USC-SIPI REPORT #400

Information Theoretic Measures for PET Image Reconstruction and Non-Rigid Image Registration

by

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December 2009

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INFORMATION THEORETIC MEASURES FOR PET IMAGE RECONSTRUCTION AND NON-RIGID IMAGE REGISTRATION

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A Dissertation Presented to the FACULTY OF THE USC GRADUATE SCHOOL UNIVERSITY OF SOUTHERN CALIFORNIA In Partial Fulfillment of the Requirements for the Degree DOCTOR OF PHILOSOPHY (ELECTRICAL ENGINEERING)

December 2009

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Dedication

To my parents Seshu and Venkata Lakshmana Rao Somayajula
Acknowledgments

I thank Prof. Richard Leahy for being an excellent advisor and teacher. His guidance at every step of my research work has been invaluable, and I am grateful for the encouragement he provided when I needed it. He has been an inspiring role model for me. I thank him for providing financial support, for helping me form productive collaborations, and for making my experience as a graduate student truly valuable. I also thank my guidance committee Dr. Manbir Singh, Dr. Krishna Nayak, Dr. Antonio Ortega, and Dr. Quanzheng Li for their time and feedback. I thank my collaborators Christos Panagiotou, Dr. Simon Arridge, and Dr. Anand Rangarajan for contributing to the PET work through their valuable suggestions. Additionally, I thank Dr. Richard Laforest, Dr. Quanzheng Li, and Dr. Giselle Petzinger for providing data to validate my PET work. I am grateful to the Women in Science and Engineering program for awarding me a top-off scholarship for the first two years of my PhD studies, and for providing travel grants that helped me attend major conferences in my field.

It has been a pleasure to work with my colleagues in the USC Biomedical Imaging group. It was an enriching experience to discuss my work with them, and I thank them for being a helpful and collaborative research group. I thank Dr. Sangtae Ahn, Dr. Anand Joshi, Dr. Quanzheng Li, and Dr. Sanghee Cho for always having the time to answer
my questions, and Dr. Evren Asma for his help early in my research work. I thank Dr. Abhijit Chaudhari, Joyita Dutta, and Dr. Belma Dogdas for being good friends besides being great colleagues. I thank Dr. Zheng Li, Dr. Dimitrios Pantazis, Yu-teng Chang, Yanguang Lin, Ran Ren, Syed Ashrafulla, Juan Poletti Soto, Hua Hui, and David Wheland for being great colleagues, and for the fun times I had while being a part of the research group.

I am grateful to my dear friends without whom I would be lost. I thank Kavitha Buddhharaju for the faith she has in me, and for not letting astronomical international call rates stop her from always being there for me. I thank Lalit Varma for teaching me to believe in myself. Though he left this world early in my PhD program, he was and will continue to be a source of inspiration for me. I thank Suma Mallavarapu for being my sounding board, though she had her own PhD and her baby to take care of. I thank Gautam Thatte, Michelle Bardis, Laura Woo, Ashok and Amola Patel, Manoj Gopalkrishnan, Krishna and Sandhya Gogineni, and Ratnakar Vejella for being strong sources of support throughout my graduate student years. I thank Srikanth Pinninti, Joaquin Rapela, Vijay Ponduru, Pradeep Natarajan, and Haritha Bodugam for making my years at USC an enjoyable experience.

Finally, I thank my family for being pillars of support throughout my graduate studies. I thank my grandmother Eswaramma Somayajula for taking immense pride in everything I do, and my sister Sunitha Mukkavalli for being my friend for as long as I can remember. My parents Seshu and Venkata Lakshmana Rao Somayajula cheered me on every step of the way, and I thank them for their constant love and support.
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Abstract

We explore the use of information theoretic measures for positron emission tomography (PET) image reconstruction and for multi-modality non-rigid registration.

PET is a functional imaging modality based on imaging the uptake of a radioactive tracer. PET images are typically of low resolution and are accompanied by high resolution anatomical images such as CT or MR for localization of activity. PET tracer uptake typically results in a spatial density that is highly correlated to the anatomical morphology. The incorporation of this correlated anatomical information can potentially improve the quality of low resolution PET images. We propose using priors based on information theoretic similarity measures to incorporate anatomical information in PET reconstruction. We compare and evaluate the use of mutual information (MI) and joint entropy (JE) between feature vectors extracted from the anatomical and PET images as priors in PET reconstruction. These feature vectors are defined using scale-space theory such that they emphasize prominent boundaries in the anatomical and functional images, and attach less importance to detail and noise that is less likely to be correlated in the two images. We present an efficient method of computing these priors and their derivatives based on fast Fourier transforms that reduce the complexity of their convolution-like
expressions. Through simulations and clinical data reconstructions, we evaluate the performance of MI and JE based priors in comparison to a quadratic prior, which does not use any anatomical information. We also apply these priors to the problem of positron range correction. Positron range is the distance traveled by a positron before annihilation, thereby causing a blurring effect in the reconstructed image and limiting its resolution. We use information theoretic priors in conjunction with a system model that incorporates positron range by modeling it as a spatially invariant blurring that is truncated at the boundary of the imaging volume. We present phantom simulation and real data results comparing these priors to the range corrected system model with quadratic prior.

Small animal non-rigid registration is especially challenging because of the presence of rigid structures like the skeleton within non-rigid soft tissue. We present two approaches to multi-modality imaging that can be applied to clinical as well as pre-clinical images. First, we describe a non-parametric scale-space approach to MI based non-rigid small animal image registration. In this application, the goal is to simultaneously align global structure as well as detail in the images. We present results based on CT images obtained from two time points of a longitudinal mouse study that demonstrate that this approach aligns both skeleton and soft tissue better than a commonly used hierarchical approach. Second, we explore an alternative formulation that uses the log likelihood of the reference image (target) given the image to be registered (template) as a similarity metric wherein the likelihood is modeled using Gaussian mixture models (GMM). Using the GMM formulation reduces the density estimation step to that of estimating the GMM parameters. These parameters are small in number for images that have few regions of distinct intensities, such as brain or microCT images. This approach is more robust than
the non-parametric MI based approach because of reduced overall dimensionality of the problem and more robust and accurate density estimation. We present comparisons of the non-parametric MI based approach to that of the GMM conditional likelihood based approach through intra-modality brain and multi-modality mouse images.

Finally, we present future directions for the work described in this dissertation.
Chapter 1

Introduction

This dissertation explores the use of information theoretic measures for two biomedical imaging applications - positron emission tomography (PET) image reconstruction, and non-rigid image registration. In both applications, the goal is to maximize similarity in the information theoretic sense between the image to be reconstructed or registered, and a correlated image.

1.1 Positron Emission Tomography Image Reconstruction

Positron emission tomography is a non-invasive biomedical imaging tool that yields the density distribution of a radioactive positron emitting tracer. It is a functional imaging modality that can be used to track metabolic and hemodynamic processes in vivo. The radioactive tracer is injected into or inhaled by the subject, after which it undergoes decay, emitting positrons. Each emitted positron is annihilated by an electron, producing two photons that travel in opposite directions by the law of conservation of momentum. These photons are detected by a circular ring of detectors, and a pair of photons striking detectors opposite to each other within a narrow timing window is called a coincidence.
Figure 1.1: PET/CT imaging: Whole body PET image (left), its corresponding CT image (middle), and their overlay (right). Image was obtained from the website http://medical.siemens.com

The data obtained is the number of coincidences in each angular direction around the subject. From this data, an image representing the spatial density of the tracer can be reconstructed, which gives insight into the physiological process that is being studied.

PET images visualize physiological processes but do not provide anatomical context, thus making it necessary to complement them with a CT or MR scan. Modern day PET scanners have CT scanners built into them, and can output PET images that are overlaid on the CT. Multi-modality systems including combined PET/MRI scanners is a growing area of research, a review of which is given in [1], [2]. Thus, PET images which are of low spatial resolution (2 mm at best for clinical PET) are usually accompanied by high resolution CT or MR (resolution < 1mm) images of the same subject. An example of PET with corresponding CT image, and their overlay is shown in Figure 1.1.

PET tracer uptake has been observed to be highly correlated with the underlying anatomical morphology of the subject. This can be seen in Figure 1.1, where the PET
activity is within regions that can be clearly delineated in the corresponding CT image. Thus, the incorporation of anatomical information from correlated MR/CT images into PET reconstruction algorithms can potentially improve the quality of low resolution PET images. Quantitative inaccuracies due to partial volume effects in the PET image could be corrected during the reconstruction process, or distinct structures that cannot be resolved in the PET image from the data alone could be visible, because of the high resolution anatomical information introduced into the reconstruction algorithm. In this dissertation, we will explore a method of incorporating anatomical information into PET reconstruction through a Bayesian framework wherein information theoretic measures of similarity are used as priors on the PET image. The goal is to reconstruct a PET image that fits the data and is most similar to the anatomical image in the information theoretic sense.

Previous work on the use of anatomical priors can be broadly classified into: (i) Methods based on anatomical boundary information, which encourage boundaries in functional images that correspond to anatomical boundaries [3] - [6], and (ii) methods that use anatomical segmentation information, which encourage the distribution of tracers in regions corresponding to anatomical regions [7]- [11]. In [12] a method that did not use boundary or segmentation information was proposed, wherein the prior aimed to produce homogeneous regions in the PET image where the anatomical image had an approximately homogeneous distribution of intensities. These approaches rely on boundaries, segmentation, or the intensities themselves to model the similarity in structure between the anatomical and functional images. In this work, we use information theoretic measures
such as mutual information (MI) and joint entropy (JE) to model the similarity between images in the information theoretic sense.

Mutual information (MI) between two random vectors is a measure of the amount of information contained by one random vector about the other and can be used as a similarity metric between two multi-modality images [13]. In [11], a Bayesian joint mixture model was formulated such that the solution maximizes MI between class labels. In [14, 15] we described a non-parametric method that uses MI between feature vectors extracted from the anatomical and functional images to define a prior on the functional image. This approach did not require explicit segmentation or boundary extraction, and aimed to reconstruct images that had a distribution of intensities and/or spatial gradients that matched that of the anatomical image. MI achieves this by minimizing the joint entropy (JE), while maximizing a marginal entropy term that models the uncertainty in the PET image itself. In [16] it was shown through anecdotal examples that due to the marginal entropy term, MI tends to produce biased estimates in the case where there are differences in the anatomical and functional images, and that joint entropy is a more robust metric in these situations.

In this work, we extend the non-parametric framework of our MI based priors to explore the use of both MI and JE as priors in PET image reconstruction. Since both these information theoretic metrics are based on global distributions of the image intensities, spatial information can be incorporated by adding features that capture the local variation in image intensities. We define features based on scale space theory, which provides a framework for the analysis of images at different levels of detail. It is based on generating a one parameter family of images by blurring the image with Gaussian kernels of increasing
width (the scale parameter) and analyzing these blurred images to extract structural features [17]. We evaluate the performance of both MI and JE priors in quantifying uptake in specific regions as well as overall, through simulations and clinical data. We also present an efficient method of computing the priors and their derivatives, which makes these priors practical to use in 3D image reconstruction. The obtained results indicate that using JE based priors in PET reconstruction can improve the resolution of reconstructed PET images when there is reasonable agreement between the anatomical and PET images.

One of the fundamental resolution limiting factors for PET is positron range [18]. In PET the positron emitting source is assumed to lie on the line of response (LOR) that joins the detectors at which the the photon pair is detected after positron-electron annihilation. However, a positron that is emitted from a radioactive source travels a certain distance called the positron range, after which it annihilates with an electron and emits a photon pair. A diagram showing this process is shown in Figure 1.2. The point of annihilation is not the same as the source position, thus limiting the resolution of the reconstructed PET image. For most popular PET isotopes such as F-18, C-11, etc., positron range is less than 1 mm, which is within the detector resolution of present day scanners. However, for some high energy isotopes such as Cu-60, Br-13, etc., positron range can be long (around 3mm), hence limiting their use as PET tracers, especially for small animal imaging [19]. Correcting this positron range effect might open up the use of new long range tracers that can possibly visualize physiological processes that cannot be imaged using present day tracers, or give a different perspective on those that can be monitored today. Additionally, with ongoing research in improving PET detector
technology [20] it is expected that detector resolution will improve in the future, making it important to address this fundamental limitation in reconstruction.

Previous work in positron range correction involved obtaining a positron range blur kernel from phantom experiments or Monte Carlo simulations, and using this kernel for correction. The approaches that have been used for positron range correction can be broadly classified into (i) methods that involve deconvolution of the range kernel from the reconstructed image or projection data at each iteration of filtered back projection [21] and (ii) methods that incorporate the range kernel into the system model and apply the correction during statistical reconstruction [22], [23] - [25]. The second approach was taken in [23] where the range kernel was assumed to be spatially invariant, and was truncated at the object boundary to reflect the assumption that a positron that escapes the object does not contribute to the sinogram counts. However, this method did not correct for blurring of the internal boundaries within the object and produced
ring artifacts at boundaries, where the assumption of spatial invariance of blur kernel is not valid. We apply the information theoretic anatomical priors to the positron range problem with the goal of utilizing the available anatomical information to correct for the positron range blurring effect. We employ the truncated homogeneous model in conjunction with the anatomical priors to correct for range blurring at the object as well as internal boundaries. We present results obtained through phantom simulations using the Cu-60 isotope (range 3.09 mm), and real data obtained by imaging mice injected with the Br-76 isotope (range 3.47 mm).

1.2 Non-rigid image registration

The second aspect of this work is related to image registration. Longitudinal and inter-subject scans are often performed in imaging studies to study changes in the anatomy and function of a subject over a period of time, or across populations. In longitudinal studies, changes may occur in subject posture, tissue growth, organ movement, etc. In inter-subject studies, anatomical variability across populations needs to be normalized for accurate analysis.

Image registration finds a mapping between images such that homologous features in the images have the same coordinates. The mapping that describes the relation between the images can be a low dimensional rigid or affine registration that is described by rotations, translations, and shear, or can be a high dimensional non-rigid registration. In longitudinal and inter-subject studies, the changes in anatomy and function mentioned above require non-rigid registration to align the images. Examples of imaging studies
that require non-rigid registration are shown in Figure 1.2 for mouse imaging. Several non-rigid registration algorithms have been developed, a review of which can be found in [26], [27]. A large subset of these algorithms use voxel based similarity metrics that measure similarity between the intensities of the images to be registered such as sum of squared difference (SSD), or some statistical measure of the intensities, such as cross correlation or MI.

MI based registration aims to find a deformation field that maximizes MI between the intensities of the images to be registered. Since MI between two images is a function of the distribution of intensities of the images rather than their actual values, it can be used to register multi-modal images. MI based registration has been successfully applied to rigid registration [13], and some approaches to non-rigid registration using MI or similar information theoretic measures have also been proposed [28]-[30]. It has been observed that though the existing non-rigid registration algorithms have been applied to brain imaging with good results, these methods do not perform well for small animal registration [31]-[33].
Small animal imaging is an important tool for pre-clinical studies. The registration of small animal images is challenging because of the presence of rigid structures like the skeleton within non-rigid soft tissue. In [31] and [32] mouse registration was performed using piece-wise rigid registration of anatomical structures, which were defined in a segmentation step prior to registration. In [33] a fully automated method was proposed for whole body registration, where they first aligned the skeleton using a point based method, after which they imposed stiffness constraints at the skeleton to align the whole body images using intensity based non rigid registration with mutual information as the similarity metric.

In this work, we describe two approaches to multi-modality non-rigid registration that can potentially be used for pre-clinical as well as clinical applications. First, we use MI as a similarity metric in a scale-space based framework similar to that proposed for PET reconstruction. The goal is to find a common mapping that aligns the fine structure that appears in the images at lower scales, as well as the global structure that remains in the images at higher scales. For small animal imaging, the skeletal structure is visible in the lower scales and is blurred out at the higher scales, while the global structure remains in the images at the higher scales. Hence this scale-space approach can potentially simultaneously align the skeleton as well as soft tissue. This multi-scale approach has the additional advantage of having a cost function that is less prone to local minima [27], and hence is able to perform large non-linear deformations more accurately than a hierarchical approach that aligns each scale hierarchically. We evaluate our scale-space method on a longitudinal mouse CT study by comparing it to a hierarchical approach. The results indicate that the scale-space approach is able to align both the skeleton and
soft tissue more accurately than the hierarchical approach. In our MI based scale-space registration, we used a non-parametric approach to estimate the unknown joint intensity distribution of the reference image (target) and the image to be registered (template) by Parzen windowing technique [34] to avoid making any assumptions about the joint distribution. However, we note that this non-parametric approach is sensitive to small changes in the images, and requires a large number of samples for accurate estimation. Additionally, we note that using a non-parametric approach to estimate the entire joint density based on the given images, in conjunction with estimation of the high dimensional non-rigid deformation field from the non-convex MI cost function could increase the total dimensionality of the problem, thus making the problem more ill-conditioned.

As an alternative approach, we use the log likelihood of the target given the template as a similarity metric, wherein the joint distribution of the target and template intensities is modeled by a Gaussian mixture model (GMM). The goal is to find a deformation field that maximizes the conditional probability of the target image given the deformed template image. Modeling the probability density using a GMM reduces the density estimation step to that of estimating the parameters of the GMM. We choose the number of components in the GMM from the joint histogram of the target and template images, and keep this number constant throughout the registration. This can hence be considered as a semi-parametric approach to density estimation. For images that have a few distinct regions of intensity such as brain or mouse CT images, the number of parameters to be estimated is small, and can be robustly estimated from the images.

Maximizing MI is closely related to maximizing the joint probability of the target and template images, or the conditional probability of the target given the template images.
An interpretation of MI as a special case of maximum likelihood estimation is given in [36]. In [37] a MAP framework for non-rigid registration is used wherein a conditional density estimate is computed using a Parzen-like density estimator, and used as the likelihood term. It was shown that under certain conditions, the optimum of this MAP objective function also maximizes the true MI. In [35] two different approaches to modeling the joint intensity distributions were explored - the Parzen density estimation, and the Gaussian mixture model (GMM). The distributions using these two approaches were estimated using a registered training set for the same modality as that of the images to be registered, and these estimated distributions were used to perform rigid registration by maximizing the joint probability between the target and template image intensities.

Our approach of using the log-likelihood of the target given the template in conjunction with a GMM can be viewed as a semi-parametric approximation to MI based registration. However, we expect our GMM based approach to perform better than the non-parametric MI based approach because of (1) reduced overall dimensionality of the registration problem through the parametrization of the density estimation step, and (2) more robust and accurate estimation of the joint density. We compare the performance of the conditional likelihood metric with GMM parametrization, with that of MI with non-parametric density estimation approach. We evaluate these methods using intra-modality brain MR images, as well as inter-subject, inter-modality mouse images. Through these datasets, we test the performance of our approach for clinical as well as pre-clinical imaging applications.
1.3 Contributions

The contributions of this dissertation are as follows:

1. Developed priors based on information theoretic measures of similarity to incorporate anatomical information in PET reconstruction. These priors take advantage of the available high resolution anatomical images to improve the reconstruction of PET images.

2. Used scale-space based feature vectors to incorporate spatial information into the global distribution based measures of MI and JE. We found that using these feature vectors improved the stability of the anatomical priors based on these measures.

3. Applied the information theoretic anatomical priors to positron range correction, a fundamental resolution limiting factor in PET.

4. Applied the scale-space feature vectors to MI based non-rigid mouse registration to simultaneously align fine skeletal as well as global soft tissue structure in the images.

5. Developed a non-rigid registration method that uses the log likelihood of the target given the deformed template image as a similarity metric wherein the distribution is modeled as a GMM. This approach reduces the dimensionality of the density estimation step, and shows improved robustness and accuracy compared to the non-parametric MI based approach.
1.4 Organization of the dissertation

Chapter 2 describes PET image reconstruction using information theoretic anatomical priors. It presents the details of reconstruction using mutual information and joint entropy based priors, and the simulation and clinical data results comparing the two metrics. The application of these priors to positron range correction is also described with phantom simulations and real data results. Chapter 3 describes MI based non-rigid mouse registration using the scale-space approach. Results are presented for a longitudinal CT study of a mouse. Chapter 4 describes the log-likelihood based non-rigid registration using a Gaussian mixture model for the probability densities. Evaluations using clinical and pre-clinical data are presented. Chapter 5 summarizes the work described in this dissertation and proposes future directions for this work.
Chapter 2

PET Image Reconstruction Using Information Theoretic Anatomical Priors

The uptake of positron emission tomographic (PET) tracers typically results in a spatial density that reflects the underlying anatomical morphology. This results in a strong correlation between the structure of the anatomical and functional images. The incorporation of anatomical information from co-registered MR/CT images into PET reconstruction algorithms can potentially improve the quality of low resolution PET images. This anatomical information is readily available from multimodality imaging equipment that is often used for acquiring data and can be incorporated into the PET reconstruction algorithm in a Bayesian framework through the use of priors.

We describe a non-parametric framework for incorporating anatomical information in PET image reconstruction through priors based on mutual information and joint entropy. Since both these information theoretic metrics are based on global distributions of the image intensities, spatial information can be incorporated by adding features that capture the local variation in image intensities. We define features based on scale space theory, which provides a framework for the analysis of images at different levels of detail.
It is based on generating a one parameter family of images by blurring the image with Gaussian kernels of increasing width (the scale parameter) and analyzing these blurred images to extract structural features [17]. We define the scale-space features for information theoretic priors as the original image, the image blurred at different scales, and the Laplacians of the blurred images. These features reflect the assumption that boundaries in the two images are similar, and that the image intensities follow similar distributions within boundaries. By analyzing the images at different scales, we aim to automatically emphasize stronger boundaries that delineate important anatomical structures, and attach less importance to internal detail and noise which is blurred out in the higher scale images.

We evaluate the performance of both MI and JE as priors through simulations as well as clinical data. The simulations explore both the ideal case of perfect agreement between the anatomical and functional images, as well as a more realistic case of the images having some differences. The simulations are based on brain imaging with $^{18}$FDG, which has uniform uptake in the three main anatomical regions of the brain, gray matter (GM), white matter (WM), and cerebrospinal fluid (CSF). We perform Monte Carlo simulations to evaluate quantitation accuracy in specific regions of interest, as well as overall. Additionally, we compare reconstructions of clinical data obtained from a PET scan using [F-18]Fallypride, which shows uptake mainly in the striatum region of the brain. This clinical data serves not only as a validation of the simulation results, but also explores the behavior of the information theoretic priors when the uptake is the same in regions corresponding to different anatomical image intensities. We present an efficient method of computing the priors and their derivatives, which reduces the complexity from
\(O(MN_s)\) to \(O(M \log M + N_s)\), where \(N_s\) is the number of voxels in the image and \(M\) is the number of points at which the distribution is computed. This makes these priors practical to use for 3D images where \(N_s\) is large.

We apply the information theoretic priors to a fundamental resolution limiting problem of PET, positron range. Positron range is the distance traveled by a positron before annihilation. For some high energy isotopes such as Cu-60, Br-13, etc, this distance can be long and causes a blurring in the reconstructed PET image, thus limiting its resolution \([19]\). In \([23]\) a truncated homogeneous system model was used to correct the positron range effect by modeling it as a spatially invariant blurring that is truncated at the object boundary to reflect the assumption that a positron that escapes the object does not contribute to the sinogram counts. We use the truncated homogeneous system model in conjunction with information theoretic priors to correct for positron range by taking advantage of anatomical information to improve the resolution of the reconstructed image. We present simulation results based on the Cu-60 isotope (range 3.07 mm) and real data results based on the Br-76 isotope (range 3.47 mm) and compare reconstructions without range correction, and with range correction in conjunction with quadratic and information theoretic priors.

2.1 Methods

2.1.1 MAP estimation using information theoretic priors

We represent the PET image by \(f = [f_1, f_2, \cdots, f_N]^T\), where \(f_i\) represents the activity at voxel index \(i\), and the co-registered anatomical image by \(a = [a_1, a_2, \cdots, a_N]^T\). Let
\( g \) denote the sinogram data, which is modeled as a set of independent Poisson random variables \( g_i, i = 1, 2, \cdots, K \). The maximum a posteriori estimate of \( f \) is given by

\[
\hat{f} = \arg \max_{f \geq 0} \frac{p(g|f)p(f)}{p(g)}, \tag{2.1}
\]

where \( p(f) \) is the prior on the image, and \( p(g|f) \) is the Poisson likelihood function given by,

\[
p(g|f) = \prod_{i=1}^{K} \frac{\exp\left(-\sum_{j=1}^{N} P_{ij} f_j \right) \left(\sum_{j=1}^{N} P_{ij} f_j \right)^{g_i} \frac{g_i!}{g_i!}}{g_i!} . \tag{2.2}
\]

\( P_{ij} \) represents the \( i, j^{th} \) element of the \( K \times N \) forward projection matrix \( \mathbf{P} \). Here, \( \mathbf{P} \) not only models the geometric mapping between the source and detector but also incorporates detector sensitivity, detector blurring, and attenuation effects [38].

The prior term \( p(f) \) can be used to incorporate a priori information about the PET image. We utilize information that is available through co-registered anatomical images, by defining the prior in terms of information theoretic measures of similarity between the anatomical and functional images. Through this framework, we aim to reconstruct a PET image that maximizes the likelihood of the data, while also being maximally similar in the information theoretic sense to the anatomical image.

To define the prior, we extract feature vectors that can be expected to be correlated in the PET and anatomical images. Let the \( N_s \) feature vectors extracted from the PET and anatomical images be represented as \( \mathbf{x}_i \) and \( \mathbf{y}_i \) respectively for \( i = 1, 2, \cdots, N_s \). These can be considered as independent realizations of the random vectors \( \mathbf{X} \) and \( \mathbf{Y} \). Let \( m \) be the number of features in each feature vector such that \( \mathbf{X} = [X_1, X_2, \cdots, X_m]^T \). If
$D(X, Y)$ is the information theoretic similarity metric that is defined between $X$ and $Y$, the prior is then defined as,

$$p(f) = \frac{1}{Z} \exp(\mu D(X, Y))$$

(2.3)

where $Z$ is a normalizing constant and $\mu$ is a positive hyperparameter.

Taking log of the posterior and dropping constants, the objective function $L(f)$ is given by,

$$L(f) = \sum_{i=1}^{K} - \sum_{j=1}^{N} P_{ij} f_j + g_i \log(\sum_{j=1}^{N} P_{ij} f_j) + \mu D(X, Y),$$

(2.4)

and the MAP estimate of $f$ is given by,

$$\hat{f} = \arg \max_{f \geq 0} L(f).$$

(2.5)

### 2.1.2 Information theoretic priors

We use mutual information and joint entropy as measures of similarity between the anatomical and functional images. Mutual information between two random variables $X$ and $Y$ with marginal distributions $p(x)$ and $p(y)$, and joint distribution $p(x, y)$ is defined as [39],

$$I(X, Y) = H(X) + H(Y) - H(X, Y),$$

(2.6)
where the entropy $H(X)$ is defined as

$$H(X) = -\int p(x) \log p(x) dx,$$  \hspace{1cm} (2.7)

and the joint entropy $H(X, Y)$ is given by

$$H(X, Y) = -\int p(x, y) \log p(x, y) dxdy.$$  \hspace{1cm} (2.8)

Mutual information can be interpreted as the reduction in uncertainty (entropy) of $X$ by the knowledge of $Y$ or vice versa. When $X$ and $Y$ are independent $I(X, Y)$ takes its minimum value of zero, and is maximized when the two random variables are maximally correlated. The distribution $p(x, y)$ that maximizes MI is sparse with localized peaks. This also corresponds to a reduction in the joint entropy $H(X, Y)$, from equations 2.6 and 2.8. We therefore define $D(X, Y)$ in terms of MI as,

$$D(X, Y) = I(X, Y),$$  \hspace{1cm} (2.9)

or in terms of JE as

$$D(X, Y) = -H(X, Y).$$  \hspace{1cm} (2.10)

For continuous random variables, the entropy $H(X)$ (called differential entropy) can go to $-\infty$ unlike the Shannon entropy of a discrete random variable which has a minimum value of zero. Differential joint entropy approaches $-\infty$ as the joint distribution
approaches a delta function. Mutual information, however, has a minimum value of zero for continuous and discrete random variables.

In Figure 2.1, the pdfs (estimated from joint histograms) of a reference image with some example reconstructed PET images are shown. The x axis of the pdfs shown denotes PET image intensities, and y axis denotes the reference image intensities. The examples shown are of a blurred reconstruction $X_{blur}$ and a noisy reconstruction $X_{noisy}$, of a true image $X_{true}$ that is identical in structure to a reference image $Y$. It can be seen that the joint histogram of the true image is more clustered than that of the blurred or noisy reconstructed image. The joint entropy of these pdfs is decreasing from left to right, $H(X_{blur}, Y) > H(X_{noisy}, Y) > H(X_{true}, Y)$. Similarly, their MI is increasing from left to right, $I(X_{blur}, Y) < I(X_{noisy}, Y) < I(X_{true}, Y)$. Thus, by using these priors, we can potentially reconstruct PET images that are less noisy, and have higher resolution due to the structural information that is available from the anatomical image.

Though both MI and JE have an optimum that is characterized by a clustered joint pdf, the additional marginal entropy terms in MI cause a difference in the set of solutions that can be obtained using these metrics. Examples of this are shown in Figure 2.2 through two test cases, the first where the functional image $X_{arbit}$ has the same structure as the reference image but with arbitrary intensities, and the second where the functional image $X_{hom}$ has two anatomically distinct regions combined into one homogeneous functional region. The joint entropy $H(X_{arbit}, Y) = H(X_{true}, Y) \approx H(X_{hom}, Y)$, whereas the mutual information $I(X_{hom}, Y) < I(X_{arbit}, Y) = I(X_{true}, Y)$. Thus JE can possibly have more than one anatomical region map to a homogeneous functional region without change in its value, while MI remains the same for all images that have the same structure as
Figure 2.1: Log of estimated joint probability distribution functions: Top: Anatomical image, Middle: Example reconstructed images (L to R): Blurred, Noisy, and true. Bottom: Estimated joint pdf of the images in the middle row and anatomical image.

the reference image. This causes JE to be more robust than MI to differences between the PET and anatomical images, where the anatomical image has structures that are not present in the PET image [16]. However, this property could also lead to distinct regions in the PET image that correspond to clusters that are close to each other in the joint density to be combined into one region. It should also be noted that since the optimization is with respect to image intensities, the marginal entropy term in MI may cause the PET image intensities to have high variance, thereby increasing the uncertainty or entropy in the image. Both these metrics measure similarity in distributions rather than actual values of intensities, and rely on the likelihood term to converge to the correct solution. We explore the properties and performance of these metrics through simulations and clinical data in the later sections.
2.1.3 Extraction of feature vectors

Mutual information and joint entropy between images are functions of the global distribution of intensities in the images. In PET images reconstructed using MI or JE where only intensity is taken as a feature, intensities in neighboring voxels of the PET image can be different as long as its global distribution matches that of the anatomical image. This can lead to voxels that have higher or lower intensities than their neighbors, which is undesirable since PET images are usually homogeneous within regions. We introduce spatial information into the information theoretic priors by defining feature vectors that capture the local morphology of the images. The feature vectors should be chosen such that they are correlated in the anatomical and functional images, and should reflect the morphology of the images. Since the information theoretic priors defined in this work are naturally insensitive to differences in intensity between the two images, the combination
Figure 2.3: Scale-space features of a coronal slice of an MR image of the brain

of local measures of image morphology and information theoretic measures should facilitate reconstruction of PET images whose structure is similar to that of the registered anatomical image.

Scale space theory provides a framework for the extraction of useful information from an image through a multi-scale representation of the image. In this approach, a family of images is generated from an image by blurring it with Gaussians of increasing width (scale parameter), thus generating images at decreasing levels of detail. Relevant features can then be extracted by analyzing this family of images rather than just the original image. Scale-space theory has a well defined mathematical framework that is described in [17].

We use this approach to define the feature vectors as:

1. The original image

2. The image blurred by a Gaussian kernel of width $\sigma_1$

3. Laplacian of the blurred image

By analyzing the image at two different scales (the original and image at scale $\sigma_1$), we are giving more emphasis to those boundaries that remain in the image at the higher
scale, and attach less importance to those that will be blurred out at the higher scale. A coronal slice of an MR image of the brain and its scale-space features are shown in Figure 2.3. More features could be added by using different sizes of kernels, but we restrict our analysis to two scales to limit complexity. We further make the simplifying assumption of independence between features so that

$$D(X, Y) = \sum_{i=1}^{m} D(X_i, Y_i).$$

Though the scale-space images are correlated, we choose to make this independence assumption to reduce the computational cost of the prior term.

### 2.1.4 Computation of prior term

The information theoretic priors defined in section 2.1.2 are expressed in terms of the pdfs of the images, which are not known. We use a non-parametric approach to estimate the joint density using the Parzen window method [34]. Given the samples $f_1, f_2, \ldots, f_{N_s}$ of the random variable $X$, the Parzen estimate $\hat{p}(x)$ of $p(x)$ is given by,

$$\hat{p}(x) = \frac{1}{N_s} \sum_{j=1}^{N_s} \phi\left(\frac{x - f_j}{\sigma}\right),$$

where $\phi(\frac{x}{\sigma})$ is a windowing function of width $\sigma$. The windowing function should integrate to one, and is commonly chosen to be a Gaussian of mean zero and standard deviation $\sigma$. For estimating the joint density between $X$ and $Y$ corresponding to the intensities of PET and anatomical images $f$ and $a$, we use a Gaussian window function with a diagonal
covariance matrix given by \( \Sigma = \begin{pmatrix} \sigma_x^2 & 0 \\ 0 & \sigma_y^2 \end{pmatrix} \). Thus, the joint density estimate is given by

\[
\hat{p}(x, y) = \frac{1}{N_s} \sum_{k=1}^{N_s} \phi\left( \frac{x - f_k}{\sigma_x} \right) \phi\left( \frac{y - a_k}{\sigma_y} \right).
\]  

(2.13)

The variance of the Parzen window is taken as a design parameter. It has been shown in [34] that as \( N_s \to \infty \), the Parzen estimate converges to the true pdf in the mean squared sense. For PET image reconstruction (especially for the 3D case) \( N_s \) is large, so it can be assumed that there are sufficient number of samples to give an accurate estimate of the pdf. Hence, we replace \( p(x, y) \) with \( \hat{p}(x, y) \) in equations 2.6 or 2.8 to compute the prior term.

2.1.5 Computation of the MAP estimate

The objective function given in Equation 2.4 can be iteratively optimized by a preconditioned conjugate gradient algorithm with an Armijo line search technique [40]. Let the estimate of the PET image at iteration \( k \) be \( \hat{f}^{(k)} \), the gradient vector be \( \mathbf{d}^{(k)} = \nabla L(f)|_{f=\hat{f}^{(k)}} \), the preconditioner be \( \mathbf{C}^{(k)} \), the conjugate gradient direction vector be \( \mathbf{b}^{(k)} \), and the step size computed from the line search be \( \alpha^{(k)} \). The PCG update equations are given by

\[
\hat{f}^{(k+1)} = \hat{f}^{(k)} + \alpha^{(k)} \mathbf{b}^{(k)} \]  

(2.14)

\[
\mathbf{b}^{(k)} = \mathbf{C}^{(k)} \mathbf{d}^{(k)} + \beta^{(k-1)} \mathbf{b}^{(k-1)} \]  

(2.15)

\[
\beta^{(k-1)} = \frac{(\mathbf{d}^{(k)} - \mathbf{d}^{(k-1)})^T \mathbf{b}^{(k)}}{\mathbf{d}^{(k-1)^T} \mathbf{b}^{(k-1)}}. \]  

(2.16)
The algorithm is initialized by $b^{(0)} = C^{(0)} d^0$. We use the EM-ML preconditioner, which is defined by $C^{(k)} = \text{diag}(\frac{f^{(k)}}{\sum_i f^{(k)}_i})$.

The $k^{th}$ element of the gradient vector $d$ can be computed from Equation 2.4 as,

$$\frac{\partial L(f)}{\partial f_k} = \sum_{i=1}^{N} (-P_{ik} + \frac{g_i}{\sum_{j=1}^{N} P_{ij} f_j}) + \mu \frac{\partial D(X, Y)}{\partial f_k}.$$  (2.17)

To compute the derivative of $D(X, Y)$, we first define the derivative of marginal and joint entropy terms corresponding to intensity of the original image. Replacing the integration in Equation 2.7 with its numerical approximation, the derivative of marginal entropy $H(X)$ with respect to $f_k$ is given by

$$\frac{\partial H(X)}{\partial f_k} = \frac{\Delta x}{N_s} \sum_{i=1}^{M} (1 + \log p(x_i)) \phi'(\frac{x_i - f_k}{\sigma_x})$$  (2.18)

where $\phi'(\frac{x_i - f_k}{\sigma_x}) = \phi(\frac{x_i - f_k}{\sigma_x})(\frac{x_i - f_k}{\sigma_x^2})$, and $M$ is the number of points at which the pdf is computed. Similarly, the gradient of joint entropy is

$$\frac{\partial H(X, Y)}{\partial f_k} = -\frac{\Delta x \Delta y}{N_s} \sum_{i,j=1}^{M} (1 + \log p(x_i, y_j)) \frac{\partial p(x_i, y_j)}{\partial f_k}$$

$$= -\frac{\Delta x \Delta y}{N_s} \sum_{i,j=1}^{M} (1 + \log p(x_i, y_j)) \times$$

$$\phi(\frac{y_j - a_k}{\sigma_y}) \phi'(\frac{x_i - f_k}{\sigma_x}).$$  (2.19)

The gradient of the MI prior is then

$$\frac{\partial I(X, Y)}{\partial f_k} = \frac{\partial H(X)}{\partial f_k} - \frac{\partial H(X, Y)}{\partial f_k}.$$  (2.20)
The gradients corresponding to the other scale-space features can be computed easily from these equations since the Laplacian and blurring are linear operations on \( f \) and \( a \). For an interpretation of these gradients through an alternative formulation, please refer to the Appendix.

We perform an approximate line search using the Armijo rule since it does not require the computation of the second derivative of \( L(f) \), which is computationally expensive for the information theoretic priors. Whenever the non-negativity constraint is violated, we use a bent line search technique by projecting the estimate from the first line search (with negative values) onto the constraint set, and computing a second line search between the current estimate and the projected estimate. The objective function is a non-convex function of \( f \), so optimization of the function using gradient based techniques require a good initial estimate to converge to the correct solution.

### 2.1.6 Efficient computation of entropy and its gradient

The computation of the Parzen window estimate \( p(x) \) at \( M \) locations using \( N_s \) samples is \( O(MN_s) \). In addition, the gradient in Equation 2.18 is also of complexity \( O(MN_s) \), since for each \( f_k \), we compute a summation of \( M \) multiplications. This can be expensive for large \( N_s \), which is the case in PET reconstruction. We take an approach similar to [41]
to compute the entropy measures as well as their gradients efficiently through the use of FFTs. The Parzen window estimate in Equation 2.12 can be rewritten as a convolution

\[
p(x) = \frac{1}{N_s} \sum_{j=1}^{N_s} \int \phi \left( \frac{x - s}{\sigma} \right) \delta(s - f_j) ds
\]

\[
= \phi \left( \frac{x}{\sigma} \right) \ast \frac{1}{N_s} \sum_{j=1}^{N_s} \delta(s - f_j). \tag{2.21}
\]

Here \( x \) is at continuous values, and the impulse train \( h(x) = \frac{1}{N_s} \sum_{j=1}^{N_s} \delta(x - f_j) \) has impulses at non-equispaced locations.

Efficient fast Fourier transforms can be used to compute convolution on a uniform grid. To take advantage of these efficient implementations, we interpolate the continuous intensity values \( f_j \) onto a uniform grid with equidistant discrete bin locations \( \tilde{x}_j \), where \( j = 1, 2, \cdots, M \), with spacing \( \Delta \tilde{x} \). Let the image with the quantized intensity values be \( \tilde{f} \). The impulses in \( h(x) \) are now replaced by triangular interpolation functions thus giving

\[
h(\tilde{x}) = \frac{1}{N_s} \sum_{j=1}^{N_s} \wedge(\tilde{x} - f_j), \tag{2.22}
\]

where

\[
\wedge(u) = \begin{cases} 1 - |u|, & \text{if } |u| < \Delta \tilde{x} \\ 0, & \text{otherwise} \end{cases} \tag{2.23}
\]
Thus each continuous sample \( f_j \) contributes to the bin it is closest to, as well as its neighboring bins. The Parzen estimate at the equispaced locations is given by

\[
p(\bar{x}) = \phi\left(\frac{\bar{x}}{\sigma}\right) \ast h(\bar{x}).
\] (2.24)

This convolution can now be performed using fast Fourier transforms with complexity \( O(M \log M) \). We can then compute the approximate marginal entropy \( H(\tilde{X}) \) by replacing \( p(x) \) with \( p(\bar{x}) \) in Equation 2.7.

The gradient equation in Equation 2.18 also has a convolution structure, and can be rewritten as

\[
\frac{\partial H(X)}{\partial f_k} = -\frac{\Delta x}{N_s} \left(1 + \log p(x)\right) \ast \phi'\left(-\frac{\bar{x}}{\sigma}\right) \bigg|_{\bar{x} = f_k}.
\] (2.25)

If we replace \( p(x) \) in the equation with \( p(\bar{x}) \), we can compute the gradient corresponding to the binned intensity values \( \tilde{f}_k \), \( k = 1, 2, \cdots, M \) using convolution with FFTs. We represent this binned gradient as \( \frac{\partial H(\tilde{X})}{\partial \tilde{f}_j} \). To retrieve the gradient with respect to original intensity values, we use similar interpolation as for binning giving,

\[
\frac{\partial H(X)}{\partial f_k} = -\sum_{j=1}^{M} \wedge (f_k - \tilde{f}_j) \frac{\partial H(\tilde{X})}{\partial \tilde{f}_j}.
\] (2.26)

This means we take the gradients at neighboring bins and interpolate to get the true gradient. Thus the complexity of computing the binned gradient is still \( O(M \log M) \) followed by interpolation which is of complexity \( O(N_s) \). Additionally, the bin locations corresponding to each \( f_k \) can be stored in the pdf estimation step, and retrieved in the gradient estimation step, thus giving further computational savings. This method
can be extended to the 2D case by using bilinear interpolation instead of the triangular windowing function to compute the binned joint density and to interpolate the gradient at the binned locations to the continuous intensity values.

We compare the Parzen window method with the FFT based method in terms of run times and accuracy of the marginal pdf and gradient estimates. For both methods we compute the pdf \( p(\tilde{x}) \) at the same locations \( \tilde{x}_i, i = 1, 2, \cdots, M \). To compute the Parzen gradient, we expressed Equation 2.18 as the inner product of the vectors given by

\[
L_p = -\frac{\Delta x}{N_s} [(1 + \log(p(x_1)), 1 + \log(p(x_2)), \cdots, 1 + \log(p(x_M))]^T
\]

\[
\Phi_k = \left[ \phi'(\frac{x_1 - f_k}{\sigma_x}), \phi'(\frac{x_2 - f_k}{\sigma_x}), \cdots, \phi'(\frac{x_M - f_k}{\sigma_x}) \right]^T.
\] (2.27)

This vectorized form enables us to use efficient linear algebra libraries for evaluating the summation for each \( f_k \).

The run-times for the combined computation of pdf \( p(x) \) and the gradient \( \nabla H(X) \) using the Parzen and FFT based methods are shown in Table. 2.1 for different sizes of image. The reported run-times were obtained on an AMD Opteron 870 cluster with 8 dual core CPUs, using a single processor. The estimated pdfs and gradients using the two methods are shown in Figure 2.4. The norm of the difference between the gradients of the two methods, normalized by the Parzen gradient was less than 1% for all the sizes of image considered. Since the main computation cost for the FFT based method is the FFT based convolutions that depend on \( M \) rather than image size, we see large speed-up factors for 3D images compared to the Parzen method for which the complexity increases linearly with image size.
Figure 2.4: Comparison of Parzen and FFT-based methods: a) Image for which pdf and gradients were estimated, b) pdf estimates using $M = 351$, c) gradient of marginal entropy using the Parzen method (left) and the FFT-based method (right). The normalized difference in the shown gradient images is 0.14%.

Table 2.1: Run-times for combined computation of pdf and gradient for different image sizes, for the Parzen and FFT based methods

<table>
<thead>
<tr>
<th>Image Size (No. of Bins)</th>
<th>Parzen (sec)</th>
<th>FFT-based (sec)</th>
<th>Speed-up Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>$128 \times 128(M = 351)$</td>
<td>1.01</td>
<td>0.11</td>
<td>9.1630</td>
</tr>
<tr>
<td>$128 \times 128 \times 111(M = 351)$</td>
<td>117.44</td>
<td>0.88</td>
<td>133.63</td>
</tr>
<tr>
<td>$256 \times 256 \times 111(M = 351)$</td>
<td>470.64</td>
<td>3.45</td>
<td>136.41</td>
</tr>
</tbody>
</table>
2.2 Results

We evaluate the performance of MI and JE priors through a) simulations that model the best case scenario of the anatomical image having identical structure as the PET image, b) simulations of a more realistic case where there are some differences between the MR and PET regions, and c) clinical data obtained from brain imaging.

We consider four different information theoretic priors:

1. JE-Intensity : Joint entropy between the intensities of anatomical and PET images
2. MI-Intensity : Mutual information between the intensities of anatomical and PET images
3. JE-scale : Joint entropy between scale-space feature vectors (original image, image blurred by Gaussian of width $\sigma_1$, and Laplacian of blurred image) of the anatomical and PET images
4. MI-scale : Mutual information between scale-space feature vectors of anatomical and PET images

We compare the performance of these priors relative to each other, and to a quadratic prior (QP), which does not use any anatomical information. QP penalizes large differences in intensities within a neighborhood $\eta_i$ defined by 8 nearest neighbors of each voxel $i$, and is given by,

$$p(f) = -\beta \sum_{i=1}^{N_s} \sum_{j \in \eta_i} \frac{(f_i - f_j)^2}{\kappa_{ij}}$$

(2.28)

where $\kappa_{ij}$ is a weight assigned to each voxel pair $f_i$, and $f_j$. 

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The weight on the prior is controlled by varying the hyperparameter $\mu$ for the information theoretic priors, and $\beta$ for the QP prior.

### 2.2.1 Simulation Results

The simulations are based on the $^{18}$FDG tracer that shows a uniform uptake in the gray matter (GM), white matter (WM), and cerebrospinal fluid (CSF) regions of a normal brain. For all the information theoretic priors considered, we computed the densities at bin centers that we keep constant throughout the optimization, the range of which we set to 2.5 times the range of intensities in the initial image. We chose the Parzen standard deviation to be about 5% of the range of the initial image. To form the scale-space features, we used a Gaussian kernel of standard deviation $\sigma_1 = 2.0$ pixels to generate the blurred image, and a $3 \times 3$ Laplacian kernel given by

$$
\begin{bmatrix}
0 & 1 & 0 \\
1 & -4 & 1 \\
0 & 1 & 0
\end{bmatrix}.
$$

#### 2.2.1.1 Anatomical and PET images with identical structure

We used a $128 \times 128$ slice of the Hoffman brain phantom as our functional image and scaled the three different regions (GM, WM, and CSF) differently to generate our anatomical image. Both images are shown in Figure 2.5. The simulations are based on a single ring of the microPET Focus 220 scanner, for which the sinogram dimensions are $288 \times 252$. The sinogram data had approximately 300,000 counts, corresponding to 30 counts per line of response (LOR). We reconstructed all images using 30 iterations of PCG algorithm, initialized by the image reconstructed after 2 iterations of ordered subset expectation maximization (OSEM) algorithm using 6 subsets. The reconstructed images using QP
and the four information theoretic priors are shown in Figure 2.6 along with the true image. The images shown for each prior are at values of hyperparameter that gave the least error between true and reconstructed images for that prior. The normalized error values as a function of iteration are shown in Figure 2.7. To evaluate the performance of these priors in quantifying uptake, we perform Monte Carlo simulations. Bias and variance of the estimated image were computed from 25 datasets, and are shown in Figure 2.8.

It can be seen that the images reconstructed using information theoretic priors have less overall error, sharper boundaries, and less noise than that of QP. MI-Intensity prior produces images that follow the structure of the anatomical image, but have several high intensity pixels distributed within the GM region. This is also reflected in the Monte Carlo results, where the MI-Intensity prior has high variance in the GM region, and less bias along the boundaries compared to QP. Adding spatial information through the MI-scale prior reduces the occurrence of these high intensity pixels, and hence decreases the variance in the estimate and also the overall reconstruction error. The JE-Intensity prior reconstructs a near perfect image, with normalized error of about 2% with respect to the true image. The bias and variance are both close to zero, except for a few pixels. This
Figure 2.6: True and reconstructed images: Top: True PET image (left), QP reconstruction with $\beta = 0.5$ (right). Middle: MI reconstruction for $\mu = 10000$ (left), JE reconstruction for $\mu = 5000$ (right). Bottom: MI-scale reconstruction for $\mu = 2500$ (left), JE-scale reconstruction for $\mu = 2500$.

Figure 2.7: Normalized error vs. iteration for best value of $\mu$ and $\beta$ of the information theoretic and QP priors respectively.
dataset illustrates the best case scenario for the JE prior, since the images are piecewise constant and the JE prior prefers piecewise homogeneous solutions. The JE-scale prior gives biased intensity values, and some smoothing along the boundaries. However, for a more realistic case where the images are not piecewise constant, we expect the scale-space features to help the JE reconstruction by providing local spatial information.

This simulation should also be the best case scenario for the MI prior, since the two images are identical except in the values of intensity, to which it is robust. However, we see that the MI priors do not reconstruct an image close to the true image. This can be explained from the joint pdfs of the anatomical image and the intensity based MI and JE reconstructions shown in Figure 2.6. These pdfs are shown in Figure 2.9, and the variation of the terms in the MI and JE priors as a function of iteration number are shown in Figure 2.10. For the same initial joint density, MI is optimized by initially increasing the marginal entropy term by increasing the variance in the intensities of the GM region, while decreasing the joint entropy by clustering the high intensity pixels together as isolated peaks in the joint histogram. This caused MI to converge to a local minimum in which the isolated peaks manifested as high intensity pixels that were distributed throughout the GM. On the other hand, minimizing the JE term alone gave a steeper decrease in JE and lower final value of JE than that of MI. However, even the JE prior has an isolated peak in its joint pdf, associated with the background/CSF of the anatomical image. Introduction of the scale-space features helps the priors avoid these local minima by adding terms that penalize the formation of these isolated peaks.
Figure 2.8: Results of Monte Carlo simulations for (L to R) QP ($\beta = 0.5$), MI ($\mu = 2500$), JE($\mu = 5000$), MI-scale($\mu = 2500$), JE-scale($\mu = 2500$)

Figure 2.9: Joint pdfs of the anatomical image and a) true PET image, b) OSEM estimate used for initialization, c) image reconstructed using MI-Intensity prior, d) Image reconstructed using JE-Intensity prior
2.2.1.2 Anatomical and PET images with differences

To simulate a more realistic scenario, we used an MR image of the brain and its corresponding manually segmented image from the IBSR dataset provided by the Center for Morphometric Analysis at Massachusetts General Hospital and available at the website http://www.cma.mgh.harvard.edu/ibsr/. We resized the MR image to size $256 \times 256 \times 111$ and generated the PET image from the segmented MR image by setting the activity in all voxels corresponding to GM as four times that corresponding to the WM, and CSF activity as zero. A coronal slice of both images is shown in Figure 2.11. Though this PET image was generated from the MR image, its structure is not identical to the MR image since the labels assign some non-homogeneous anatomical regions to a homogeneous functional region. The 3D simulations are based on a 55 ring Biograph scanner with sinogram dimensions $336 \times 336 \times 559$. The datasets simulated each had approximately 26 counts per LOR. All images were reconstructed by initializing with 2 iterations of OSEM with 21 subsets, followed by 20 iterations of MAP.
Reconstructed images using the QP and information theoretic priors are shown along with the true image in Figure 2.12. Normalized error between true and reconstructed images are shown in Figure 2.13 for the value of hyperparameter that gave the least error for each prior. Bias and standard deviation images computed from 25 datasets are shown in Figure 2.14 for the values of hyperparameter that gave the least overall error. We compute the normalized bias and variance in mean uptake in regions of interest (ROIs) by,

\[ Bias = \frac{1}{N_d} \frac{\sum_{i=1}^{N_d} \bar{\hat{f}}_{ROI} - \bar{f}_{ROI}}{\bar{f}_{ROI}}, \]

\[ SD = \sqrt{\frac{\sum_{i=1}^{N_d} (\bar{\hat{f}}_{ROI} - \frac{1}{N_d} \sum_{i=1}^{N_d} \bar{\hat{f}}_{ROI})^2}{\bar{f}_{ROI}}}, \]  

(2.29)

where \( \bar{f}_{ROI} \) is the mean true activity in the region of interest, \( \bar{\hat{f}}_{ROI} \) is the mean activity in ROI of the estimated image from the \( i^{th} \) dataset, and \( N_d \) is the number of datasets.

We draw ROIs in the thalamus, caudate, putamen, and WM, and the plots of bias versus
Figure 2.12: True and reconstructed images: Top: True PET image (left), QP reconstruction with $\beta = 0.5$ (right). Middle: MI reconstruction for $\mu = 1e5$ (left), JE reconstruction for $\mu = 150000$, Bottom: MI-scale reconstruction for $\mu = 1e5$ (left), JE-scale reconstruction for $\mu = 1e5$.

standard deviation as the hyperparameter is varied are shown in Figure 2.15 for these regions.

The reconstructed images using the information theoretic priors show higher contrast than that of QP, with the JE priors giving the best performance in terms of overall error metric. MI-Intensity prior has isolated voxels with high intensity, and high variance in the GM as expected. The MI-scale prior reduces the occurrence of these hot spots, and the variance in the GM. However, the overall error for the MI priors is now higher than that of QP, possibly both because of the differences in the anatomical and PET images, and
the higher ratio of volume of GM to WM compared to the Hoffman phantom simulations (2.15 for this dataset compared to 0.7 for Hoffman phantom). In this realistic simulation, the JE-Intensity prior has some isolated hot and cold spots within regions of homogeneous activity, which is undesirable for any application of PET. The JE-scale prior reduces these inhomogeneities and the variance in the estimate of the image relative to the JE-Intensity prior.

The ROI curves shown in Figure 2.15 indicate that the information theoretic priors do not offer a traditional trade-off between bias and variance. These priors are non-linear functions of the image that is being estimated, and exhibit a non-monotonic variation as the hyperparameter is changed. However, the ROI plots still offer an insight into the relative performance of these priors with respect to QP and to each other. In the WM, all the information theoretic priors have lesser bias and variance compared to QP. In all the GM regions considered (putamen, caudate, thalamus), the JE-Intensity prior gave the least bias and variance compared to all the priors considered, but has an unpredictable

<table>
<thead>
<tr>
<th>Iteration number</th>
<th>Normalized error</th>
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<tbody>
<tr>
<td></td>
<td>QP</td>
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<td>MI</td>
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<td>MI-scale</td>
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<td>JE-scale</td>
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Figure 2.13: Normalized error as a function of iteration.
Figure 2.14: Coronal slice of bias and SD images: Top (L to R): QP ($\beta = 0.5$), MI-Intensity($\mu = 1e5$), JE-Intensity($\mu = 150000$), Bottom (L to R): MI-scale($\mu = 1e5$), JE-scale ($\mu = 1e5$).
behavior as the hyperparameter is varied. The JE-scale prior is better behaved, showing a more stable decrease in variance with corresponding increase in bias.

Overall, we see that JE based priors outperform QP and MI based priors, and that introducing spatial information through the scale-space features mitigates the instability produced due to the lack of spatial correlations inherent in these global distribution based information theoretic metrics. We therefore prefer the JE-scale prior despite its increased bias compared to JE-Intensity prior.
2.2.2 Clinical data

PET data was acquired on a Siemens Biograph True Point HD scanner from a Parkinsons disease patient who was injected with [F-18]Fallypride. This tracer has a high affinity to D2 receptors, which have a high density in the striatal regions of the brain. PET images using this tracer show uptake mainly in the caudate and putamen subcortical nuclei (jointly referred to as the striatum), and therefore this data would serve to both validate the simulation results, and to investigate the behavior of the information theoretic priors when the PET image intensities do not uniformly follow the variation in the anatomical image intensities.

A T1 weighted MRI scan of the subject was performed on the same subject and is used as the anatomical image for reconstruction. The MR image was registered to a MAP with QP reconstructed image using MI based rigid registration in RView [42]. The QP image along with the co-registered MR image is shown in Figure 2.16. Histogram equalization was then performed on the MR image to improve contrast.

We initialized the MAP algorithm for all priors by the result of 2 iterations of OSEM with 21 subsets, and ran 20 iterations of MAP. We used similar parameters for the bin widths and Parzen variance as for the simulations. However, for the JE-scale prior, we used a scale parameter of $\sigma_1 = 3.0$ pixels, and dropped the Laplacian term. The Laplacian of these PET images is non-zero in few voxels (mainly in the striatal regions), which produces a joint density of the Laplacian images consisting of a large peak around zero, and a few peaks corresponding to the edges. Since the JE metric can produce homogeneous solutions, it tends towards a Laplacian joint density which has one large
peak around zero, producing undesirable results. The marginal entropy term in the MI-scale prior prevents this from occur.

The images reconstructed using the different priors are shown in Figure 2.17 for the slice corresponding to that of the anatomical image shown in Figure 2.16. The QP and JE-scale reconstructions are overlaid on the anatomical image and the coronal, sagittal, and transaxial views are shown in Figure 2.18. It can be seen that for QP, the activity is often spread outside the anatomical region to which it belongs, while this spreading is reduced for the priors using anatomical information.

We quantify the performance of these priors in terms of their contrast recovery coefficient (CRC) as a function of noise. We define CRC between the uptake in a region of the striatum and a region in the WM close to it as,

$$CRC = \frac{\bar{f}_{Str}}{\bar{f}_{WM}},$$  \hspace{1cm} (2.30)

where $\bar{f}_{Str}$ is the mean activity in a region defined in the striatum, and $\bar{f}_{WM}$ is the mean activity in a WM region. We considered ROIs in the caudate region of the striatum, and the WM that separates the caudate from the putamen on 11 contiguous slices of the brain. One transaxial slice showing the ROIs for CRC computation are shown in Figure 2.19(a), with red and blue representing the caudate and WM ROIs. We computed the noise variance from regions in the brain where we do not expect uptake of tracer, such as the white matter in the posterior of the brain. The noise variance ROIs for one transaxial slice is shown in Figure 2.19(b). The CRC vs noise plots are shown in Figure 2.20.
These results demonstrate that the information theoretic priors can reconstruct high contrast images that take advantage of anatomical image information, in a real world scenario where the anatomical and functional images are independently acquired and then registered to each other. Additionally, the results indicate that these priors can be used even for tracers that bind non uniformly within regions of similar intensities in the anatomical image (e.g., GM, WM, CSF for the brain), as long as the uptake is homogeneous within some clearly delineated anatomical structure that is large enough to form a prominent peak in the joint density.
Figure 2.18: Overlay of PET reconstruction over co-registered MR image for QP (top) and JE-scale (bottom) priors.

Figure 2.19: Regions of interest for computing CRC vs noise curves: a)ROIs in caudate (red) and WM separating it from putamen (blue), (b) ROIs for computing noise variance
The CRC curves show that the information theoretic priors reconstructed images with higher contrast than that of QP. While QP has a decreasing contrast and noise variance as the hyperparameter is increased, the MI and JE based priors have increasing contrast as \( \mu \) is increased. An increasing \( \mu \) would give larger weight on the prior, thus causing the joint density to be more clustered, which would correspond to a higher contrast. We see that consistent to our simulation results, the scale-space priors have reduced noise variance compared to the intensity based priors, without appreciable loss in contrast.
The JE-scale prior gives the best contrast to noise performance among all the priors considered.

However, we note that the behavior of the information theoretic priors is unpredictable at high values of $\mu$, where the prior dominates the likelihood term. MI tends to produce images that spuriously follow the anatomical regions, whereas JE prefers a piecewise homogeneous image, and may join regions that are close to each other in intensity. This is illustrated in Figure 2.21, which shows the MI-scale and JE-scale reconstructions for high values of $\mu$.

This instability makes the choice of hyperparameter especially important for these priors. We have observed that the knee of the L curve between the log likelihood and log prior terms yields reasonable values of $\mu$, but for clinical use, these results should always be verified with the OSEM or MAP with QP results to ensure that no misleading artifacts are produced.

### 2.3 Application of Information Theoretic Priors to Positron Range Correction

In this section, we apply JE with scale-space features as a prior in conjunction with the truncated homogeneous system model for positron range correction. The motivation for using JE-scale priors for this application is to take advantage of anatomical information to correct for the positron range effect where the truncated homogeneous system model is insufficient. Using these priors can reduce blurring due to positron range outside regions where uptake is expected, by using a co-registered anatomical image that delineates these
regions. Thus we aim to correct for positron range such that the PET image not only fits the data, but also is most similar to the anatomical image in the information theoretic sense.

The likelihood $p(g|f)$ in Equation 2.1 is a Poisson likelihood function with mean $Pf$, where $P$ is the system matrix that models the mapping from the source voxels to the detectors. In [43] a factored system model is used, where this $P$ matrix incorporates detector blurring, detector sensitivity, and attenuation effects. Positron range effect can also be included in this factored system model through a matrix $P_{\text{range}}$ and the complete system model can be written as

$$P = P_{\text{sens}}P_{\text{blur}}P_{\text{attn}}P_{\text{geom}}P_{\text{range}},$$

(2.31)

where $P_{\text{geom}}$ represents the geometric projection from source to detector and $P_{\text{attn}}, P_{\text{blur}}, P_{\text{sens}}$, and $P_{\text{range}}$ model the attenuation, detector blurring, sensitivity, and positron range effects respectively. We use the truncated homogeneous model of [23] which can be represented as

$$P_{\text{range}} = TP_{\text{hom}}T,$$

(2.32)

where $P_{\text{hom}}$ is a convolution matrix representing the spatially invariant blurring assumed within the subject and $T$ is a diagonal truncation matrix that truncates at the object boundary.

We use the $P$ matrix as defined in Equation 2.31 in the likelihood function to obtain the objective function as defined in Equation 2.4, with the prior $D(X, Y)$ being the JE-scale prior defined as $-H(X, Y)$, where $X$ and $Y$ are the scale-space features.
2.3.1 Positron Range Simulations

We used a 3D Hoffman brain phantom resampled to size $128 \times 128 \times 63$ as our functional image and scaled the three different regions (gray matter, white matter, and CSF) differently to generate our anatomical image. A truncation mask for the homogeneous truncated model was generated from the anatomical image, one transaxial slice of which is shown along with the PET and anatomical images in Figure 2.22. For simulating positron range blurring, we used the range kernel obtained from Monte Carlo simulations of Cu-60 isotope in water, which has a range of 3.08mm \cite{19} and is shown in Figure 2.23. The same kernel is used for generating the data and for reconstruction. The simulations are based on a Concorde R4 scanner, and the sinogram data obtained had 15 counts per line of response. We used 30 PCG iterations to reconstruct all images at a voxel size of $0.4 \times 0.4 \times 1.211$ mm.

We compare three methods for reconstruction:

1. NRC-QP: No range correction and quadratic prior.

2. RC-QP: Range correction using the truncated homogeneous model, and quadratic prior.
Figure 2.23: Positron range kernel for Cu-60 isotope vs distance (mm)

Figure 2.24: Reconstructed images (L to R): No range correction with QP (NRC-QP), range correction with quadratic prior (RC-QP) with $\mu = 0.01$, range correction with JE prior (RC-JE), scale $\sigma = 0.5$ voxels and $\mu = 5\epsilon4$.

3. RC-JE: Range correction using truncated homogeneous model, and JE-scale prior.

The reconstructed images using NRC-QP, RC-QP, and RC-JE with scale $\sigma = 0.5$ voxels are shown in Figure 2.24. A profile drawn through the reconstructed images is shown in Figure 2.25.

It can be seen that without range correction, the reconstructed image is blurred and the boundaries between GM/WM and WM/CSF are not clearly delineated. The RC-QP images demonstrate improvement compared to the NRC-QP reconstruction, but we see
Figure 2.25: Profile drawn through true and reconstructed images

that the activity is overestimated close to the object boundary, and the internal structures are still blurred. The RC-JE reconstruction is able to take advantage of the anatomical information to provide sharper boundary definition even for the smaller internal structures in the image. The overall normalized quantitation error between true and reconstructed images is 0.6467 for NRC-QP, 0.5232 for RC-QP, and 0.1498 for RC-JE. Thus the JE prior based reconstruction yields images with improved boundary definition and lower quantitation error.

2.3.2 Real data results

PET data was acquired on a Focus 220 scanner from two mice that were injected with Br-76 isotope. This isotope has a positron range of 3.47 mm. A CT image of these mice was acquired on a microCT scanner at a resolution of 0.197 × 0.197 × 0.197 mm. We reconstructed a PET image of size 256 × 256 × 95 at a resolution of 0.4 × 0.4 × 0.8 mm, without range correction and using a quadratic prior. The CT image was then registered to the reconstructed PET image, and used as the anatomical image for the
range corrected reconstruction. A truncation mask is obtained from the CT image for truncated homogeneous range correction. The CT image registered to the PET image is shown in Figure 2.26 along with the truncation mask. The positron range kernel for Br76 was obtained using Monte Carlo simulations and is shown in Figure 2.27.

We reconstruct PET images from the data using NRC-QP, RC-QP, and RC-JE methods. We initialized the MAP algorithm for all three methods by the result of 2 iterations of OSEM with 21 subsets, and ran 50 iterations of MAP. For the RC-JE method we used the JE-scale prior with a scale parameter of 2 pixels. A coronal view of the images reconstructed using these methods is shown in Figure 2.28 along with their overlay on the CT image.

It appears from these figures that the range corrected images are less blurred and have better boundary definition than the NRC-QP image. The RC-QP image produces ring artifacts around region boundaries by overestimating the activity within the region and compensating it by undershoots around the region. The RC-JE prior takes advantage of the available anatomical information to produce apparently better boundary definition with activity that is contained within anatomical regions, and without the ring artifacts. These results are preliminary, but they are sufficiently encouraging to warrant further study of range correction using the approach described here.

### 2.4 Conclusions

We described a scale-space based framework for incorporating anatomical information into PET image reconstruction using information theoretic priors. Through idealized
Figure 2.26: (a) CT image registered to reconstructed PET image and (b) truncation mask obtained from the CT image for truncated homogeneous model of range correction.
Figure 2.27: Positron range kernel for Br-76.

Figure 2.28: Reconstructed images: Top row: Coronal slice of PET images reconstructed using NRC-QP (left), RC-QP (middle), and RC-JE (right). Bottom row: Overlay on CT of the images shown in the top row.
simulations, realistic simulations, and clinical data, we evaluated the performance of mutual information and joint entropy as priors in PET reconstruction. We presented an efficient FFT based approach for the computation of these priors and their derivatives, making it practical to use these priors for 3D image reconstruction.

The reconstructed images using the information theoretic priors have higher contrast than those obtained using the quadratic prior, and look realistic without the mosaic-like appearance that results from anatomical priors that impose piecewise smoothness. Through simulation studies we showed that the marginal entropy term in MI tends to increase the variance in the estimate of the image. We noted that testing the information theoretic priors on piecewise constant images can give misleadingly accurate results while using only intensity as a feature, especially for JE, since it prefers piecewise constant images. The more realistic simulations that we performed show that using JE-Intensity is prone to isolated hot and cold spots, and varies unpredictably with the hyperparameter. It is therefore important to incorporate spatial correlations into these priors. The scale-space based approach described in this dissertation improved the behavior of the information theoretic priors, and the JE-scale prior gave better quantitation compared to QP while using only two additional features, the coarser scale image and its Laplacian. The clinical data results were consistent with the simulations, and show that these priors can be used even for tracers that do not have homogeneous uptake in regions corresponding to similar intensities in the anatomical image, provided the activity corresponds to a prominent anatomical region.

It should however be noted that both mutual information and joint entropy priors in PET reconstruction rely on the likelihood term to ensure convergence to an image with
the correct intensity. When the prior term dominated the objective function, MI tended to produce an image that had a structure identical to the anatomical image, whereas JE tended towards a piecewise homogeneous solution. This suggests that a weighting between zero (JE) and one (MI) on the marginal entropy term of mutual information could be used depending on the application.

We applied these information theoretic priors to a fundamental resolution limiting problem in PET, positron range. These priors were used in conjunction with a truncated homogeneous model for positron range that is incorporated into the system model. We performed simulations that modeled the best case scenario where the PET and anatomical images have identical structure, and the range kernel is the same for forward projection and reconstruction. The results of these simulations indicate that using the JE-scale prior with truncated homogeneous system model for range correction yielded images that have better boundary definition and lower quantitation error than those obtained using no range correction with QP or range correction with QP. We also performed reconstructions from real data obtained by imaging mice that were injected with a long range isotope. The results indicate that using the JE-scale prior along with a homogeneous system model incorporating positron range is able to produce reconstructions with apparently better boundary definition and without the ring artifacts that are produced by the RC-QP method at regions where the system model is not accurate. The positron range modeling used in these results is simple and does not consider the effects of spatially variant range resulting from differences in electron density within the imaging volume. However, it is promising that the JE based prior appears to perform well with experimental data, even with this imperfect range model.
This framework of information theoretic priors can also be applied to other functional imaging modalities where the data is severely ill-conditioned and the use of anatomical information may improve the accuracy of reconstruction. Promising results were obtained for diffuse optical tomography using this framework of information theoretic anatomical priors [44]. For PET, we expect these priors to be most useful in applications involving quantification of uptake in specific organs, where the uptake is within a region that is clearly discernible in the anatomical image.

2.5 Appendix: Alternative formulation to the JE prior

Joint entropy between two random variables $X$ and $Y$ is defined as

$$
H(X, Y) = -\int p(x, y) \log p(x, y) dx dy.
$$

$$
= -E(\log p(X, Y)), \quad (2.33)
$$

This expectation can be approximated by the sample mean. If the intensities of image $f$ and $a$ are realizations of $X$ and $Y$ respectively,

$$
H(X, Y) = -\frac{1}{N_s} \sum_{i=1}^{N_s} \log p(f_i, a_i) \quad (2.34)
$$

If we use Parzen windows to estimate $p$ as described in the chapter, then,

$$
H(X, Y) = -\frac{1}{N_s} \sum_{i=1}^{N_s} \log \left( \sum_{j=1}^{N_s} \phi \left( \frac{f_i - f_j}{\sigma_x}, \frac{a_i - a_j}{\sigma_y} \right) \right) \quad (2.35)
$$
The derivative of $H(X,Y)$ with respect to $f_m$ is then,

\[
\frac{\partial H(X,Y)}{\partial f_m} = -\frac{1}{N_s} \sum_{i=1, i \neq m}^{N_s} \frac{\phi'(\frac{f_i-f_m}{\sigma_x}) \phi'(\frac{a_i-a_m}{\sigma_y})}{p(f_i, a_i)} + \frac{1}{N_s} \sum_{i=1}^{N_s} \frac{\phi'(\frac{f_m-f_i}{\sigma_x}) \phi'(\frac{a_m-a_i}{\sigma_y})}{p(f_m, a_m)} \tag{2.36}
\]

Since we are using a symmetric Gaussian kernel with standard deviation $\sigma_x$ for the functional image, $\phi'(\frac{f_i-f_m}{\sigma_x}) = \phi'(\frac{f_m-f_i}{\sigma_x}) = \phi'(\frac{f_i-f_m}{\sigma_x}) \frac{f_i-f_m}{\sigma_x^2}$, we get

\[
\frac{\partial H(X,Y)}{\partial f_m} = -\frac{1}{N_s} \sum_{i=1}^{N_s} \left( \frac{\phi'(\frac{f_i-f_m}{\sigma_x}) \phi'(\frac{a_i-a_m}{\sigma_y})}{p(f_i, a_i)} + \frac{\phi'(\frac{f_m-f_i}{\sigma_x}) \phi'(\frac{a_m-a_i}{\sigma_y})}{p(f_m, a_m)} \right) \left( \frac{f_i-f_m}{\sigma_x^2} \right). \tag{2.37}
\]

If we let

\[
\alpha_{im} = \frac{\phi'(\frac{f_i-f_m}{\sigma_x}) \phi'(\frac{a_i-a_m}{\sigma_y})}{p(f_i, a_i)}, \quad \text{and}
\]
\[
\beta_{im} = \frac{\phi'(\frac{f_m-f_i}{\sigma_x}) \phi'(\frac{a_m-a_i}{\sigma_y})}{p(f_m, a_m)}, \tag{2.38}
\]

then Equation 2.37 can be rewritten as,

\[
\frac{\partial H(X,Y)}{\partial f_m} = -\frac{1}{N_s} \sum_{i=1}^{N_s} \left( \alpha_{im} + \beta_{im} \right) \left( \frac{f_i-f_m}{\sigma_x^2} \right),
\]

\[
= -\frac{1}{N_s} \sum_{i=1}^{N_s} W_{im} \left( \frac{f_i-f_m}{\sigma_x^2} \right). \tag{2.39}
\]
The weights $\alpha_{im}$, and $\beta_{im}$ are both between 0 and 1, since the denominator in each of these terms is obtained by a summation of the numerator over all $m$ or $i$. The weight $W_{im}$ approaches 0 when $f_i$ and $a_i$ are distant from $f_m$ and $a_m$ respectively, and approaches 2 when they are close. Hence, $W_{im}$ selects the voxels with intensities that are within a few standard deviations of $f_m$ in both the anatomical and functional images, and updates $f_m$ such that the intensities in these voxels are clustered closer together.

The gradient in Equation 2.39 can be further approximated by considering only voxels that are spatially close to the voxel $m$, i.e., in a neighborhood $\eta_m$, giving

$$
\frac{\partial \tilde{H}(X,Y)}{\partial f_m} = -\frac{1}{N_s} \sum_{i \in \eta_m} W_{im} \left( \frac{f_i - f_m}{\sigma_i^2} \right). \tag{2.40}
$$

This expression is similar to that of the quadratic prior, except that the weight $W_{im}$ selects only those voxels within the neighborhood $\eta_m$ that have intensities close to $f_m$. In this case, the voxels that are spatially close to each voxel influence the gradient rather than the more global interaction described in Equation 2.39.
Chapter 3

Mutual Information Based Non-Rigid Mouse Registration

Using A Scale-Space Approach

Longitudinal and inter-subject studies are often performed in small animal imaging in order to study changes in mouse anatomy and function over a period of time, or across populations. Changes in animal posture, tissue growth, organ movement and other anatomical changes during longitudinal studies require non-rigid registration of the acquired images for accurate analysis. Normalization of anatomical variability across populations in inter-subject studies also requires non-rigid registration.

In this chapter, we describe a Gaussian scale-space theory based approach to simultaneously align global structure such as overall shape, as well as detail such as the skeleton, in small animals. We extract scale-space feature vectors at each voxel of the target and template images to be registered. Each feature vector is defined as a vector of the intensities of its scale-space images at that voxel. Detail such as the skeletal structure is visible in the lower scales and is blurred out at the higher scales, while the global structure such as overall shape remains in the images at the higher scales. The scale-space feature vectors are aligned with the goal of finding a common mapping that aligns the
skeleton as well as overall shape in the mouse images. We do not explicitly impose any rigidity constraints near the skeleton. We use mutual information (MI) as a similarity metric between the target and template images since it measures the similarity between distributions of intensities rather than actual values of intensities in the image, thus being more robust to intensity differences in these images [13]. This multi-scale approach has the additional advantage of having a cost function that is less prone to local minima [27], and hence is able to perform large non-linear deformations accurately.

We parameterize the displacement field using the DCT basis, and use the Laplacian of the field as a regularizing term. The DCT bases are eigen functions of the discrete Laplacian, so using the DCT representation of the displacement field in conjunction with Laplacian regularization simplifies the regularization term to a diagonal matrix. Moreover, the DCT basis gives an efficient representation of the displacement field, allowing us to represent the field using only a few coefficients. Fast implementations of DCT are readily available, and can be used to further reduce the computation time. In [45] a similar parametrization is used for non-rigid registration using SSD metric with Gauss Newton optimization, which is fast but can be used only for the SSD metric. The Gauss-Newton optimization also restricts the number of bases that can be used in practice. The paper recommends the use of less than $8 \times 8 \times 8$ coefficients to avoid problems. We use a conjugate gradient optimization with Armijo line search to optimize our cost function, and there is no constraint on the number of bases that can be used.

We apply the scale-space registration algorithm to CT images from a longitudinal mouse study. We compare the results of this scale-space approach with those obtained by a hierarchical multi-scale approach that is commonly used in brain registration, where
images at each scale are aligned hierarchically, starting at the highest scale and ending at the lowest scale.

### 3.1 Methods and Results

#### 3.1.1 Mutual information based non-rigid registration

Let $i_1(x)$ and $i_2(x)$ represent the intensities of target and template images respectively at position $x$. Let the transformation that maps the target to the template be $T(x) = x - u(x)$, where $u(x)$ is the displacement field. Let the feature vectors extracted from the target and template images at voxel $x$ be $i_1(x)$ and $i_2(x)$ respectively. These can be considered as realizations of the random vectors $I_1$ and $I_2$. Let $N_s$ be the number of features in each feature vector such that $I_1 = [I_1^1, I_1^2, \cdots, I_1^{N_s}]^T$, where $I_1^j$ is the random variable corresponding to feature $j$ of the target image. Then, the mutual information between feature vectors of the target and deformed template, $D_u(I_1, I_2)$ is defined as [39],

$$
D_u(I_1, I_2) = \int p_u(I_1, I_2) \log \frac{p_u(I_1, I_2)}{p(I_1)p_u(I_2)} dI_1 dI_2. \tag{3.1}
$$

A differentiable estimate of the density can be obtained using Parzen windows given by [34],

$$
p_u(I_1, I_2) = \sum_x \phi \left( \frac{I_1 - i_1(x)}{\rho} \right) \phi \left( \frac{I_2 - i_2(x - u(x))}{\rho} \right), \tag{3.2}
$$

where $\phi$ is a Gaussian window and $\rho$ determines the width of the window.
We use the Laplacian of the displacement field as a regularizing term. The objective function is given by

$$\max_u D_u(I_1, I_2) - \mu \| \nabla^2 u(x) \|^2,$$  \hspace{1cm} (3.3)

where $\mu$ is a hyperparameter that controls the weight on the regularizing term.

For simplicity, we assume independence between features so that MI between the feature vectors simplifies to the sum of MI between the individual features.

$$D_u(I_1, I_2) = \sum_{j=1}^{N_s} D_u(I_1^j, I_2^j).$$ \hspace{1cm} (3.4)

### 3.1.2 Scale-space feature vectors

We use a Gaussian scale-space theory based approach to define feature vectors that are extracted from the images. We generate images at different scales by blurring the target and template with Gaussian kernels of increasing widths. Let the width of the Gaussian kernel at each scale be $\sigma_j$, $j = 1, 2, \ldots, N_s$, such that $\sigma_j > \sigma_k$ for $j > k$. Let the smoothing kernel at scale $j$ be represented by $S_j(x)$. We define the scale-space based feature vector as $i_1(x) = [i_1(x)^1, i_1(x)^2, \ldots, i_1(x)^{N_s}]$, where $i_1(x)^j$ is the intensity of the target image at scale $j$ and position $x$, and is given by,

$$i_1(x)^j = S_j(x) * i_1(x).$$ \hspace{1cm} (3.5)

In this work, we use $N_s=2$, where the first scale corresponds to no smoothing and $\sigma_2 = 3$. A coronal slice of a mouse image at these scales is shown in Fig. 3.1.
Figure 3.1: Coronal slice of scale-space images of a mouse for $N_s = 2$, with scale 1 corresponding to no smoothing (left) and $\sigma_2 = 3$ (right).

We generate the feature vector of the deformed template $i_2(x - u(x))$ by first applying the displacement field to the original template, and then generating the scale-space images of that deformed image. So the feature $i_2(x - u)^j$ is given by,

$$i_2(x - u)^j = S_j(x) \ast (i_2(x - u)).$$  \hspace{1cm} (3.6)

We observed that this performs better than deforming the original scale-space images of the template. This could be because in our approach, we are retaining the original relationship between intensities as a function of position in the template image while computing $i_2(x - u(x))$, and then computing the similarity metric between target and a linear transformation (smoothing) of the deformed template. In contrast, applying the displacement field to the original scale-space images will mean that we are computing the similarity metric between the target and a new function $(S_j \ast i_2)(x - u)$ of the template. Though generating the scale-space feature vectors at every iteration increases the computation, we take this approach for better accuracy.
3.1.3 Displacement field and regularization

The displacement field $u(x)$ can be represented in terms of the DCT basis. Let $b_i(x)$ and $\beta_i(x)$, $i = 1, 2, \cdots, N_b$ represent the DCT basis vectors, and the DCT coefficients respectively. Then, the displacement field is given by,

$$u(x) = \sum_{i=1}^{N_b} \beta_i b_i(x) = B(x) \beta,$$

(3.7)

where $B(x) = [b_1(x), b_2(x), \cdots, b_{N_b}(x)]$, and the coefficient vector $\beta = [\beta_1, \beta_2, \cdots, \beta_{N_b}]^T$.

The DCT basis vectors are eigen functions of the discrete Laplacian. Let $L$ be the discrete Laplacian matrix and $\gamma_i$, $i = 1, 2, \cdots, N_b$ be the eigen values of $L$ corresponding to the basis $b_i(x)$. Then,

$$Lu(x) = L \sum_{i=1}^{N_b} \beta_i b_i(x) = \sum_{i=1}^{N_b} \gamma_i \beta_i b_i(x)$$

(3.8)

Hence, the discrete approximation of the norm of the Laplacian, $\|\nabla^2 u(x)\|^2$ is given by,

$$\|Lu(x)\|^2 = \sum_{i=1}^{N_b} \gamma_i^2 \beta_i^2.$$  

(3.9)

Thus we save the computation cost of the matrix vector multiplication required by the left hand side of Equation 3.9, by reducing it to the vector norm on the right hand side. Additionally, since the DCT gives a sparse representation, we can model the displacement field with $N_c < N_b$ coefficients to reduce computation time further.
Finally, the objective function is given by

$$\max_\beta \sum_{j=1}^{N_s} D_{B,\beta}(I_1^j, I_2^j) - \mu(\sum_{j=1}^{N_c} \gamma_j \beta_j^2).$$

(3.10)

We optimize this objective function using a gradient descent algorithm with Armijo line search [40].

### 3.1.4 Results

We use CT images from a longitudinal mouse study. We use two time points of that study that are two months apart. The CT images were acquired using the microCT system at a resolution of $0.2 \times 0.2 \times 0.2$ mm. We first perform a MI based rigid registration on the two images using the RView software [42]. The rigidly registered images are shown in Figure 3.2. It can be seen that there are considerable differences in the skeleton, soft tissue, and overall shape of the images. We downsampled the CT images to size $128 \times 128 \times 64$, corresponding to a resolution of $0.4 \times 0.72 \times 0.51$ mm to reduce computation. We used these downsampled images as the target and template images in the non rigid registration.
algorithm. We use two scales to form our feature vectors—the unsmoothed image, and the image smoothed by a Gaussian kernel of size $[7\times7\times7]$ and width $\sigma = 3$. We use $N_c = 30$ DCT bases and $\mu = 5e - 6$. We first perform the non-rigid registration at only the higher scale ($\sigma = 3$), and we use this to initialize the scale-space non-rigid registration. Registration at only the lower scale (unsmoothed images) gave a local minimum that would not be a good initialization.

We compare the scale-space registration results to those of hierarchical multi-scale registration. We used the same number of DCT bases and the same scales as the scale-space registration, with $\mu = 2e - 5$. We first registered the smoothed images, and used the resulting displacement field to initialize the registration for the unsmoothed images. The overlay on the target image of the template image, and images registered using both approaches are shown in Fig. 3.3. We applied the displacement field resulting from both registration algorithms to the higher resolution (0.2 mm) images for display purposes. The displacement field obtained from the scale-space registration is shown in Fig. 3.4, for the coronal and sagittal slices shown in Fig. 3.3.

It can be seen that the scale-space non-rigid registration algorithm gives better skeletal, limb, and soft tissue (such as liver and heart) alignment than the hierarchical multi-scale algorithm.

### 3.2 Discussion

We used a scale-space theory based approach for non-rigid mouse registration using mutual information. We used the DCT basis to efficiently represent the displacement field,
Figure 3.3: Comparison of registered images: Coronal and sagittal views of overlay on target image of template (top row), hierarchical multi-scale registered image (middle row), and scale-space registered image (bottom row).
as well as to simplify the Laplacian regularization term. We generated the scale-space feature vectors at each iteration after applying the displacement field to the original images, since this approach gave better accuracy. We applied this algorithm to CT images of a mouse in a longitudinal study.

By using images at different scales simultaneously, we obtained better alignment of the global structure such as the overall shape of the mouse as well as improved alignment of detail such as the skeleton, compared to the hierarchical multi-scale approach. Additionally, using multiple scales simultaneously gives a cost function that is less prone to local minima. Though these results are encouraging, we note that this approach is computationally intensive. We also note that using non-parametric density estimation in conjunction with non-rigid MI based registration might pose problems because of the high dimensionality of the problem coupled with the non-convexity of the cost function. In the next chapter, we explore an alternative formulation for non-rigid registration that deals
with the high dimensionality of the combined density and displacement field estimation problems.
Chapter 4

Non-Rigid Registration Using Gaussian Mixture Models

Registration techniques that use mutual information as a similarity metric typically use a non-parametric approach for estimating the unknown distribution of intensities in the images [13], [28] - [30], [Chapter 3]. This framework has some limitations when dealing with the high dimensional non-rigid registration problem. Firstly, mutual information is a non-convex function of the registration parameters and the registration could converge to an inaccurate local optimum. This problem is exacerbated for the non-rigid registration case due to the increase in dimensionality of the deformation field to be estimated compared to the rigid registration case, for which this framework has been successfully applied [13]. Secondly, the non-parametric Parzen based approach that is typically used in MI based registration estimates the entire joint density from the images, and this joint density is then used to estimate the deformation field. Using these two high dimensional estimation steps in conjunction with each other could result in a registration that is sensitive to small changes in the images, thus making it more susceptible to local optima. Additionally, the Parzen window method requires appropriate choice of the Parzen window width, which is usually taken as a design parameter, and is kept fixed over the entire
sample [34]. This has the drawback that for long-tailed distributions the density estimate tends to be noisy at the tails, and increasing the window width to deal with this might lead to oversmoothing the details in the distribution [46]. This would mean that the former scenario results in a cost function that has more local optima, while the latter could lead to inaccurate registration results. This approach also requires a large number of samples to accurately estimate the distribution.

In this chapter, we describe an alternative approach for multi-modality non-rigid registration that uses the log-likelihood of the target image given the deformed template as a similarity metric for non-rigid registration, wherein the log-likelihood is modeled using Gaussian mixture models (GMM). Gaussian distributions are commonly used in image segmentation to represent the distribution of intensities corresponding to a particular tissue type in MR or CT images [47]. The joint distribution of two MR or CT images is characterized by localized blobs that can also be modeled as a mixture of Gaussians. Using GMMs reduces the density estimation step to that of estimating the parameters of the GMM, which consist of the mean, covariance, and weight of each Gaussian. Since the estimated parameters can be different for each Gaussian, the density estimates obtained using this approach are smoother than the non-parametric approach while retaining the detail in the distribution. We choose the number of components in the GMM from the joint histogram of the target and template images, and keep this number constant throughout the registration. This can hence be considered as a semi-parametric approach to density estimation. For images that have a few distinct regions of intensity such as brain or mouse images, the number of parameters to be estimated is small, and can be robustly estimated from fewer samples compared to the non-parametric approach.
Our approach of using the log-likelihood of the target given the template in conjunction with a GMM can be viewed as a semi-parametric approximation to MI based registration. However, we expect our GMM based approach to perform better than the non-parametric MI based approach because of (1) reduced overall dimensionality of the registration problem through the parametrization of the density estimation step, and (2) more robust and accurate estimation of the joint density. Additionally, we parameterize the displacement field using the DCT basis, and use the Laplacian of the field as a regularizing term as described in Chapter 3.

We compare the performance of the conditional likelihood metric with GMM parametrization, with that of MI with non-parametric density estimation approach. We will henceforth refer to these methods as the GMM-Cond and the Parzen-MI approaches respectively. We evaluate these methods using intra-modality brain MR images, as well as inter-subject, inter-modality mouse images. Registration of mice and other small animals is challenging because of the presence of rigid skeleton within non-rigid soft tissue, and it has been observed that though the existing non-rigid registration algorithms have been applied to brain imaging with encouraging results, they do not perform well for small animal images [31] -[33]. We test the performance of our approach for both brain as well as mouse imaging to evaluate potential in clinical as well as pre-clinical imaging applications.
4.1 Methods and Results

Let the target and template images be $I_1$ and $I_2$, and their intensity at position $x$ be $i_1(x)$ and $i_2(x)$ respectively. Let the transformation that maps the template to the target be $T(x) = x - u(x)$, where $u$ is the displacement field. The deformed template is then represented by $I_2^u$, whose intensities are given by $i_2(x - u(x))$. We define the similarity metric $D_u(I_1, I_2)$ between the target and deformed template as the log likelihood of the target given the deformed template. Assuming that the voxel intensities in $I_1$ and $I_2$ are independent identically distributed random variables with joint density $p(i_1, i_2)$, the similarity metric is given by,

$$D_u(I_1, I_2) = \log p(I_1|I_2^u) = \sum_x \log \frac{p(i_1(x), i_2(x - u(x)))}{p(i_2(x - u(x)))}. \quad (4.1)$$

We assume a Gaussian mixture model for the joint and marginal densities $p(i_1, i_2)$ and $p(i_2)$. Let the number of components of the Gaussian mixture model be $K$, the mixing proportions be $\pi_k$, and $g(i_1, i_2|m_k, \Sigma_k)$ be a Gaussian with mean $m_k$ and covariance $\Sigma_k$, where $k = 1, 2, \cdots, K$. Let the unknown GMM parameters for each component $k$ be represented as $\theta_k = (\pi_k, m_k, \Sigma_k)$, and let $\Theta = [\theta_1, \theta_2, \cdots, \theta_K]$ be the vector of all unknown parameters. Then, the joint density is given by

$$p(i_1, i_2|\Theta) = \sum_{k=1}^K \pi_k g(i_1, i_2|m_k, \Sigma_k), \quad (4.2)$$

where $\pi_k > 0$ and $\sum_{k=1}^K \pi_k = 1$. 

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We use the Laplacian of the displacement field as a regularizing term $R(u)$ to penalize deformations that are not smooth. Let $L$ be the discrete Laplacian matrix. The objective function is then given by,

$$\max_{u, \Theta} \sum_x \log p(i_1(x)|i_2(x - u(x)), \Theta) - \mu ||Lu||^2,$$

(4.3)

where $\mu$ is a hyperparameter that controls the weight on the regularizing term.

To simplify the problem, we replace the combined optimization with respect to the deformation field and GMM parameters with an iterative two step procedure. Here, the GMM parameters are first estimated from the target and deformed template images through maximum likelihood estimation, and the deformation field is then estimated given the estimated GMM parameters. The two step optimization is given by

\[
\hat{\Theta}(\hat{u}^m) = \arg \max_{\Theta} \sum_x \log p(i_1(x), i_2(x - \hat{u}^m(x))|\Theta) \\
\hat{u}^{m+1} = \arg \max_u \sum_x \log p(i_1(x)|i_2(x - u(x)), \hat{\Theta}(\hat{u}^m)) - \mu ||Lu||^2,
\]

(4.4)

where $\hat{u}^m$ represents the estimated deformation field at overall optimization iteration $m$.

### 4.1.1 Estimation of parameters of Gaussian mixture model

The maximum likelihood estimate of the GMM parameters $\hat{\Theta}$ in equation 4.4 can be obtained by the EM algorithm [48]. Let the data sample at voxel $j$ corresponding to
the position $x_j$ be $S^u_j = [i_1(x_j), i_2(x_j - u(x_j))]^T$, where $j = 1, 2, \cdots, N$, and $N$ is the number of voxels in each image. The component of the GMM from which $S_j$ arises is taken as the hidden variable in the EM algorithm. The EM update equations are given by,

$$
\tau_{jk}^i = \frac{\pi_k^i g(S^u_j, m_k^i(u), \Sigma_k^i(u))}{\sum_{h=1}^K \pi_h^i g(S^u_j, m_h^i(u), \Sigma_h^i(u))},
$$

(4.6)

$$
\pi_k^{i+1}(u) = \frac{1}{N} \sum_{j=1}^N \tau_{jk}^i,
$$

(4.7)

$$
m_k^{i+1}(u) = \frac{\sum_{j=1}^N \tau_{jk}^i S^u_j}{\sum_{j=1}^N \tau_{jk}^i},
$$

(4.8)

$$
\Sigma_k^{i+1}(u) = \frac{\sum_{j=1}^N \tau_{jk}^i (S^u_j - m_k^{i+1}(u))(S^u_j - m_k^{i+1}(u))^T}{\sum_{j=1}^N \tau_{jk}^i},
$$

(4.9)

where $\pi_k^i(u), m_k^i(u)$, and $\Sigma_k^i(u)$ are the GMM parameter estimates at EM iteration $i$ and deformation field $u$.

The objective function to be optimized in equation 4.4 is a non-convex function of $\Theta$, so a good initial estimate is needed to converge to the correct local optimum. We use the k-nearest neighbors algorithm [34] to identify cluster centers in the joint histogram of the target and template images, and the number of samples that fall into a particular cluster. The cluster centers and the proportion of samples in a cluster relative to the total number of samples were used as the initializations $m_k^0$ and $p_k^0$ respectively, and $\Sigma_k^0$ was assumed to be identity for all $k$. The number of clusters was chosen to match the initial histogram of the two images, after which it was kept constant throughout the registration process. Figure 4.1 shows an example of a pdf estimated using GMM. Two brain images were considered, transaxial slices of which are shown in Figure 4.1(a) and
in Figure 4.1(b). The joint histogram of the intensities of these two images is shown
in Figure 4.1(c), and the pdf estimated using GMM is shown in Figure 4.1(d) with the
component means overlaid on it. The number of components was chosen to be \( K = 6 \) to
match the joint histogram.

In the GMM based approach for probability density estimation, the number of pa-
rameters to be estimated is \( K + KD + KD(D + 1)/2 \), where \( D \) is the dimensionality of
the sample, and \( K \) is the number of components in the GMM. For registration of two
images \( D = 2 \), so the number of parameters is \( 6K \), which is small for small values of
\( K \). On the other hand, the non-parametric Parzen based estimation computes the en-
tire \( N_{bin} \times N_{bin} \) pdf from the samples, where \( N_{bin} \) is the number of bins at which the
pdf is computed. We expect to gain computationally as well as in robustness, from this
reduction in dimensionality of the problem. However, if the joint density does not fit a
GMM, the number of mixture components might be large, approaching a Parzen window
estimate.

### 4.1.2 Estimation of displacement field

Let the objective function in equation 4.5 be represented by \( J^m(u) \), which can be rewritten
as

\[
J^m(u) = D_u(I_1, I_2|\hat{\Theta}(\hat{u}^m)) - \mu R(u)
\]  

We use conjugate gradient (CG) optimization to optimize this objective function with
an Armijo line search [40]. Let \( \hat{u}^{(i)} \) be the estimate of deformation field at CG iteration
\( i \). Let the gradient vector be \( d^{(i)} = \nabla J^m(u)|_{u=u^{(i)}} \), the conjugate gradient direction
Figure 4.1: Estimation of GMM parameters: (a) Target image, (b) Template image, (c) Joint histogram of intensities of target and template images, (d) Joint pdf of target and template images computed using GMM (the component means shown with 'x' marks).
vector be $b^{(i)}$, and the step size computed from the line search be $a^{(i)}$. The CG update equations are given by

$$\hat{u}^{(i+1)} = \hat{u}^{(i)} + a^{(i)} b^{(i)}$$
(4.11)

$$b^{(i)} = d^{(i)} + \beta^{(i-1)} b^{(i-1)}$$
(4.12)

$$\beta^{(i-1)} = \frac{(d^{(i)} - d^{(i-1)})^T b^{(i)}}{d^{(i-1)^T} b^{(i-1)}}$$
(4.13)

The algorithm is initialized by $b^{(0)} = d^0$. If the CG algorithm converges in $r$ iterations, the estimated deformation field in equation 4.5 is $\hat{u}^{m+1} = \hat{u}^{(r)}$.

The gradient vector $d$ is given by

$$d = \frac{\partial}{\partial u} D_u(I_1, I_2) - \mu \frac{\partial}{\partial u} R(u),$$
(4.14)

where $\frac{\partial}{\partial u} D_u(I_1, I_2)$ and $\frac{\partial}{\partial u} R(u)$ are the gradients of the similarity metric and the regularizing terms respectively. The gradient of the similarity metric with respect to the displacement field at voxel position $x_l$ is given in equation 4.15, where $i_1(l)$ and $i_2^{\text{u}}(l)$ are shorthand notations for $i_1(x_l)$ and $i_2(x_l - u(x_l))$ respectively, the intensities of the target and deformed template at $x_l$ are represented by $S_l^u = [i_1(l), i_2^{\text{u}}(l)]^T$, and $\nabla i_2^{\text{u}}(l)$ is the spatial gradient of the deformed template at $x_l$. The gradient of the regularizing term is given in equation 4.16.
\[ \frac{\partial D_u(i_1, i_2)}{\partial u(x_i)} = \frac{\nabla i_2^u(l)}{p(i_1(l), i_2^u(l))} \sum_{k=1}^{K} \pi_k g(S_1^u|\hat{\Theta}(\hat{u}^m)) \left( \begin{bmatrix} 0 & 1 \end{bmatrix} \Sigma_k^{-1} \begin{bmatrix} i_1(l) - m_{k,1} \\ i_2^u(l) - m_{k,2} \end{bmatrix} \right) - \frac{\nabla i_2^u(l)}{p(i_2^u(l))} \sum_{k=1}^{K} \pi_k g(i_2^u(l)) m_{k,2,2} \frac{(i_2^u(l) - m_{k,2})}{\Sigma_{k,2,2}}, \] (4.15)

\[ \frac{\partial}{\partial u} R(u) = 2 \Lambda^T \Lambda u. \] (4.16)

We use a DCT parametrization of the displacement field as described in Chapter 3.

The objective function is finally given by

\[ \max_{\beta} D_B(\beta) = \max_{\beta} \left( \frac{\partial}{\partial \beta} D_B(\beta) \right) = \mu \sum_{i=1}^{N_b} \gamma_i^2 \beta_i^2, \] (4.17)

where \( B, \beta \) represent the DCT bases and coefficients respectively, and \( \gamma_i, i = 1, 2, \cdots, N_b \) represents the eigen values of \( \Lambda \) corresponding to the \( i^{th} \) basis vector.

Since the DCT is a linear transformation of \( u \), the gradient with respect to \( \beta \) can be obtained in terms of the gradient with respect to \( u \) as

\[ \frac{\partial}{\partial \beta} D_B(\beta) = B^T \frac{\partial}{\partial u} D_u(I_1, I_2) \] (4.18)

Thus the gradient can be computed by using equation 4.15 and then taking its DCT .

The gradient of the regularizer is

\[ \frac{\partial}{\partial \beta} R(B\beta) = 2\Lambda \beta, \] (4.19)
where $\Lambda$ is a diagonal matrix with elements $[\gamma_1^2, \gamma_2^2, \cdots, \gamma_N^2]$.

### 4.1.3 Relation to Mutual Information

Let the random variables corresponding to the intensities of $I_1$ and $I_2$ be $\zeta_1$ and $\zeta_2$ respectively. Mutual information between $\zeta_1$ and $\zeta_2$ is defined as [39],

$$
D(\zeta_1, \zeta_2) = \int p(z_1, z_2) \log \frac{p(z_1, z_2)}{p(z_1)p(z_2)} \, dz_1 \, dz_2
$$

$$
= E(\log \frac{p(z_1, z_2)}{p(z_1)p(z_2)}).
$$

(4.20)

MI between two random variables can be interpreted as the reduction in uncertainty of one random variable given the other. Using MI as a similarity metric for registration aims to find a deformation that makes the joint density of the target and deformed template images maximally clustered, thus implying that the uncertainty of one image given the other is minimized.

An alternative formulation can be obtained by approximating the expectation in equation 4.20 by a sample mean where the intensity at each voxel in the target and deformed template images constitutes the random sample. Hence we get

$$
\hat{D}(\zeta_1, \zeta_2) = \frac{1}{N} \sum_x \log \frac{p(i_1(x), i_2(x - u(x)))}{p(i_1(x))p(i_2(x - u(x)))}
$$

$$
= \frac{1}{N} \sum_x \log \frac{p(i_1(x), i_2(x - u(x)))}{p(i_2(x - u(x)))}
$$

$$
- \frac{1}{N} \sum_x \log p(i_1(x)).
$$

(4.21)
Since the target is fixed and independent of $u(x)$, dropping the terms containing the marginal density $p(i_1)$, we get the approximate MI based similarity metric as

$$\hat{D}(\zeta_1, \zeta_2) = \frac{1}{N} \sum_x \log \frac{p(i_1(x), i_2(x - u(x)))}{p(i_2(x - u(x)))}.$$  

(4.22)

Thus, computing the deformation field that maximizes mutual information is approximately equivalent to maximizing the conditional density of the target given the template image as defined in Equation 4.1. In [36] a similar relationship between maximum likelihood and conditional entropy was derived. The pdf $p(i_1, i_2)$ is unknown, and needs to be estimated. The pdf can be estimated using a non-parametric approach such as Parzen windowing, or a GMM based approach can be taken to parametrize the pdf and estimate those parameters. The approach taken to estimate the pdf causes differences in the behavior of these metrics, and we will explore these differences by looking at the gradient equations of these two methods.

The Parzen window estimate of a pdf at random variable values $z_1, z_2$ is defined by [34]

$$p(z_1, z_2) = \frac{1}{N} \sum_{j=1}^N g\left(\frac{z_1 - i_1(j)}{\sigma}\right)g\left(\frac{z_2 - i_2(u)(j)}{\sigma}\right),$$  

(4.23)

where $g(\frac{z}{\sigma})$ is a Gaussian window of width $\sigma$, which is usually taken as a design parameter.

Note that this can be considered as a Gaussian mixture model with the fixed parameters

$$K = N, \quad \pi_k = \frac{1}{N}, \quad m_k = [i_1(k), i_2(u)(k)]^T, \quad \Sigma_k = \begin{bmatrix} \sigma^2 & 0 \\ 0 & \sigma^2 \end{bmatrix}.$$  

The gradient of Parzen based MI cost function with respect to the displacement field at voxel $l$ is then given by
\[
\frac{D(\zeta_1, \zeta_2)}{\partial u(l)} = -\frac{1}{N} \sum_{z_1, z_2} (1 + \log p(z_1, z_2)) g\left(\frac{z_1 - i_1(l)}{\sigma}\right) \frac{g'(z_2 - i^u_2(l))}{\sigma} \nabla_i u_2(l) \\
+ \frac{1}{N} \sum_{z_2} (1 + \log p(z_2)) g'(z_2 - i^u_2(l)) \nabla_i u_2(l),
\]

where \(g'(z_2 - i^u_2(l)) = g\left(\frac{z_2 - i^u_2(l)}{\sigma}\right) \frac{z_2 - i^u_2(l)}{\sigma^2}\).

We can rewrite the gradient equation for the GMM-Cond approach in Equation 4.15 as

\[
\frac{\partial D_u(i_1, i_2)}{\partial u(x_l)} = \frac{\nabla_i u_2(l)}{p(i_1(l), i^u_2(l))} \sum_{k=1}^K \pi_k g'(S_i^u | \hat{\Theta}(\hat{u}^m)) - \frac{\nabla_i u_2(l)}{p(i^u_2(l))} \sum_{k=1}^K \pi_k g'(i^u_2(l)|m_{k,2}, \Sigma_{k,2,2})
\]

where \(g'(S_i^u | \hat{\Theta}(\hat{u}^m)) = \begin{bmatrix} 0 & 1 \end{bmatrix} \Sigma_{k,1}^{-1} \begin{bmatrix} i_1(l) - m_{k,1} \\ i^u_2(l) - m_{k,2} \end{bmatrix}\). Thus, in both Parzen-MI as well as the GMM-Cond approaches, the gradient expressions have the derivative of a Gaussian weighting the summand. In Parzen-MI case, the Gaussian that is being differentiated has a mean given by the data sample at which it is centered, and a fixed variance over the entire sample. In the GMM case, the Gaussian has a mean and covariance given by the estimated GMM component mean and covariance. The Gaussian derivative has a large magnitude for values that are close to but not equal to the mean, and low everywhere else. A plot of \(g'(z_2)\) is shown for a Gaussian window of width \(\sigma = 2\) in Figure 4.2. Thus, for MI, the gradient selects the voxels that are close (within a few standard deviations (SDs) of Parzen window width) in intensity to a pdf bin value, and in order
to increase MI, deforms the image such that these voxel intensities get closer to that bin.
The GMM based approach on the other hand, selects voxels that are within a few SDs of the estimated GMM mean, and tries to get the voxel intensities closer to the mean. The GMM approach can thus be more robust to local minima, since the Parzen approach might produce deformations that have regions of misalignment because of splitting a large cluster in the joint density into multiple localized peaks around certain bin values.

We therefore expect the GMM-Cond approach to have the following two advantages over the Parzen-MI approach.

1. The density estimation is reduced to that of estimating $6K$ GMM parameters that can be robustly estimated from the given images for small $K$.

2. Estimation of displacement field may be more robust to local minima since the gradient tries to form localized peaks around the estimated GMM means in the joint pdf, rather than around the closest pdf bin.
4.1.4 Results

We perform validation studies of our method using (1) brain images, and (2) mouse images. Through these studies, we evaluate the performance of our approach in brain imaging as well as small animal imaging. Brain images usually consist of three distinct regions of intensity, the gray matter, white matter, and cerebro-spinal fluid. Thus the distribution of intensities of these images can be modeled using a GMM with a small number of components, making the GMM-Cond approach feasible. Similarly mouse CT images typically consist of mainly soft tissue versus bone contrast, and can be assumed to follow a GMM. Though mouse MR images have a larger number of intensity levels than the CT, the number of components required in the GMM is not prohibitively large.

4.1.5 Brain Image registration

We randomly chose 9 brains from the Internet Brain Segmentation Repository (IBSR) provided by the Center for Morphometric Analysis at Massachusetts General Hospital and is available at http://www.cma.mgh.harvard.edu/ibsr/. This dataset contains T1 weighted MR images along with their manual segmentations. We skull-stripped the brains using the available manual segmentations, and performed histogram equalization on their intensities to enhance contrast. Each brain from this dataset is used as an atlas that is registered to the rest of the brains, thus giving 72 registrations in total. We first rigidly registered each atlas to each target using AIR registration [49], [50], and then performed non-rigid registration using the GMM-Cond, Parzen-MI, and SSD methods to register these images. Though SSD is not a multi-modality metric like conditional likelihood and MI, it is widely used for intra-modality registration and hence we include it in our
evaluation. For all three methods, we used $30 \times 30 \times 30$ DCT bases, and conjugate gradient optimization. Additionally, for all three methods, we check the determinant of the Jacobian of the deformation field for negative values, and choose a value of $\mu$ to weight the regularization term such that the Jacobian determinant has no negative values. This ensures a smooth deformation field for all methods considered.

For the Parzen-MI registration we followed a hierarchical approach, first aligning the images that were smoothed with a Gaussian of width 3 voxels, and using the displacement field thus obtained to initialize the registration of the original images. We observed that directly aligning the original images causes the algorithm to reach an inaccurate local minimum. A Parzen window width of $\sigma = 2$ was used to compute the distribution at every iteration. For the GMM-Cond approach, we used $K = 6$ components in the GMM, and 5 overall iterations between the density estimation and deformation field estimation. Each displacement field estimation in the GMM-Cond approach involved 50 iterations of the conjugate gradient algorithm.

The registration results for one atlas to target registration are shown in Figure 4.3. The corresponding displacement field obtained from GMM-Cond registration is shown in Figure 4.4. The joint density estimated using the Parzen window and GMM methods before and after registration of the brains is shown in Figure 4.5. It can be seen that the GMM-Cond approach is able to cluster the intensities more tightly along the diagonal than the MI-Parzen approach, implying a better alignment. The registration accuracy is evaluated based on the following metrics:
Figure 4.3: Registration results for one dataset: (a) Target, (b) Atlas, (c) SSD registered image, (d) Parzen-MI registered image, and (e) GMM-Cond registered image.
Figure 4.4: Displacement field obtained from GMM-Cond registration corresponding to the slice shown in Figure 4.3.

Figure 4.5: Comparison of Parzen-MI vs GMM-Cond based methods: Log of the estimate of initial joint density between target and template images using (a) Parzen window method, (b) GMM method (the ‘x’ marks denote the GMM means), and log of the estimate of joint density between target and registered template using (c) Parzen window method and (d) GMM method.
1. Dice coefficients: Dice coefficient is a measure of the amount of overlap in sets $A$ and $B$, and is defined as,

$$d = \frac{|A \cap B|}{|A| + |B|},$$

(4.25)

where $|A|$ denotes the number of elements in the set $A$.

We use dice coefficients to evaluate registration accuracy in terms of its ability to align various segmented regions of the brain. The deformation field obtained from each atlas-target registration is applied to the labels of the atlas to obtain the segmentation of the target image. The dice coefficients between the manual segmentation of the target image, and that obtained by atlas based segmentation is computed. The mean dice coefficients over all 72 atlas-target registrations are shown for various brain regions in Figure 4.6.

2. Normalized mean squared difference between intensities: The images used are all T1 weighted MR images, and computing the squared difference between the target and deformed atlas image intensities gives a measure of registration accuracy. Since SSD optimizes this metric, we expect this approach to have the best performance for this metric. However, for the Parzen-MI and the GMM-Cond methods, this would be an unbiased measure of registration accuracy. The MSE values are shown in a box plot for the SSD, Parzen-MI, and GMM based methods in Figure 4.7.

3. Qualitative comparison: The deformed atlases registered to each image are averaged to produce an image that would look more blurred if the registration was inaccurate. The images obtained by averaging the atlases deformed to match one target brain are shown in Figure 4.8 for the MI and GMM based methods.
Figure 4.6: Mean dice coefficients for different brain regions (Caud: Caudate, GM: Cerebral gray matter, WM: Cerebral white matter, Put: Putamen, Tha: Thalamus, HC: Hippocampus, CBGM: Cerebellar gray matter, CBWM: Cerebellar white matter)

From these figures, it can be seen that the GMM-Cond method has higher mean dice coefficients for all regions of the brain considered than the Parzen-MI and SSD methods, and lesser MSE between intensities than for the Parzen-MI approach. The average images using the GMM based approach look less blurred than that of the Parzen-MI approach, implying that the former is more consistently aligning the atlas to the target images. Thus, the GMM based approach shows higher registration accuracy than the Parzen-MI method in terms of the metrics considered.

4.1.6 Mouse image registration

Mouse image registration is a challenging problem because of the presence of rigid structures such as skeleton within non-rigid soft tissue. We apply our GMM based method to
multi-modality, inter-subject registration of mouse images. We consider two mice that were imaged using both MR and CT (referred to as MR1 and CT1, MR2 and CT2) and two other mice that were imaged using only CT (referred to as CT3 and CT4). The MR images were obtained on a Biospec 7T system at a resolution of $0.23 \times 0.17 \times 0.50$ mm. The CT images corresponding to the MR were acquired on a Siemens Inveon system and the others were obtained from a microCT system, at a resolution of $0.2 \times 0.2 \times 0.2$ mm. We first perform a 12 parameter affine registration of the CT images to the MR image using the AIR software [49], [50]. We downsampled the MR and affinely registered CT images to size $128 \times 128 \times 64$ to reduce computation. The downsampled MR and affinely registered CT images were then used as the target and template images respectively for non rigid registration (corresponding to 6 inter-subject multi-modality registrations). We
Figure 4.8: Transaxial and sagittal views of average images: Top row: Template image to which the atlases are registered. Middle row: Average of deformed atlases using Parzen-MI approach, Bottom row: Average of deformed atlases using GMM based approach.
compare the GMM-Cond and Parzen-MI approaches. SSD is not a suitable metric for multi-modality registration, and hence we do not use it for this comparison.

For both GMM-Cond and Parzen-MI methods, we used $15 \times 15 \times 15$ DCT bases to represent the displacement field. As before, we check the displacement field for foldings (negative determinant of Jacobian) and use a value of $\mu$ that gives a smooth field without foldings. We used a hierarchical approach for the Parzen-MI registration with images smoothed by a Gaussian of 3 voxels as the first step, followed by registration of the original images. A Parzen window width of $\sigma = 5$ was used to compute the distribution at every iteration. For the GMM-Cond approach, we used $K = 8$ components in the
GMM, and 5 overall iterations between the density estimation and deformation field estimation. Each displacement field estimation involved 50 iterations of the conjugate gradient algorithm. Coronal and sagittal views of the registered images for one dataset along with the target and template images are shown in Figure 4.9. The template and registered images for the same dataset are overlaid on the target MR and shown in Figure 4.10.

We quantify the performance of the registration through three measures:

1. Overall overlap: The target and template images can be segmented into mouse and background regions. The overall overlap of the target and deformed template can then be measured by computing the dice coefficients of the region labeled mouse in the two images.
2. Overlap of skeleton: The skeleton can be segmented out in the target and template images by thresholding. The dice coefficients of the skeleton in the target and deformed template images give a measure of overlap in the skeleton.

3. Mean squared difference (MSD) between intensities: The target MR image has a corresponding CT image acquired with it. The normalized mean squared difference between intensities of the CT corresponding to the target, and the deformed template images gives a measure of registration accuracy.

The values of these measures for the four CT images registered to the MR images are given in Table 4.1.

On an average, the GMM-Cond method has higher dice coefficients for the skeleton as well as overall shape, and lower normalized MSD between intensities than the MI-Parzen registration method, indicating better alignment. It is promising that the GMM-Cond shows improved performance for the inter-subject, multi-modality registration considered, since these images have considerable difference in intensities, overall shape, and skeletal structure.

4.2 Discussion

We used the conditional density of the target given the deformed template as a similarity metric for non-rigid registration, wherein the conditional density is modeled as a Gaussian mixture model. A DCT representation of the deformation field was used in conjunction with a Laplacian regularizing term to reduce computation. We compared the performance of our approach with that of Parzen-MI based approach using clinical and pre-clinical
Table 4.1: Quantitative measures of overlap

<table>
<thead>
<tr>
<th>Registration</th>
<th>Affine</th>
<th>Parzen-MI</th>
<th>GMM-Cond</th>
</tr>
</thead>
<tbody>
<tr>
<td>CT1-MR2</td>
<td>0.8953</td>
<td>0.9048</td>
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<td>CT3-MR2</td>
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<td>CT4-MR2</td>
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<tr>
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<tr>
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<td><strong>0.8892</strong></td>
<td><strong>0.9151</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Registration</th>
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<th>Parzen-MI</th>
<th>GMM-Cond</th>
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<td>CT4-MR1</td>
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<td><strong>0.3085</strong></td>
<td><strong>0.3404</strong></td>
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</table>

<table>
<thead>
<tr>
<th>Registration</th>
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<th>Parzen-MI</th>
<th>GMM-Cond</th>
</tr>
</thead>
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<tr>
<td>CT1-MR2</td>
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<tr>
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<td>CT3-MR1</td>
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<td>0.5383</td>
<td>0.6518</td>
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<tr>
<td>CT4-MR1</td>
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<td><strong>0.5027</strong></td>
<td><strong>0.4378</strong></td>
</tr>
</tbody>
</table>
images of both intra modality brain MR images as well as multi-modality MR/CT mouse images.

The GMM-Cond approach showed higher registration accuracy than the Parzen-MI approach in terms of dice coefficients and mean squared difference between intensities of the target and registered images. The GMM parametrization is not only computationally more efficient than the Parzen method, but also improves performance by reducing the overall dimensionality of the estimation problem, and through more robust and accurate density estimation. Additionally, the only design parameter that needs to be chosen is the number of clusters in the GMM, which can be obtained from the initial joint histogram.

The consistent performance of the GMM-Cond method for brain as well as mouse images, and for intra as well as multi-modality images indicates that this is a robust approach that can potentially be applied to multimodality registration applications. This approach can be used as an alternative to MI based registration when dealing with high dimensional deformation fields. It can be viewed as a general framework that can be used in conjunction with other models for the deformation field. It should be noted however, that if the joint density of the images does not follow a GMM, a large number of clusters would be required to fit the data, thus increasing the number of parameters to be estimated and might not perform better than Parzen-MI in that case. We expect this approach to be useful in applications where the images have a few distinct regions of intensity, such as brain or mouse CT images.
Chapter 5

Summary and Future Work

We explored the application of information theoretic measures to PET image reconstruction and non-rigid registration. For PET imaging, we described an approach that uses mutual information and joint entropy as anatomical priors on the PET image. We used a scale-space framework to incorporate spatial information into these global distribution based metrics. The novelty in this approach to using anatomical information in PET reconstruction is in aiming to reconstruct a PET image that is similar to the anatomical image in the information theoretic sense rather than defining explicit correlations between the boundaries or regions in the images. We presented an efficient FFT based method to compute these priors and their derivatives, making it practical to use these priors for 3D image reconstruction. The simulations and clinical data results indicate that joint entropy with scale-space information reconstructs images with more accurate quantitation and sharper boundaries than that of the commonly used quadratic priors, which do not use anatomical information. We applied these anatomical priors to positron range correction, which is a fundamental resolution limiting problem in PET. A truncated homogeneous model was used to incorporate range blurring into the system model, and
the joint entropy prior with scale-space features was used to correct for blurring outside regions where uptake is expected. Promising results were obtained from simulations as well as pre-clinical data, which indicate that joint entropy based priors can be used in conjunction with a system model that incorporates positron range, to improve the resolution of PET even when long range isotopes are used.

We described two approaches to non-rigid registration that can be applied to small animal images. First, we apply the mutual information based scale-space framework to non-rigid registration of mouse images. The results indicate that the scale-space based registration gives better skeletal as well as soft tissue alignment than a hierarchical multiscale approach. We used a non-parametric approach to estimating the distribution of intensities that defines the mutual information cost function, to avoid making assumptions about the form of the distribution. However, using this approach in conjunction with estimation of the non-rigid deformation field increases the dimensionality of the problem. We therefore describe an alternative approach that uses the log likelihood of the target given the deformed template wherein the likelihood is described by a Gaussian mixture model. This reduces the density estimation step to that of estimating the GMM parameters, thus reducing the overall dimensionality of the problem for images that have a few regions of distinct intensities. We evaluated the GMM approach using intramodality brain MR images, and multi-modality mouse MR/CT images. The results indicate that this GMM based approach is robust and yields more accurate registration accuracy compared to the non-parametric MI based approach. The GMM based approach can therefore be potentially applied for clinical as well as preclinical imaging.
In summary, we find that the information theoretic measures considered in this dissertation have potential for application in multi-modality imaging studies. Since there is ongoing research in constructing multi-modality imaging systems, the approaches described in this dissertation are relevant and could be used as a general framework for reconstruction and registration utilizing the available multi-modality information.

5.1 Future Work

The work presented in this dissertation can be extended to various problems in reconstruction and registration. A few of the possible extensions are:

1. Atlas based mouse segmentation: Mouse atlases such as the Digimouse described in [51] have a mouse CT or MR image with the various organs labeled by an expert. Automatic segmentation of a target image can be performed by registering the atlas to it, and applying the obtained deformation field to the labels thus obtaining the target image segmentation. This requires non-rigid registration, possibly of images from different modalities. The GMM based non-rigid registration described in this dissertation can be used to perform this registration of different subjects to the atlas for segmentation.

2. Positron range: The positron range model used in this work assumes that the range blurring is spatially invariant, which is not accurate especially at organ boundaries. Though the JE based approach is able to improve the resolution of the reconstructed image even with this assumption, using a more accurate range model in conjunction with the JE prior might improve the results further.
3. Application of information theoretic anatomical priors to diffuse optical tomography (DOT): We obtained some preliminary phantom results for application of the information theoretic anatomical priors described in this dissertation to the ill-conditioned problem of diffuse optical tomography [44]. This application can be explored further to improve the resolution of DOT.
References


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mouse: a 3d whole body mouse atlas from ct and cryosection data. *Physics in