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Abbreviations

GM	Grey matter
WM	White matter
CSF	Cerebrospinal fluid
PD	Proton density
FLAIR	Fluid attenuation inversion recovery
WML	White matter lesion
WMH	White matter hyperintensity
DAWM	Dirty appearing WM
MS	Multiple Sclerosis
AD	Alzheimer's Disease
PVE	Partial volume effect
VLR	Voxel-Wise Logistic Regression
MLE	Maximum likelihood estimation
MAP	Maximum a posteriori
SI	Similarity index
ICC	Interclass Correlation Coefficient
LL	Lesion load
CV	Cross validation
LOO-CV	Leave-one-out CV
KF-CV	K-fold CV
LOSO-CV	Leave-one-source-out
SVM	Support vector machine
K-NN	K-nearest neighbours
MRF	Markov random field
FPR	False positive reduction
PMF	Probability mass function
CDF	Cumulative density function

Notation

Variables		
y	feature	$\in \mathbb{R}$
\tilde{y}	standardized feature	$\in \mathbb{R}$
c	true class	$\in \{0, 1\}$
\hat{c}	estimated class	$\in [0, 1]$
β	logistic model feature weight	$\in \mathbb{R}$
γ	synthetic feature	$\in \mathbb{R}$
ρ	prior probability	$\in [0, 1]$
Indexing – e.g. arbitrary variable a		
k	feature index	$\in \{1, \dots, K\}$
n	subject index	$\in \{1, \dots, N\}$
t	iteration index	$\in \{1, \dots, X\}$
x	spatial location	$= [x_1, x_2, x_3]$
$a_n^{k(t)}(x)$	k^{th} feature; n^{th} subject; t^{th} iteration; location x	
Images & sets – e.g. arbitrary variable a		
a	one voxel, one feature, one subject	
\mathbf{a}	one voxel, all features, one subject	
$A(x)$	image in native space	
$A(x)$	image in standard space (MNI)	
$\mathbf{A}(x)$	image set: all features, one subject	
$\mathcal{A}(x)$	image set: one feature, all subjects	
$\mathcal{A}(x)$	image superset: all features, all subjects	
\mathbb{A}	full dataset: all features, subjects, voxels	

Chapter 1

Introduction

Digitization of medical imaging has facilitated innumerable advances in disease understanding and treatment. From multi-modal image fusion to image guided therapy, software tools now underpin research and clinical workflows in almost every domain of medical imaging.

This work concerns an unsolved segmentation problem in 3D brain magnetic resonance imaging (MRI), in which the objective is to automatically predict the class, or label, of every voxel (“volume pixel”) in the image. The objects of interest are white matter hyperintensities (WMH), non-cancerous brain lesions which are correlated with several neurodegenerative diseases. This chapter presents the motivation for automated WMH segmentation, gives a problem definition, and explores the previously proposed solutions.

1.1 Background

The brain is composed of three major classes of tissue: grey matter (GM), white matter (GM), and cerebrospinal fluid (CSF). Grey matter constitutes the peripheral surface of the brain – the cortex, approximately 5mm thick – as well as some deeper structures called the basal ganglia. It contains neuronal cell bodies, and performs the bulk of neural processing. The white matter is composed primarily of myelinated axons, and functions to relay information between different GM structures in the brain. The brain is surrounded by CSF, which provides mechanical and immunological defence. It is produced by the choroid plexuses in the ventricles of the brain – a series of 4 connected cavities.

1.1.1 Magnetic Resonance Imaging

Magnetic resonance imaging (MRI) provides superior and flexible brain tissue contrast versus computed tomography (CT) imaging, and is the primary modality for imaging brain disease. Whereas CT measures tissue density via attenuation of transmitted X-rays, which does not vary significantly among brain tissues, MRI measures a mutable combination of 3 tissue characteristics: the proton density (PD)¹, and T1 and T2 relaxation constants [1]. The physics of signal generation are described below.

In an MR scanner, a powerful magnetic field induces alignment of proton dipoles with the field. Only a tiny fraction of the total protons align, but they create a small magnetic field M_z which is distinct from the main field [2]. The aligned protons also rotate about the axis of alignment, imperfectly, like a spinning top; this is called precession, and the frequency of rotation is roughly homogeneous and proportional to the main field strength [2]. If a second magnetic field is applied which is 90° perpendicular to the first, and rotating at the precession frequency, the aligned protons can be forced into temporary alignment with this transverse rotating field, before decaying back towards their original state, as illustrated in Figure 1.1 [2]. This transient applied magnetic field is induced by a radio frequency (RF) pulse, and the rate at which the original magnetization M_z is regained is described by the tissue-specific T1 relaxation constant,

$$M_z = M_0 \left(1 - e^{-\left(\frac{t}{T_1}\right)} \right). \quad (1.1)$$

The T1 constant is dictated by the ability of protons in the tissue to transfer energy to bonded atoms and surrounding molecules, since this energy transfer defines the transition from the high energy transverse state to the low energy original state [2, 3]. Large macromolecules, membranes, and lipids are generally able to facilitate this energy transfer more effectively than small molecules like water, producing a shorter

¹ MRI can be used to image any nucleus with a net nuclear dipole, but proton (hydrogen) imaging is most common since hydrogen is biologically abundant and gives a strong signal intensity.

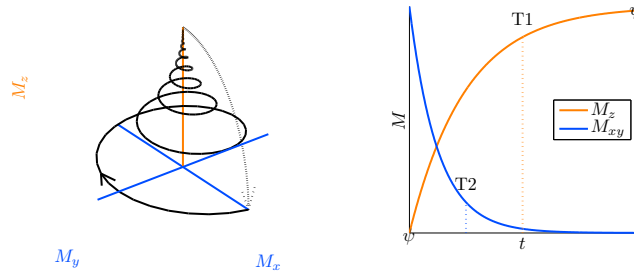


Figure 1.1: Visualization of T1 and T2 relaxation.

Table 1.1: T1 and T2 constants for brain tissues at 1.5 Tesla.

Tissue	T1 (ms)	T2 (ms)	$K[H]$ (a.u.)	Ref
WM	719 ± 33	73 ± 6	0.81 ± 0.03	[6]
GM	1165 ± 88	92 ± 11	0.98 ± 0.07	[6]
CSF	3337 ± 111	2562 ± 123	1.00 ± 0.07	[6]
WML	1124 ± 372	136 ± 79	—	[7] ^a

^a Estimated from Fig 1 supratentorial data (numerical results not given); \pm IQR, not SD; cf. § 1.1.2 for definition.

T1 [4]. For this reason, myelinated WM has a shorter T1 than GM, which in turn has a shorter T1 than CSF, which is mostly water [5].

The rate of decay of the transverse moment M_{xy} is actually not equal to the rate of regeneration of M_z . Rather, this is governed by the T2 relaxation constant,

$$M_{xy} = M_0 \left(e^{-\left(\frac{t}{T_2}\right)} \right), \quad (1.2)$$

which is always shorter than T1. This is because, in addition to T1 effects, the net rotating moment M_{xy} is eroded by proton dephasing. When precessing protons, having a net dipole, interact with other dipoles or charged particles, their rotational frequency can be increased or decreased, but overall less coherent, reducing the perceptible net magnetization M_{xy} [2]. In highly structured tissues like GM and WM, these interactions are more variable, dephasing is faster, and T2 is shorter [5]. In fluid environments like CSF, proton interactions are more homogeneous, yielding longer T2 [5]. For this reason, T2-weighted images are especially useful in identifying pathologies which degrade tissue structure, since they will have abnormally high T2 [5]. Both relaxation constants depend in a small way on the main magnetic field strength, measured in Tesla (T); T1 and T2 values for various brain tissues at 1.5T are summarized in Table 1.1.

Image acquisition involves sensing the transverse magnetization M_{xy} following proton excitation by an RF pulse. The problem is that this small signal decays very quickly due to proton dephasing, which occurs even faster than T_2 would predict due to a third factor, inhomogeneity in the main magnetic field [8]. The time constant for this decays is termed T_2^* , and its effects are usually undesirable [8]. As a result, M_{xy} is easily overpowered by the magnetic moment from the RF pulse, even after it is turned off, due to resonance. An important solution to this, called the spin-echo, was proposed by Erwin Hahn in 1950 [9]. If T_2^* for each proton is assumed to be constant, then reversing the direction of rotation at a time t should cause all protons to align again at exactly $2t$. Therefore, at $2t$ the transverse magnetization M_{xy} – the image signal – manifests again for sensing, no longer confounded by RF coil resonance [9].

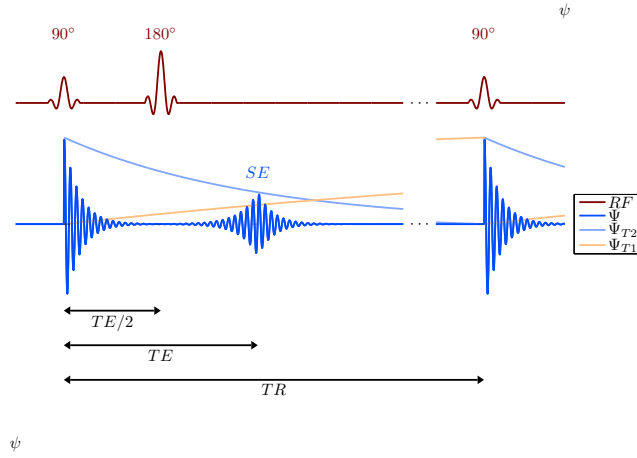


Figure 1.2: RF and MR signal for a basic Spin Echo sequence.

This second signal is called the Spin Echo (SE), and the interval $2t$ is termed the echo time (TE). Reversing the direction of rotation can be achieved by a 180° RF pulse at time $TE/2$, in the same way the original excitation is achieved using a 90° RF pulse (amount of rotation is proportional to the energy of the pulse). Acquisition of an entire image requires repetitions of this sequence with an interval called the repetition time (TR). An example spin echo sequence, showing TE and TR , as well as $T1$ and $T2$ decay, is illustrated in Figure 1.2. Spatial encoding for creation of 2D and 3D images requires the use of additional electromagnetic gradients; however this topic is omitted here since it is quite involved, and not essential to the current work².

Using these principals, the nature of MR image contrast can finally be understood. That is, the signal intensity Ψ for a spin echo sequence at location x can be described by the following 3-term equation,

$$\Psi_{SE}(x) = \left[K[H](x) \right] \left[e^{-\left(\frac{TE}{T2(x)}\right)} \right] \left[1 - e^{-\left(\frac{TR}{T1(x)}\right)} \right], \quad (1.3)$$

where K is scaling factor, and $[H]$ denotes the proton density. If TR is chosen to be relatively long, then the longitudinal magnetization M_z is allowed to recover completely after each repetition, the third term tends towards 1 for all tissues, and differences in tissue specific $T1$ are nullified. Similarly, if TE is relatively short, then M_{xy} has little time to dephase, the second term is maintained close to 1, and differences in $T2$ are nullified. In order to emphasize differences in $T1$, therefore, TR can be chosen shorter; for $T2$ -weighted contrast, TE can be chosen longer; and if differences in $[H]$ (proton density, PD) are to be emphasized, TR can be kept long and TE short. An example MRI slice using each of these image sequences is shown in Figure 1.3a, 1.3b, and 1.3c.

² The interested reader is directed to this comprehensive resource on the topic: <http://mri-q.com/>

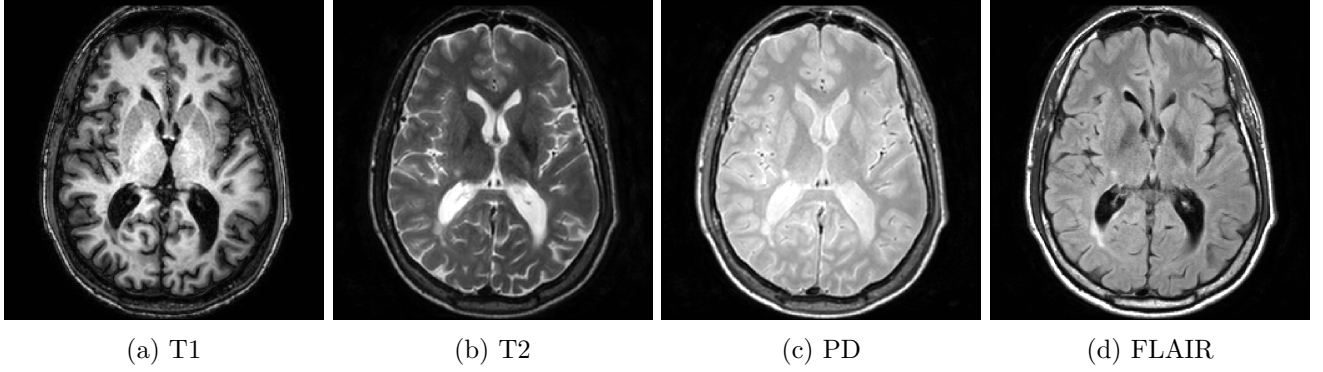


Figure 1.3: Example MRI image set with WMH pathology; from [12].

For identifying WML, T2-weighted images were conventionally used, since the lesions appear bright. However, CSF in the sulci and ventricles also appears bright on T2 images, making delineation of lesions – especially periventricular ones – difficult in T2 images (Figure 1.3b). To solve this problem, an adaptation of the spin echo RF pulse sequence can be used, called an inversion recovery (IR) [10]. In this sequence, an additional 180° inverting RF pulse is added before the 90° pulse, so that the longitudinal magnetization M_z is inverted, then recovers to the original state, passing for a brief moment through zero net magnetization. The rate of recovery is governed by $T1$, so it is tissue specific. Furthermore, if the 90° pulse is applied at the instant of zero net magnetization, no transverse moment will develop, nor the subsequent spin echo. Therefore this time interval, called the inversion time (TI), can be chosen to null the signal from any tissue with a unique $T1$. The equation governing the image signal simply adds an inversion term,

$$\Psi_{IR}(x) = \left[K[H(x)] \right] \left[e^{-\left(\frac{TE}{T2(x)}\right)} \right] \left[1 + e^{-\left(\frac{TR}{T1(x)}\right)} - 2e^{-\left(\frac{TI}{T1(x)}\right)} \right]. \quad (1.4)$$

This inversion principal is now often used to null the signal from CSF, especially for delineation of WMH, in a sequence called FLuid Attenuation Inversion Recovery (FLAIR) [11]. FLAIR images are usually T2-weighted. Figure 1.3d shows an example FLAIR image, where a WMH can be seen, posterior to the occipital horn of the left lateral ventricle, much more clearly than in the T2 image.

1.1.2 White Matter Disease

“Normal” ageing of the brain is characterized by a variety of physical and cognitive changes. Memory, synaptic plasticity, and brain volume decline, with observable effects on cognitive function [13, 14]. Brain ageing is also expedited in many patients by neurodegenerative diseases targeting the white matter, including Alzheimer’s disease (AD), cerebrovascular disease, and in rare cases Multiple Sclerosis (MS). While the etiologies of these diseases are not yet fully understood, there is considerable evidence to suggest

that they are intertwined [15, 16, 17, 18].

Cerebrovascular disease describes changes to blood vessels in the brain which increase the risk of ischemic injury – a reduction in blood flow due to vessel occlusion or hemorrhage. Ischemic injuries include major events (stroke) [19], transient ischemic attacks [20], and chronic hypoperfusion due to small vessel disease [21]. In all such events, neuronal death occurs from insufficient nutrient supply [19]. Strokes involving major cerebral arteries can be fatal, and post-event quality of life in survivors is highly variable [22]. In the less dramatic courses, clinically quiet disease progression can lead to personality changes, memory loss, and reduced cognitive ability; such changes are termed vascular dementia [23].

Alzheimer’s Disease is another subclass of dementia with similar symptoms; in fact it is the most common type, affecting about 6% of the population over age 65 [24]. The cause of Alzheimer’s disease is hotly debated. Two 30-year-old theories linking the disease to the build up of amyloid β protein and misfolded protein τ have been widely supported by correlational studies [25, 26, 27], but have lacked clear mechanisms of injury until recently. It is now thought that amyloid β oligomers interfere with neuronal mitochondria and synapse function, leading to cell death [28, 29], while aberrant τ proteins disrupt microtubules necessary for intraneuronal transport [27]. During the search for these mechanisms, competing theories implicating vascular injury [18], immune response [17], and blood brain barrier disruption [30] have emerged, painting the picture of a more complex disease.

The pathophysiology of multiple sclerosis is similarly unclear, though genetics are a necessary factor, and it is known that symptoms arise from erosion of myelin – a fatty insulating layer surrounding axons which is critical for normal neuron firing [31]. Several theories hypothesize either that this damage is driven by autoimmune attack, followed by neuronal dysfunction and death, or that neurodegenerative changes stimulate recruitment of immune cells as part of the usual response to injury [32, 31]. Recent evidence favours the former mechanism, particularly with inflammatory injury as the initiating event [33, 34].

Connecting all these diseases are white matter lesions (WML, AKA Leukoariosis), which represent the macroscopic changes to brain tissue in regions of white matter damage [15, 35, 36]. WML are very common in elderly populations, and a small volume of lesion does not necessarily implicate one of the above diseases; in one study of 1077 subjects aged 60-90, 95% had at least one WML [37]. WML appear as bright tissue regions in T2-weighted MRI due to some combination of inflammatory injury and degradation of tissue structure [35, 36]; in this imaging context, WML are often called white matter hyperintensities (WMH). Lesions are often focal, as opposed to diffuse, but there is evidence to suggest that surrounding regions of moderate hyperintensity, sometimes called “dirty appearing white matter (DAWM)”, are also related to the diseases [38]. As biomarkers of the most common WM diseases – conditions with unsolved

etiologies and inadequate treatments – WML are of special interest to many brain researchers. The next section discusses how they are used.

1.1.3 MRI in White Matter Disease

MR imaging plays important roles in diagnosis and research of white matter diseases. Typical MRI protocols include T1, T2, and FLAIR sequences, though only the latter two sequences depict WML as hyperintense [39, 40]. Depending on the disease and context, WMH can be quantified in several ways, including binary criteria (e.g. is there a lesion in a specific location) [41, 23], rating scales (e.g. a summary of several criteria) [42], or explicit manual segmentation of the lesions by an expert [43].

WMH are arguably most important in MS. Particularly since WMH are more specific to this disease in younger patients, WMH have long been used in the diagnosis of MS, and can even be used to replace some clinical criteria, as in the 2010 McDonald Criteria [41]. MRI can also be used to discriminate between MS subtypes, which stratify disease aggression and course [41, 44, 45]. In numerous clinical trials, WMH have also been used as biomarkers of treatment efficacy [46, 47, 48], since WMH have been shown to be more sensitive to disease progression than clinical features in certain subtypes [49]. In fact, despite the central role of MRI in management and research of MS, there exists a so-called “clinico-radiological” paradox, which is the surprisingly limited correlation between WMH and clinical MS symptoms like physical and cognitive impairment [50]. However, this only strengthens the case for continued WMH research, particularly considering the recommendations by Mollison et al. in [50] to standardize image analysis in order to better understand the paradox.

In dementia (including vascular and AD), WMH are used to discriminate between disease subtypes during diagnosis. For example, the presence of at least one WML was deemed *necessary* for diagnosis of vascular dementia in 1993 [23], and subsequent revisions to these widely used criteria (NINCDS-ADRDA) have added this feature as an *exclusionary* criteria for AD [51]. While diagnosis of additional dementia subtypes may be improved using imaging [52, 53], diagnosis of the most prevalent – AD – continues to be based on clinical features alone [54]. As a result, WMH have not been used as an endpoint to any AD clinical trial. In fact, only recently have specific standards for use of WMH in vascular dementia studies been outlined [40, 36], with some subsequent uptake [55]. And yet, a 2010 meta-analysis found that WMH in brain MRI were independently correlated with stroke risk, dementia (including AD) and death [15], suggesting that much more can be done to make use of WMH as hallmarks of neurodegenerative disease.

1.2 Problem Statement

White matter hyperintensities, as ubiquitous biomarkers of several diseases with unsolved pathophysiology, are of great interest to brain researchers. Segmentation of WMH, compared to visual rating scales, provides a finer resolution for quantification of lesion load, and gives the explicit spatial distribution of pathology. This spatial information can be very useful, since diagnostic criteria often consider lesion location [52] and there are several correlations between lesion location and suspected etiology of WMH [56, 36].

Unfortunately, manual segmentation of WMH is laborious, and subject to large inter- and intra-rater variability, as reported in several works. Table 1.2 summarizes these reports, where similarity index ($SI \in [0, 1]$) is a measure of voxel-wise agreement, and interclass correlation coefficient ($ICC \in [0, 1]$) measures total volume agreement (cf. § 2.6 for definitions). Table 1.2 also gives the results using four semi-automated approaches, since these methods are reported to reduce variability and task time over strictly manual segmentation. Yet, for very large scale research studies, any approach requiring human intervention would be prohibitively time consuming and subjective.

Therefore, a fully automated algorithm to segment WMH in MRI is required. Such an algorithm would have, by construction, perfect repeatability, and consistent bias – which is especially important for perceiving small changes in longitudinal studies [57]. Additionally, while an automated approach may not necessarily be faster than manual or semi-automatic segmentation on a per-case basis, it could be run on several computers in parallel, yielding significant overall speed up.

Furthermore, while T1, T2, and FLAIR sequences are typically recommended for both MS and dementia investigations [39, 40, 45], FLAIR sequences are at least as sensitive as T2 images for the detection of WML³. As noted above, FLAIR images also have the advantage of easily distinguishing WMH from confounding CSF hyperintensity, which is important for highly prevalent periventricular lesions, and also for excluding lacunar infarcts [60, 61]. Consequently, it should be feasible to detect WMH using FLAIR MRI alone. This has several advantages, including minimizing the required MR sequences available during retrospective analyses, decreasing cost and scan time in prospective studies, and eliminating the need for intra-subject image registration if sequences are acquired at different resolutions (as is often the case).

³ Early studies exploring the utility of FLAIR sequences may contradict this claim [58, 59], but FLAIR imaging has since improved [36].

Table 1.2: Mean inter-rater agreement measures for manual and semi-automated WMH segmentation reported in previous works.

	Ref	Raters	Data	SI	ICC
Manual	[62]	5	10 images	0.64	—
	[63]	2	6 images	0.75	—
	[64]	2	120 slices	0.83	0.96
	[43]	3	50 images	0.66	0.97
Semi-Automated	[65]	1	16 images	—	0.99
	[66]	1	2 images	0.70	—
	[67]	1	33 slices	0.78	—
	[68]	2	30 images	0.78	—

1.2.1 Objective

The primary objective of this thesis is to develop an algorithm for fully automatic segmentation of WMH, using FLAIR MRI alone. Secondary objectives include:

- analysis of the limitations of prior work in this area;
- exploration and definition of appropriate cross validation techniques for the task;
- validation of the proposed algorithm on a large and heterogeneous database of FLAIR images.

1.2.2 Challenges to Automatic Segmentation

While fully automated segmentation of WMH is attractive, translation of expert knowledge into algorithmic constructs is difficult, and often requires assumptions which induce sensitivity of the model to seemingly extraneous image features. Moreover, human understanding of MR acquisition physics help radiologists to distinguish WML from image artifacts. Thus, there are several challenges to automatic segmentation. These can be summarized as follows:

1. Overlapping distributions:

Using the relaxation constants in Table 1.1 in conjunction with the FLAIR signal equation (1.4), the predicted intensity distributions of GM and WMH can overlap, depending on the choice of TE, TR, and TI (cf. Table 3.2 for typical values and § A.1 for modelling). As a result, the intensity of image voxels alone cannot be used to determine their class [69].

2. Bias field:

The most common image artifact in MRI is due to inhomogeneity in the main magnetic field or RF coil during acquisition, which is difficult to eliminate in strong electromagnets; this creates a low

frequency variation in signal intensity over the imaged volume [70, 71]. The overall effect is that the same tissues may have different graylevels in different locations, further confounding the uniqueness of WMH graylevels [36].

3. **DAWM:**

Most of the inter-rater disagreement in manual segmentation of WMH is arguably due to ambiguity of pathological extent at the lesion borders, where the core lesion meets so-called DAWM [38]. If human judgement of this boundary is difficult, then programmatic definitions could be expected to be similarly challenged.

4. **Partial volume effect:**

With finite image resolution, voxels located on tissue boundaries will inevitably contain tissues of two or more tissues. This is known as partial volume effect (PVE), and the resulting signal intensity can be modelled as a linear mixture of the components [72]. Niessen et al. [73] show that inadequate modelling of PVE can result in significant errors in tissue segmentation, though the widely reported 30% figure from this work is derived from unrealistic conditions.

5. **Artifacts:**

Due to the complexity of signal acquisition, there are several artifacts which can manifest in MR images. Artifacts which appear hyperintense in T2 images (including FLAIR) are of particular importance to the current work, since these confound bright pathologies, and must therefore be excluded using other features; the most notable artifacts include [36]:

- CSF flow artifacts – ventricular hyperintensities resulting from movement of magnetically polarized CSF fluid during the inversion interval (cf. § 1.1.1) [74];
- Perivascular spaces – minuscule spaces adjacent to cerebral vessels whose properties differ from ventricular and sulcal CSF, and are therefore not attenuated in FLAIR images [36];
- Motion artifacts – artifacts which originate during frequency-domain encoding of spatial image content with subject motion, which is more common in MRI due to long acquisition times (several minutes); these typically manifest as high frequency “ringing” artifacts [75].

6. **Image variability:**

There are a large number variable characteristics of MR images; some of these can be selected at acquisition time based on time constraints, and physician preferences, while others are immutable. “Image variability” is taken to comprise:

- differences in image contrasts (and tissue graylevel distributions), due to selection of MRI parameters;

- differences in image resolution (voxel size);
- differences in MRI scanner, including field strength and proprietary image reconstruction;
- inter-subject anatomical variability and lesion heterogeneity.

Modelling this immense gamut of possible image characteristics (e.g. using parametric distributions or task-specific assumptions) represents perhaps the most challenging aspect to automated image analysis. Some specific impacts will be further discussed in § 1.3.3.

An optimal WMH segmentation algorithm will therefore consider and address each of these challenges.

1.3 Prior Work

This endeavour is far from original. Efforts to automate segmentation of WMH date back to 1990 [76] and the task has been the subject of several major reviews in 2012 [77, 69], 2013 [78], and 2015 [79]. The task has also been featured in four international competitions at the MICCAI (Medical Image Computing and Computer Assisted Intervention) Conference – 2008 [80], 2016 [81], and 2017 [12] – and the ISBI (International Symposium on Biomedical Imaging) Conference – 2015 [57] – in which researchers vie to produce the best segmentation algorithms (for more on competitions, cf. § 3.1.2). A summary of many of the proposed approaches is given in Table 1.3⁴. Below, the most important contributions in this area are introduced and summarized.

1.3.1 Segmentation Models & Features

Segmentation models represent a mapping from the content of an observed image to an image of labels or classes – in this case, tissues. The output class image comprises an estimated label for each observed voxel, or, in probabilistic models, the probability of each class for each voxel. As in many classification problems, models can be described as either supervised or unsupervised. Supervised models have relatively large capacity to model arbitrary mappings, but learn a mapping relevant to the current task using feedback from labelled examples (i.e. by a human). Unsupervised models, by contrast, are usually problem-specific, and leverage prior knowledge and the image features to predict the label image; they do not require labelled data for optimization, at least in principle.

Features used for segmentation can be derived from individual voxels (e.g. graylevel), groups of voxels

⁴ A more detailed and interactive version of this table is available at www.uoguelph.ca/~jknigh04/wmlseg/table.html

Table 1.3: Summary of previous approaches to WMH segmentation with respect to image variability and reported performance (SI).

#	Ref.	Year	Authors	MRI Sequences	I	S	SI
1	[82]	2001	Van Leemput et al.	T1, T2, PD	20	1	0.51
2	[83]	2001	Jack et al.	FLAIR	39	1	
3	[84]	2002	Zijdenbos et al.	T1, T2	10	1	0.6
4	[85]	2004	Anbeek et al.	T1, T2, PD, FLAIR, IR	20	1	0.61
5	[86]	2005	Anbeek et al.	T1, T2, PD, FLAIR, IR	10	1	0.78
6	[87]	2005	Admiraal-Behloul et al.	T2, PD, FLAIR	100	1	0.75
7	[88]	2006	Lao et al.	T1, T2, PD, FLAIR	45	1	
8	[89]	2006	Wu et al.	T1, T2, PD	12	1	
9	[90]	2006	Sajja et al.	T2, PD, FLAIR	23	1	0.78
10	[62]	2006	Harmouche et al.	T1, T2, PD	10	1	0.61
11	[91]	2008	Khayati et al.	FLAIR	20	1	0.75
12	[92]	2008	Wels et al.	T1, T2, FLAIR	6	1	0.57
13	[93]	2008	Herskovits et al.	T1, T2, PD, FLAIR	42	2	0.6
14	[94]	2008	Bricq et al.	T2, FLAIR	25	2	
15	[95]	2008	Dyrby et al.	T1, T2, FLAIR	362	10	0.56
16	[96]	2008	Souplet et al.	T1, T2, FLAIR	25	2	
17	[63]	2009	Boer et al.	T1, PD, FLAIR	20	2	0.72
18	[97]	2009	García-Lorenzo et al.	T1, T2, PD	10	1	0.63
19	[98]	2009	Akselrod-Ballin et al.	T1, T2, PD, FLAIR	41	1	0.53
20	[99]	2009	Schwarz et al.	T1, T2, PD	165	2	
21	[100]	2010	Gibson et al.	T1, T2, FLAIR	18	1	0.81
22	[101]	2010	Shiee et al.	T1, FLAIR	10	1	0.63
23	[102]	2010	Scully et al.	T1, T2, FLAIR	17	1	
24	[103]	2011	García-Lorenzo et al.	T1, T2, FLAIR	10	1	0.65
25	[104]	2011	Geremia et al.	T1, T2, FLAIR	20	2	
26	[105]	2011	Smart et al.	T1, FLAIR	30	1	
27	[106]	2012	Samaille et al.	T1, FLAIR	67	6	0.72
28	[107]	2012	Khademi et al.	FLAIR	24	1	0.83
29	[108]	2012	Schmidt et al.	T1, FLAIR	53	1	0.75
30	[109]	2012	Abdullah et al.	T1, T2, FLAIR	61	3	
31	[110]	2013	Sweeney et al.	T1, T2, PD, FLAIR	111	1	0.61
32	[111]	2013	Datta et al.	T1, T2, FLAIR	90	3	
33	[64]	2013	Steenwijk et al.	T1, FLAIR	40	2	0.8
34	[112]	2014	Khademi et al.	FLAIR	25	1	0.78
35	[113]	2014	Ithapu et al.	T1, FLAIR	38	1	0.67
36	[114]	2014	Yoo et al.	FLAIR	32	2	0.76
37	[115]	2015	Harmouche et al.	T1, T2, PD, FLAIR	100	35	0.56
38	[116]	2015	Guizard et al.	T1, T2, PD, FLAIR	108	32	0.6
39	[117]	2015	Jain et al.	T1, FLAIR	20	1	0.67
40	[118]	2015	Tomas-Fernandez et al.	T1, T2, FLAIR	51	2	
41	[119]	2015	Wang et al.	T1, T2, FLAIR	70	2	0.84
42	[120]	2015	Roy et al.	FLAIR	38	3	0.56
43	[121]	2015	Brosch et al.	T1, T2, FLAIR	20	2	0.36
44	[122]	2015	Fartaria et al.	FLAIR	39	1	0.55
45	[123]	2015	Deshpande et al.	T1, T2, PD, FLAIR	52	1	0.5
46	[124]	2015	Roura et al.	T1, FLAIR	20	2	0.34
47	[125]	2016	Knight et al.	FLAIR	15	3	0.7
48	[126]	2016	Mechrez et al.	T1, T2, FLAIR	20	2	0.31
49	[127]	2016	Strumia et al.	T1, FLAIR	20	3	0.52
50	[128]	2016	Griffanti et al.	T1, FLAIR	130	2	0.76
51	[129]	2017	Valverde et al.	T1, FLAIR	33	2	
52	[130]	2017	Dadar et al.	T1, FLAIR	80	3	0.62
53	[131]	2017	Zhan et al.	T1, T2, FLAIR	50	2	0.76

Abbreviations. I: number of MR image sets used for validation; S: number of MRI scanners used for validation; SI: reported validation similarity index.

(e.g. local mean graylevel), the entire image (e.g. a histogram feature), spatial location (e.g. coordinates in a standardized space), or prior knowledge (e.g. class prior probability). It is often useful to imagine the space spanned by all possible values of all features; this is called the feature space, Each observed voxel, having a unique value for each feature, therefore represents a unique location in this space, The task of segmentation is therefore to divide the feature space into subspaces corresponding to each class, In probabilistic models, these subspaces are better described as distributions of each class over the features.

Previous approaches to WMH segmentation have generally employed three types of features:

- **Graylevel:** graylevels of MRI sequences, often following standardization (e.g. T1, T2, PD, FLAIR);
- **Prior:** prior tissue probability, often derived from a coregistered prior image (e.g. ICBM [132]);
- **Spatial:** spatial location, often normalized to a common space (e.g. x_1, x_2, x_3).

Additional features types are rarely used, since the combination of the above features are typically the only features employed by human raters, At least one graylevel feature is always used, since it is the only image-specific information (i.e. the evidence).

1.3.2 Proposed Methods

The specific methods proposed for WMH segmentation are now reviewed.⁵

Thresholding Techniques

Since WMH are brighter than healthy brain tissue in FLAIR images, many unsupervised works have used thresholding of FLAIR intensities as the initial lesion segmentation. For example, in the works by Jack et al. [83], Boer et al. [63], and Smart et al., [105] optimal FLAIR thresholds are empirically estimated relative to histogram statistics, though Boer et al. use only estimated GM voxels in the histogram. Gibson et al. use a conservative FLAIR threshold initially, but then classify the remaining voxels using Fuzzy C Means clustering [100]. Samaille et al. use nonlinear diffusion filtering and watershed segmentation, before classifying candidate regions based on a FLAIR image threshold. Yoo et al. estimate the optimal threshold for FLAIR images using histogram statistics, derived from a regression model primarily considering the total lesion load [114]. In works by Khademi et al., a peak in the conditional probability of edge content on graylevel is used to predict the transition between healthy tissue and lesion [112, 134, 125].

⁵ This section copied verbatim from a paper in submission [133]

Mixture Models

Most other unsupervised approaches are probabilistic models, often framed as a mixture model. The work by Van Leemput et al. [82] uses a similar framework as the early work by Ashburner and Friston [135], later incorporated into the SPM “segment” tool [136], which jointly estimates Gaussian graylevel distributions for each tissue class, and also bias field, using expectation maximization. In the model by Van Leemput et al., distribution parameters are estimated using outlier-insensitive estimators, and WMH are derived from model outliers using heuristic rules. The predicted classes are also smoothed spatially using a Markov Random Field (MRF).

Similar works by Bricq et al. [94], Schmidt et al. [108], Jain et al. [117], and Roura et al. [124] use parametric mixture models to predict WMH as model outliers, and all but [124] embed the model in a MRF. Khayati et al. [91] and Subbanna et al. [137] also use MRF-constrained mixture models, but model WMHs as a Gaussian-distributed tissue class, rather than as outliers. In the works by Harmouche et al., parametric distributions are also used to model lesions, but such distributions are parameterized independently per brain region, in order to reflect lobe heterogeneity; a MRF is again used for regularization [62, 115]. Schwarz et al. again employ a Bayesian MRF model, but use lognormal distributions for WM and WMH [99]. Souplet et al. use an augmented mixture model which includes partial volume averaging classes and an outlier class to perform initial brain tissue segmentation; WMH are subsequently classified using a FLAIR intensity threshold after contrast enhancement [96]. The work by Herskovits et al. is much the same, but uses statistical information from training data to classify lesions (i.e. it is supervised) [93]. More recently, Graph-Cuts have been used in conjunction with mixture models, as in the works by García-Lorenzo et al. [97], Tomas-Fernandez and Warfield [118], and Strumia et al. [127].

The Lesion-TOADS method by Shiee et al. [101], a lesion-specific adaptation of the TOADS algorithm [138], presents an entirely new non-Gaussian paradigm for modelling class distributions, and incorporates topological energies in the objective function. Other proposed unsupervised methods have used clustering by Fuzzy C-Means, including the works by Admiraal-Behloul et al. [87], Gibson et al. [100], and Valverde et al. [129].

Classic Supervised Methods

Many early supervised methods used K-Nearest Neighbours (K-NN) for voxel-wise WMH classification. Anbeek et al. used a K-NN model with features derived from spatial coordinates and voxel intensities from several modalities [85, 86]. In the works by Wu et al. [89], Steenwijk et al. [64], and Fartaria et al. [122],

spatial coordinates are substituted for tissue priors as K-NN features. In the recently proposed BIANCA algorithm by Griffanti et al. [128], spatial coordinates are added back, along with some patch-based features.

Other works have also explored Support Vector Machines (SVM) for classification. The works by Lao et al. [88], Abdullah et al. [109], and Scully et al. [102] each use a selection of intensity features, neighbouring intensities, tissue priors, morphological, and texture features with an SVM classifier. Several more recent works have used decision tree-based classifiers, including Random Forest (RF) and AdaBoost. Akselrod-Ballin et al. [98] employ over 30 features for multi-scale image representation and classify voxels using RF. Both Geremia et al. [104] and Roy et al. [120] use a combination of intensity and tissue prior features to train a RF classifier, whereas Wels et al. [92] use a large number of Haar-like features to train an AdaBoost model. Ithapu et al. [113] explore the use of texton features in both SVM and RF models.

Logistic regression models have also gained popularity recently. In the OASIS model by Sweeney et al. [110], image intensities from T1, T2, PD, and FLAIR sequences are used individually, in multiplicative combination, and with Gaussian blurring as predictors for a global set of logistic regression parameters. In the work by Zhan et al. [131], a similar logistic model is fitted using only the raw T1, T2, and FLAIR intensities, while bias correction is performed as preprocessing and spatial smoothness using MRF post processing. In the work by Dadar et al. [130], spatial and intensity features from a flexible selection of MR sequences are used to train a linear regression model, the results of which are thresholded to give the lesion prediction. Still more works have proposed other supervised models, including nonparametric Parzen classifiers [90].

Deep Learning

A number of deep learning approaches have also been proposed, though their permeation in this problem space is surprisingly limited. Both Zijdenbos et al. [84] and Dyrby et al. [95] train fully-connected voxel-wise Neural Networks with a selection of intensity, spatial, and tissue prior features to predict the lesion class. In contrast, Brosch et al. [121] construct a more modern deep convolutional model, which is capable of capturing both local and global dependencies.

External Toolboxes

Many of the proposed methods use registration, brain extraction, bias field correction, and segmentation tools available in freely available toolkits; these include the SPM⁶ toolkit [90, 95, 98, 105, 108, 114, 113, 120, 129] and the FSL⁷ toolkit [93, 100, 111, 64, 110, 120, 119, 128, 131], as well as bias correction by the N3/4 [139] algorithm [84, 62, 122, 116, 115, 126, 129, 130, 131].

1.3.3 Limitations

Despite over 50 proposed algorithms and several competitions, no WMH segmentation algorithm has clearly emerged the superior method, nor has any been taken up for use in the wider research community. This is contrasted with other neuroimaging tasks, where several robust tools noted above are now regularly used in analysis pipelines – e.g. N3/4 [139] for bias field correction, BET for brain extraction [140], SPM Segment [136] / FSL FAST [141] for healthy brain segmentation, SPM Normalize [136] / FSL FNIRT [142] for registration. There are two hypothesized reasons for this gap in WMH segmentation tools.

First, very few of the proposed methods have been released as either open-source code or compiled applications. Researchers may not want to release source code for reasons related to intellectual property, or the additional work of ensuring robustness and writing documentation. Yet the field of deep learning illustrates how these practices can accelerate progress in the field enormously. Similarly, compiling applications for cross-platform compatibility is no small feat, though there are many examples for SPM extensions⁸, as well as events by NA-MIC (National Alliance for Medical Image Computing) for development of 3D Slicer modules⁹.

Second, very few of the WMH segmentation methods have been validated on large, multi-centre databases: of the 53 works reviewed (Table 1.3), less than half use more than one scanner for validation, and only 4 use more than three. As noted in § 1.2.2, there are several sources of image variability in MRI, and both supervised and unsupervised methods can be sensitive to these factors, as noted by several authors [77, 110, 130]. Therefore, while many of the proposed methods may be of use for in-house work (i.e. with images from a consistent source), there can be little confidence that they will perform as reported on data from new sources (generalization performance).

⁶ <http://www.fil.ion.ucl.ac.uk/spm/>

⁷ <https://fsl.fmrib.ox.ac.uk/fsl/>

⁸ <http://www.fil.ion.ucl.ac.uk/spm/ext/>

⁹ <https://na-mic.org/wiki/Events>

In supervised models, graylevel features must be standardized, since the MRI intensity scale is not consistent across scanners or scan parameters, due to the complexity of signal acquisition [143]. However, this is not an easy task. For example, Steenwijk et al. validate a supervised WMH segmentation algorithm using same-scanner training and testing for two different scanners independently (mean SI = 0.75, 0.84), after variance scaling of intensity features [64]. Yet, a follow-up experiment which saw the method trained on one scanner and tested on the other showed a precipitous drop in performance to mean SI = 0.50.

Unsupervised models also have parameters which can be inadvertently over-tuned to data from one or two sources. For example, mixture models which classify lesions as outliers often employ an outlier definition which depends on mixture model parameters (e.g. tissue graylevel mean and variance), which in turn are subject to MR slice thickness, noise level, and contrast [82, 96, 103, 124]. Graylevel thresholding techniques [83, 105, 106, 108, 112] are similarly affected by changes in image properties.

It is worth noting three works¹⁰ which run counter to this trend, demonstrating robust validation of their proposed methods. These works are summarized in Table 1.4. Perhaps not surprisingly, these papers report lower performance (mean SI ≤ 0.60) than other works; as a result, these algorithms would not likely be used for clinical research. Yet, even these works do not optimally estimate the model generalization performance for data from new scanners, as will be discussed in § 3.1. Moreover, the proposed algorithms in these papers require at least three MRI sequences, which fails the objectives of the current work.

Table 1.4: Works demonstrating excellent validation of a WMH segmentation algorithm.

Ref.	Year	Authors	I	S	SI
[95]	2008	Dyrby et al.	362	10	0.56
[116]	2015	Guizard et al.	108	32	0.60
[115]	2015	Harmouche et al.	100	35	0.56

Abbreviations. I: number of MR image sets used for validation; S: number of MRI scanner-parameter combinations used for validation; SI: reported validation similarity index.

1.4 Contributions

This thesis therefore aims to produce a WMH segmentation algorithm which can be used on FLAIR MRI from any source, and to characterize the expected performance on unseen data. The major contributions are as follows:

¹⁰The work by Samaille et al. (2012) [106] is also a good candidate, having used 6 scanners for validation; however, 43 of the 67 images (64%) come only from one scanner, reducing the robustness of generalization results.

1. A review and critique of the previously proposed WMH segmentation algorithms, especially with respect to expected performance on unseen data;
2. Voxel-Wise Logistic Regression (VLR): a new FLAIR-only WMH segmentation algorithm;
3. Leave-One-Source-Out Cross Validation (LOSO-CV): a validation framework which accurately characterizes the generalization performance of medical image analysis methods;
4. Extensive validation of the proposed method and its components.

The remainder of this thesis is organized as follows: Chapter 2 motivates and develops the voxel-wise logistic regression model, including expected challenges and solutions with this approach; Chapter 3 explores optimization of model components through experiment, and then presents segmentation performance results under various cross validation schemes; Chapter 4 discusses the performance results, summarizes the contributions, and highlights avenues of future work.

Chapter 2

Methodology

In this section we motivate and introduce the proposed model, and explore its parametrization. Wherever possible, concepts related to segmentation and classification will be described in their most general sense, since many of these may also apply to other image analysis and machine learning tasks.

2.1 Model Design

As suggested in § 1.3.2, three general paradigms have emerged for segmentation of brain MRI. The first and most popular is the pipeline, in which sub-tasks are completed in sequence, such as: pre-processing, classification, post-processing. This approach permits a flexible model definition which can incorporate existing algorithms for individual steps. Most thresholding and classic supervised models fall under this category. The main drawback of this approach is that some steps could be improved by the results of downstream steps. For example, tissue classifying modules typically assume that the bias field is already corrected, but bias field estimation can be more accurate if the tissue segmentation is known.

The second paradigm, a unified generative model, aims to solve this chicken-egg problem. The segmentation is parameterized in one integrated probabilistic model, which often combines the input images with tissue prior probability images, a bias field model, and smoothness terms. Parameters of each sub-model are estimated using several expectation maximization (EM) iterations before the final segmentation is inferred. The challenges to this approach include balancing model complexity with estimability, robust convergence issues, and reduced ability to include external tools.

The final paradigm, deep learning, uses error back-propagation to update thousands of parameters in a large, relatively unstructured model mapping the input MRI to the output segmentation image. Using

so-called “end-to-end” training, there is no guarantee that the usual sub-components (e.g. bias field, tissue graylevel distributions, etc.) will be estimated, though such elements could be expected to develop in the internal relationships if they are relevant to the task at hand. Most deep learning models still require standardization in space and graylevel, and more importantly, large training datasets – which are rare in medical imaging.

For several reasons, the supervised pipeline approach was selected for this work. First, preliminary work drawing on existing algorithms [112, 144] showed the feasibility of several relatively simple thresholding methods, which confer robustness through simplicity. Second, the parametric assumptions required by unified probabilistic models may be challenged by data from multiple sources [82]. Third, only a small number of training cases were available during initial development, which limited the feasibility of deep learning approaches. Finally, time constraints favoured the incorporation of existing tools to address challenges like bias correction and image registration, which would have been otherwise difficult to develop in a unified model.

2.1.1 Features

Many FLAIR-only WMH segmentation algorithms use a thresholding technique: map a single graylevel feature directly to a class or class probability (e.g. “healthy” or “lesion”). Such models often have complex methods of deriving this mapping (e.g. using mixture model parameters [124], histogram features [114], or conditional edge probability [145]), but the final rule is applied equally to the entire graylevel image, usually the FLAIR sequence. The most common challenge for these methods is a high number of false positives (cf. § 2.6 for definitions), since several artifacts and GM can overlap the WMH intensity distribution. As a result, nearly all previously proposed thresholding algorithms employ heuristic “false positive reduction” (FPR) rules as post-processing to remove these errors.

Preliminary investigations sought to characterize the spatial distribution of false positives, in order to explore possible FPR rules. Using a database of 96 FLAIR images in MNI space (cf. § 3.1.3 for details), the optimal threshold for each image was calculated¹ and used to generate binary class images (0 = healthy, 1 = lesion). Comparing these results with the manual segmentations, the average distribution of true positives (TP), false positives (FP), and false negatives (FN) was computed, as shown in Figure 2.1. The median optimal similarity index (SI) was 0.36.

Evidently, even if an optimal threshold could be estimated for these data, a high proportion of FP and FN

¹ The `fminsearch` function in Matlab maximized the similarity index with respect to the manual segmentation.

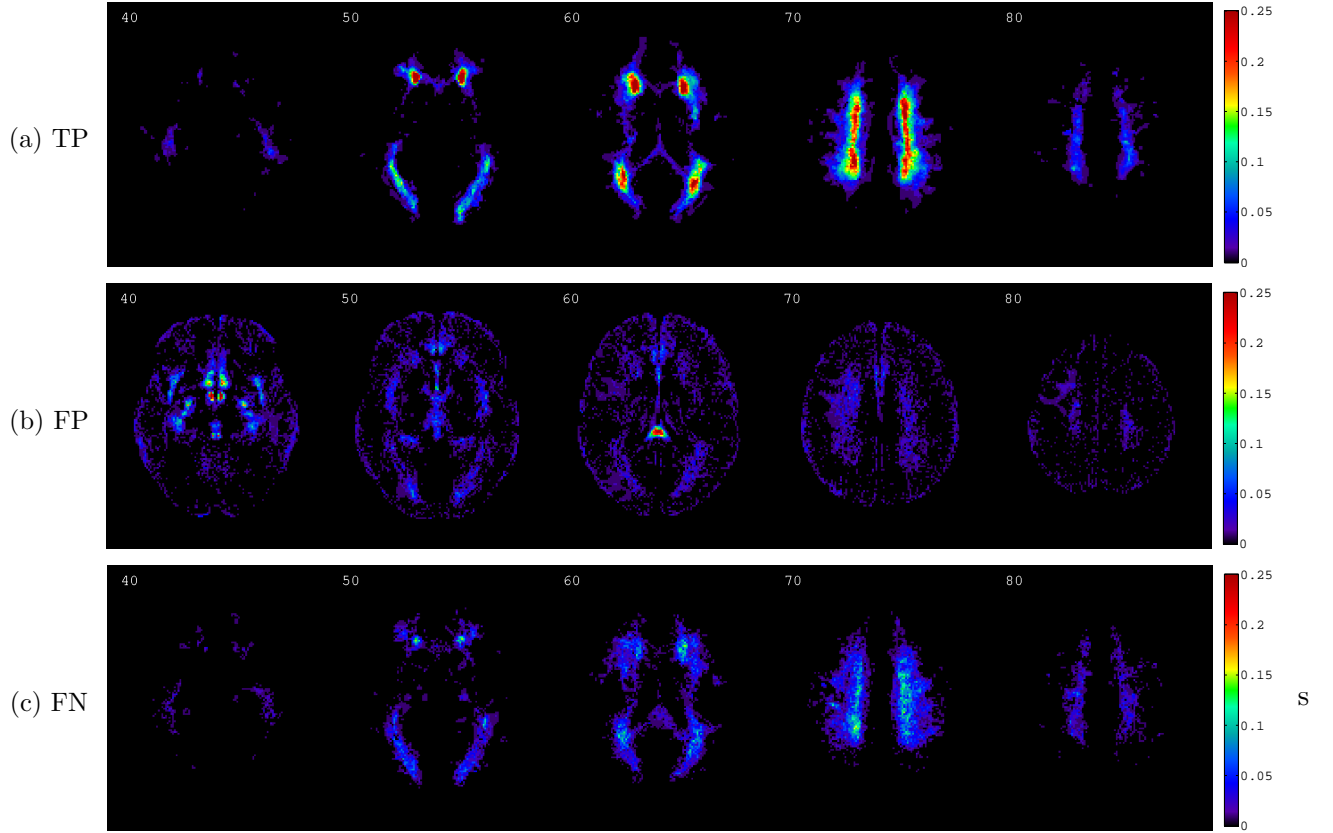


Figure 2.1: Distributions of TP, FP, and FN in 96 FLAIR MRI, following supervised optimal thresholding in MNI space.

would be incurred. Yet in Figure 2.1 there are regular and distinct spatial distributions of these errors. Moreover, it has been suggested that there is regional heterogeneity in relaxation rates of brain tissues [146], and that WML intensity depends in part on location [7, 115]. This implies that spatial coordinates would be useful features for WMH segmentation, especially when incorporated into the main classification model (i.e. not as post-processing). Indeed such features have previously been used [85, 86, 95, 128, 130].

Spatial features could be used with FLAIR intensities in supervised classification models like K-NN, SVM, RF, etc. However, such models treat all features equally, which can lead to artifacts in the decision boundary that contradict prior knowledge. For example, in spatial locations which do not observe any lesions during training, the optimized model may learn to never predict the lesion class, regardless of graylevel features. Similarly, there is no guarantee that FLAIR graylevels will map monotonically to the lesion class, as would be expected. Therefore, spatial features should be treated differently.

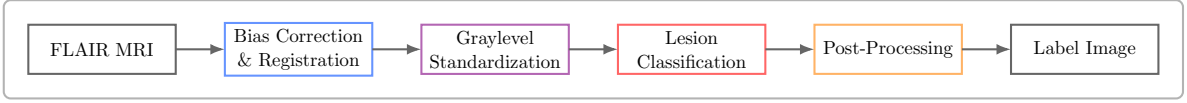


Figure 2.2: Overview of the necessary processing steps.

2.1.2 Overview

The proposed pipeline can be summarized as follows. Pre-processing steps will aim to correct any bias field effect and standardize spatial coordinates. The classification model will employ FLAIR intensities and spatial features to give the initial segmentation. Since most classification models assume features are drawn from a consistent distribution, a graylevel standardization step is appended to the pre-processing. Post-processing steps will generally aim to improve segmentation performance. This pipeline is illustrated in Figure 2.2. Next, the proposed classification model is developed.

2.2 Voxel-Wise Logistic Regression

This section presents the proposed classification model: Voxel-wise Logistic Regression (VLR). The VLR model solves the problem of differential treatment for spatial and graylevel features, and produces uniquely interpretable parameters following training. The model is inspired by the LPA algorithm in the LST toolbox for SPM by Schmidt [144, 147]², however, novel methods of model parametrization and estimation are presented here, in addition to differences in pre- and post-processing from the LPA algorithm.

Logistic regression models have recently gained popularity for WMH segmentation [148, 110, 147, 131], and have several advantages. First, model parameters are generally more interpretable than those in other models, permitting better design of regularizations. Second, the simplicity of the model reduces its capacity for over-fitting. Third, model foundations in statistical theory allow probabilistic interpretations of the outputs, which may be helpful for quantifying marginally pathological tissues like DAWM. The major drawback of logistic models for WMH segmentation is that standardization of graylevels is required.

In the two class formulation, the lesion class is denoted $c = 1$, and the healthy class $c = 0$. The probability of the lesion class, given a set of features $\mathbf{y} = [1, y^1, \dots, y^K]^T$ is modelled by a logistic function,

² This algorithm was originally understood through the Matlab code, after downloading the toolbox. Section 6.1 of [147], published in 2017, provides more details.

parameterized by a vector of feature weights $\boldsymbol{\beta} = [\beta^0, \beta^1, \dots, \beta^K]^T$,

$$P(c = 1 \mid \mathbf{y}, \boldsymbol{\beta}) = \frac{1}{1 + e^{-\eta}}, \quad \eta = \boldsymbol{\beta}^T \mathbf{y}. \quad (2.1)$$

This probability – the estimated lesion class label – is denoted $\hat{c} = P(c = 1 \mid \mathbf{y}, \boldsymbol{\beta}) \in [0, 1]$. Considering the spatial location $x = [x_1, x_2, x_3]$, the estimated label image is defined as $\hat{C}(x) = P(C(x) = 1 \mid \mathbf{Y}(x), \boldsymbol{\beta})$. Figure 2.3 shows an example class prediction for an arbitrary input graylevel.

Previous works have not included spatial dimensions in the logistic feature set \mathbf{y} , since \hat{c} is not expected to be monotonically related to any spatial dimension. The circumvention proposed by Schmidt is to parameterize the feature weights themselves spatially – that is $\boldsymbol{\beta} \rightarrow \boldsymbol{\beta}(x)$,

$$\hat{C}(x) = \frac{1}{1 + e^{-\eta(x)}}, \quad \eta = \boldsymbol{\beta}(x)^T \mathbf{Y}(x). \quad (2.2)$$

In this way, the characteristics of the logistic regression model are maintained with respect to graylevel features, but spatial features can have more complex relationships with the output. A second pre-processing step, however, is now required: standardization of spatial dimensions – i.e. registration of the input image to a consistent target – so that the same voxel in every input image corresponds to the same brain region (cf. § ?? for more details). Training the VLR model then yields a unique vector $\boldsymbol{\beta}$ for each voxel, or equivalently, one complete image for each parameter. Such “parameter images”, as they will be called, are then readily interpretable and can be regularized intuitively.

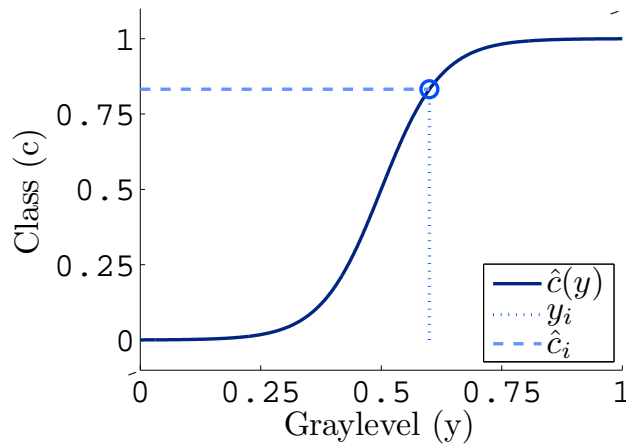


Figure 2.3: The logistic regression model.

2.2.1 Model Fitting

Fitting the VLR model involves estimating β for each voxel x . This requires some training data: feature vectors from a population of N observations $\mathcal{Y}(x) = \{\mathcal{Y}_1(x), \dots, \mathcal{Y}_N(x)\}$, and the corresponding labels $\mathcal{C}(x) = \{C_1(x), \dots, C_N(x)\}$. Typically, parameter estimation involves maximizing the likelihood of the model, given this data – i.e. maximum likelihood estimation (MLE).

Three major challenges emerge during model fitting. These challenges involve contradictions between prior knowledge and the fitted model using the available training data. That is, these challenges could all be overcome by a more complete training set, but this is rarely available. The three challenges are:

1. **Separable classes:** When data from two classes are perfectly separable, the MLE-fitted logistic model can approach a step-function – i.e. $\beta^k \rightarrow +\infty$. This implies that on either side of a specific graylevel threshold, the model is either 100% confident in predicting the healthy class, or 100% confident in predicting the lesion class. In fact, no threshold is ever so perfect, and instead a level of uncertainty should be maintained around the decision boundary. These two cases are illustrated in Figure 2.4a.
2. **Sparsely observed lesion class:** Since WMH are often distributed in consistent locations, many brain regions contain no lesions across the entire training dataset. These voxels will be termed “healthy training” voxels, and denoted \mathcal{X}_h . In some locations, this is expected (e.g. the GM, since by definition WMH manifest in the WM), while in others, prior knowledge predicts lesions will eventually be observed (e.g. the rest of the WM). As illustrated in Figure 2.4b, the MLE-fitted model may not maintain the ability to predict $\hat{c} = 1$ in such locations, regardless of the features. However, the ability to predict lesion should be maintained in many of these locations.
3. **Smooth parameter images:** It is assumed that similar locations will contain similar training data, yielding smooth parameter images. If this assumption is sometimes invalid, parameter images could contain noise or discontinuities, creating artifacts in estimated lesion class images.

Joint Estimation

In the LPA algorithm, the logistic feature weights for each voxel are not considered independent. The FLAIR graylevel effect is tied across all voxels – i.e. $\beta^1(x) \rightarrow \beta^1$, so $\eta(x) = \beta^0(x) + \beta^1 Y(x)$ – and a Gaussian MRF model is used to estimate the spatially varying intercept $\beta^0(x)$. This design addresses the aforementioned challenges by greatly increasing the number of observations used to estimate each

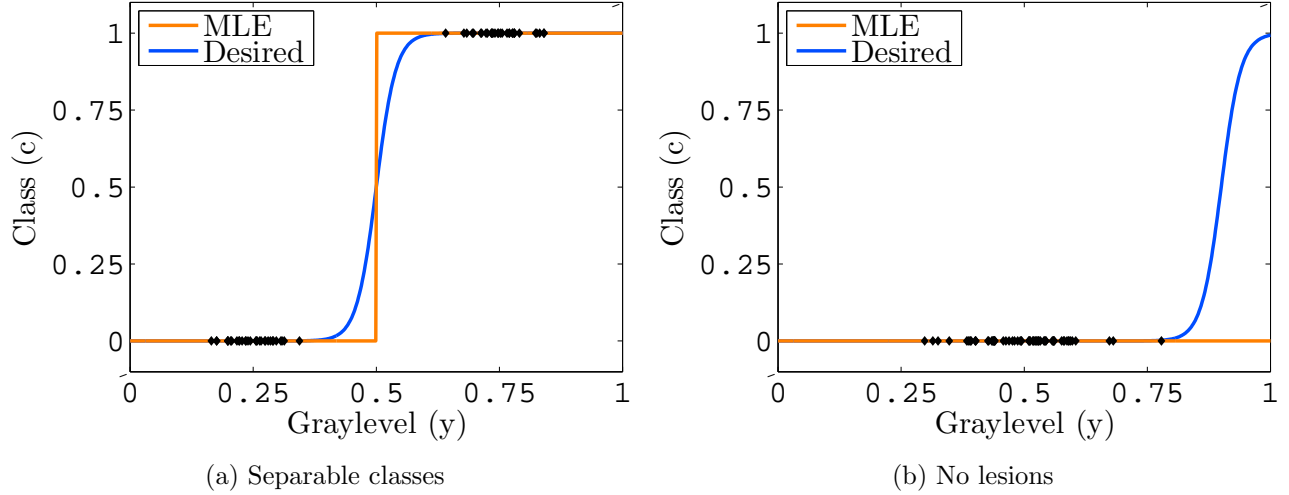


Figure 2.4: Challenges encountered during estimation of a logistic model.

parameter. However, it requires joint estimation of all parameters simultaneously, and dramatically increases the complexity of estimation [147].

Efficient fitting of such models was the subject of major works by Schmidt et al. [149, 147], but several drawbacks remain. First, Markov Chain Monte Carlo estimation of the model appears to create discontinuity artifacts in the spatial effect image $\beta^0(x)$, as shown in Figure 3.2. Second, MRF modelling of the parameter images assumes that the missing data (i.e. WMH training examples in the more superficial brain regions) can be interpolated spatially, but this may not be justified. Third, the proposed joint estimation procedures are computationally expensive (versus the methods proposed here), requiring approximately two hours to estimate $\beta(x)$ [147]. Finally, it is not clear whether any tied β are necessary or advantageous in this context. These deficiencies then motivate investigations into alternative solutions to the above challenges.

In addition to these potential modelling weaknesses, there were also several limitations to the validation methodology for the LPA algorithm, worth noting here. First, the “ground truth” segmentations were generated using an automated algorithm – the Lesion Growth Algorithm (LGA) [108] of the same toolbox – rather than a human expert. Second, the graylevel standardization procedure employed does not consider the variance of image graylevels (only the mean is subtracted); this strongly assumes that user images will have graylevels spanning a similar range. Third, all 53 training cases were obtained on the same MRI scanner, which may limit generalization performance. Finally, no segmentation performance results are given in either of the associated publications [149, 147]. Therefore, while the open-source release of the LPA algorithm is greatly appreciated, significant improvements can be made to this algorithm.

Independent Estimation

An alternative approach is to consider each parameter vector $\beta(x)$ independent. Doing so greatly simplifies model fitting, but does not address the challenges described above. These must instead be addressed using regularization strategies, as discussed below in § ???. In this section, ML estimation of independent parameter vectors β is developed. For clarity, only a single voxel is considered – i.e. y from $Y(x)$, etc.

As note above, the optimal β for each independent voxel can be resolved using MLE. If the training data are also assumed to be independently observed, then the likelihood (conditioned on the data) is defined from binomial theory as

$$\begin{aligned} L(\beta \mid \mathcal{C}, \mathcal{Y}) &= \prod_{n=1}^N P(c = 1 \mid \mathbf{y}_n, \beta)^{c_n} (1 - P(c = 1 \mid \mathbf{y}_n, \beta))^{1-c_n} \\ &= \prod_{n=1}^N \left[\hat{c}_n^{c_n} (1 - \hat{c}_n)^{1-c_n} \right]. \end{aligned} \quad (2.3)$$

For computational reasons, it is simpler and asymptotically equivalent to maximize the log-likelihood,

$$\begin{aligned} \mathcal{L}(\beta) &= \log \prod_{n=1}^N \left[\hat{c}_n^{c_n} (1 - \hat{c}_n)^{1-c_n} \right] \\ &= \sum_{n=1}^N \left[c_n \log \hat{c}_n + (1 - c_n) \log(1 - \hat{c}_n) \right] \\ &= \sum_{n=1}^N \left[c_n \beta^T \mathbf{y}_n - \log(1 + e^{\beta^T \mathbf{y}_n}) \right]. \end{aligned} \quad (2.4)$$

The optimal β is therefore resolved by maximizing the log-likelihood,

$$\begin{aligned} \beta^* &= \arg \max_{\beta} \mathcal{L}(\beta) \\ &= \arg \max_{\beta} \sum_{n=1}^N \left[c_n \beta^T \mathbf{y}_n - \log(1 + e^{\beta^T \mathbf{y}_n}) \right] \end{aligned} \quad (2.5)$$

Iterative Updates

Estimation of β^* can be performed using iterative optimization, using an initial estimate $\beta^{(0)}$ and an update term $\Delta\beta^{(t)}$,

$$\beta^{(t+1)} \leftarrow \beta^{(t)} + \alpha \Delta\beta^{(t)}, \quad (2.6)$$

where α is a small valued learning rate parameter. There are many possible definitions of $\Delta\beta$, including simply the gradient of $\mathcal{L}(\beta)$, denoted $\nabla_{\beta}\mathcal{L}$. However, it can be shown that $\mathcal{L}(\beta)$ is convex, so higher order update equations can be used. The work by Minka [150] compares several options, including Newton’s method (and variants), conjugate gradient, iterative scaling (and variants), and dual optimization³. For small feature dimensionality (K), performance differences among the options were small. Classic Newton updates gave a good balance between memory requirements and computational order, so they are used.

If the gradient $\nabla_{\beta}\mathcal{L}$ and Hessian matrix $\nabla_{\beta}^2\mathcal{L}$ are defined as

$$\nabla_{\beta}\mathcal{L} = \begin{bmatrix} \frac{\partial L}{\partial \beta^1} \\ \vdots \\ \frac{\partial L}{\partial \beta^K} \end{bmatrix}, \quad (2.7)$$

$$\nabla_{\beta}^2\mathcal{L} = \begin{bmatrix} \frac{\partial^2 L}{\partial \beta^1 \partial \beta^1} & \cdots & \frac{\partial^2 L}{\partial \beta^1 \partial \beta^K} \\ \vdots & \ddots & \vdots \\ \frac{\partial^2 L}{\partial \beta^K \partial \beta^1} & \cdots & \frac{\partial^2 L}{\partial \beta^K \partial \beta^K} \end{bmatrix}, \quad (2.8)$$

then the Newton update is given by

$$\Delta\beta = -\nabla_{\beta}^2\mathcal{L}^{-1}\nabla_{\beta}\mathcal{L}. \quad (2.9)$$

In the current model, the gradient is given by

$$\nabla_{\beta}\mathcal{L} = \sum_{n=1}^N \mathbf{y}_n (c_n - \hat{c}_n), \quad (2.10)$$

and the Hessian by

$$\nabla_{\beta}^2\mathcal{L} = \sum_{n=1}^N \mathbf{y}_n \mathbf{y}_n^T (c_n - \hat{c}_n). \quad (2.11)$$

Substituting (2.10) and (2.11) into (2.9), the explicit update $\Delta\beta$ for (2.6) is obtained. At each iteration, $\Delta\beta^{(t)}$ is re-computed, and the process continues until some convergence criterion is satisfied.

Simplification

It is not necessary to complete the above procedure for all voxels in the standardized space. Instead, only voxels in the expected location of the brain need to be computed; such voxels can be selected using a binary “brain mask”, denoted $M(x)$. More details about the brain mask used in this work can be found

³ Matlab code available at <https://github.com/tminka/logreg/>

in § B.3.2. Moreover, since the parameters of each voxel are estimated independently, this can also be computed in parallel. The details of this implementation are presented in § B.2.1, after incorporation of the regularizations described in the next section.

Finally, the model has so far been derived in general terms, so that any choice of feature set \mathbf{y} can be used. However, with only one feature – the FLAIR graylevel – it is possible to reparameterize the sigmoid argument as

$$\begin{aligned}\beta^T \mathbf{y} &= \beta^0 + \beta^1 y^1 \\ &= s(y - \tau) \quad \begin{cases} s = \beta^1 \\ \tau = -\frac{\beta^0}{\beta^1} \end{cases} .\end{aligned}\tag{2.12}$$

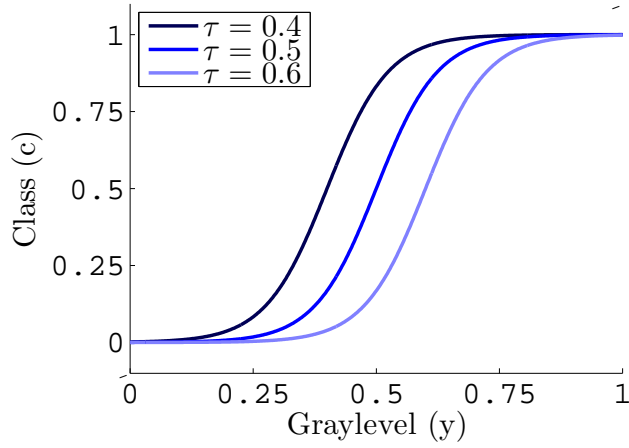
In this form, the parameters τ and s emerge as a graylevel threshold and a slope parameter, respectively. Specifically, when $y = \tau$, the predicted probability of lesion is $\hat{c} = \frac{1}{1+e^0} = 0.5$, so τ controls the location of the class discrimination, as shown in Figure 2.5a. Similarly, the s parameter defines the sensitivity of the logistic function to y , as shown in Figure 2.5b. By contrast, varying the original parameter β^1 with β^0 constant (Figure 2.5d) results in correlation of these characteristics. These new parameters, and the corresponding images $\mathcal{T}(x)$ and $\mathcal{S}(x)$, are therefore salient descriptors of the predictive model.

2.3 Regularization

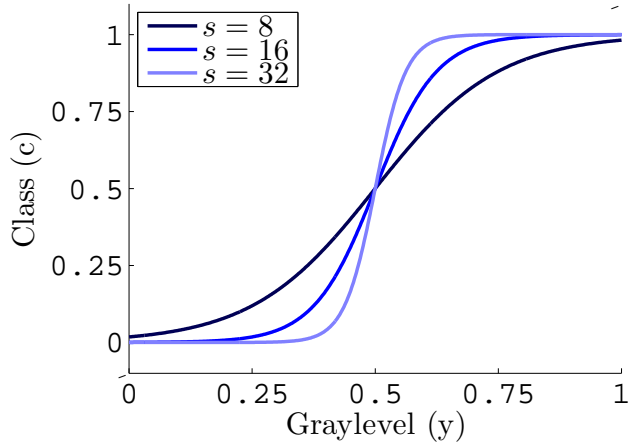
Regularizations are methods of injecting prior knowledge about the expected model into the optimization. Assuming voxel-wise independence of model parameters requires the use of regularization strategies to solve the challenges outlined in § 2.2.1. Several regularization methods are explored below.

2.3.1 Data Augmentation

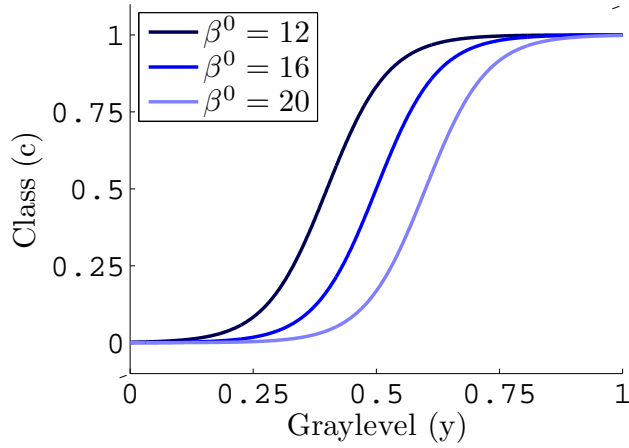
Noting the central role of training data in each of the challenges, methods of artificially increasing the training dataset size may be particularly useful in solving them. Data augmentation has long been used in machine learning tasks with limited training data, and there are several methods of generating synthetic data. In low dimensional input/output spaces, random sampling of fitted class-conditional posterior distributions can produce reasonable samples with known labels [151]. In higher dimensional problem spaces, however, imputation is more difficult [152]. For example, the space of potential $100 \times 100 \times 100$ -sized images has 100^3 dimensions (one per voxel), yet only a small subspace represents plausible images.



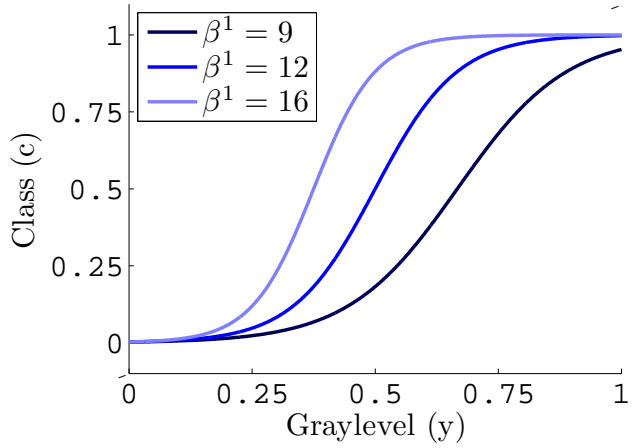
(a) Vary τ with $s = 16$ constant.



(b) Vary s with $\tau = 0.5$ constant.



(c) Vary β^0 with $\beta^1 = 16$ constant.



(d) Vary β^1 with $\beta^0 = -6$ constant.

Figure 2.5: Effect of varying the logistic model parameters.

Generating synthetic examples in this space is therefore challenging, especially for segmentation tasks, where the outputs have dimensionality roughly equal to the input.

Alternatively, simple image manipulations can still afford model improvements [153]. In segmentation tasks, both the input image(s) and the corresponding label images can be translated, reflected, rotated, and perhaps resized, thereby avoiding the generation of genuinely synthetic examples. In the current work, reflections and small (one-voxel) translations can be applied to the label and FLAIR images following registration to the MNI brainspace. The potential benefits of this augmentation are explored in § ??.

2.3.2 Classic Regularization

The separable classes challenge is well-known in regression problems, and a good solution is to penalize the magnitude of model parameters using the L_p -norm: $\lambda ||\boldsymbol{\beta}||_p$ [154]. It can be shown that L_1 regularization corresponds to a Laplacian prior on elements of $\boldsymbol{\beta}$, with scale parameter inversely proportional to λ (equivalently, this assumes that the model error follows this distribution). Similarly, L_2 regularization implies a Gaussian prior, with standard deviation inversely proportional to λ [154]. Model fitting which includes this prior-derived term is called maximum a posteriori (MAP) estimation, and the penalty can be appended to the objective function (2.5), as in

$$\begin{aligned}
\boldsymbol{\beta}^* &= \arg \max_{\boldsymbol{\beta}} \mathcal{J}(\boldsymbol{\beta}) \\
&= \arg \max_{\boldsymbol{\beta}} \mathcal{L}(\boldsymbol{\beta}) - \lambda ||\boldsymbol{\beta}||_p \\
&= \arg \max_{\boldsymbol{\beta}} \sum_{n=1}^N \left[c_n \boldsymbol{\beta}^T \mathbf{y}_n - \log(1 + e^{\boldsymbol{\beta}^T \mathbf{y}_n}) \right] - \lambda ||\boldsymbol{\beta}||_p
\end{aligned} \tag{2.13}$$

Due to its relatively large gradient near zero, L_1 regularization is typically used to encourage sparsity in the feature weights (i.e. $\beta^k \rightarrow 0$) [155]. This is not desirable in the current model, since the feature (FLAIR graylevel) is known to be discriminative. Moreover, the expansion of the $||\boldsymbol{\beta}||_1$ term in the gradient of the objective function is not straightforward, since it is non-differentiable at zero [155, 156]. Conversely, L_2 regularization is more effective at limiting parameter magnitude – which is the current aim – and the first and second order gradients of (2.13) derive easily [150]. For these reasons, only L_2 regularization is considered, yielding the following change to the Newton update expression (2.9),

$$\begin{aligned}
\Delta \boldsymbol{\beta} &= -\nabla_{\boldsymbol{\beta}}^2 \mathcal{J}^{-1} \nabla_{\boldsymbol{\beta}} \mathcal{J} \\
&= -(\nabla_{\boldsymbol{\beta}}^2 \mathcal{L} - \lambda I)^{-1} (\nabla_{\boldsymbol{\beta}} \mathcal{L} - \lambda \boldsymbol{\beta})
\end{aligned} \tag{2.14}$$

What remains is to select an appropriate value of λ . This is explored experimentally in § 3.3.1 using a toy model.

2.3.3 Pseudo-Lesions

The sparsely observed lesion class challenge is less common, since discriminative models are rarely fit in the absence of one class altogether. This occurs here because all voxels are modelled independently. It is therefore tempting to simply sample features from the lesion class at other spatial locations in order to fit the logistic model in the healthy training voxels, similar to the approach by Schmidt. However, as noted in § 1.2.2 and § 2.1, WML are thought to have different intensities in different brain regions [146, 7], and some locations will likely never contain any WMH. Considering these facts, the use of deterministic synthetic lesion-class samples, or “pseudo-lesions”, could instead permit better use of prior knowledge about WMH. These synthetic observations could be appended to the training data for each voxel so as to minimally balance the training classes, and act as a prior on the distribution of lesion-class features.

If the same number of synthetic observations are appended to the training data for each voxel, this is equivalent to appending a number of synthetic images to the training set. The synthetic feature data are denoted $\mathbf{V}(x) = \{\gamma_1(x), \dots, \gamma_v(x)\}$. It is assumed that the labels of all synthetic data are “lesion”, so the set of synthetic label images is simply denoted $\mathbf{1}(x)$. The updated training set is therefore $\mathcal{Y}_\gamma(x) = \{\mathcal{Y}(x), \mathbf{V}(x)\}$, and $\mathcal{C}_\gamma(x) = \{\mathcal{C}(x), \mathbf{1}(x)\}$.

Design of the synthetic image set $\mathbf{V}(x)$ should be guided by prior knowledge. For the same reasons as described above, it is not possible to derive this knowledge from the training set. Unfortunately, few other sources of structured information are available. One reasonable approach could make use of healthy tissue prior probability images (Figure 2.6), denoted $\rho(x)$. Specifically, the expected intensity for the lesion class in each tissue can be multiplied by the tissue probability image, and the results summed to give the overall synthetic image,

$$\gamma(x) = \gamma_{\text{GM}} \cdot \rho_{\text{GM}}(x) + \gamma_{\text{WM}} \cdot \rho_{\text{WM}}(x) + \gamma_{\text{CSF}} \cdot \rho_{\text{CSF}}(x) \quad (2.15)$$

While WMH are not possible in either the GM or the CSF, it is necessary to select a FLAIR graylevel – perhaps the maximum possible intensity – to complete this model. Additionally, such parameters will inevitably play a role for subjects with imperfect registration or outlier anatomy. The contributions of pseudo-lesions to model fitting are explored experimentally in § 3.3.1.

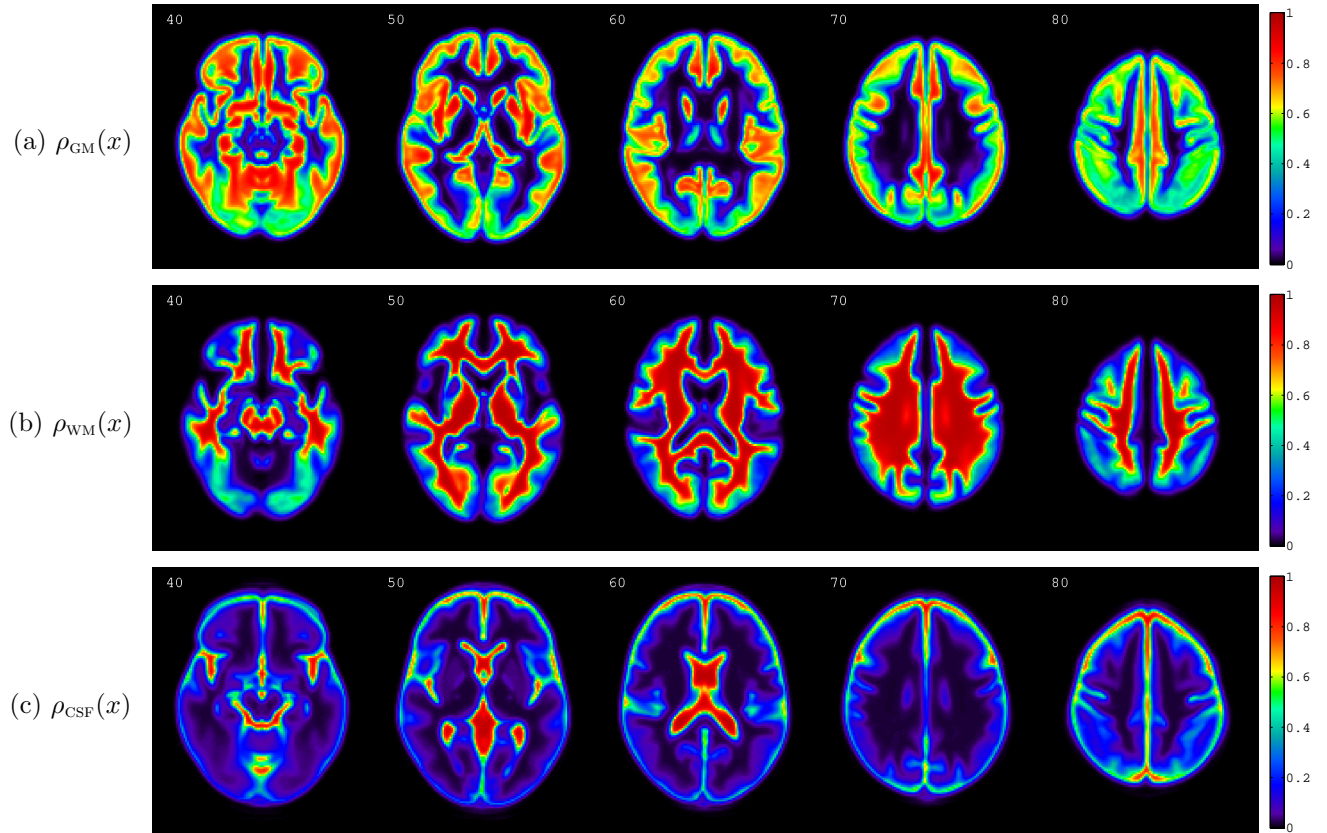


Figure 2.6: Tissue prior probability images in MNI space. Derived from [132].

Table 2.1: Image filters considered for smoothing the estimated parameter images.

Name	Parameters	Advantages	Disadvantages	Ref.
Gaussian	width σ	no artifacts	blurs edges	
Median	width w	preserves edges	square artifacts	
Anisotropic Diffusion				
Bilateral				

2.3.4 Parameter Image Smoothing

Finally, independent model fitting in every voxel risks yielding noisy parameter images. The simplest solution to this problem involves filtering the reconstructed parameter images after estimation. A wide range of possible filters for this task exist, and a selection of these are summarized in Table 2.1, including the mutable parameters for each.

An alternative solution might involve modelling the parameter images as a spatial function (e.g. band-limited discrete cosine / Fourier transform). However, there are two challenges with this approach. First, deriving the update gradients for such a model would be challenging, and their computation could significantly increase training time. Second, such an encoding may introduce artifacts in the resulting parameter images. Moreover, it is well known that frequency domain band-limiting can be equivalently

achieved by convolution (i.e. filtering) in the spatial domain [157]. Therefore, only conventional filtering is explored in this work (cf. § ??)

2.4 Pre-Processing

The proposed VLR classification model addresses several of the challenges outlined in § 1.2.2. The problem of overlapping tissue graylevel distributions is mostly solved through expansion of the feature space to include spatial features. CSF flow-through artifacts, which appear in roughly consistent locations, are similarly managed. Heterogeneity in the appearance of lesions is also considered by the spatial parametrization of logistic parameters. Finally, ambiguity regarding moderately hyperintense DAWM, and voxels affected by partial volume effect, is captured in the probabilistic output.

Several challenges, however, still remain. In particular, a number of assumptions were made about the input data for the VLR model which are likely invalid for raw images. These assumptions are that: 1) input MRI images are free of bias field artifact; 2) feature intensities are consistent across different subjects; 3) images are consistently sized and voxels represent the same anatomical regions across different subjects. Solving these challenges must therefore be accomplished by one or more pre-processing steps.

2.4.1 Registration

Image registration is the process of geometrically transforming a source image so that the image content is aligned per-voxel with a target image of the same subject. This process facilitates voxel-wise analysis of MRI from different subjects, such as “voxel-based morphometry” [158] and analysis of functional MRI data [159]⁴. Source images from multiple subjects are usually registered to the same target image; in this context, it is useful to define a “native space” and a “standardized space”, denoting the original, subject-specific geometry, and the standardized target geometry. By convention, the target brain space is usually either the Talairach space [160] or the Montreal Neurological Institute (MNI) space [161], though any reasonable target image could be used.

Parameters defining registration transforms are fitted by maximizing some measure of overlap between the images [162]. Simple registration methods employ only affine transformations, comprising a combination of translation, scaling, and shear transformations (e.g. the FLIRT tool [163] in FSL). The utility of these

⁴ Incidentally, investigation of these topics were the motivations for developing of the SPM and FSL software packages, respectively.

methods for neuroimage analysis is limited, since there are often significant differences in brain anatomy between subjects. Rather, most tools parametrize a spatial warping model, permitting local nonlinear deformations to better match brain structures (e.g. cubic B-splines in FSL FNIRT [142], discrete cosine transform in SPM Normalize [135, 136], a general diffeomorphism in the SyN algorithm [164]). While these models are more difficult to fit, several robust algorithms have been released for general use, with widespread acceptance (e.g. the toolboxes mentioned above). A 2009 comparison of 14 different methods is a good resource on the subject [165], while a more recent (2014) study ranks the popular toolboxes SPM, FSL, and Brainsuite [166].

In the current work, registration is required during both training and testing. During training, the model parameters $\beta(x)$ are estimated using mutually aligned segmentation examples $\{\mathcal{Y}, \mathcal{C}\}$ in a standard brain space. At test time (or for actual use), these parameter images are transformed in the opposite direction from the standard space to the native space of the current subject. Since registration transforms are typically bijections – i.e. invertible – the second registration case can be estimated using the same method as the first, minimizing bias.

Unlike other applications, it is not essential that perfect image registration is achieved here. As noted by Harmouche et al. [115], in smoothly varying models, small registration errors can be expected to have a negligible impact. Therefore, registration was not a primary focus of optimization in this work.

While the registration component of the SPM8 “Segment” feature [136] was not among the top performing in the 2009 study [165], subsequent implementation revisions in SPM12⁵ called “New Segment” have apparently improved results. In the 2014 study [166], the SPM12 New Segment method achieved the highest Similarity Index of all methods on real (IBSR [167]) data.

Furthermore, the SPM module has several other features amenable to this work. First, unlike many other registration algorithms, the objective function does not require source and target images to have the same contrast. This is helpful, since no suitable FLAIR template image is available for use as a target.⁶ Second, it is simple to invoke the SPM modules via the command line or Matlab scripts; this facilitates smooth integration of this tool in the pipeline. Third, it is possible to save previously estimated registration transformations, and apply them in the forward or reverse directions efficiently. This can save significant time during cross validation, since the estimated parameter images $\beta(x)$ must eventually be transformed to the native space of every subject. Finally, the SPM “New Segment” model additionally estimates the

⁵ http://www.fil.ion.ucl.ac.uk/spm/software/spm12/SPM12_Release_Notes.pdf

⁶ One FLAIR template is available in [168]; however, this was generated using SPM5, so using it would compound any registration biases associated with the older method.

bias field during execution with high accuracy, saving an additional step (cf. § 2.4.2, below). For all these reasons, the registration performed by SPM New Segment was used throughout this work. After satisfactory visual inspection of all training set images, no other registration tools were investigated. The default brain space of this module is MNI.

2.4.2 Bias Correction

As noted in § 1.2.2, bias field (AKA intensity inhomogeneity), is a smoothly varying intensity variation artifact common in MRI. The sources of bias field artifact include inhomogeneities in the magnetic field and RF coils used for pulse transmission and signal sensing, as well as non-ideal magnetic properties of the imaged object [71]. The field and coil related sources are more significant at clinical field strengths. These can be corrected prospectively, though techniques for doing so are limited, and often a small bias field continues to corrupt acquired images [71]. Therefore, retrospective correction has been the subject of much research.

Similar to image registration, several widely accepted algorithms for estimating and correcting bias field have emerged. The N3 algorithm [169], subsequently updated to N4(ITK) [139] and integrated in the FreeSurfer toolbox⁷ is perhaps the most popular, it makes minimal assumptions about the image. This method aims to sharpen the image histogram by dividing the image by an estimated bias field, parameterized by B-splines for smoothness. As noted above, the SPM Segment model [136] includes integrated bias field estimation and correction. The bias field model in this work is parameterized by the discrete cosine transform, as described in in a previous work by Ashburner and Friston [170]. The FSL Segment feature also estimates bias field in a similar overall model to SPM Segment, except the tissue prior probability maps in SPM are replaced with a Hidden Markov Random Field Model, as described in [141]. Again, two reviews with quantitative performance comparisons provide a good reference of other proposed algorithms, including a comprehensive review in 2006 [171] and a comparison of mainly popular methods in 2016 [172].

In the 2016 comparison [172], the authors note that the SPM and FSL models – which include segmentation – outperformed the N3 algorithm [169] and another non-segmenting method [173]. These results are consistent with the advantages of unified generative models described in § 2.1 and discussed in “Unified segmentation.” [136]. Specifically, the estimation of both bias field and tissue segmentation could each be improved if the other was already known. Rolling these tasks into a single EM-fitted model allows

⁷ <https://surfer.nmr.mgh.harvard.edu/>

alternating conditional estimates to converge on better results overall.

Bias field estimation was not a primary focus of this work. Therefore, due to the better performance over N3/4, and the advantages already afforded by SPM Segment for registration noted above, this model was employed for bias correction throughout this work. No other bias field correction tools were investigated.

2.4.3 Graylevel Standardization

The flexibility of image contrast in MRI is a double-edged sword. This feature, in addition to properties of spatial encoding during acquisition, preclude direct interpretation of image graylevels as a tissue property. As a result, the same brain region in the same subject may be assigned a different graylevel depending on the MRI sequence time constants, scanner, and spatial acquisition protocol. For automated analysis of MRI images, therefore, standardization of image graylevels is required.

Graylevel standardization can be achieved using a univariate transformation $\tau : y \mapsto \tilde{y}$, defined as

$$\tilde{y} = \tau(y), \quad (2.16)$$

where τ is monotonic, and considers characteristics of the input image, such as basic graylevel statistics or the histogram. The histogram of an image represents the number of occurrences of each graylevel in the image. Normalizing the histogram by the total number of voxels X yields the probability mass function (PMF) $f_Y(y)$,

$$\begin{aligned} f_Y(y) &= \frac{1}{X} h_Y(y) \\ &= \frac{1}{X} \sum_{i=1}^X \begin{cases} 1 & Y(x_i) = y \\ 0 & Y(x_i) \neq y \end{cases} \end{aligned} \quad (2.17)$$

The cumulative density function (CDF) $F_Y(y)$ is the cumulative sum of $f_Y(y)$,

$$F_Y(y) = \sum_{\gamma=y_{\min}}^y f_Y(\gamma) \quad (2.18)$$

The PMF of an MRI can be decomposed into the contributions of each constituent tissue, as shown in Figure A.2. This is the fundamental principle underlying mixture models (cf. § 1.3.2). The goal of standardization is therefore to align these sub-distributions as closely as possible.

Previously proposed methods of standardizing MRI intensities include the following:⁸

- **Range Matching:** The simplest approach to standardization involves rescaling the data using the minimum and maximum intensities,

$$\tau(y) = \frac{y - y_{\min}}{y_{\max} - y_{\min}}. \quad (2.19)$$

Naively, this method is very susceptible to corruption by outliers. For more robustness, y_{\min} and y_{\max} can be defined using intensity quantiles – e.g. $[\epsilon_1, 1 - \epsilon_2]$. However, selection of an appropriate ϵ_2 for WMH segmentation is difficult, since WMH typically constitute only the top 1% of the total brain volume. Furthermore, differences in image contrasts are not considered by this approach.

- **Statistical standardization:** Another simple but popular approach uses the first and second order moments of the PMF,

$$\tau(y) = \frac{y - \mu_Y}{\sigma_Y}. \quad (2.20)$$

As with range matching, variable image contrasts are not well modelled by this method.

- **Histogram equalization:** Histogram equalization transforms the image PMF to a uniform distribution, thereby distributing image intensities equally across the available range. The desired transform is defined as the CDF of the input image (cf. [157] for derivation),

$$\tau(y) = F_Y(y). \quad (2.21)$$

The chief assumption of histogram equalization for standardization is that input images contain consistent amounts of each tissue class. In MRI with WMH, this assumption may not be valid; however, this technique may still have value.

- **Histogram matching:** Histogram matching is similar to histogram equalization, except that the output PMF is not uniform, but some other specified distribution, $f_{\tilde{Y}}$. This transform is defined as the function composition of the input CDF and the inverse target CDF,

$$\tau(y) = F_{\tilde{Y}}^{-1}(F_Y(y)) \quad (2.22)$$

Histogram matching (cf. A.2.1 for an illustration and experimental evidence.)

⁸ This section considers only univariate standardization methods, since only FLAIR intensities are used in this work.

- **Nyul standardization:** In [143, 174], Nyul and Udupa proposed a method for intensity standardization which has subsequently been used in other works. This method defines τ with piecewise linear segments connecting the Q quantiles of the input PMF q , with quantiles of a target PMF r ,

$$\tau(y) = r_i + (y - q_i) \left(\frac{r_{i+1} - r_i}{q_{i+1} - q_i} \right), \quad y \in [q_i, q_{i+1}] \quad (2.23)$$

However, it can be shown that this transformation is a non-uniform trapezoidal Riemann approximation of true histogram matching, which performs worse in terms of intensity standardization.⁹

- **Regional characteristics:** Decorrelating variation in intensities from variability in anatomical content is a central challenge in intensity standardization. One solution is to define the image-specific transformation using characteristics from a more anatomically consistent brain region. This is the approach employed by Shinohara et al. in the so-called white stripe method [176]. In the current work, this region, denoted \mathcal{X}_τ , can be defined in MNI space using tissue priors (Figure 2.6), anatomical label maps, or any other method of selecting a subset of voxels.

Quantifying Standardization

While the major advantages and challenges to several graylevel standardization methods have been briefly noted, it remains to explore the utility of each experimentally. In the current work, the goal of this step is to maximize the separation of the two classes. This can even be maximized voxel-wise, due to the characteristics of the VLR model. Therefore, an intermediate objective function \mathcal{Z} should be defined which quantifies the degree of separation of the two classes.¹⁰ This objective function can then be used to optimize any tunable parameters in each of the transforms – e.g. ϵ , Q , \mathcal{X}_τ – and also to select the best overall method.

Two such functions are proposed. The first is discrete, and inspired by the Zero-Crossing Rate [177]. It measures the number of class transitions in the sorted feature data $\tilde{\mathcal{Y}}_s$, as shown in Figures 2.7a and 2.7c. With \mathcal{C}_s as the class labels after sorting by the feature $\tilde{\mathcal{Y}} = \tau(\mathcal{Y})$, the objective function \mathcal{Z}_Δ is defined as

$$\mathcal{Z}_\Delta = \sum_{n=1}^{N-1} \begin{cases} 1 & \mathcal{C}_s^n \neq \mathcal{C}_s^{n+1} \\ 0 & \mathcal{C}_s^n = \mathcal{C}_s^{n+1} \end{cases}. \quad (2.24)$$

⁹ This result is presented and supported with experiments in [175].

¹⁰ Alternatively, the entire pipeline can be executed under cross validation and overall performance compared between standardization methods. However this does not consider potential interactions between the standardization method and tunable downstream parameters.

This function is discrete and bounded, as in $\mathcal{Z}_\Delta \in \mathbb{Z} [1, \lfloor \frac{N}{2} \rfloor]$ ¹¹ and the lower bound is optimal – i.e. \mathcal{Z}_Δ should be minimized.

The second function is continuous, and inspired by probability theory. It measures the relative overlap of class distributions, $p(\tilde{y} \mid c = 1)$ and $p(\tilde{y} \mid c = 0)$, estimated using kernel smoothing, as shown in Figures 2.7b and 2.7d. This objective function \mathcal{Z}_\star can be defined as

$$\mathcal{Z}_\star = \int_{\tilde{y}_{\min}}^{\tilde{y}_{\max}} \frac{\min\{p(\zeta \mid c = 1), p(\zeta \mid c = 0)\}}{\max\{p(\zeta \mid c = 1), p(\zeta \mid c = 0)\}} \partial\zeta, \quad p(\zeta \mid c) \approx \sum_{\tilde{y} \in \{\mathcal{Y} \mid c\}} \delta(\zeta - \tilde{y}) \star G_\sigma(\zeta) \quad (2.25)$$

where $G_\sigma(\zeta)$ is a Gaussian convolution kernel with width σ . This function is continuous and bounded, as in $\mathcal{Z}_\Delta \in [0, 1]$, and the lower bound is again optimal.

Supervised Standardization

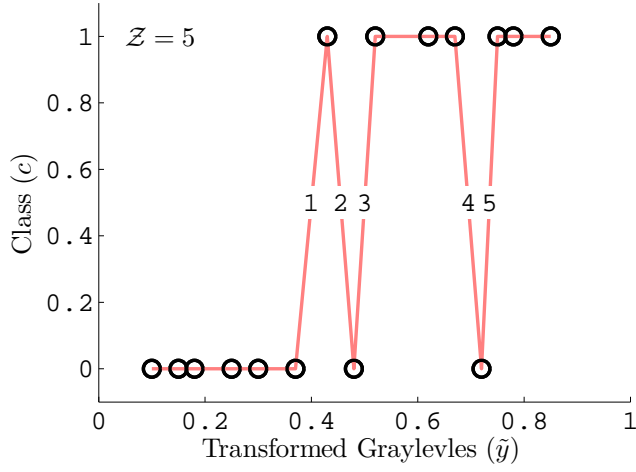
It is not hard to see that it should be possible to estimate an optimal graylevel standardization transform using the training data. That is, a *supervised graylevel standardization*.¹² If \mathcal{Z} is differentiable, then this optimization can be performed using gradient descent, or similar methods. Unfortunately, neither of the above objective functions \mathcal{Z} were reasonably differentiable, but many other optimization paradigms which do not require gradients could be used.

2.5 Post-Processing

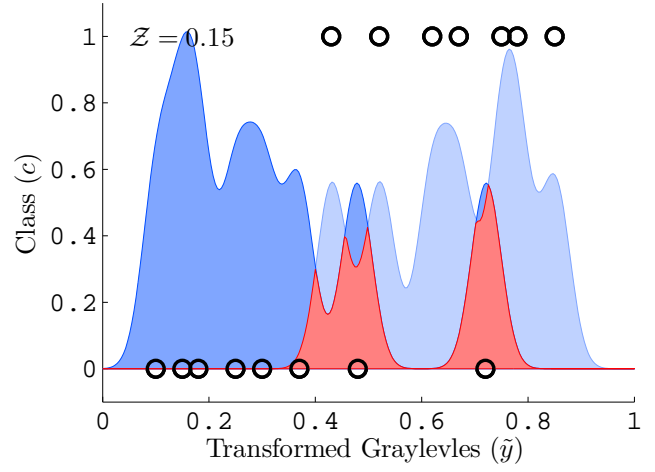
At this point, the motivation and details of the proposed VLR model have been presented, in addition to the preprocessing steps required to satisfy its assumptions. The last component of a segmentation pipeline typically includes post-processing. In principle, this step aims to incorporate any additional knowledge of the problem which has not been considered in upstream elements. Here, these include the connected morphology of WML, and the minimum lesion size. Discussions of these topics, however, would assume that the label image is already binary, whereas the output from the VLR model is probabilistic. Therefore, the first post-processing step thresholds the WMH class probability to give a “hard” classification: $\hat{c} \rightarrow \hat{c}^\circ$. This also facilitates comparison with manual segmentation masks, which are usually binary.

¹¹ The lower bound can be zero if one class is not observed.

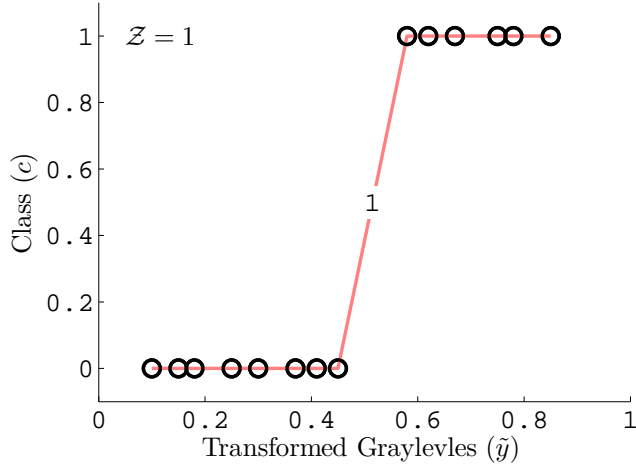
¹² To the best of this author’s knowledge, such a technique has never been proposed.



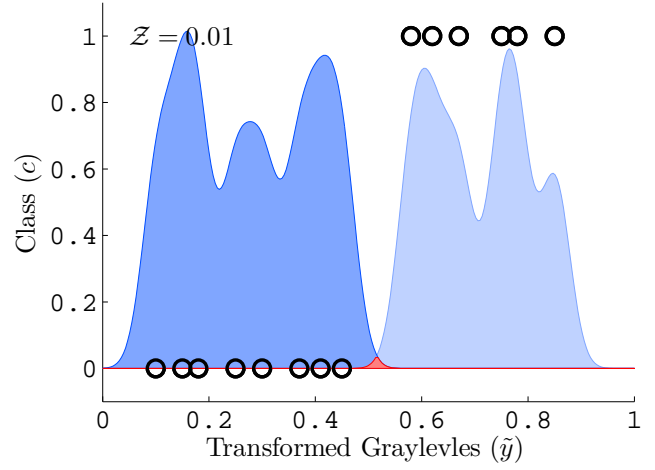
(a) \mathcal{Z}_Δ with overlapping classes



(b) \mathcal{Z}_\star with overlapping classes



(c) \mathcal{Z}_Δ with separable classes



(d) \mathcal{Z}_\star with separable classes

Figure 2.7: Illustration of potential separability objective functions.

2.5.1 Thresholding

If the assumptions of any probabilistic model are valid, then the “hard” classification is straightforward:

$$\hat{c}^\circ = \arg \max_{\omega} p(\omega \mid \mathbf{y}, \boldsymbol{\beta}) \quad \forall \omega. \quad (2.26)$$

In a 2-class logistic regression model, this simplifies to thresholding:

$$\hat{c}^\circ = \begin{cases} 0 & \hat{c} < \pi_c \\ 1 & \hat{c} \geq \pi_c \end{cases}, \quad (2.27)$$

with $\pi_c = 0.5$. However, since these assumptions are often only partially true, most models are able to achieve better agreement with manual segmentations using a different threshold π_c for the WMH class. In fact several of the freely available toolboxes (cf. § ??) permit a user-specified threshold which can be optimized for the user’s data. During model validation, this parameter should be optimized using the training data for each cross validation fold. It is also prudent to illustrate the sensitivity of the model to this parameter, using either a plot of performance versus threshold [64], or a precision-recall (PR) curve [178]. In the current work, both these techniques are employed: π_c is optimized on the fly, and a PR curve is given after.

2.5.2 Minimum Lesion Size

With finite image resolution and appreciable noise in MRI, lesions appearing as only a few connected voxels are indistinguishable from image noise, even by human experts. Such potential lesions are therefore not included in radiologists assessment of WML. Accordingly, most WMH segmentation algorithms employ a minimum-connected-voxels exclusion criterion during post-processing. Connectedness can be defined in 2D or 3D, and consider only direct adjacency or diagonal connections too. Most works employ the most liberal definition: 26-connectedness, which considers all $3 \times 3 \times 3 - 1 = 26$ candidates surrounding a given voxel in 3D. Ideally, the number of required connected voxels will adapt to the image resolution, and correspond to a minimum lesion volume. Typical volumes range from about $x_{\min}^c = 3.5 \text{ mm}^3$ in [64, 122], to 9.0 mm^3 in [114, 179].

In the current work, the inclusion of a minimum lesion size rule is explored. The optimal value for x_{\min}^c is resolved experimentally during each cross validation fold, and the resulting values compared with the above conventions. Additionally, the gains in performance afforded by this step are quantified.

2.6 Segmentation Performance Metrics

Quantifying the performance of a model is essential to optimizing its design. Typically, WMH segmentation performance is characterized in two respects: voxel-wise agreement and total lesion load (LL) volume agreement. When comparing the estimated class \hat{c} to the ground truth class c , each individual voxel can occupy one of four states (colours shown for future reference):

- True Positive (TP): $c = 1$ and $\hat{c} = 1$, correctly predicted “lesion”.
- False Positive (FP): $c = 0$ and $\hat{c} = 1$, incorrectly predicted “lesion”.
- False Negative (FN): $c = 1$ and $\hat{c} = 0$, incorrectly predicted “healthy”.
- True Negative (TN): $c = 0$ and $\hat{c} = 0$, correctly predicted “healthy”.

Summing the number of voxels in each state over an entire image volume, voxel-wise agreement can then be quantified using the following measures:

- **Similarity Index (SI)** (AKA Dice Similarity Coefficient, F1-Score)

Measures overall segmentation performance.

$$SI = \frac{2TP}{2TP + FP + FN} \quad (2.28)$$

- **Precision (Pr)** (AKA Overlap Fraction, Positive Predictive Value)

Fraction of predicted positives which are true positives.

$$Pr = \frac{TP}{TP + FP} \quad (2.29)$$

- **Recall (Re)** (AKA Sensitivity, True Positive Rate)

Fraction of true positives which are predicted positive.

$$Re = \frac{TP}{TP + FN} \quad (2.30)$$

Each measure is $\in [0, 1]$, where higher is better. Note that typical performance metrics like accuracy and specificity are avoided, since they include the TN count in the numerator, which is typically much larger than $TP + FP + FN$ combined – i.e. $c = 1$ is a rare event.

Overall volume agreement between segmentations is characterized using the 2-way mixed-effects single-

rater absolute intraclass correlation coefficient (ICC)¹³ [180], while trends in over/undersegmentation with lesion load are illustrated using Blant-Altman plots [181].

2.7 Model Summary

In summary, the proposed algorithm uses graylevel features to train a logistic regression model for each voxel independently – Voxel-Wise Logistic Regression. In order to train the VLR model, a set of labelled training images must first be registered to a standard brain space (MNI). This is achieved using the SPM Segment tool, which also produces bias corrected images. Next, image intensities are standardized using a graylevel transformation, to be determined in the next section. The parameter images $\beta(x)$ are then computed using iterative MAP estimation, with Newton updates and an augmented dataset. These images are smoothed to reflect prior knowledge. Finally, the optimal probability threshold π_c and minimum lesion size x_{\min}^c are estimated using the training data. This completes the training phase.

At test time, SPM Segment is used again to correct the bias field and estimate the registration to MNI space for a given input image. However, the inverse transform is now used to warp the parameter images $\beta(x)$ from MNI space to the native space. This transformation of the smooth parameter images prior to inference is preferable to transforming the detailed label image afterwards. The probability of the WMH class is computed by evaluating the independent logistic models at every voxel. This initial estimate $\hat{C}(x)$ is then thresholded using π_c , and binary objects smaller than x_{\min}^c are removed. The resulting label image is the final output. These training and testing phases are illustrated in Figure 2.8.

2.7.1 Tunable Parameters

In order to achieve the best possible model performance, it is prudent to track tunable model parameters (AKA hyperparameters) which are distinct from those fitted during each cross validation fold – i.e. $\beta(x)$ and π_c . Considering both the main VLR model and the pre- and post-processing aspects, the parameters of the proposed algorithm are summarized in Table 2.2 and also programmatically in `hypdef.m`. The optimization of these model components will be the subject of the next chapter.

¹³ Option ‘A-1’ in the Matlab function ICC from <https://www.mathworks.com/matlabcentral/fileexchange/22099>

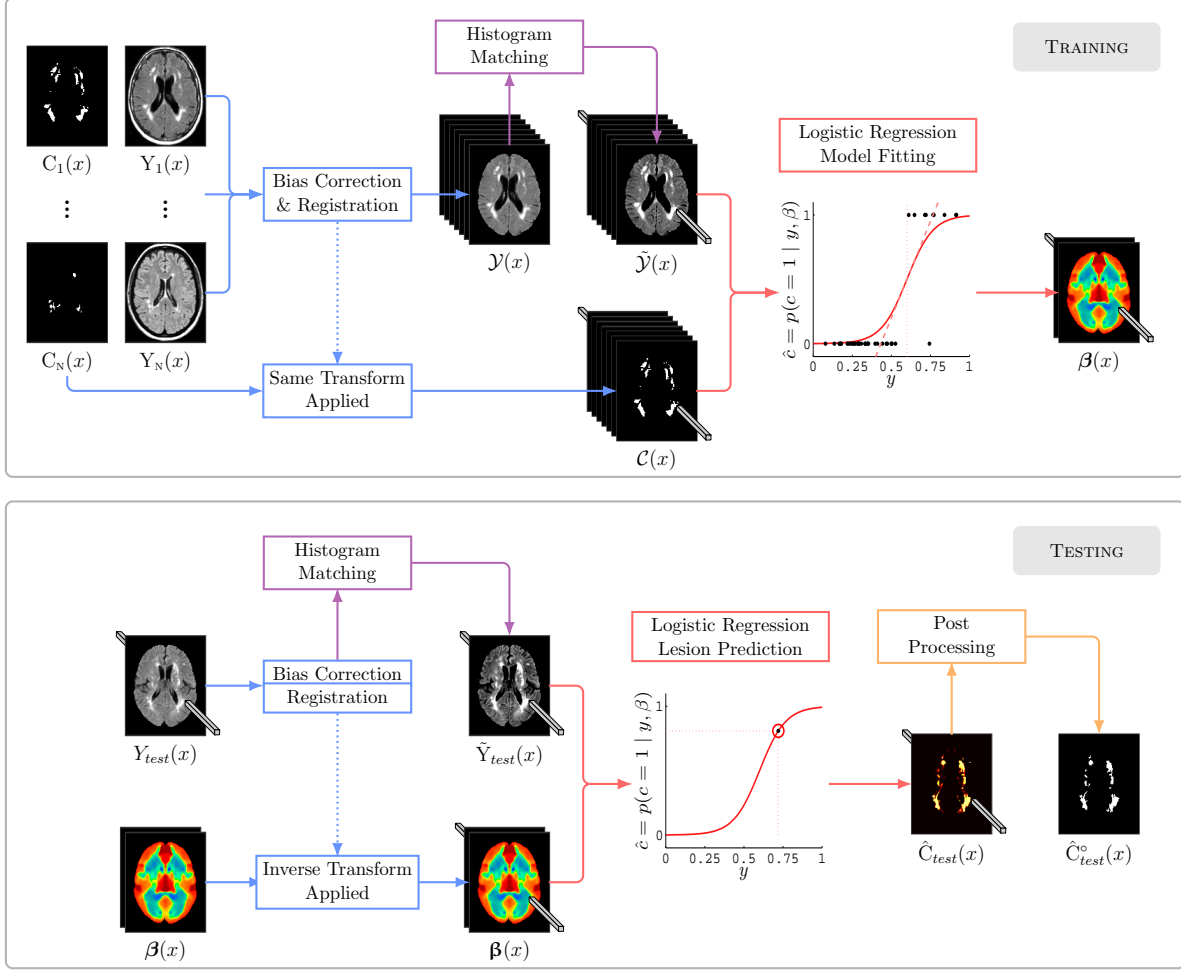


Figure 2.8: Overview of the proposed algorithm. Typefaces – upright Roman: images in native space; italic Roman: images in standard (MNI) space; calligraphic: a set of images from several patients; bold: a set of images corresponding to different features; Variables – $C(x)$: manual segmentation; $Y(x)$: FLAIR image, $\beta(x)$: parameter image; $\hat{C}(x)$: estimated lesion segmentation.

Table 2.2: Model hyperparameters.

Stage	Parameter	Notation	Type	Naive Default
Pre-Processing	Reflect Augmentation	a_R	\mathbb{B}	0
	Shift Augmentation	a_S	\mathbf{N}_p	\mathbf{N}_0
	Graylevel Transform	τ_y	$f : \mathbb{R} \mapsto \mathbb{R}$	τ_{he}
	Transform Mask	\mathcal{X}_τ	$\mathbb{B}(x)$	$\mathcal{X}_{\text{brain}}$
VLR Fitting	Iterations	T	\mathbb{Z}	30
	Initial β	$\beta^{(0)}$	\mathbb{R}^2	$[0, 0]$
	Learning Rate	α	\mathbb{R}	1
	Regularization	λ	\mathbb{R}	0
	Pseudo-Lesions	$\mathcal{V}(x)$	$\{\cdot \in \mathbb{R}\}$	$\{\}$
	β Filter	F_β	$f : \mathbb{R}(x) \mapsto \mathbb{R}(x)$	$\tilde{\beta}(x) = \beta(x)$
Post-Processing	Min Lesion Size	x_{\min}^c	$\mathbb{R} \text{ (mm}^3\text{)}$	0

Notation. \mathbb{B} : boolean value; \mathbb{Z} : integer value; \mathbb{R} : real value; \mathbb{R}^n : vector; $\mathbb{R}(x)$: image; \mathbf{N}_p : nearest p voxel neighbourhood.

Chapter 3

Experiment & Results

This section explores model validation. Performance of model components is characterized with respect to intermediate objectives, including graylevel standardization and regularization in toy scenarios. The segmentation performance of the full model is then presented under several cross validation frameworks, and compared to a similar algorithm.

3.1 Cross Validation Frameworks

Supervised segmentation models require the capacity to model complex relationships between the input image(s) and output label images. When models with large capacity are trained on a dataset which does not represent the full gamut of potential input data, they risk *overfitting*: acquiring a bias towards the training data [182]. The main problem associated with overfitting is decreased performance on new data (AKA generalization performance) [182]. Popular techniques for characterizing this expected decrease include cross validation (CV) procedures. These involve splitting the N available examples into training (t) and testing (e) subsets, where the training data are used to fit the model parameters, and the test data are used to approximate the expected generalization performance; the data splits are usually repeated, randomly or exhaustively, to ensure robust results [183]. The most popular CV frameworks include:

- **LOO – Leave-One-Out:** Withhold one example from the training set, and use it as the test case ($N_t = N - 1$; $N_e = 1$); repeat N times.

Benefit: Close approximation of the expected generalization performance

Drawback: Expensive to compute – $\mathcal{O}(N)$

- **KFCV – K-Fold Cross Validation:** Withhold a random batch of $B = N/K$ examples from the training set, and use it as the test set ($N_t = N - B$; $N_e = B$); repeat K times (without replacement).
Benefit: Less expensive to compute – $\mathcal{O}(N/K)$
Drawback: Worse approximation of the expected generalization performance

3.1.1 Leave-One-Source-Out CV

The choice of cross validation framework can have significant impacts on the reported model performance (see [183] for an in-depth review), and there is at least one assumption of the above methods which is not always valid: that examples are independent and identically distributed (iid). This is not true for data originating from multiple sources with different underlying distributions (e.g. MRI with different scan-parameter combinations) [184]. In fact, Geras and Sutton show that in multi-source problems where the expected use case involves data from entirely new sources, random KFCV (and therefore also LOO, as a special case of KFCV with $B = 1$) significantly overestimates the generalization performance. This is because random training fold selection allows the model to perceive source-specific characteristics of the test examples, which cannot be repeated for truly new examples. In such scenarios, the authors propose the following:

- **LOSO – Leave-One-Source-Out¹:** Withhold all examples from source $s \in 1 \dots S$ from the training set, and use these as the test set ($N_t = N - N_s$; $N_e = N_s$); repeat S times.
Benefit: Best approximation of the expected generalization performance in multi-source problems
Drawback: Still only an approximation

As noted in the introduction (cf. § 1.3.3), there has been surprisingly limited use of data from multiple sources for validation of WMH segmentation algorithms. Moreover, CV frameworks vary widely among papers, and to the best of this author’s knowledge, no WMH algorithm has yet been validated using LOSO CV. This represents a significant caveat to reported performances, since MRI have many sources of variability (cf. § 1.2.2), including scanner manufacturer, field strength, sequence parameters, resolution, anatomical and disease variability. As the aim of this work is to develop a segmentation algorithm which will perform well on any given FLAIR MRI, the LOSO framework was initially developed without knowledge of the work by Geras and Sutton. However, this paper happily corroborates the importance of LOSO CV to the current work. In this case, one data source is defined as a unique scanner-parameter combination.

¹ Authors’ original name was “Multi-Source Cross Validation”

3.1.2 Competition Frameworks

One notable exception to this trend in WMH algorithms have been segmentation competitions [80, 57, 81, 12]. These generally provide both multi-source datasets and a robust validation framework. Table 3.1

Table 3.1: Summary of competition image databases.

Database	Ref.	Training		Testing	
		I	S	I	S
WMH 2017	[12]	60	3	110	5
MS 2016	[81]	15	3	38	4
MS 2015 ISBI	[57]	21	1	61	1
MS 2008	[80]	20	2	32	2

3.1.3 Data

For the reasons outlined above, it was important to collect a large and diverse database of FLAIR images for model validation. We use 129 FLAIR images from 10 different scanners. The number of images and scan parameters are summarized in Table 3.2. With the exception of the MS 2008² and In-House datasets, all of the data are freely available as part of the segmentation competitions. Since direct comparison of results on equal datasets is important for establishing state-of-the-art, we present results using only these freely available data (“Dataset A”) in addition to all .

Regarding simulated MRI data, none are used in this work. While such data (e.g. BrainWeb [186]) are useful for evaluation of whole-brain segmentation methods, only three examples of WMH are currently available, and there is little documentation as to how these were generated³. Furthermore, several works [187, 188] have noted significant discrepancies in estimated segmentation performance between BrainWeb and real data (ISBR [167]).

² The manual segmentations used in this dataset were generated in-house, as described in § B.3.1.

³ Documentation found here: http://brainweb.bic.mni.mcgill.ca/brainweb/selection_ms.html.

Table 3.2: Summary of experimental image database.

Img (#)	Database	Ref.	Scanner	TE (ms)	TR (ms)	TI (ms)	Voxel Size (mm)	Manuals (#)
20	WMH 2017 (1) ■	[12]	3T Philips Achieva	125	11000	2800	$0.96 \times 0.96 \times 3.00$	1 ^a
20	WMH 2017 (2) ■	[12]	3T Siemens TrioTim	82	9000	2500	$1.00 \times 1.00 \times 3.00$	1 ^a
20	WMH 2017 (3) ■	[12]	3T GE Signa HDxt	126	8000	2340	$0.98 \times 1.20 \times 3.00$	1 ^a
5	MS 2016 (1) ■	[81]	3T Philips Ingenia	360	5400	1800	$0.50 \times 1.10 \times 0.50$	7 ^b
5	MS 2016 (2) ■	[81]	1.5T Siemens Aera	336	5400	1800	$1.04 \times 1.25 \times 1.04$	7 ^b
5	MS 2016 (3) ■	[81]	3T Siemens Verio	399	5000	1800	$0.74 \times 0.70 \times 0.74$	7 ^b
21	MS 2015 ISBI ■	[57]	3T Philips	68	11000	2800	$0.43 \times 0.43 \times 3.00$	2 ^c
13	In-House ■	—	3T Philips Achieva	125	9000	2800	$1.00 \times 1.00 \times 1.00$	1 ^d
10	MS 2008 CHB ■	[80]	—	—	—	—	$0.50 \times 0.50 \times 0.50$	1 ^e
10	MS 2008 UNC ■	[80]	3T Siemens Allegra	125	9000	2800	$0.50 \times 0.50 \times 0.50$	1 ^f

^a Manuals were generated following the standards outlined in [79], and were subsequently reviewed by a second rater, only WMH labels were included; ^b Manuals were fused using the LOP-STAPLE method [185]; ^c Manuals were fused using logical ‘and’; ^d Manuals were generated in-house; ^e REVIEW; ^f REVIEW.

3.2 Graylevel Standardization

3.3 Regularization

3.3.1 Toy Model

In order to investigate the effect of different regularization strategies, a toy model is used. The model represents a single voxel during training, with synthetic observations. Regularizations are then chosen by hand to maintain desired characteristics in the fitted functions.

It is also possible to plot the MAP objective function $\mathcal{J}(\beta)$ in the 2D plane composed of $\beta = [\beta^0, \beta^1]$.

3.4 Full Model

3.4.1 Convergence

The rate of convergence in each voxel will be unique. During parallel fitting, it is prudent to stop training after a reasonable number of iterations, rather than wait for all voxels to achieve a certain stopping criterion, in case a few aberrant voxels do not converge. In order to determine this number, the model was fitted using [X] selection(s) of the training data, and the value of the MAP objective

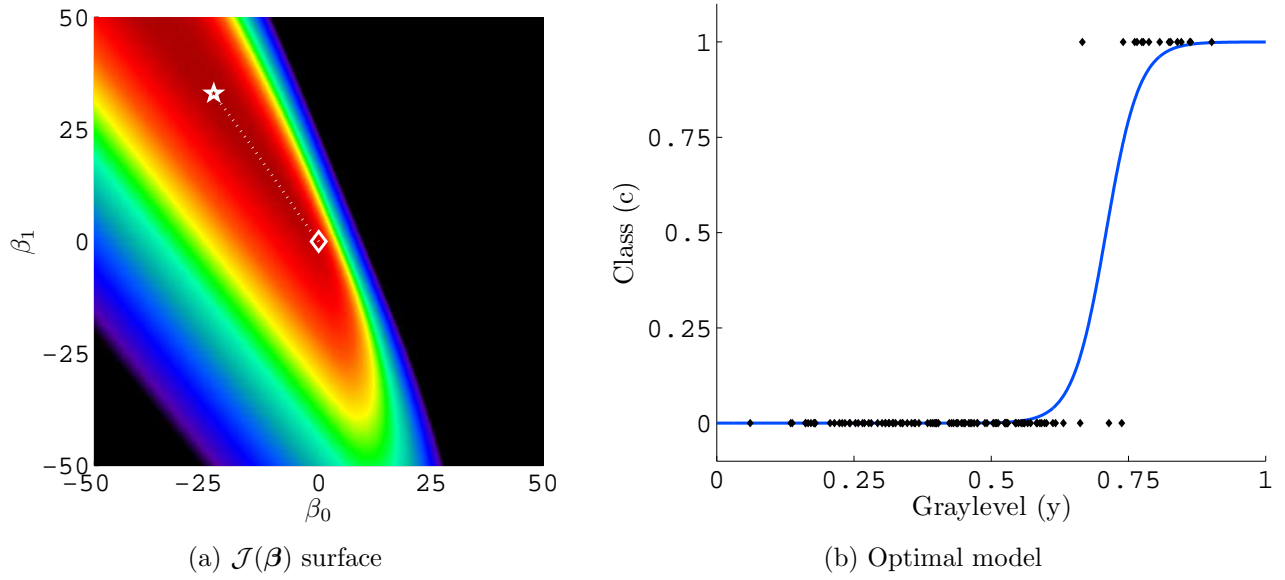


Figure 3.1: MAP objective function in 2D; initialization (diamond), fitting (dotted), optimum (star).

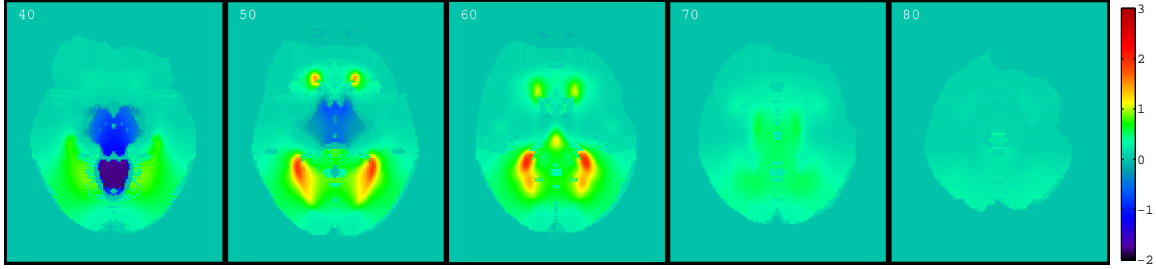


Figure 3.2: Spatial effect parameter image from the LPA toolbox after registration to MNI space.

3.4.2 Parameter Images

For comparison with the LPA algorithm, the

3.4.3 Quantitative Performance

Regularization

3.4.4 Comparison with Another Method

3.5 Discussion

Chapter 4

Conclusion

4.1 Summary

4.2 Future Work

4.2.1 Other Features

4.2.2 Supervised Graylevel Standardization

4.2.3 Integrative Models

4.2.4 Open Sourcing

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Appendix A

Maths

This section presents various mathematical results which are not essential to the thesis.

A.1 FLAIR MRI Intensity Modelling

While MR imaging is both complex and mutable, simulation of the expected signal intensities is possible using the relaxometry data in Table 1.1 and sequence signal equations – e.g. (1.3) and (1.4). In practice, this simulation helps select appropriate acquisition parameters $TE/TR/TI$ for the desired contrast; however, these characteristics can also be later considered as covariates in performance analysis of segmentation tools. To this end, Equation (1.4) – repeated here for reference,

$$\Psi_{IR}(x) = \left[K [H(x)] \right] \left[e^{-\left(\frac{TE}{T2(x)}\right)} \right] \left[1 + e^{-\left(\frac{TR}{T1(x)}\right)} - 2e^{-\left(\frac{TI}{T1(x)}\right)} \right] \quad (\text{A.1})$$

– was used with the acquisition parameters of the FLAIR image database (Table 3.2) and the relaxometry data from Table 1.1 to calculate the expected tissue intensities and WMH contrasts for the experimental data. These results are summarized in Table A.1. In Figure A.1, an example image is shown for each parameter set using the tissue maps from the BrainWeb database [186]¹, while in Figure A.2, the PMF for each tissue class from the same data are given.

¹ <http://brainweb.bic.mni.mcgill.ca/>

Table A.1: Simulated FLAIR tissue intensities and WMH contrasts using scan parameters from the experimental database. Tissue intensities are normalized to the WM value.

Scanner	GM	WM	CSF	WMH	$\frac{\text{WMH}}{\text{GM}}$	$\frac{\text{WMH}}{\text{WM}}$	$\frac{\text{WMH}}{\text{CSF}}$
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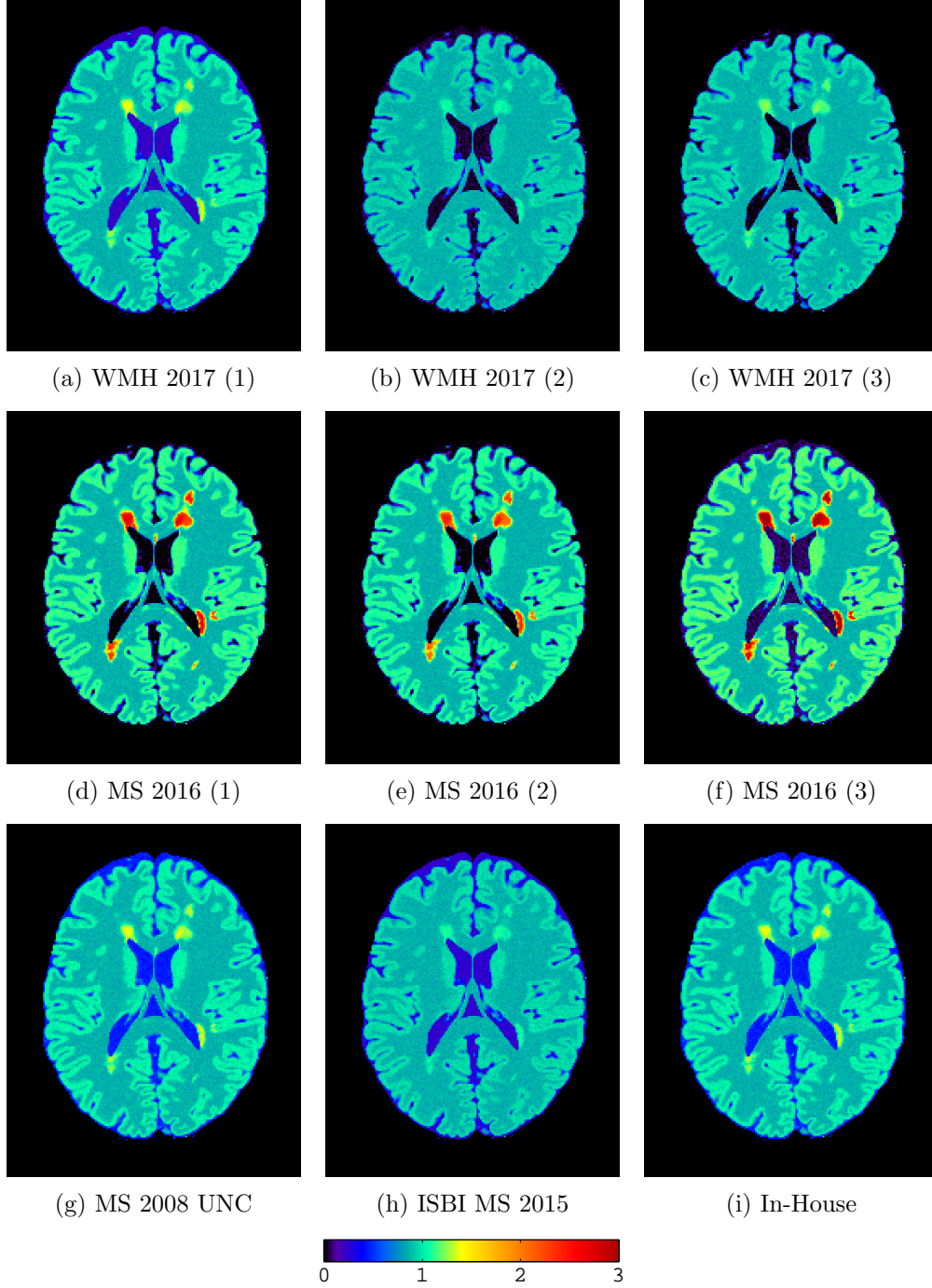


Figure A.1: Simulated FLAIR images using scan parameters from the experimental database. Colourmap is arbitrary but consistent.

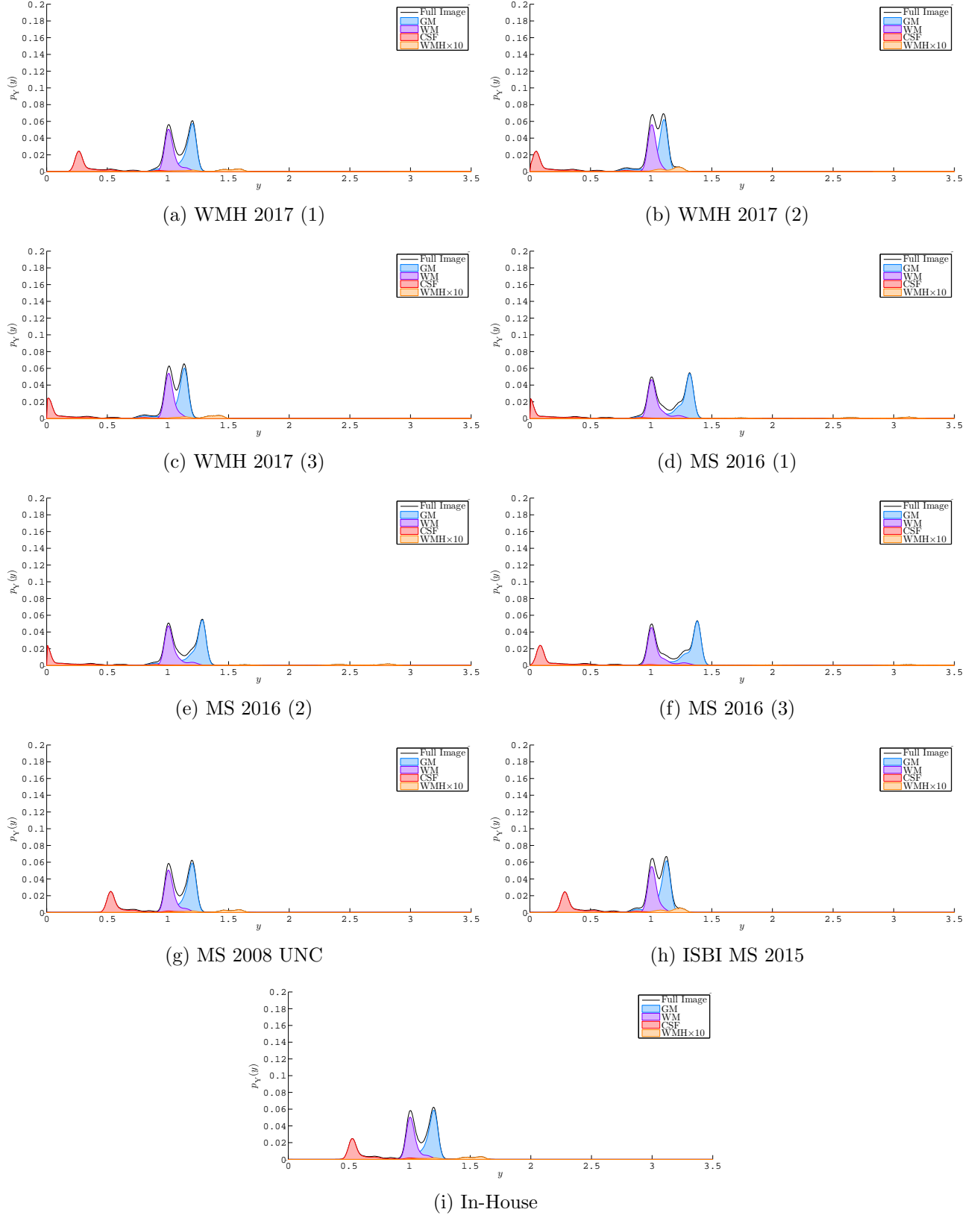


Figure A.2: PMF of each tissue from simulated FLAIR images. The PMF of the WMH class is scaled by 10 for visibility.

A.2 Graylevel Standardization

A.2.1 Histogram Matching vs Histogram Equalization

In § 2.4.3 it was argued that histogram matching is equivalent to histogram equalization in terms of effectiveness at standardizing graylevels in heterogeneous input images. This is because the histogram matching operation is defined as the function composition of the histogram equalization transform of the input image, F_Y , and the inverse equalization transform for the desired output histogram, $F_{\tilde{Y}}^{-1}$,

$$\tau(y) = F_{\tilde{Y}}^{-1}(F_Y(y)) \quad (\text{A.2})$$

The second transformation in the cascade does not depend on Y , and so it is applied equally to images. Therefore the choice of target PMF is not important for the objective of graylevel standardization. This result is verified experimentally using synthetic images. Four $100 \times 100 \times 100$ images were generated to have the following density functions f_Y ,

- Uniform: $y \sim \mathcal{U}(y_{\min} = 0, y_{\max} = 1)$
- Unimodal: $y \sim \mathcal{N}(\mu = 0.5, \sigma = 0.08)$
- Bimodal: $y \sim (0.5\mathcal{N}(\mu = 0.3, \sigma = 0.05) + 0.5\mathcal{N}(\mu = 0.7, \sigma = 0.05))$
- Trimodal: $y \sim (0.3\mathcal{N}(\mu = 0.25, \sigma = 0.05) + 0.4\mathcal{N}(\mu = 0.5, \sigma = 0.05) + 0.3\mathcal{N}(\mu = 0.75, \sigma = 0.05))$

All four images were then histogram-matched to each of the respective density functions, with the aim of increasing agreement of image intensities. This agreement can be approximated by the alignment of intensity quantiles, since this is the target of histogram matching operations. As shown in Figure A.3, the quantiles agree almost perfectly in every output image, regardless of the choice of output PMF.

A.2.2 Nyul Approximation of Histogram Matching

This topic is treated in [175].

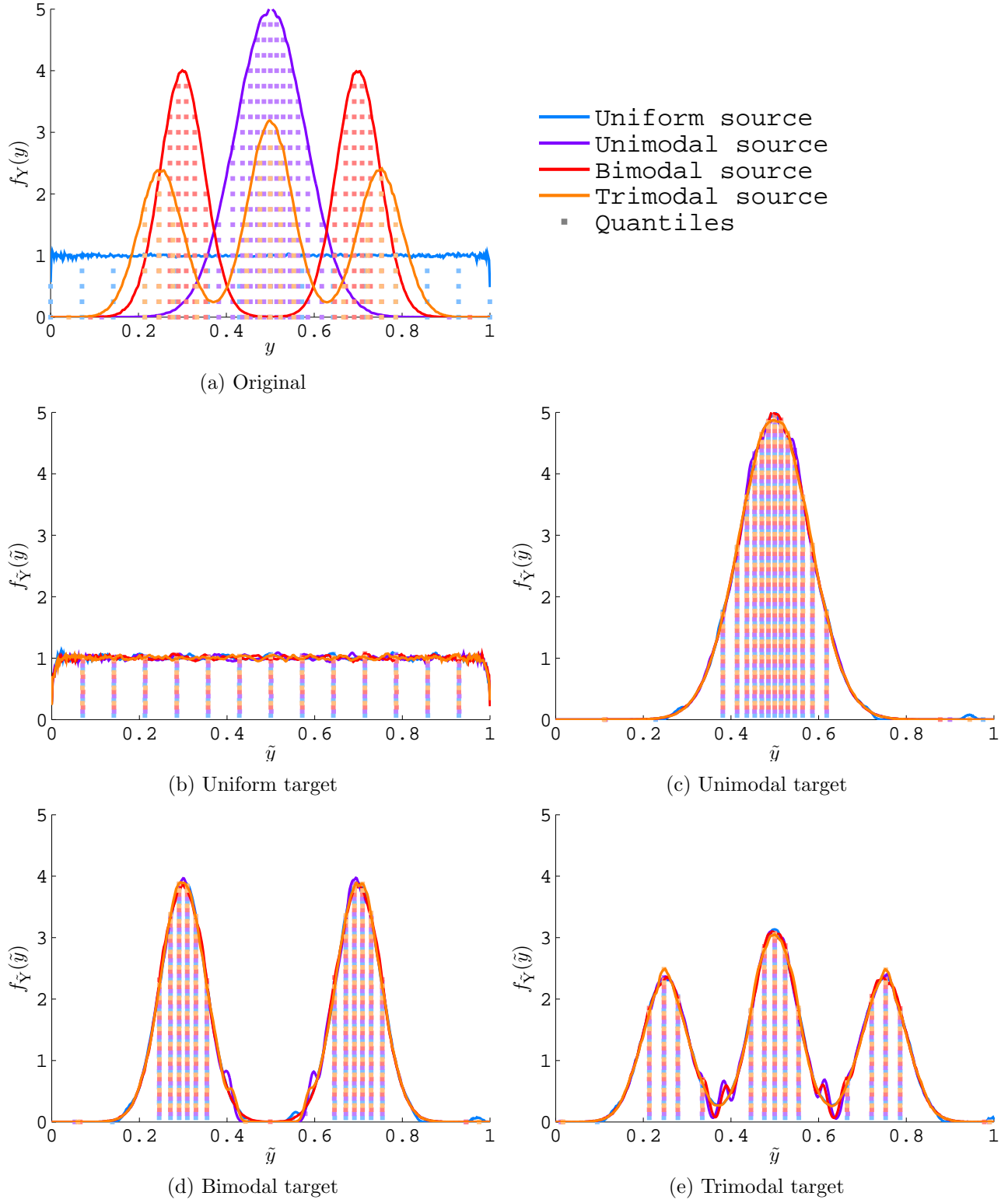


Figure A.3: Histogram matching of synthetic data to different target histograms. Quantiles show high agreement regardless of the target histogram.

Appendix B

Implementation

This appendix contains implementation details.

B.1 Computing

All computation in the current work was performed using the following workstation and software:

- **CPU:** Intel Core i7-6700K 4.00 GHz
- **RAM:** 16.0 GB DDR4
- **GPU:** NVIDIA GeForce GTX 980 Ti
- **OS:** Windows 10 Pro
- **Code:** MATLAB R2011a

B.2 Parallelization

Speed of model fitting is a significant factor during development, particularly considering optimization of hyperparameters. Faster training yields more model iterations, which inevitably bear improvements.

B.2.1 Model Estimation

While the estimation procedure outlined in § [2.2.1](#) must be repeated for all standardized voxels in the brain mask, it is possible to do this in parallel, since every estimation is independent. To do so, the

training data must first be vectorized with respect to spatial location x , and matrix operations expanded explicitly to accommodate the new dimension.

To begin, the standardized training data from all subjects – features $\tilde{\mathcal{Y}}_\gamma(x)$, with $\tilde{\mathcal{Y}}_\gamma^0 = 1$, and labels $\mathcal{C}_\gamma(x)$ – are sampled from nonzero locations in the brain mask $M(x)$. These data are stored in two matrices \mathbb{Y} and \mathbb{C} , with dimensions $[X, N, K + 1]$ and $[X, N, 1]$, respectively, where X is the total number of nonzero voxels in the brain mask, N is the number of subjects, and K is the number of features. A similar matrix is constructed for the initial parameters $\beta^{(0)}(x)$, denoted $\mathbb{B}^{(0)}$, with dimensions $[X, 1, K + 1]$. Let \mathbb{Y}_n^k denote the vector of data from all voxels for the k^{th} feature from the n^{th} subject, and so on for \mathbb{C} and \mathbb{B} .

In order to simplify subsequent calculations, the feature data are rectified according to the class labels, before the first iteration, as in

$$\mathbb{Y}_n^k = \begin{cases} +\mathbb{Y}_n^k, & \mathbb{C}_n \geq 0.5 \\ -\mathbb{Y}_n^k, & \mathbb{C}_n < 0.5 \end{cases}, \quad \forall k \in \{1, \dots, K\}. \quad (\text{B.1})$$

Next, for a given iteration t , the following vector-compatible expansions of Equations (2.10), (2.11), and (2.14) yield the desired update matrix $\Delta\mathbb{B}^{(t)}$. These calculations are performed in ???. Regarding notation: 1) the iteration index $^{(t)}$ is omitted for clarity, 2) element-wise multiplication is denoted by \circ , and 3) the variable K is now defined as 1, since this is essential to the simplification.

$$\mathbb{S} = \frac{1}{1 + e^{+\eta}}, \quad \eta = \mathbb{B}^0 + (\mathbb{B}^1 \circ \mathbb{Y}^1) \quad (\text{B.2})$$

$$\mathbb{A} = \mathbb{S} \circ (1 - \mathbb{S}) \quad (\text{B.3})$$

$$\begin{aligned} \mathbb{G} &= \nabla_{\mathbb{B}} \mathcal{L} - \lambda \mathbb{B} \\ &= \begin{bmatrix} \mathbb{G}^0 \\ \mathbb{G}^1 \end{bmatrix} \\ &= \begin{bmatrix} \sum_{n=1}^N (\mathbb{Y}_n^0 \circ \mathbb{S}) \\ \sum_{n=1}^N (\mathbb{Y}_n^1 \circ \mathbb{S}) \end{bmatrix} - \lambda \begin{bmatrix} \mathbb{B}^0 \\ \mathbb{B}^1 \end{bmatrix} \end{aligned} \quad (\text{B.4})$$

$$\begin{aligned} \mathbb{H} &= \nabla_{\mathbb{B}}^1 \mathcal{L} - \lambda \mathbb{I} \\ &= \begin{bmatrix} \mathbb{H}^{0,0} & \mathbb{H}^{0,1} \\ \mathbb{H}^{1,0} & \mathbb{H}^{1,1} \end{bmatrix} \\ &= \begin{bmatrix} \sum_{n=1}^N (\mathbb{A} \circ \mathbb{Y}_n^0 \circ \mathbb{Y}_n^0) & \sum_{n=1}^N (\mathbb{A} \circ \mathbb{Y}_n^1 \circ \mathbb{Y}_n^0) \\ \sum_{n=1}^N (\mathbb{A} \circ \mathbb{Y}_n^0 \circ \mathbb{Y}_n^1) & \sum_{n=1}^N (\mathbb{A} \circ \mathbb{Y}_n^1 \circ \mathbb{Y}_n^1) \end{bmatrix} - \lambda \begin{bmatrix} 1 & \\ & 1 \end{bmatrix} \end{aligned} \quad (\text{B.5})$$

$$\begin{aligned} \mathbb{D} &= \det \mathbb{H} \\ &= (\mathbb{H}^{0,0} \circ \mathbb{H}^{1,1}) - (\mathbb{H}^{0,1} \circ \mathbb{H}^{1,0}) \end{aligned} \quad (\text{B.6})$$

$$\begin{aligned} \Delta \mathbb{B} &= -\mathbb{H}^{-1} \mathbb{G} \\ &= \frac{1}{\mathbb{D}} \begin{bmatrix} (\mathbb{H}^{1,1} \circ \mathbb{G}^0 - \mathbb{H}^{1,0} \circ \mathbb{G}^1) & (\mathbb{H}^{0,1} \circ \mathbb{G}^1 - \mathbb{H}^{0,0} \circ \mathbb{G}^0) \end{bmatrix}^T \end{aligned} \quad (\text{B.7})$$

B.2.2 Image Deformations

Following training, each indi

While these deformations are saved after initial estimation, their application is computationally expensive.

B.3 Manual Segmentations

It was necessary to create and edit a small number of binary segmentation masks during this work. To do this, the Editor module from the 3D Slicer imaging platform [189] was used¹, including the Wand, Paint, and Erase functions. Figure B.1 shows the user interface during a lesion segmentation.

¹ 3D Slicer Editor tool documentation is available here: <https://www.slicer.org/wiki/Documentation/4.6/Modules/Editor>.

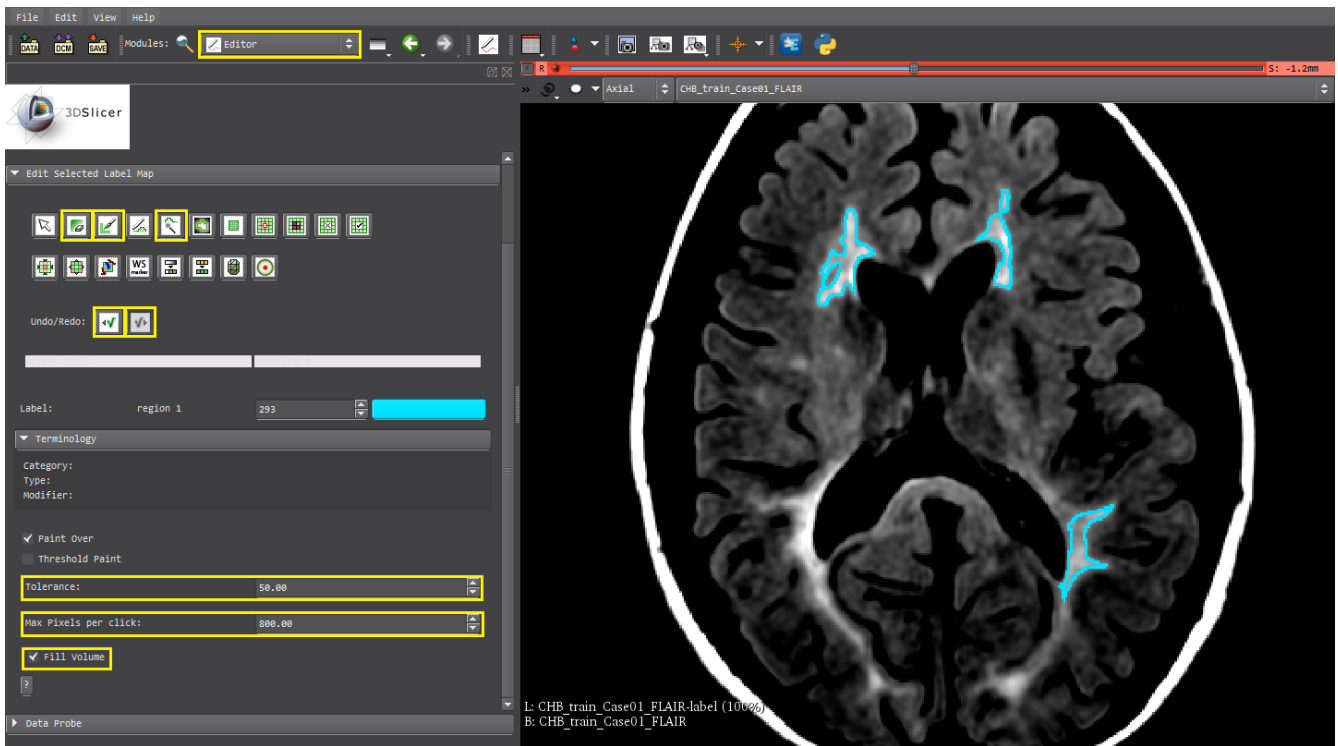


Figure B.1: 3D Slicer user interface for performing in-house manual segmentations and revisions. The tools used are highlighted in yellow, while the segmentation (in-progress) is shown in blue.

B.3.1 MS 2008 WMH Masks

Since the reported performance of an automatic segmentation algorithm depends on the manual segmentations to which it is compared, it is important to obtain good manual segmentations. Unfortunately, the original manuals in the MS 2008 Segmentation Challenge contained obvious artifacts and inconsistencies, as shown at left in Figure B.2. Therefore, it was deemed necessary to redo these manuals. The resulting revisions are shown at right in Figure B.2.

B.3.2 Brain Mask

In order to vectorize image data for parallel processing, a binary mask selecting voxels of interest in standardized space is also required. Since only voxels in the brain are of interest, this mask is called a “brain mask”. The brain mask used here was derived from the ICBM tissue prior images [132] in MNI space: after initial thresholding of the combined GM + WM + CSF probabilities at 0.5, manual refinements were completed and symmetry was enforced. The mask is slightly small on purpose, since tissues outside the brain are frequently bright in FLAIR images, and can be mistaken for lesions by naive models. The resulting mask is shown in Figure B.3.

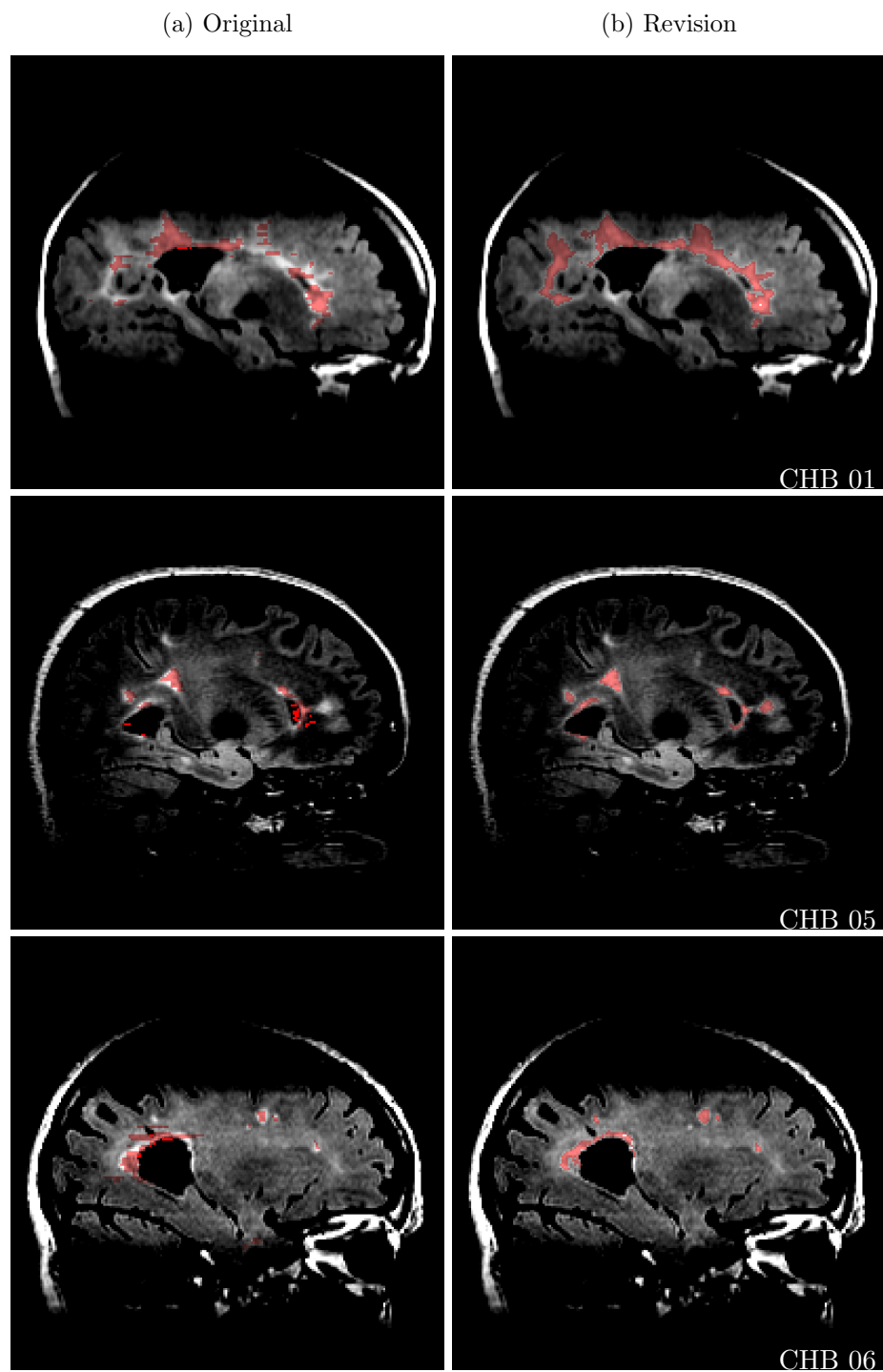


Figure B.2: Example revisions to the manual segmentations for the MS 2008 challenge dataset.

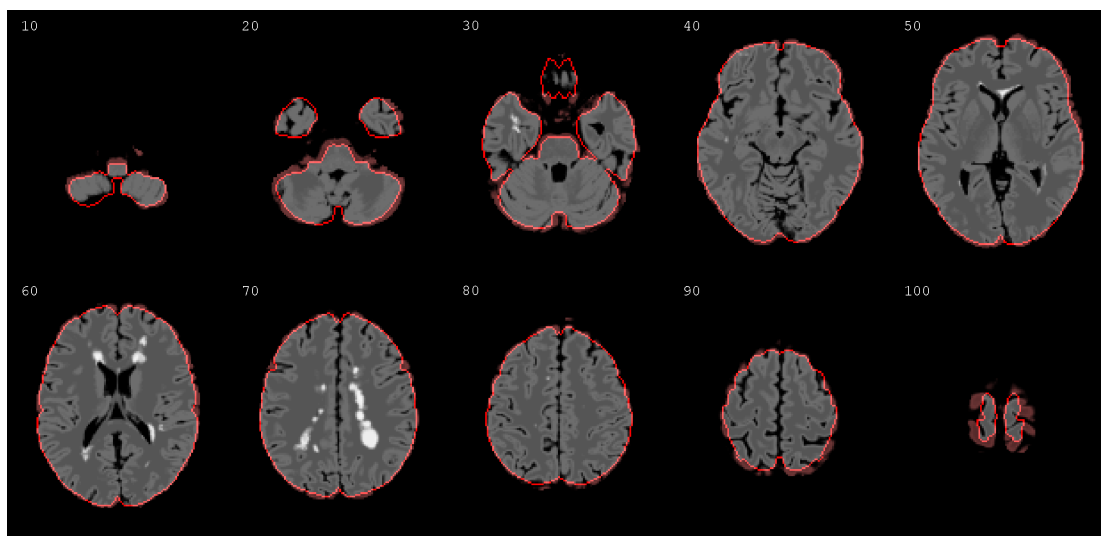


Figure B.3: Manually refined brain mask in MNI space, overlaid on a simulated BrainWeb FLAIR image. Mask outline is highlighted in red; inclusions are shown in grayscale; exclusions are tinted red.

Appendix C

Code

>> hypdef.m

```
function [h] = hypdef(h)
% flag-like names
h.name.key      = 'WMH17';    % h.name.key      = 'LPA';
h.name.data     = 'mni96';    % h.name.data     = 'mni109';
h.name.cv       = 'loso';     % h.name.cv       = 'nocv';
% scanner parameters
h.cmap          = NIH3;
h.Ni            = [];
h.scan.idx      = [1,2,3,4,5,6,9]; % h.scan.idx      = [1,2,3,4,5,6,9,7,8,10];
h.scan.clr      = rainbow7;    % h.scan.clr      = rainbow10;
% sampling parameters
h.sam.fresh     = 0;
h.sam.resize    = 0.5;
h.sam.dx        = kernelshifts(binsphere(1));
h.sam.flip      = 1;
% grey standardization parameters
h.std.type      = 'pdfeq';
h.std.pdf       = 'gauss';
h.std.mask      = 'brain';
% logistic regression parameters
h.lr.Nit        = 30;
h.lr.B          = [0,0];
h.lr.alpha      = 1;
h.lr.reg.la     = 1e-3;
h.lr.reg.py     = [0,1];
h.lr.reg.pc     = [0,1];
h.lr.pad        = [-1.5;1];
h.lr.pp.filter  = @(B)(op23(B,@(B)medfilt2(B,[3,3])));
% post processing parameters
h.pp.saveles    = 'les';
h.pp.thr.def    = 0.5;
h.pp.thr.Nit    = 30;
h.pp.minmm3     = 5;
h.pp.mrf.Nit    = 0;
h.pp.mrf.w      = 0.5;
h.pp.mrf.q      = 0.8;
% cross validation and
h = hypfill(h);
```