on transfer learning paradigm discussed in Sections 4.2 4.3 on STT and MTT. Specifically, for STT, the pre-training dataset is defined as the PICo modality of PPMI.
- Non-CL-based (NCL): methods with cost functions regularized by co-occurrence agreement or link reconstruction, including Node2Vec [25], DeepWalk [65], and VGAE [46].
• Single-scale CL (SCL): methods utilizing either node- or graph-level representations in the CL optimization, including GBT [5], ProGCL [85], and GraphCL [93].

• Multi-scale CL (MCL): methods whose CL optimization utilizes both nodes- and graph-level representations, including DGI [82] and InfoG [75].

• Ego-graph sampling (EGS): methods whose contrastive samplings consider $k$-hop ego-networks as discriminative instances, which are the most similar to the proposed framework, including GCC [67] and EGI [99].

The experiments reveal the following insights:

• The proposed framework of ours consistently outperforms all the baselines, achieving a relative improvement of 7.34%-13.30% over the best-performing baselines and 31.80%-38.26% over the NPT setting. The reported results of the selected baselines are also statistically compared against that of the proposed framework under the paired $t$-test. With the significance level set to 0.05, the largest two-tailed $p$ value is reported at 0.042, which means that the proposed framework demonstrates statistically significant performance increase over other selected methods.

• The transfer learning based pipeline, including STT and MTT improve over the NPT baseline with a relative gain 8.27% and 16.67% respectively across both metrics, suggesting the relative benefit of model pre-training and knowledge transfer. However, the transfer learning setting still suffers high variance in performance results and inferior overall performance compared to the proposed full framework.

• Compared with the transductive methods of Node2Vec and DeepWalk, the GNN pre-trained by VGAE learns structure-preserving representations and achieves
the best results in the NCL-type methods. This indicates the potential benefit of the locality-preserving regularizer design in the proposed framework.

- Maximizing mutual information between augmented instances may hinder GNNs from learning a shared understanding of the entire dataset. For baselines belonging to the categories of SCL, MCL, and EGS, pre-training with non-augmented CL (InfoG, EGI) generally results in a 4.36% relative improvement across both metrics and a 7.63% relative decrease in performance variance compared to their augmentation-based counterparts (GBT, GraphCL, ProGCL, DGI, GCC). This explains why the proposed framework does not employ data augmentation.

- Multi-scale MI promotes the capture of effective local (i.e., node-level) representations that can summarize the global (i.e., graph-level) information of the entire network. The MCL-type methods typically outperform the SCL-type ones by a relative gain of 2.68% in ACC and 3.27% in AUC.

- The group of baselines considering $k$-hop neighborhoods (EGS) presents the strongest performance, indicating the importance of local neighborhoods in brain network analysis. The proposed the proposed framework, which captures this aspect through both node- and graph-level CL, is the only one that comprehensively captures the local neighborhoods of nodes.

- The added component of task adaptive reweighing demonstrates a promising working effectiveness by bringing over the non reweighed training with a relative improvement of 4.29% in accuracy score and 5.80% in AUC metric. This shows that the issue of task-biased convergence during multi-task pre-training exists and can be mitigated by additional handling through reweighing mechanisms.
8.2 Ablation Studies (RQ2)

This ablation studies examine two key components of the proposed framework - (1) the two-level contrastive sampling and (2) the atlas mapping regularizers. The best contrastive sampling configuration is fixed when examining the atlas regularizers, and all regularizers are equipped when examining the contrastive samplings. The results, shown in Figure 8.1 (with additional DTI version in Appendix B.2), are analyzed based on the four possible variants of contrastive sampling listed in Table 8.2. The analyses yield the following observations:

- leveraging $k$-hop neighborhood (i.e., positive $S_2$) MI maximization brings visible performance gain, confirming its benefit in brain structure learning.

- The extension to multi-graph CL (i.e., consideration of $S_3$) facilitates the extraction of unique ROI knowledge, leading to improved results in Var. 3/4.

- Var. 4 outperforms Var. 3 as it effectively summarizes of global (i.e., graph-level) information in local node representations.

- The full implementation of the proposed framework brings a relative gain of 4.27% in both metrics on top of Var. 4, highlighting the significance of considering shared substructure knowledge across multiple graphs (i.e., through the inclusion of $S_4$).
Figure 8.1: Ablation comparisons on contrastive sampling choices (top two) and atlas mapping regularizers (bottom two). The $y$-axis refers to the numeric values of evaluated metrics (in %).
The bottom two sub-figures examine the impact of the atlas mapping regularizers by comparing the results of the full framework to those without the sparsity regularizer (w/o SR), the locality regularizer (w/o LR), and the community regularizer (w/o CR). Two key observations are made:

- The removal of SR leads to the greatest performance drop, emphasizing its crucial role in learning robust projections that can effectively handle noise and prevent over-fitting.
- The inferior results when LR and CR are absent emphasize the importance of spatial sensitivity and blockwise feature information in brain network analysis. This supports our intuition to consider the relative positioning of ROIs in the 3D coordinate as well as knowledge on community belongings based on modularity measures.

### 8.3 Analysis of Two-level Contrastive Sampling (RQ3)

Figure 8.2 offers insight into the pre-training convergence, target adaptation progression, and pre-training runtime consumption of the four sampling variants and the full framework. Key observations include:

- As seen in Figure 8.2(a), all variants demonstrate efficient pre-training convergence due to the multi-dataset joint optimization inspired by MAML. The full model demonstrates the most optimal convergence, highlighting the advantage of learning shared neighborhood information in brain network data through two-level node contrastive sampling.
- Figure 8.2(b) shows the superiority of our design in terms of downstream adaptation performance compared to other variants.
Figure 8.2: In-depth comparison among the four variants and the full model. The $x$-axis is epochs.
- Figure 8.2(c) reveals that the more sophisticated the sampling considerations result in greater computational complexity for mutual information evaluation, leading to longer runtime for each pre-training epoch. However, the total time consumptions are all on the same scale.

8.4 Analysis of ROI Alignment (RQ4)

To further validate the variance-based virtual ROI sorting, this experiment selects the top 2 virtual ROIs with the highest sample variances for each atlas template (i.e., dataset) and backtrack to locate their corresponding projected ROIs. The results are illustrated in Figure 8.3, which shows a 3D brain surface visualization highlighting the original ROIs. From this, one can draw two main conclusions:
Figure 8.3: The virtual ROI mapping across the three investigated datasets. Overlapping regions are highlighted with colored boxes. In particular, the annotation use gold-colored boxes for the PPMI and BP atlases; blue-colored boxes for the BP and HIV atlases; and purple-colored boxes for the PPMI and HIV atlases.

- There exists multiple regional overlaps between pairs of two atlas templates, reflecting some working effectiveness of our proposed solution as well as confirming the feasibility of converting between atlas templates.

- It is relatively harder to find regions that overlap across all three atlas templates which shows a limitation of the proposed unsupervised ROI alignment scheme, suggesting a need to modify against the current heuristic that considers beyond mere variance measures which may inspire further study and research opportunity.
Chapter 9

Conclusions and Future Directions

This work focuses on data efficient learning on small-sized brain network datasets through leveraging meta learning techniques, self-supervised contrastive pre-training objectives, and brain network oriented design considerations. The experiments have demonstrated the effectiveness of the proposed framework in the application of brain network based disease predictions. This work is also the first to discover the inherent challenges in learning on small-sized brain network datasets and formulate this problem into a data-efficient learning objective, where the goal to find a generalizable model initialization that achieves efficient adaption on target tasks. To this end, the proposed framework leverages transfer learning and, more importantly, meta-learning strategies to serve as backbone frameworks for model pre-training. In addition, the framework also features a novel two-level contrastive sampling strategies to enable unsupervised model pre-training. Considering the special properties of brain networks from traditional graphical data, the framework proposes an automated atlas transformation design and variance-based sorting to help address the incompatibility challenge of cross-dataset brain network ROI template dimensions. Besides, the proposed framework also introduces an adaptive task reweighing algorithm that helps resolve biased learning issues in the conventional meta-training pipeline. Extensive ex-
perimentation demonstrated the effectiveness of proposed methodologies. It is worth noting that the proposed framework is naturally generic and can be easily scaled to other types of neuroimaging datasets. The training pipeline can also be generalized to any parameterized model that is optimized on any customizable objectives and data sampling strategies.

However, Learning on brain network data is still prescribed to various challenges. First, most brain networks are expressed by multiple views and modalities, in which to achieve a comprehensive feature extraction, would require GNN models to capture complex inter-relations within graph modalities. Simply applying multi-facet meta-learning and separately optimizing on individual views fail to consider the intricacies of some shared and complementary knowledge underneath the multi-view datasets. Second, the target performance on supervised disease classification still suffers from relatively high data variance under the $k$-fold evaluation scheme. This suggests that, assuming given a balanced dataset, the current GNN models are sensitive to batch effects, which would require additional handling of data noise and further development of GNN models that achieve good out-of-distribution performance. For future investigation, my research will primarily focus on addressing the aforementioned challenges by performing theoretical and empirical analyses on GNN architectures for brain network learning. To tackle the data scarcity issue, exploration of data augmentation and synthetic generation techniques [26, 101] are also advised to expand available training samples with artificially constructed, domain- and distribution-aware data instances. Since the raw neuroimaging signals are represented in time-series, it is also worth investigating methods of learning over time-series data and dynamic structures which can further reduce the cost of data collection by removing the need to pre-process these signals into brain networks which are known to be time consuming.
Appendix A

Autoencoder Structure Analysis

A.1 Bridging Reconstruction Minimization and Variance Maximization

In this section, we briefly discuss how the reconstruction minimizing objective in one-layer AE can be cast to a variance-maximizing objective in PCA. Assume given a data matrix $X \in \mathbb{R}^{n \times d}$, its covariance matrix $\Sigma = X^T X \in \mathbb{R}^{n \times n}$, and a single-layer AE projection matrix $W \in \mathbb{R}^{d \times m}$ with parameters randomly initialized from the continuous uniform distribution $\mathcal{U}(0,1)$, the reconstruction objective is:

\[
\frac{1}{n} \|X - XWW^\top\|^2 = \frac{1}{n} \text{tr}((X - XWW^\top) \\
\cdot (X - XWW^\top)^\top) \\
= \frac{1}{n} \text{tr}((X - XWW^\top) \\
\cdot (X^\top - WW^\top X^\top)) \\
= \frac{1}{n} [\text{tr}(XX^\top) - \text{tr}(XWW^\top X^\top) \\
- \text{tr}(XWW^\top X^\top) \\
+ \text{tr}(XWW^\top WW^\top X^\top)]
\]
\[
\frac{1}{n}[c_1 - 2 \cdot \text{tr}(XW^T X^T)] + \text{tr}(\hat{X}\hat{X}^T)] \\
= \frac{1}{n}[c_1 - 2 \cdot \text{tr}(XW^T X^T) + c_2] \\
= c_3 - c_4 \cdot \text{tr}(W^T X^T XW) \\
= c_3 - c_4 \cdot \text{tr}(W^T \Sigma W)
\]

Notice that \(c_1, c_2, c_3, c_4\) are non-negative scalar constants that do not influence the overall optimization trajectory. Hence, alternatively, the optimal AE projection also maximizes the sample variance \(\text{tr}(W^T \Sigma W)\), achieving an identical end goal of PCA transform. Specifically, according to PCA, variance maximization is realized by constructing the projection \(W\) to contain the set of orthonormal eigenvectors of \(\Sigma\) that gives the largest eigenvalues [35]. That is, there is an orthogonality constraint on \(W\). Minimizing the MSE reconstruction also results in an orthogonal \(W\):

\[
\frac{1}{M} \|X - XW^T\|^2 = 0 \Rightarrow WW^T = I
\]

Therefore, the optimal AE projection \(W\) is also capturing a set of variance-maximizing orthogonal vectors. Note that the AE optimized \(W\) is theoretically equivalent to the eigendecomposition of \(\Sigma\) if and only if the reconstruction loss is 0. Therefore, in practice, the AE is, at best, an approximate solution to variance maximization.

### A.2 Variance-based Sorting Procedure

Following the discussion in A.1, assuming a perfect optimization, the linear one-layer AE behaves similarly to PCA, and there is an equivalence relation between their respective objective functions. Notice that in PCA, the eigenvalue of the covariance matrix \(\Sigma\) signifies the intensity of data variation along the direction of its
corresponding eigenvector, which is essentially a column entry of the transformation matrix. Then intuitively, given an optimized AE projection $W$, we can examine, for each column of $W$, its representativeness (i.e., data variance) of the data covariance with a scalar estimate (i.e., an eigenvalue-like scoring). Inspired by the properties of eigendecomposition, we can approximate these estimates by measuring the distance of $W$ w.r.t to the product of linearly transforming $W$ through $\Sigma$ by a scaling factor of $\lambda$. More specifically, we want to solve for $\lambda$ such that $\Sigma w = \lambda w$ for every column vector $w \in W$. Under the PCA perspective, $\lambda$ contains the variance estimate for each column-wise individual projection of $W$. To this end, we detail the sorting procedure in Algorithm 4.

**Algorithm 4** Overview procedure for variance-based sorting

**Input:** Original feature matrix $X \in \mathbb{R}^{M \times M}$; AE optimized projection matrix $W \in \mathbb{R}^{M \times D}$

**Initialize:** Scalar vector $\lambda \in \mathbb{R}^D$; Small positive float $\epsilon$

**Output:** Sorted AE projection matrix $\tilde{W}$

1: Normalize the feature matrix: $X_n \leftarrow X / \|X\|
2: Compute data covariance matrix: $\Sigma \leftarrow X_n^\top X_n$
3: Solve for $\lambda$ such that $|\Sigma W - W \odot \text{diag}(\lambda)| \leq \epsilon$
4: Sort column vectors $w \in W$ according to (sorted) decreasing order of $\lambda$ to obtain $\tilde{W}$
Appendix B

Additional Experiments

B.1 Performance with GAT and GIN

Table B.1: Disease prediction performance of our framework using GAT and GIN. The best performer is highlighted in bold.

<table>
<thead>
<tr>
<th>Method</th>
<th>BP-IMRI</th>
<th>BP-DTI</th>
<th>HIV-IMRI</th>
<th>HIV-DTI</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ACC</td>
<td>AUC</td>
<td>ACC</td>
<td>AUC</td>
</tr>
<tr>
<td>Ours w/ GCN</td>
<td>68.84</td>
<td>±8.26</td>
<td>68.45</td>
<td>±8.96</td>
</tr>
<tr>
<td>Ours w/ GAT</td>
<td>66.96</td>
<td>±9.71</td>
<td>69.68</td>
<td>±9.61</td>
</tr>
<tr>
<td>Ours w/ GIN</td>
<td>66.30</td>
<td>±8.77</td>
<td>68.92</td>
<td>±9.41</td>
</tr>
</tbody>
</table>

Table B.1 reports the downstream performance of the proposed full framework using GAT and GIN as backbone encoders. In general, the two encoders deliver inferior performance compared to GCN, which suggests that complex GNN convolutions (e.g., GAT and GIN) might not be as effective as they seem when learning on brain network datasets.

B.2 Additional Ablation Studies on DTI

Figure B.1 presents the ablation studies on the DTI view following the same setup as discussed in Section 8.2. One can draw similar conclusions from the DTI-based
Figure B.1: Additional ablation comparisons on DTI views. The top two subfigures refer to contrastive sampling considerations and the bottom two subfigures refer to atlas mapping regularizers. The $y$-axis refers to the numeric values of evaluated metrics (in %). This Appendix benchmarks results on the DTI modality of the BP and HIV dataset.

analysis where each constituent component of the two-level sampling consideration as well as the atlas mapping mechanism has proven positive contribution and significance towards the overall performance and robustness.
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