

Application of Graph Learning to inverse problems

Master Thesis Preperation

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Table of Contents

1	Intr	oduction	1
2	Imaging methods		
	2.1	Computed tomography	2
	2.2	Cryo-EM	3
	2.3	Abstract form	4
3	Gra	ph Denoising	6
	3.1	Graph Foundations	6
		3.1.1 Graph definition	6
		3.1.2 Graph Construction	7
	3.2	Graph Denoising definition	8
		3.2.1 Non-local means:	8
	3.3	Graph Laplacian	9
		3.3.1 Manifolds	9
		3.3.2 Connection to machine learning	11
		-	11
4	Mas	ter Thesis project	13
	4.1	Problem conclusion	13
	4.2	Work packages	14
	4.3	Gantt chart	15
Bi	bliog	raphy	16
\mathbf{A}	ppen	dix A Mathematical tools	18
	A.1	Power Iterations	18
	Δ 2	Wasserstein metric	18

Introduction

Inverse problems aim to estimate an original signal that went through a system, based on the output signal observation. Machine learning (ML) is a tool to model and solve such inverse problems. They are widely used throughout different science directions, such as ML, signal processing, computer vision, natural language processing and many more.

In recent years, graphs got a lot of attention in ML and are one of the most promising research areas. Graphs are a well suited data structure, simple but with high expressiveness. For some specific scenarios ordinary ML algorithm fail but Graph ML approaches have great success, e.g dimensionality reduction for high-dimensional data. Data can be in a graph structure already, like social networks, or they can be constructed for arbitrary datasets.

Cryo-electron microscopy (cryo-EM), where molecules are imaged in an electron microscope, gained a lot of attention in recent years. Due to ground-breaking improvements regarding hardware and data processing, the field of research has highly improved. In 2017, the pioneers in the field of cryo-EM got the Nobel Prize in Chemistry¹. Today, using cryo-EM many molecular structures can be displayed with near-atomic resolution. The big challenge with cryo-EM is enormous noise, which makes calculation challenging.

The following report resulted as the Master Thesis Preparation report. During the six week project, the aim was to familiarize with the research area, build up some mathematical foundation and define the project content as well as a project plan. The report is structured the following:

In chapter 2, the main motivation of the Master Thesis is given. The two imaging methods computed tomography and cryo-EM are introduced and an abstract model for dealing with both methods is defined. Chapter 3 is dedicated to graphs, were the connection of graphs to computed tomography and cryo-EM is established. Further, the problem of "Graph Denoising" is defined and methods like graph construction and Graph Laplacian is introduced. Finally, in chapter 4 the project content in shortly concluded, work packages are defined and the project schedule is given. Throughout the report, red boxes are used to denote important statements and connections regarding the Master Thesis.

¹ https://www.nobelprize.org/prizes/chemistry/2017/press-release/

Imaging methods

In the current chapter, imaging methods computed tomography and cryo-electron microscopy (cryo-EM) will be introduced. Further, their observation model is defined in a mathematically way and their reconstruction is observed. Application of cryo-EM is the major motivation for the Master Thesis, as the problem is not easy to solve due to dealing with enormous noise and other difficulties.

2.1 Computed tomography

Computed tomography is a well established imaging method. Using X-ray source, fan shaped beams are produced which scan the imaging object, resulting in many measurements taken over straight lines [3].

Tomography reconstruction: Tomographic reconstruction[6] is a popular inverse problem. The aim is to reconstruct an imaged object from observed measurements. The reconstruction object can be in two-dimension (2D) or in three-dimension (3D).

In the Master Thesis, the focus on computed tomography will be on 2D case, which is called *classical tomography reconstruction*.

2D tomographic reconstruction: Mathematically, the observed measurements can be defined as follows:

$$y_i[j] = R(x, \theta_i, s_j) + \eta_i[j], \text{ with } 1 \le i \le N \text{ and } 1 \le j \le M,$$
 (2.1)

where N is number of observations and M the observation dimension. Then, $x \in L^2(\Omega)$ is the original object with $\Omega \subset \mathbb{R}^2$ and L^2 the Lebesgue space. Further, $y_i \in \mathbb{R}^M$ is the *i*-th observation with $y_i[j] \in \mathbb{R}$ *j*-th element of the observation.

 $R(\cdot; \theta, s) : L^2(\Omega) \to L^2(\tilde{\Omega}), x \mapsto R(x; \theta, s)$ refers to the Radon Transform[18] with $\tilde{\Omega} \subset \mathbb{R}$, θ as the observation angle from the x-axis and s_j as the sampling point. η refers to noise and is defined as $\eta_i[j] \sim \mathcal{N}(0, \sigma^2)$.

Imaging methods 3

Filter Backprojection: Filter Backprojection [6] is a reconstruction method, typically used in classical tomography. It allows to inverse the Radon Transform and enables reconstruction of the original object x. The algorithm fails when working with noisy data [15].

2.2 Cryo-EM

Cryo-EM is another imaging method, that enables the view of molecules in near-atomic resolution. In the Master Thesis, only single-particle cryo-EM[9] is considered, when writing about cryo-EM it always refer to single-particle cryo-EM. During imaging process molecules are frozen in a thin layer of ice, where they are randomly oriented and positioned. Random orientation and positioning of molecules makes reconstruction challenging but the freezing process allows to observe molecules in a stable state where they are not moving. With an electron microscope, two-dimensional tomographic projection images of the molecules are observed in the ice, which are called *micrograph*. The frozen molecules are fragile and the electron microscope needs to work with very low power (electron dose), resulting in highly noisy images. The resulting signal-to-noise ration (SNR) is typically smaller than 1, which indicates that there is more noise than signal [15]. Further, observed molecules are not equal in the sense that there are some structural varieties between the molecules within a observation.

3D cryo-EM reconstruction: Similar to tomographic reconstruction, cryo-EM reconstruction problem [2] is defined. It can be seen as a 3D reconstruction problem as the original object $x \in L^2(\Omega)$ to be reconstructed is in 3D. To keep the notation from previous section, now $\Omega \subset \mathbb{R}^3$ and $\tilde{\Omega} \subset \mathbb{R}^2$.

Mathematically, the observed measurements can be defined as follows:

$$y_i = \Pi_z(Rot(x; \theta_i)) + \eta_i$$
, with $1 \le i \le N$ (2.2)

where $\Pi: L^2(\Omega) \to L^2(\tilde{\Omega}), x \mapsto \int x(\cdot, \cdot, z) dz$ is the projection operator and $Rot: L^2(\Omega) \to L^2(\Omega), Rot_{\theta}(x) = ((x_1, x_2, x_3) \mapsto x(x_1R^1, x_2R^2, x_3R^3))$ is the rotation operator modelling the rotation during freezing. Further, $\theta_i = [\theta_i^1, \theta_i^2, \theta_i^3]$ where entries $\theta_i^1, \theta_i^2, \theta_i^3 \in \mathbb{R}$ and $R = [R^1, R^2, R^3] \in SO(3)$. $\eta_i[j, k] \sim \mathcal{N}(0, \sigma^2 I)$ corresponds again to the noise of the observation.

As y_i is not observable directly, discretization is needed:

$$y_{i} = (\Pi_{z}(Rot(x;\theta_{i})) + \eta_{i})(\Delta), \text{ with } 1 \leq i \leq N$$

$$y_{i}[j,k] = \Pi_{z}(Rot(x;\theta_{i}))_{j,k} + \eta_{i}[j,k], \text{ with } 1 \leq i \leq N \text{ and } 1 \leq j,k \leq M$$

$$(2.3)$$

where $\Delta \subset \tilde{\Omega}^{M^2}$ is the sampling grid and M is the first and second dimension of the sampling grid.

Extended formula: Equation $\ref{eq:condition}$ is a simplified version of cryo-EM. First of all, point spread function (PSF) of the microscope is not taken into account. Secondly, structural variety is not taken into account, the underlying object x is not the same for every observation

Imaging methods 4

as modelled in the equation. Precisely, it can be seen as a random signal from an unknown distribution defined over all possible molecules structures.

The equation can be extended and defined as the following:

$$y_i = h_i \circ \Pi_z(Rot(x_i; \theta_i)) + \eta_i, \text{ with } 1 \le i \le N$$
 (2.4)

where h_i is the PSF of the microscope and \circ defines the convolution.

During Master Thesis, equation 2.3 is used, not the extended version.

Difference to tomographic reconstruction: The two problems are highly related, but the cryo-EM reconstruct is more challenging. During CT observation, the patient is asked to not move and therefore, the angles of projection are known. Whereas, in cryo-EM this information will be lost during the freezing process. Secondly, the high level of noise makes cryo-EM much more challenging regarding tomographic reconstruction.

2.3 Abstract form

As the tomographic reconstruction and the cryo-EM reconstruction are rather similar, the aim of the Master Thesis will be to design an algorithm, that can be applied in both scenarios. Therefore, an abstract form of the problems will be defined in the following. First of all, a similar notation as before is used, but in a more general way $x \in L^2(\Omega)$ where $\Omega \subset \mathbb{R}^D$ with D as the dimension of the space and $\tilde{\Omega} \subset \mathbb{R}^{D-1}$.

$$y_i = (A(x, \theta_i) + \eta_i) (\Delta)$$
(2.5)

where $y_i \in \tilde{\Omega}^M$ is the observed measurements, M the measurement dimension, $x \in L^2(\Omega)$ our original object, A a non-linear operator $A: L^2(\Omega) \to L^2(\tilde{\Omega}), x \mapsto A(x; \theta)$ and $\eta \sim \mathcal{N}(O, \sigma^2 I)$ gaussian noise. $\Delta \subset \tilde{Omega}^{M^2}$ is a term for discretization.

Classical tomography reconstruction: For classical tomography parameters are defined defined with D=2 and $\theta \in \mathbb{R}$. Further, $A(\cdot)$ is the Radon Transform, defined in equation 2.1. A distance measure between measurements can be set up by using the ℓ 2-norm $||y_i - y_j||$.

Cryo-Em reconstruction: For cryo-EM parameters are defined with D=3 and $\theta \in \mathbb{R}^3$. Further, $A(\cdot)$ can be defined as Π_z ($Rot(x;\theta)$) where Rot is the 3D rotation and Π_z the tomographic projection.

As measurements are drawn with some random 3D rotation and projection, it can happen that two samples are equivalent up to 2D rotation. Consider a first example y_1 , which has no 3D rotation and a second sample y_2 with a rotation only in in x-y plane by 45°. The two samples have a defined in-plane rotation g, such that $gy_1 = y_2$. Therefore, in our distance measure we add this term of in-plan rotation: $\min_{g \in SO(1)} \|g * y_i - y_j\|$, which is inspired by the work of [10].

Imaging methods 5

High noise regime: Cryo-EM measurements are highly noisy, which makes reconstruction challenging. There are different ways to reduce noise from measurements, most of them are related to averaging. Averaging need to consider similar measurements and ignore diverse ones. In the defined abstract model, averaging over paired measurements from θ should be a good averaging model. But how can it be achieved?

One idea would be to measure distances between observation (therefore introduced above). Another way is to find a low-dimensional embedding which maps our measurements y to some θ . When talking from low-dimensional embeddings, there is no way around Graph Learning, which will be introduced in the following chapter.

During the Master Thesis, high-noise regime is the domain of interest. The main practical application is cryo-EM, where an algorithm for denoising is expected to boost quality of the overall 3D-reconstruction. As cryo-EM is a 3D problem, computed tomography will be considered as well which allows to test on a corresponding 2D problem. The aim of the Master Thesis is to introduce a denoising algorithm, which is able to work well even on highly noisy data, where cryo-EM is major field of interest.

Graph Denoising

In the following chapter, the connection between graphs and the domain of denoising in high-noise domains, such as cryo-EM, is established. First, a broad definition of graphs is given and further, the term "Graph Denoising" is introduced and explained. Finally, the connection to Graph Laplacian is established and different opportunities exploiting for a good denoising algorithms are shown.

3.1 Graph Foundations

Real world data can be in graph structure, like social networks, citation networks, protein interaction networks or google search. If data is not available in graph structure, a graph can be artificially constructed with methods like k-nearest neighbours (k-NN) or others.

Graph Learning: Graph Learning is a hot research area and got a lot of attention in recent years. It is a way of applying ML on graphs and algorithms emerged from ML but also other fields. When a graph is available, one can start using Graph Learning algorithms for solving tasks. Popular tasks are node classification or link prediction within a graph, where the model is learned from node and edge features as well as the topology of the graph. The model can than be used for prediction or classification. Another common task is community detection, where the aim is to identify cluster of nodes within the input graph. Further, graphs are highly favoured for dimensionality-reduction, where graph algorithms provide a helpful tool, as ordinary algorithms like principle component analysis fail.

3.1.1 Graph definition

A graph is defined as $G = \langle V, E \rangle$, where V is a set of vertices (or nodes) and E is a set of edges (or links). Edges are defined as a set of tuples (i, j), where i and j determine the index of vertices in the graph.

Graph properties: A graph can be either *directed* or *undirected*. In a directed one, an edge connects explicitly from one node to another, which means that edge $(i, j) \neq (j, i)$. In undirected graphs ordering does not matter and (i, j) = (j, i).

The neighbourhood, denoted by $\mathcal{N}(i)$, of a node i is defined as all adjacent nodes, or in other words, there is an edge between the neighbourhood nodes and i. Further, edges can have weights, which is a method to define importance to neighbours of a node. If edges are dealing with weights, the term weighted graph is used. The degree of a node are the number of incoming edges.

Adjacency matrix: To do calculations with graphs, it is common to translate graphs in a well suitable mathematically form, which are matrices. The adjacency matrix can be seen as a way of representing graphs as a matrix. The (binary) adjacency matrix of graph G is defined as follows:

$$A_{ij} = \begin{cases} 1 & \text{if } (i,j) \in E \\ 0, & \text{otherwise} \end{cases}$$
 (3.1)

The matrix A has dimension $\mathbb{R}^{N\times N}$ with N as number of nodes, and indices of A correspond to nodes of the graph. If there exists an edge between two nodes, the entry in A will be set to 1, otherwise to 0. This leads to an unweighted graph, as weight of all edges will be 1, but could easily be extended by assigned not just values of 1 or 0. When the graph is undirected, the corresponding adjacency matrix will be symmetric. Eigenvalues of A are called *spectrum* of the graph.

3.1.2 Graph Construction

When data is not available as a graph, it can be easily constructed. Consider data from space $\Omega \subset \mathbb{R}^M$, but could basically be any arbitrary space. Then, each node is associated with some element $x \in \Omega$. Further, the graph G can be constructed by using:

$$A_{ij} = \begin{cases} 1 & \text{if } d(x_i, x_j) < \tau \\ 0, & \text{otherwise} \end{cases}$$
 (3.2)

where n_i , n_j are nodes from indices i, j, d is corresponds to a similarity measure between two nodes and τ is a threshold, when to consider two nodes to be adjacent. K-NN is one possible implementation of a graph construction algorithm, where for every node, k neighbours will be defined. The neighbourhood of node i is defined as \mathcal{N}_i and consists of the nodes, with the k smallest similarity measure.

Noise regime In the case of noise, observation of x it not possible directly. Measurements will give access to $y = x + \eta$ where $y, x \in \Omega$ and the noise η is assumed to be drawn from gaussian distribution $\mathcal{N} \sim (0, \sigma^2)$. The noisy graph G_0 can be constructed as in equation 3.2, but replacing y with x:

$$A_{ij} = \begin{cases} 1 & \text{if } d(y_i, y_j) < \tau \\ 0, & \text{otherwise} \end{cases}$$
 (3.3)

3.2 Graph Denoising definition

First of all, $Graph\ Denoising$ is not a common term in literature. In the previous section, noisy graph G_0 was introduced and the aim is to denoise this graph, which means to estimate the original graph G from a given noisy graph G_0 . This is what is meant with Graph Denoising, which is rather related to signal or image denoising. Reconstruction of a true signal given noisy observation signal is done via averaging, that can be performed locally, by the calculus of variations or in the frequency domain[4].

Noisy Graph: For every noisy graph there exists an original graph $G = \langle V, E \rangle$. The noisy graph G_0 can further be defined as $G_0 = \langle V, E_0 \rangle$, where $E_0 = E \setminus E^- \cup E^+$ with $E^- \subseteq E$ and $E^+ \cap E = \emptyset$.

 G_0 consists of same nodes V as the original graph G. From the edges of G denoted by E, some are removed (denoted by E^-) and some are added (denoted by E^+), which results is the edges E_0 of G_0 .

The adjacency matrix of G_0 is denoted by A_0 . The task of Graph Denoising, can therefore be written as $GD: A_0 \mapsto \tilde{A} \approx A$, where A_0 , \tilde{A} , A denotes the adjacency matrix from noisy input graph, denoised graph and original graph respectively.

The goal of the Master Thesis is to introduce a method to estimate the original graph G with corresponding A based on an observed noisy graph G_0 with A_0 .

Connection to link prediction Link prediction is a task in Graph Learning. The idea is to predict existence of a link (edge) between two nodes. The task can be formulated as a missing value estimation task. A model M_p is learned from a given set of observed edges. The model finally maps links to probabilities $M_p: E' \to [0,1]$ where E' is the set of potential links.

Further, U determines the set of all possible vertices of G, therefore $E \subseteq U$. Obviously, Graph Denoising can be seen as a link prediction problem. The difference is, that in link prediction a model from a set of observed links is learned $E_{observed} \subseteq E$ and in Graph Denoising the model is learned from $E_{observed} \subseteq U$.

3.2.1 Non-local means:

In the following section, a short introduction to the state-of-the-art image denoising method non-local means is given[4]. For a given noisy image v, the denoised image is defined as $NL[v](i) = \sum w(i,j) \ v(j)$. where w(i,j) is the weight between pixel i and j. Weight can be seen as similarity measure of pixels. Moreover, similarities are calculated over square neighbourhoods of pixels, where $\ell 2$ -norm of the neighbourhood is used. Similar pixel neighbourhoods have a large weight and different neighbourhoods have a small weight. More general, the denoised image pixel i is computed as an weighted average of all pixels in the image, therefore, in a non-local way.

Non-local means is not a denoising algorithm, which works with graph as a data structure. But, it uses a neighbourhood for averaging, which shows great potential of graphs as a data

structure as graphs can represent neighbours really well.

3.3 Graph Laplacian

The Graph Laplacian is a matrix that represents the graph and can be used to find many important properties. It is a very powerful tool and therefore, a complete section is dedicated to it. A good introduction and overview can be found by [17, 20].

The matrix is defined as follows:

$$L = D - A, (3.4)$$

where A is the adjacency matrix and D the degree matrix (diagonal matrix with degree of nodes as entries).

3.3.1 Manifolds

In high-dimensional data euclidean distances are not meaningful, in the sense that they will not capture similar data points well. Graph Laplacian can be used to compute a Manifold, which can help in such a scenario. In the manifold space, euclidean distances make sense again. Let manifold M be defined as $\mathcal{M} = \{f(x), f \in C^K, f : \mathbb{R}^D \to \mathbb{R}^d\}$. Manifolds are a well established mathematical concept. In the Master Thesis, only C^k differentiable d-dimensional manifold defined by \mathcal{M} are considered. When $d \ll D$, manifold defines a low-dimensional embedding, which maps from high-dimensional space \mathbb{R}^D to low-dimensional space \mathbb{R}^d .

Lets give two popular examples of such manifolds, namely the *circle* and the *sphere*. The circle is a 1D manifold, where d=1 and D=2. A sphere is a 2D manifold, with d=2 and D=3. In figure 3.1(a), 200 samples are drawn from a uniform distribution of circle manifold and in figure 3.1(b), 400 samples are drawn from a uniform distribution of sphere manifold.

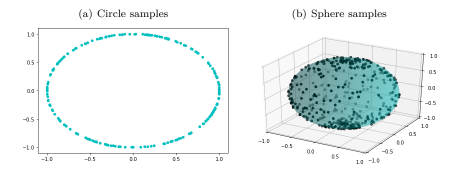


Figure 3.1: Samples drawn from 1D and 2D manifold.

One popular algorithm for calculating manifolds is Diffusion Maps[7], which is a non-linear approach for calculating low-dimensional manifolds for (high-dimensional) datasets, using Graph Laplacian. Vector Diffusion Maps[16] generalize the concept of Diffusion Maps for vector fields. Multi-Frequency Vector Diffusion Maps (MFVDM)Fan and Zhao [10] can be seen as an extension to Vector Diffusion Maps [16], which works well even on highly

noisy environments. [10] was successfully applied in cryo-EM setting, where it was used for denoising purpose[11].

Manifold assumption: The manifold assumption is a popular assumption for high-dimensional datasets. For a given dataset in high-dimension, one can assume that data points are samples drawn from a low-dimensional manifold, that embeds the high-dimensional space. Therefore, if underlying manifold can be approximated, a dimensionality reduction is established as one can embed the data points in the low-dimensional manifold space. There is a complete area of research devoted to this manifold assumption called Manifold Learning[5], but it is not only used there.

Manifold estimation: The manifold, and therefore, a low-dimensional embedding, can be calculated the following:

- 1. Construct k-NN graph from observations (see section 3.1.2).
- 2. Calculate the (normalized) Graph Laplacian.
- 3. Extract the second, third (and fourth) smallest eigenvectors.

Therefore, it can be observed how the manifold of classical tomography and cryo-EM looks like. In the following, the Shepp-Logan phantom is used as an example of a classical tomography image.

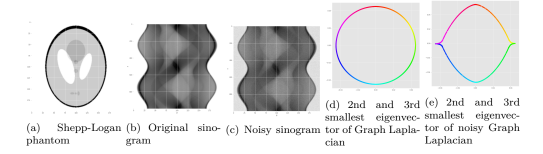


Figure 3.2: Shepp-Logan phantom manifold

In Figure 3.2(d) the manifold calculated from the original image Graph Laplacian can be seen and it is a perfect circle. Further, noise was added to the image and in Figure 3.2(e) the noisy version with $\sigma=2$ is plotted and the manifold is circle like but not a perfect circle. The more noise is added to the original image, the less the manifold looks like a circle. In Figure 3.3(b) the manifold for $\sigma=100$ is plotted. In all plots, knn-graph have been constructed with k=10.

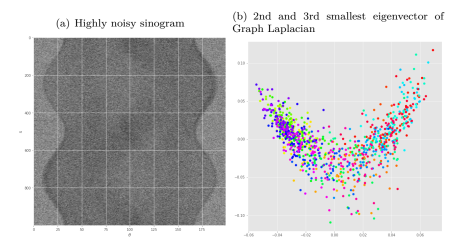


Figure 3.3: Shepp-Logan phantom manifold for high noise level

In the field of classical tomography and cryo-EM, the manifold is well defined for none-noisy data. In the 2D case of classical tomography, the underlying manifold is a circle, whereas in 3D case of cryo-EM the manifold if defined as a sphere. This fact can be exploited during learning, by using wasserstein loss function (see A.2).

3.3.2 Connection to machine learning

Graph Laplacian is used for dimensionality reduction for high-dimensional data, as well as spectral clustering and semi-supervised learning. Coifman et al. [8] used Graph Laplacian in a complete other domain, namely in tomography. They showed that Graph Laplacian approximates the Laplace-Beltrami operator. Further, Graph Laplacian is depended on the adjacency matrix A, if A is noisy, Graph Laplacian will be noisy as well.

In the Master Thesis, further experience of Graph Laplacian in the domain of tomography and denoising should be gained. Additionally, connection with machine learning will be explored, hopefully allowing to learn a denoised adjacency matrix to fully enable the power of Graph Laplacian.

3.3.3 Graph Deep Learning

As already mentioned, Graph Denoising can be seen as a way of link prediction. The state-of-the-art method for solving link prediction are graph deep learning approaches. Graph deep learning is a fast evolving field in research. With Graph Neural Networks (GNN)[13] the framework for GNN has been established.

Using Graph Convolutional Networks (GCN) [14] for graph feature extraction is a popular way. Basically, with GCN a new feature representation is iteratively learned for the node features (edges features are not taken into account). It can be seen as an averaging of nodes over their neighbourhood where all the neighbours get the same weight combined with some

non-linear activation. To consider the node itself in the averaging process they apply to so-called "Renormalization trick", where self-loops are added to the adjacency matrix and after every layer, a normalization step is applied. The topology of the graph will not be adjusted during the learning process.

Veličković et al. [19] extended the concept of GCN with attention and not all the neighbouring nodes get the same weight (attention). Simple Graph Convolutional Network (SGC) [22] proposed a simplified version of GCN. They could verify their hypothesis that GCN is dominated by the local averaging step and the non-linear activation function between layers do not contribute to much to the success of GCN. Therefore, it can be seen as a way of power iteration (see A.1 for further information) over the adjacency matrix with normalization in every layer. Wang et al. [21] proposed a extended version of GCN by not operating on the same graph in every layer but adopting the underlying graph topology layer by layer.

In the Master Thesis, connection between Graph Laplacian and GNNs should be further studied. The fact, that Wu et al. [22] could simplify the existing GCN algorithm is motivation enough that similar connections can be drawn in other fields of Graph Learning.

Master Thesis project

In the last chapter of the report, a short conclusion of the Thesis problem is given. Further, the project plan will be introduced as well as a broad overview of different work packages. Finally, the project timeline can be seen as a Gantt chart.

4.1 Problem conclusion

Cryo-Em and computed tomography within the high noise regime is the field of interest of the Master Thesis. In both imaging methods, the final goal is to reconstruct an original object from observed noisy data. It is expected, that better denoising will boost the overall quality of reconstruction.

The Master Thesis will therefore introduce a new method for denoising. We believe, that graphs are a well suited data structure for denoising. Additionally, we hope to explore further connection of the powerful Graph Laplacian and machine learning and use it to estimate a denoised graph from a noisy one. Further, the idea is to exploit the fact, that the manifold of the noise-less input data is known.

Evaluation: The introduced denoising method will be evaluated during the Master Thesis. During evaluation, 2D and 3D scenario will be considered. A first evaluation will be done on artificial constructed toy-dataset. If time allows, real dataset from classical tomography and/or cryo-EM² can be evaluated as well.

During evaluation, two baselines are considered which already solved part of the problem. The first one is a multi-frequency diffusion map approach[10, 11], which aims to denoise cryo-EM images. Secondly, [8] a Graph Laplacian approach solving classical tomography with random projection angles will be compared against.

The evaluation process is a first broad idea. Any adjustments in baseline papers or dataset are possible during the Master Project.

² https://www.ebi.ac.uk/emdb/

4.2 Work packages

In the upcoming section, some work packages are defined. Probably, there are some parts of the project which will not work out as expected and adjustments are needed throughout the Thesis. The project plan can be seen as a rough guideline.

Implement algorithm for 2D case: The first step will be, to familiarize with the problem and implement the algorithm for 2D.

Evaluate 2D case on toy dataset and implement baselines: As a second step, the implemented 2D algorithm will be tested on a toy dataset, where noise is added to images by hand. As the goal is to work with highly noisy images, the noise level can be selected and increased when working with toy datasets. The evaluation in 2D is crucial and needs to in a satisfying matter. It does not make sense to continue with 3D implementation, when the simple 2D case is not handled well enough. Therefore, if evaluation results are not satisfying, the algorithm needs to be iteratively adjusted, such that the evaluation will be in a good enough quality.

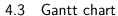
Implement algorithm for 3D case: After successfully evaluating the algorithm in 2D, the aim is to extend the algorithm to work in 3D as well.

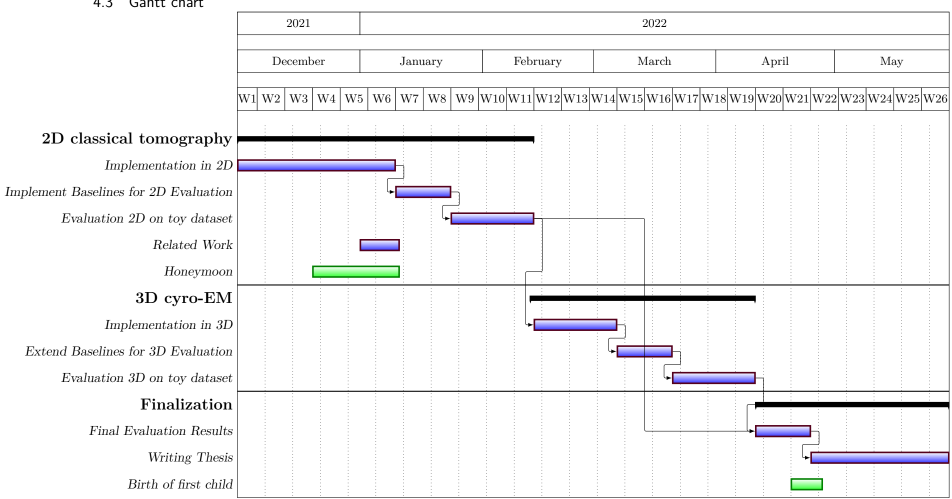
Evaluate 3D case on toy dataset and adjust baselines: Again, the implementation will be evaluated on a toy dataset, where noise can be adjusted by hand.

Nice to have: Evaluate of real dataset If time allows and the 2D and 3D implementation are evaluated successfully on toy datasets, real data can be used for further evaluation. This step will only be done, if time allows.

Evaluate related work: As cryo-EM reconstruction is a hot research topic, related work can not only be considered during the start of the Thesis and needs to be evaluated throughout the Thesis.

Writing Thesis: Document implementation and evaluation result.





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Mathematical tools

A.1 Power Iterations

Power iteration (also called power method) is an iteratively method, that approximates biggest eigenvalue of a diagonalizable matrix A.

The algorithm starts with a random vector b_0 or an approximation of the dominant eigenvector.

$$b_{k+1} = \frac{Ab_k}{||Ab_k||} \tag{A.1}$$

The algorithm not necessarily converges. The algorithm will converge, if A has an eigenvalue strictly grater than its other eigenvalues and initial vector b_0 is not orthogonal to the eigenvector associated with the largest eigenvalue.

A.2 Wasserstein metric

The Wasserstein metric is a distance measure between two probability distributions and it is used in ML as a loss function[12]. Intuitively, it can can be understood as the minimum cost to transfer the mass of one distribution to the other. Therefore, it is also known as the earth mover's distance.

As Arjovsky et al. [1] could show, ordinary distance measurements like *Total Variation*, *Kullback-Leibler divergence* and *Jensen-Shannon divergence* are not sensible when learning with distributions supported by manifolds On the contrary, Wasserstein metric does a good job as loss function in such scenarios.