

DIPLOMA THESIS

LINKING CONNECTIVITIES AND GENE EXPRESSION PATTERNS IN MICE BRAINS

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Tilman Hinnerichs
Matrikelnummer: 4643427
Technische Universität Dresden

Tutor: Dr. Nico Scherf
MPI for CBS

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proper title?

Abstract

Contents

1	Introduction	5
2	Literature overview	6
2.1	Gene expression databases and prediction	6
2.2	Finding spatial patterns in gene expression in mice brains	7
2.3	Structural and functional connectivity prediction	8
3	Materials and methods	10
3.1	Problem description	10
3.1.1	Spatial gene expression prediction	10
3.1.2	Dimensionality reduction in brains	10
3.1.3	Structural and functional connectivity prediction	10
3.2	Datasets	10
3.2.1	Spatial gene expression values in mice brain	11
3.2.2	Structural and functional connectivity databases	11
3.3	Model	12
3.3.1	Feature generation	12
3.3.2	Graph convolutional neural layers	12
3.3.3	Dimensionality reduction techniques	13
3.3.3.1	Principal component analysis (PCA)	13
3.3.3.2	tSNE	13
3.3.3.3	UMAP	13
3.3.3.4	Parametric UMAP	13
3.3.4	Hyperparameter tuning	13
3.4	Evaluation and metrics	13
4	Results	14
4.1	Gene expression prediction	14
4.2	Dimensionality reduction and its combination with different graphs structures	14
4.3	On the linkage of connectivities and gene expression patterns	14
5	Discussion	15
6	Conclusion	16

To be sorted somewhere

- Variability and different interpretations of different graph convolutional neural filters [Kipf and Welling, 2016, Li et al., 2020, Feng et al., 2022] etc.
- DeepGOPlus for feature generation [Kulmanov and Hoehndorf, 2019]
- discussion of different PPI network databases [Szkklarczyk et al., 2014]
- discussion of potential databases associating gene expression data with their spatial distribution [Hawrylycz et al., 2011]
- discussion of best neural learning/graph convolutional methods [Paszke et al., 2019, Fey and Lenssen, 2019]
- how to handle highly imbalanced data, metrics, preprocessing, sampling, modification of loss function [Jeni et al., 2013] and optimization over them (with Adam [Kingma and Ba, 2015])
- maybe introduction of PhenomeNET for MP/GO for more sophisticated protein representation [Hoehndorf et al., 2011, Ashburner et al., 2000, Carbon et al., 2020, Smith and Eppig, 2009] and derive features from DL2vec [Chen et al., 2020, Mikolov et al., 2013]
- evaluation of „Using ontology embeddings for structural inductive bias in gene expression data analysis“ [Trebacz et al., 2020]
- take some ideas from Zitnik and Leskovec [2017] with title „Predicting multicellular function through multi-layer tissue networks“. (OhmNet)
- potentially group results based on InterPro [Blum et al., 2020] families eventually
- choice of model organism?!

1 Introduction

General thread for introduction and motivation:

- Gene expression patterns are difficult to analyze in humans → take mouse as model organisms
- The brain is a multi-level system in which the high-level functions are generated by low-level genetic mechanisms. Thus, elucidating the relationship among multiple brain levels via correlative and predictive analytics is an important area in brain research. Currently, studies in multiple species have indicated that the spatiotemporal gene expression patterns are predictive of brain wiring. Specifically, results on the worm *Caenorhabditis elegans* have shown that the prediction of neuronal connectivity using gene expression signatures yielded statistically significant results.
- no in-depth analysis of mouse brain genetic patterns and their relation to different connectivity patterns has been made yet
- we analyze
- studies have shown circadian patterns of gene expression in human brain and the disruption of those in depressive disorder [Li et al., 2013]
- [Twine et al., 2011] show the importance of gene expression patterns, by linking gene expression aberration with increase in Alzheimer's disease
- Guilt by association over gene networks [Oliver, 2000, Gillis and Pavlidis, 2012] in genetic networks
- protein function prediction from PPI networks [Vazquez et al., 2003]
- GCNs have been applied successfully to variety of tasks over different types of graphs.

General Introduction of the Research Study

Research problem or Questions with Sub-Questions

Reasons or Needs for the Research Study/Motivation for my research

Definition and explanation of Key Terminology

Context of Research Study within the Greater Discipline

- Introduction to mouse brains as model organisms for insights into human brain
- Works on mouse brain in general and potential tasks
- works on gene expression in mouse brains
 - traditional approaches
 - importance of gene expression patterns in mouse brains
- neural networks for this purpose
 - how were
- gene expression for general tissue

2 Literature overview

2.1 Gene expression databases and prediction

Research in gene expression prediction and profiling has a long history in bioinformatics and systems biology, but was almost exclusively linked to cancer research. Moreover, with the rise of machine learning, and more specifically (deep) neural networks and its variants, this field became increasingly data reliant. The Human Genome Project [Watson, 1990], launched in 1990 and declared finished in 2003 while the first gapless assembly was finished in 2022, also sparked various works in relating these genetic representations to other tissue- and individual-specific properties and traits.

For comparison of gene expression profiling works there exist multiple prominent variables. Most significantly, the chosen organism is a crucial choice for both data availability and predictive complexity. Second, the chosen tissue is naturally important for the proposed hypotheses, especially with respect to tissue definitive cancer research, and its potential ability to generalize without transfer learning. While gene expression pattern analysis approaches frequently focus on tissues like *mamma*, (primarily female) breast, [Herschkowitz et al., 2007], liver [Flores-Morales et al., 2002], and skeletal muscle [Lecker et al., 2004] for exploration of diseases like cancer and atrophy, respectively, in humans.

However, the nervous system is often investigated separately as it bears different molecular processes and structure, anatomy and cell life cycles, while brain and spinal cord are even based in a separate nutritional circuit for mammals. Moreover, gene expression determination in the human brain may almost certainly remain an deadly intervention for most brain tissues, hence allowing only for careful extraction of specific tissues in living organisms. Also this disallows for *in-vivo* extraction of vital brain regions and structures, e.g. the brainstem. Furthermore, the human brain’s gene expression patterns are varied and diversified [Ramasamy et al., 2014], aligning with its anatomical and embryogenesis complexity, and its compartments are exceptionally and deeply connective and collaborative [Fornito et al., 2015]. Both also hold for invertebrates, i.e. insects. Thus, full genetic profiles of expression are mandatory for a full understanding of the mammalian and invertebrate brain and primary nervous system, respectively. By the strong intervention of the tissue extraction, full genome atlases are fit together from various experiments on multiple individuals.

The human brain is among the most intricate and complicated networks we do know of, and is far from being fully understood. Additionally, full transcriptomic atlases of human brains are difficult to collect while raising decisive privacy concerns. Yet, there were multiple efforts and projects with rather small sample sizes. A detailed elaboration on dataset and organism choice, and their respective properties may be found in Section 3.2.

However, there have been works on numerous works for other tissues and other organisms. modENCODE Consortium et al. [2010] correlate activity patterns in the regulatory network within *Drosophila*, proposing their model for identification of functional elements "modENCODE". As this work was published back in 2010, the approach relies purely on statistical correlation and covariance. Chikina et al. [2009] follow a similar approach in *C. elegans* predicting tissue-specific gene expression in 2006 utilizing support-vector machines (SVM)[Noble, 2006].

More modern, data-oriented machine learning models such as (deep) neural networks (NN) were applied successfully to similar problems. Aromolaran et al. [2020] achieved to predict essential genes based on their respective sequence and functional features profiting off NNs, while transcriptomic interaction prediction was done based on functional gene data using deep learning in Yang et al. [2019] in *Drosophila* over different tissues.

Within humans, as mentioned previously, gene expression was primarily used for cancer and disease research. Schulte-Sasse et al. [2021] and Wang et al. [2021] were the first to apply graph convolutional neural networks to the task of gene expression prediction within humans. While Schulte-Sasse et al. [2021] was applied on data from The Cancer Genome Atlas (TCGA)[Tomczak et al., 2015] across multiple tissues, Wang et al. [2021]’s MOGONET is proposed as a general framework

for gene expression prediction with example computations on ROSMAP dataset and TCGA. Both approaches implement the original formulation of GCNs[Kipf and Welling, 2016], which we will discuss in more detail in Section 3.3.2, over protein-protein interaction networks and accomplish outstanding performances and both measure biomarker importance for prediction in order to leverage explainability. The authors thereby exploit the "guilt by association" principle [Oliver, 2000, Gillis and Pavlidis, 2012] over gene networks, adding background knowledge such as biological interaction and pathways.

Crucial for almost all classification tasks in machine learning is the choice of entity representation. In the mentioned works molecular [Schulte-Sasse et al., 2021, modENCODE Consortium et al., 2010, Noble, 2006] and phenotypical [Wang et al., 2021, Chikina et al., 2009] features were used for expression prediction, but never both combined. The combination of phenotypical and molecular features over GCNs was proven to raise predictive performance in drug-target interaction prediction [Hinnerichs and Hoehndorf, 2021] but remains an open challenge for this very task.

2.2 Finding spatial patterns in gene expression in mice brains

In this subsection we will constrain the issue of gene expression analysis to both "spatial patterns", mammals and the tissues of the brain, which we study in this work. The term *spatial patterns* is rather vague and allows for various interpretations, both discrete and continuous, which will form the classes for the following literature review.

Zapala et al. [2005] is among the earliest works, showing that local structures beared "transcriptional imprint" that coincide with the embryological origin of the examined regions. However, they only were able to identify up to 24 neural tissues. They further conclude that this may be important for functional collaboration within the adult mouse brain. The authors measure pairwise correlation show the existence of clusters over a heatmap.

The Allen Institute Brain Atlas (AIBA), is a collection various atlases such as Allen Mouse Brain Atlas (AMBA)[Lein et al., 2006, Daigle et al., 2018], the Allen Mouse Brain Connectivity Atlas (AMBCA) [Oh et al., 2014, Harris et al., 2019] and the Allen Mouse Brain Common Coordinate Framework (CCFv3) [Wang et al., 2020] to name only the ones related to adult mice's brains. As it was the first coherent collection of spatially resolved expression values, mapping 2D expression images consistently to 3D coordinates, the AMBA has sparked a range of publications. Within Lein et al. [2006], the Allen Institute also published the "Allen Reference Atlas"(ARA) proposing a number morphological and histologically induced sub-regions of the brain and hence a precisely defined parcelation. Moreover, they propose the ARA *ontology*, a semnatic hierarchy, providing a hierarchical cluster of all sub-structures and map them back to their coordinates with the CCF.

Takata et al. [2021] propose a flexible annotation atlas of the mouse brain, introducing a flexible ontology construction framework which may be used on the transcriptomic data such as the AMBA, leveraging anatomic structure and axonal projection data. Here, FAA focuses on consistent and reproducible regions-of-interest (ROIs) definition for other downstream tasks such as resting-state functional connectivity annotation. Further, this ontology may be seen as a pattern within mouse brain, while it may only detect connected structures.

The authors of Valk et al. [2020] analyze structural covariance of cortical thickness within primate brains, namely macaques, and its correlation to each cortical layers transcriptome. Further, transcriptomic variation was related to a continuum of functions by mapping them the brain anatomy, inducing a *continuous*, functional parcelation of the primates brain. Further, this study suggests a relation of functional and transcriptomic links.

While also focused on mouse brains, Partel et al. [2020] submits a novel database based on their own *in-situ* sequencing data, and a consecutive spatial gene expression analysis pipeline, and relates the results to tissue morphology and hence indirectly to the AMBA. Similarly to our proposed approach, brain parcelation are present as n -dimensional, continuous embeddings, representing closeness in gene expression space. Due to the similarity in the pipeline especially in their visualization utilizing UMAP,

Name a few
sparked
research
works on
this

we will use the generated images of this work for a brief comparison in Section 4.

- Sparked by the Lein et al. [2006]
-
- Have GCNs be applied before here?
- usage of ontologies?
 - functional graph
 - structural ontology?
 - developmental ontology

2.3 Structural and functional connectivity prediction

- [Lee and Lee, 2020] train classifiers using blood gene expression data in order to predict Alzheimer’s disease in Humans
- [Fornito et al., 2015] elaborate on the connectomics of brain disorders and its complexity in connectivity. Understanding how brain networks respond to pathological perturbations is crucial for understanding brain disorders and behavior

Structural/Axonal connectivity

- [Fakhry and Ji, 2015] predict axonal connectivity from gene expression patterns in mice brain with an accuracy of 93%
- [Roberti et al., 2019] use transcriptomic information to anatomical connectivity patterns and gene expression of neurons using (shallow) neural networks. Yield a 85% accuracy in prediction of unconnected and connected regions.
- experimental setup from Allen Institute for axonal projection data
- paraphrase description of „Technical tour: Explore the Allen Mouse Brain Connectivity Atlas“

Functional connectivity

- [Whitfield et al., 2003] were one of the first to link transcriptomic data with behavior and hence functional patterns in individual honey bees back in 2003. The authors show that changes in the messenger RNA were connected to behavior.
- [Rankin, 2002] first developed the idea of combining behavioral analyses of *Caenorhabditis elegans* with their genetics. Further, [Sun and Hobert, 2021] only recently described the distinct functional states and the corresponding distinct molecular states within the transcriptome. While honey bees and nematodes are rather simple model organisms, enabling both full transcriptomic analyses of the organisms, and their bearing and actions. However, "behavior" may be ambiguous and vague for such taxonomically distant animals, from the viewpoint of humans, and may only be linked to very basic meta-tasks such as basic routing, orientation and basic social interaction.
- Why do we not directly investigate gene expression patterns in human brains?
 - only few data points are given for the entirety of the human brain, while spatial decomposition and partitioning is crucially more complex.

- [Wang et al., 2022] recently proposed a novel-network based method integrating molecular-based gene association networks such as protein-protein interaction networks with brain connectome data. They further link these gene expression patterns to four brain diseases, including Alzheimer’s disease, Parkinson’s disease, major depressive disorder and autism.
- Where is data coming from? [Pallast et al., 2019]
- How to calculate functional connectivity matrix → AIDAconnect (no paper yet? cite dataset?)
- How to combine functional connectivity for multiple samples?
- [Zerbi et al., 2021]

Brief Overview of Literature Reviewed, Discussed and applied

Study Model and Process Aligning with literature reviewed

Hypotheses and justifications tied to prior sections and statements

The Scope of the study with theoretical assumptions and limitations

To be searched

- find other papers on
 - gene expression patterns within mouse brain and both possible hypothesis and tasks, and models over this
 - gene knockout models and whether they can learn propagation of those?
 - connection of FC and gene expression patterns and how to prove such interaction/correlation?
 - possible gene knockout targets within mouse brain and possible structural influences

Spatial patterns of gene expression

Data discussion, hypotheses and traditional approaches:

- [noa]
- Possible effects of rabies virus on gene expression [Prosniak et al., 2001] for potential knockout targets
- Review paper on regional variation in gene expression in mouse brain [Pavlidis and Noble, 2001]

Modern approaches on learning from gene expression patterns in mouse brain:

- Deep learning methods for capturing spatiality w.r.t. gene expression withing the brain [Zeng et al., 2015]
- R package for simulating gene expression from graph structures over general biological pathways [Kelly and Black, 2020]

[Read this](#)

3 Materials and methods

In this study, we utilized and incorporated various approaches from other works and applied them to diverse datasets. The following section will give a brief overview over all modules of the proposed model, while the combined method will be presented and described in the results section (Section 4).

Introduction and general description, study method and study design

3.1 Problem description

Here we give a brief introduction to each of the three tackled issues and further summarize data properties, challenges and goals of each problem.

3.1.1 Spatial gene expression prediction

Firstly, the issue of gene expression prediction

3.1.2 Dimensionality reduction in brains

-

3.1.3 Structural and functional connectivity prediction

-

Assumptions of study method and study design with implied

3.2 Datasets

As human brains are among the most complex in structure and connectivity within nature, a full transcriptomic atlas may be very valuable for the research community and our experiments in this work. However, full transcriptomic atlases of homo sapiens are ethically difficult to gather. Additionally, as a valuable, public genetic atlas of deceased relatives may provide highly critical information about the remaining, living ones, such as genetic diseases, genetic markers for correlating with addiction and other social behavior, or ancestry in general, this raises tremendous privacy concerns. As we want to investigate transcriptomic patterns in the brain and their relation to structural and functional connectivity as a generalized, organism-invariant methodology, we also want our experiments to be as understandable and replicable as possible. As rodents and more specifically mice are well studied in behavior and due to their taxonomic proximity to humans serve as model organisms for diverse genetic, social and medical experiments, we opted for mice as the study organism. The ultimate goal still shall be the further understanding of brains of our species.

However, there have been multiple initiatives towards collaborative and open human brain data, such as the

- Human connectome atlas
- Allen Human Brain atlas -> very small sample size

Furthermore, immense effort was put into enormous projects and databases for example invertebrates, namely *Drosophila* (specifically *Drosophila melanogaster*, also called *fruit fly*) and *Caenorhabditis elegans* (short: "*C. elegans*", colloquially also called *roundworm*) with the two projects "Virtual Fly Brain" [Milyaev et al., 2012] (<https://virtualflybrain.org/>) and "Wormbase" [Lee and Sternberg, 2003, Davis et al., 2022] (<https://wormbase.org>), respectively. Yet, we wanted to stay within the same taxonomic phylum leading our choice towards mice brains.

In literature there are several approaches, compromising sequencing depth, accuracy, throughput and spatial resolution. Generally, one can identify two classes of gene expression measurement while preserving spatial information. The first approach is to store spacial coordinates first, followed by a the sequencing of single-cell RNA where Achim et al. [2015] and Chen et al. [2017] propose the mapping and the Geo-seq protocol for this method. The second method includes the usage of "barcodes", decoded in the tissue sample, while running a parallel analysis of numerous mRNAs [Ke et al., 2013, Moffitt et al., 2016]. Samples are collected from multiple individuals.

3.2.1 Spatial gene expression values in mice brain

- include average brain and annotation brain for parcelation
- copy introduction from graph-iss
- add section on gene set enrichment analysis (GSEA)

Protein-protein interaction graph:

- CPDB78, STRING-db79, IRefIndex80, Multinet81 and PCNet82
- STRING[Szklarczyk et al., 2014] for mice

3.2.2 Structural and functional connectivity databases

in-depth description of the study design/datasets used and motivation why they were used for these experiments

Explanation of Sample used in the study

- show distribution (histo, mean, median, boxplot?) of expression densities see 'get_ge_structure_mat'
- how to normalize expression intensity (see discussion in DeepMOCCA paper, Sara Alghamdi), as there are regions with much more activity than others (e.g. bone narrow vs. bone boarder); thresholds for intensity varies across genes
 - over all intensities →
 - per structure →
 - per gene →
- why were these datasets used and not others?
- How did we achieve the matching?
- what are premises of the dataset?
- transfer learning working for other structure/regions
- dataset: Allen Mouse brain atlas vs.
 - phenoview impc data
 - mousephenotype
 - HPO/MP project expression data
- Mouse brain CCFv3
- Allen mouse brain atlas [Lein et al., 2006]

– discussion on different normalization schemes

- STRING for PPI network and how we chose suitable interactions [Szkarczyk et al., 2014]

Four graphs were used in this study:

- Protein-protein interaction graph from STRING
- structure hierarchy/ontology from [Lein et al., 2006]
- structural connectivity data from (Mouse Projection data)
- functional connectivity data from [Pallast et al., 2019]

3.3 Model

Explanation of Measurement, Definitions, Indexes, Reliability and Validity of study method and study design

Description of Analytical Techniques to be Applied and justification for them

Reliability and validity of internal/external design and related subtypes

3.3.1 Feature generation

Data preparation for regression task

- unbalanced data for prediction task

3.3.2 Graph convolutional neural layers

We include these molecular and ontology-based sub-models within a graph neural network (GNN) [Kipf and Welling, 2016]. The graph underlying the GNN is based on the protein–protein interaction (PPI) graph. The PPI dataset is represented by a graph $G = (V, E)$, where each protein is represented by a vertex $v \in V$, and each edge $e \in E \subseteq V \times V$ represents an interaction between two proteins. Additionally, we introduce a mapping $x : V \rightarrow \mathbb{R}^d$ projecting each vertex v to its node feature $x_v := x(v)$, where d denotes the dimensionality of the node features.

A graph convolutional layer [Kipf and Welling, 2016] consists of a learnable weight matrix followed by an aggregation step, formalized by

$$\mathbf{X}' = \hat{\mathbf{D}}^{-1/2} \hat{\mathbf{A}} \hat{\mathbf{D}}^{-1/2} \mathbf{X} \Theta \quad (1)$$

where for a given graph $G = (V, E)$, $\hat{\mathbf{A}} = \mathbf{A} + \mathbf{I}$ denotes the adjacency matrix with added self-loops for each vertex, $\hat{\mathbf{D}}$ is described by $\hat{D}_{ii} = \sum_{j=0} \hat{A}_{ij}$, a diagonal matrix displaying the degree of each node, and Θ denotes the learnable weight matrix. Added self-loops enforce that each node representation is directly dependent on its own preceding one. The number of graph convolutional layers stacked equals the radius of relevant nodes for each vertex within the graph.

The update rule for each node is given by a message passing scheme formalized by

$$\mathbf{x}'_i = \Theta \sum_j^N \frac{1}{\sqrt{\hat{d}_j \hat{d}_i}} \mathbf{x}_j \quad (2)$$

where both \hat{d}_i, \hat{d}_j are dependent on the edge weights e_{ij} of the graph. With simple, single-valued edge weights such as $e_{ij} = 1 \forall (i, j) \in E$, all \hat{d}_i reduce to d_i , i.e., the degree of each vertex i . We denote this type of graph convolutional neural layers with GCNCONV.

While in this initial formulation of a GCNConv the node-wise update step is defined by the sum over all neighboring node representations, we can alter this formulation to other message passing schemes. We can rearrange the order of activation function σ , aggregation AGG, and linear neural layer MLP with this formulation as proposed by [Li et al., 2020]:

$$\mathbf{x}'_i = \text{MLP}(\mathbf{x}_i + \text{AGG}(\{\sigma(\mathbf{x}_j + \mathbf{e}_{ji}) + \epsilon : j \in \mathcal{N}(i)\})) \quad (3)$$

where we only consider $\sigma \in \{\text{ReLU}, \text{LeakyReLU}\}$. We denote this generalized layer type as GENCONV following the notation of PyTorch Geometric [Fey and Lenssen, 2019]. While the reordering is mainly important for numerical stability, this alteration also addresses the vanishing gradient problem for deeper convolutional networks [Li et al., 2020]. Additionally, we can also generalize the aggregation function to allow different weighting functions such as learnable SoftMax or Power for the incoming signals for each vertex, substituting the averaging step in GCNCONV. Hence, while GCNCONV suffers from both vanishing gradients and signal fading for large scale and highly connected graphs, each propagation step in GENCONV emphasizes signals with values close to 0 and 1. The same convolutional filter and weight matrix are applied to and learned for all nodes simultaneously. We further employ another mechanism to avoid redundancy and fading signals in stacked graph convolutional networks, using residual connections and a normalization scheme [Li et al., 2019] [Li et al., 2020] as shown in Supplementary 3. The residual blocks are reusable and can be stacked multiple times.

- what is GATConv?
- what is KerGNN and what is its idea?
- add some sentences to the section above
- node vs. graph classification vs. link prediction

3.3.3 Dimensionality reduction techniques

3.3.3.1 Principal component analysis (PCA)

3.3.3.2 tSNE

3.3.3.3 UMAP

3.3.3.4 Parametric UMAP

3.3.4 Hyperparameter tuning

- RayTune[Liaw et al., 2018] for automated hyperparameter tuning

3.4 Evaluation and metrics

- AUC and AUPR for gene expression prediction
- self-built metric for evaluation of dim-red
- AUC and AUPR for conn pred

4 Results

4.1 Gene expression prediction

- We originally started from the per section prediction in order to paste its performance and results to other "related" structures within in the mouse brain. We propose multiple ideas As mentioned we used three different feature types in this study. ...(molecular features, phenotypical features, pure taxonomic features (InterPro embedding))) ...Due to the poor performance of the predictor with all three used feature types, we abandoned these plane

–

- structure specific features?
 - structural ontology / closeness
 - developmental hierarchy of tissue

Our model also allows us to test different ways of representing omics data. We tested different ways to normalize values assigned to genes as these normalizations convey different biological information; in the matrix of values assigned to genes from cancer samples, we can normalize values across the entire matrix, across each row (cancer sample), or across each column (gene). While a global normalization is more common, row-based normalization allows us to highlight values that are significantly higher or lower within one sample (e.g., which genes are expressed at high or low levels within a single sample), and column-based normalization allows us to highlight values assigned to a particular gene that are significantly higher or lower within one sample (e.g., whether a gene is expressed at higher or lower levels within one sample compared to all others). We find that column-based normalization performs better than row-based normalization, while the global normalization approach performs close to random. The best results are achieved when combining both row- and column-based normalization (Supplementary Table 2).

4.2 Dimensionality reduction and its combination with different graphs structures

- plot for showing validity of embeddings: K-means colour with respect to cluster
- plot colour parent structure all similar

4.3 On the linkage of connectivities and gene expression patterns

Brief Overview of Material

Findings (Results) of the Method of Study and Any Unplanned or Unexpected Situations that Occurred

Brief Descriptive Analysis Reliability and Validity of the Analysis

Explanation of the Hypothesis and Precise and Exact Data (Do Not Give Your Opinion)

5 Discussion

Brief Overview of Material

Full Discussion of Findings (Results) and Implications

Full Discussion of Research Analysis of Findings

Full Discussion of Hypothesis and of Findings

Post Analysis and Implications of Hypothesis and of Findings

Novelty:

- GCNs over gene expression was never applied here

6 Conclusion

Summary of Academic Study

Reference to Literature Review

Implications of Academic Study

Limitations of the Theory or Method of Research

Recommendations or Suggestions of Future Academic Study

- gene expression patterns within mouse brain and both possible hypothesis and tasks, and models over this
- gene knockout models and whether they can learn propagation of those?
- connection of FC and gene expression patterns and how to prove such interaction/correlation?
- possible gene knockout targets within mouse brain and possible structural influences

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