



"Segmentation in CT scan and multimodal MRI datasets using 3D active contours"

Vanden Bulcke, Colin

ABSTRACT

Context: Segmentation is an important tool in the diagnosis and prognosis as well as for the treatment planning of medical disorders. In hospitals, the two main departments that need segmentation are the neurosurgery and the radiotherapy departments. The first one uses currently mainly manual segmentation tools while the second one uses also AI-based and atlas-based segmentation the easiest applications.

Objective: The objective of this work is to develop and implement a user-friendly semi-automatic tool to perform segmentation on medical images that would reduce the task time taken by the doctors.

Method: The method developed uses a region-based 3D active contour algorithm using robust statistics which allows good interactions with the user. To support this algorithm, a graphical user interface was implemented to enable users to easily use this segmentation tool. This method was tested in two applications; the segmentation of femoral heads in CT scans and brain pathologies in MRI.

Results: The results show that the algorithm produces promising results with mean dice coefficients of 0.868 and 0.9016 for the two applications. This algorithm helps dividing the drawing time by a mean factor of 6.996 compared to software used in hospitals.

Conclusion: This region-based active contour algorithm is an interesting and flexible segmentation tool. It allows to have good interactions with the user and, with a good GUI, can be used by doctors in their daily work for a variety of applications. The tool could also be inserted in the workflow of an active learning segmentation scheme ...

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École polytechnique de Louvain

Segmentation in CT scan and multimodal MRI datasets using 3D active contours

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Master in Biomedical Engineering

Abstract

Context Segmentation is an important tool in the diagnosis and prognosis as well as for the treatment planning of medical disorders. In hospitals, the two main departments that need segmentation are the neurosurgery and the radiotherapy departments. The first one uses currently mainly manual segmentation tools while the second one uses also AI-based and atlas-based ~~segmentation~~ the easiest applications.

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Results The results show that the algorithm produces promising results with mean dice coefficients of 0.868 and 0.9016 for the two applications. This algorithm helps dividing the drawing time by a mean factor of 6.996 compared to software used in hospitals.

Conclusion This region-based active contour algorithm is an interesting and flexible segmentation tool. It allows to have good interactions with the user and, with a good GUI, can be used by doctors in their daily work for a variety of applications. The tool could also be inserted in the workflow of an active learning segmentation scheme to further improve the segmentation process and provide the medical staff with a new efficient and robust way to label images.

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List of Abbreviations

i.e.	id est (That is)
e.g.	exempli gratia (for example)
CT	Computerized Tomography
MRI	Magnetic Resonance Imaging
ROI	Region of Interest
VOI	Volume of Interest
CAD	Computed-Aided Diagnosis
DNA	Deoxyribonucleic Acid
CTV	Clinical Target Volume
GTV	Gross Tumor Volume
OARs	Organs At Risk
GUI	Graphical User Interface
PET	Positron Emission Tomography
RF	RadioFrequency
FOV	Field of View
LUT	LookUp Table
AI	Artificial Intelligence
ML	Machine Learning
DL	Deep Learning
ANN	Artificial Neural Network
AN	Artificial Neuron
DN	Deep Network
CNN	Convolutional Neural Network
FCN	Fully Convolutional Network
MED	Median
IQR	Inter-Quartile Range
MAD	Meadian Absolute Deviation
KDE	Kernel Density Estimation
pdf	Probability Density Function
AC	Active Contour
DC	Dice Coefficient
JI	Jaccard Index
IH	Intensity Homogeneity
CW	Curvature Weight
In	Inertia

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Chapter 1

Medical Segmentation Nowadays

Contents

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This chapter aims to describe the segmentation techniques used nowadays by doctors in the framework of their work. It will begin by giving the medical context where the segmentation are used, the most commonly used methods and software in hospitals in neurosurgery and radiotherapy and finally the objective of this thesis and how this thesis could contribute facilitating the work performed by the doctors.

1.1 Motivation

Nowadays, medical imaging is an important tool in the diagnosis and prognosis as well as for the treatment planning of medical disorders. Computed Tomography (CT) is the most common data type used in medical imaging [1] with Magnetic Resonance Imaging (MRI) closely behind [2]. With the increase in the use of medical imaging, especially 3D-scan, segmentation has undergone rapid advancements over the past decades [1]. Segmentation is a very important process and consists of dividing an image into regions with similar properties, called Regions of Interest (ROIs) in 2D or Volumes of Interest (VOIs) in 3D, with respect to certain criteria [2, 3]. The segmentation of a meningioma can be observed in Figure 1.1. The goal of generating these regions is to be able to reconstruct a 3D model of the VOI [1]. In medical imaging, the aim of these 3D models is to [2]:

- study anatomical structure
- identify ROI or VOI (tumor, lesions, abnormalities, ...)
- measure tissue volume (growth of a tumor, ...)
- help in treatment planning (radiation therapy, surgery, ...)

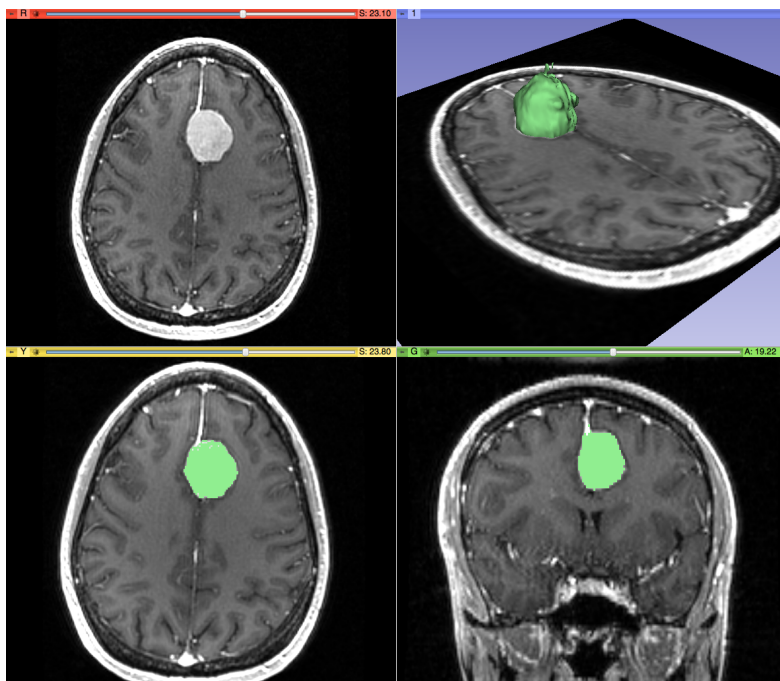


Figure 1.1: Segmentation and 3D modelling of a meningioma [4]

Segmentation can be made manually by a specialist or can be performed by a computer using an automatic or semi-automatic algorithm. Recently, with the increasing use of CT and MRI for diagnosis, treatment planning and clinical studies, it has become almost compulsory to use computers to assist radiological experts in clinical diagnosis and treatment planning. Doing segmentation manually on 3D volumes is today too time-consuming [2]. Moreover, even state-of-the-art techniques are not fast enough for real-time processing or high-resolution images [5]. Therefore, reliable algorithms are required for the delineation of VOI. The goals of Computed-Aided Diagnosis (CAD) are [2]:

- To automate the process so that a large number of cases can be handled with the same accuracy, i.e. the results are not affected as a result of fatigue, data overload or missing manual steps.
- To achieve fast and accurate results. Very high-speed computers are now available at modest costs, speeding up computer-based processing in the medical field.
- To support faster communication, wherein patient care can be extended to remote areas using information technology.

The techniques available for segmentation are often specific to the application and function on one hand of the medical imaging modality and on the other hand of the body part which is studied. Indeed, depending on the imaging modality, the contrast and the artifacts of the image between certain tissue are totally different. The texture can even vary in the same VOI. There is no universal algorithm for the segmentation of every medical image. Each imaging system has its own specific limitations [1, 2]. In the following sections, different tools and software used nowadays by doctors will be described.

1.2 Methods used in hospital

This section highlights the different segmentation techniques that are commonly used in hospitals. The two main departments in hospitals using segmentation are the neurosurgery and the radiotherapy departments. Both use segmentation as a tool for treatment planning.

1.2.1 Neurosurgery

Segmentation of relevant structure is really important in neurosurgery, especially in the treatment of brain pathology. Proper identification of pathology localization is crucial to deliver the best treatment possible. It gives information about the surrounding healthy tissue and whether or not the pathology is located in a sensitive region. For some cases, if the region is not too sensitive (e.g. in the frontal cortex), it will be preferable to enlarge a bit the region to be sure to treat correctly all the pathological tissue. On the contrary, if the region is really sensitive (e.g. in the visual cortex), then it is important to treat only the tissue that absolutely must be treated. The goal here is to cure the pathology while provoking the least damage possible. From a more technical point of view, the localization of the pathology in the brain helps the neurosurgeon to choose all the parameters of its procedure; namely where to make the incision, which angle is needed to reach the pathology, etc. All of this is chosen again in order to minimize damage to the patient [6].

Now, let us see what tools the neurosurgeons are using nowadays to perform their segmentation. In Belgium, the principal software that are used in this department are BrainLab and StealthStation by MedTronic, depending on the hospital. You can find the main working window of these software respectively in Figure 1.2 and Figure 1.3. Although slightly different, both these software use the same principle to perform segmentation. It is an enhanced manual segmentation. The neurosurgeon needs to draw manually on the different slices the region of interest but the software provides the user with an interpolation function that allows him to not draw on every slice. The interpolation function will connect the slices drawn by completing the slices in between. In practice, this allows the neurosurgeon to draw only on every 4 slices on average which is a little time-saving. Since it uses an interpolation function, it works well for regions that have a relatively smooth curvature (which is often the case). However, this principle is not using any information from the image while connecting the drawn slices which could lead to some big inaccuracy. Therefore, it can be said that neurosurgeons are performing manual segmentation [6].

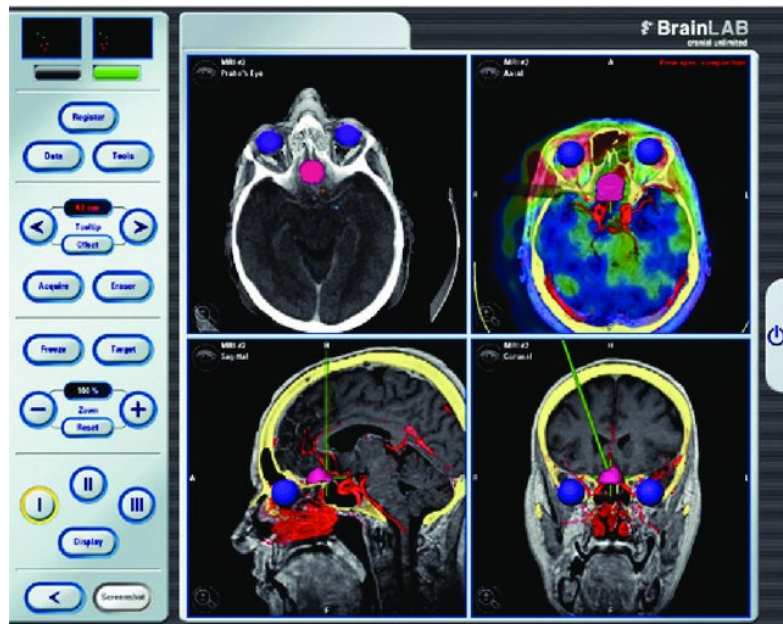


Figure 1.2: Main working window of the BrainLab Software [7]

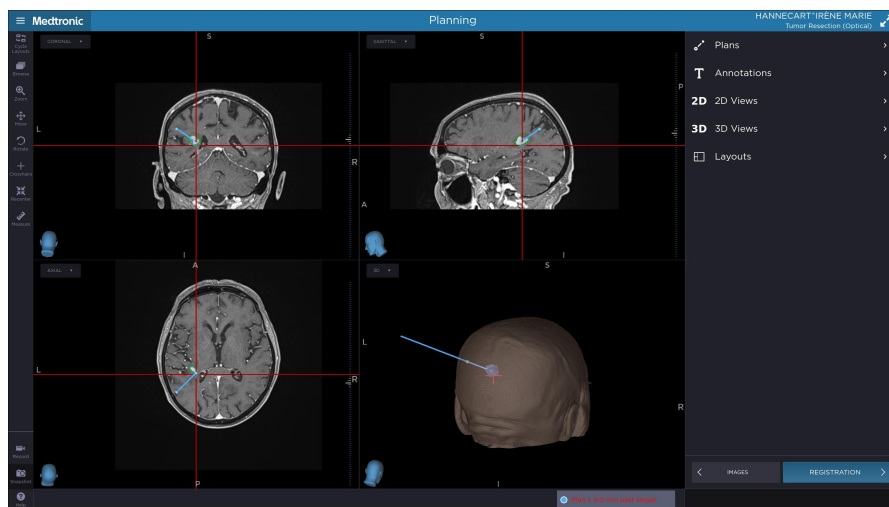


Figure 1.3: Main working window of the MedTronic Software [8]

1.2.2 Radiotherapy

Radiotherapy is a cancer treatment that uses the properties of radiation as a local tumor control. More precisely, it uses a beam of high energy X-rays, called ionizing

radiation, that carries enough energy to free electrons from atoms or molecules. These radiations aim to damage the DNA (Deoxyribonucleic Acid) of the cancerous cells in order to kill them. This is based on the fact that the mechanisms of DNA repair in cancerous cells are often altered. These radiations will therefore have a more deadly impact on the cancerous cells than on the healthy ones. The principal objective of radiotherapy is to deliver the highest dose of radiation to cancerous cells while minimizing the dose received by healthy tissue [9–11].

Figure 1.4 represents a typical treatment planning in radiotherapy. The first phase of the treatment (the top line of the Figure 1.4) corresponds to the imaging of the patient and the definition of the interesting volumes. In simple terms, the interesting volumes correspond to the Clinical Target Volume (CTV), corresponding to the Gross Tumor Volume (GTV), which is the visible cancerous cells, plus a margin to encompass the microscopic tumorous cells that lie around the GTV. This CTV corresponds to the region that needs to be treated. The other interesting volumes are the Organs At Risk (OARs), which are the surrounding vital organs that need to be clear from the radiation [9]. The delineation of these volumes is thus essential for the quality of the treatment delivery. And this is where segmentation comes in. Then, the second phase of the treatment (the bottom line of the Figure 1.4) corresponds to the planning of dose distribution to cure the cancer. This second phase does not fall into the scope of this work [11].

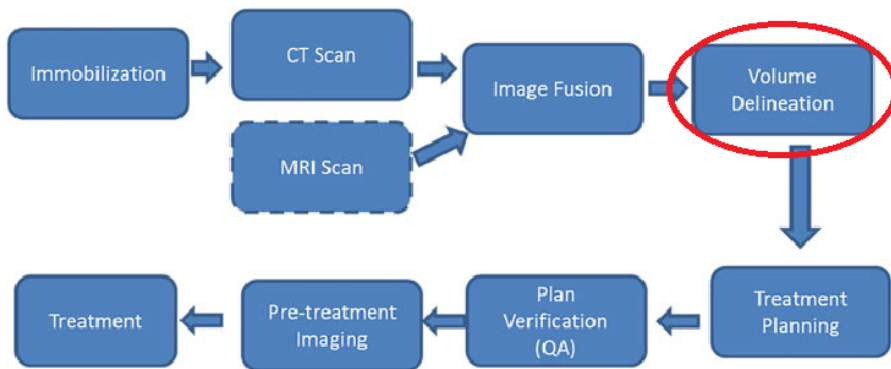


Figure 1.4: Flowchart of a classical treatment planning in radiotherapy. The red circle represents the moment where segmentation is performed [12]

In the radiotherapy field, segmentation is still principally done manually with some assistance, although automatic methods are starting to arrive. These automatic tools, composed of Atlas-based and Deep Learning methods, are used to segment anatomical volumes (OARs) but only in particular cases (mainly lungs, and head & neck). However, the clinical volumes (GTV, CTV) are always segmented manually due to its complexity. The most commonly used software in radiotherapy is RayStation from RaySearch

Laboratories [13] and its main working window can be found in Figure 1.5. This software contains all the features to plan completely a treatment; from segmentation of the volumes to the dose calculation and optimization. But, other tools exist on the market for the segmentation of the different interesting volumes; the Limbus AI [14] and the Mvision AI [15] software are using AI tools, while ABAS segmentation tool from the Elekta company [16] uses an Atlas-Based Autosegmentation scheme. [9].

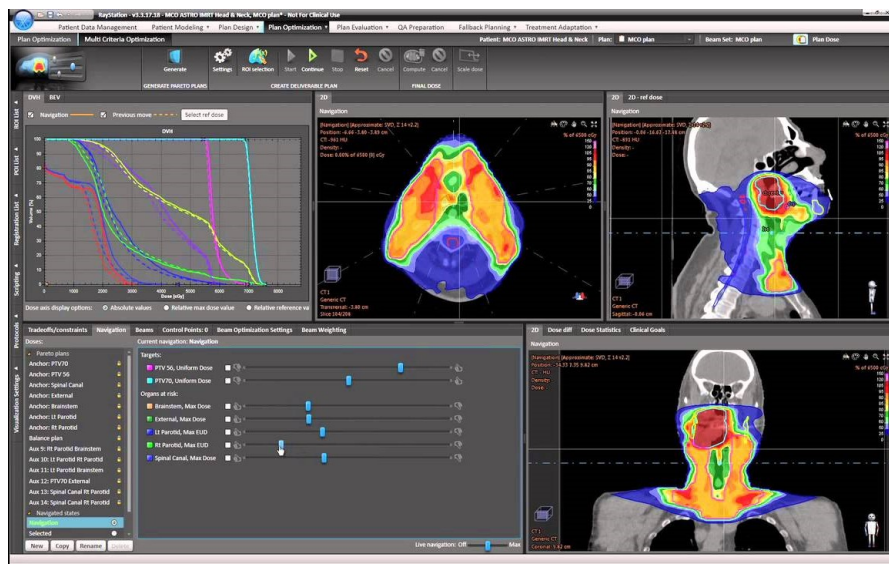


Figure 1.5: Main working window of the RayStation software [17].

1.3 Objectives of this work

This section presents the objective of this work. As a reminder, segmentation of medical images is an important part of the daily work of doctors for the diagnosis or the treatment planning of several disorders. It is especially used in the neurosurgery and radiotherapy field. Nowadays, most doctors still perform segmentation manually which takes an awful lot of time, especially for 3D images. However, a large amount of automatic segmentation tools have been developed and published for the last 15 years, but their deployment in the daily clinical practice of doctors is restrained. They are starting to arrive in radiotherapy, but for the easiest applications. The doctors seem to have a lack of confidence in these techniques when it comes to more sensitive cases such as the delineation of pathological tissue. This is mainly due to the lack of interaction between the user and these algorithms and the fact that it can produce really bad results for complex application. Although we believe that deep learning algorithms are the

future of medical segmentation, we feel that other, more interactive, techniques will help its insertion into the daily clinical practice of doctors. Active Learning is a special case of machine learning, where a learning algorithm can interactively query the user to label new data points when it faces difficulty, thus involving more the doctors. In this case, providing the user with a powerful semi-automatic segmentation tool is a good idea to accelerate this process.

Therefore, **the objective of this work is to develop and implement a user-friendly semi-automatic tool to perform segmentation on medical images that would reduce the task time taken by the doctors**. The choice of using a semi-automatic algorithm is to keep it easy to use for non-algorithm experts such as doctors. Indeed, it is known that doctors tend to dislike using nor trust automatic methods where they cannot interact with them. An emphasis is put on the intention of having, as a final result, a ready-to-use software that could be used by doctors.

Now that the main outlines have been set, a detailed list of requirements for the software developed in this thesis will be described below.

- The first requirement is for the software to be developed for medical professionals (doctors). This means that the users will be people not familiar with informatics nor algorithms. Also, that they could use this software in their everyday work. To do so, the software needs to have a well designed Graphical User Interface (GUI) that is easy to understand and to use.
- The second requirement is to implement a powerful semi-automatic algorithm using active contour that produces satisfying results such that the correction done manually by the doctor is minimal.
- The third requirement is that the algorithm needs to reduce the drawing times of doctors because segmentation is a repetitive and time-consuming task.
- The last requirement for this algorithm is to be flexible to a lot of applications/imaging modalities. Indeed, there is a lot of segmentation algorithms or software that are specific for one application only. This is not convenient at all since it would mean having a large number of segmentation methods for the different segmentation fields.

This first chapter explained the segmentation tools used in hospitals nowadays, as well as the objective of this master thesis.

Chapter 2 reviews the literature about segmentation methods. More specifically,

this chapter focuses on the two main image acquisition techniques and then, on the basic and more advanced segmentation tools.

In Chapter 3, all the methods and datasets are described. This chapter begins by describing the algorithm developed in this work through an intuitive and then detailed approach. It continues by explaining the functioning of the Graphical User Interface. Finally, the datasets and the comparison criteria are described.

Chapter 4 compiles the results about the performance and the functioning of the algorithm.

In Chapter 5, the feasibility and the utility of optimizing the choice of parameters for the algorithm are addressed.

Finally, all the results are discussed in Chapter 6.

Chapter 2

State of the art

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This chapter will describe firstly the principles of image acquisition methods used in everyday diagnosis. The second objective of this chapter is to present the state-of-the-art techniques available to perform segmentation; namely automatic methods such as deep learning and semi-automatic methods such as active contours. But before, basic principles for medical imaging segmentation will be presented.

2.1 Image Acquisition Method

Nowadays, there are a lot of different image acquisition methods such as MRI, CT, ultrasound, Positron Emission Tomography (PET), etc. Each of these methods has advantages and drawbacks and therefore are used for specific applications. For example, a PET scan is used to have a functional understanding of the human body while a CT scan helps to see the anatomy. We will consider exclusively CT and MRI imaging in the rest of this work because ~~it is~~ the two most common techniques to acquire volumetric images and there are used in a wide range of diagnosis. Furthermore, they are used to take anatomic images which are more relevant for segmentation. Indeed, interesting regions on functional imaging are already highlighted by the definition of the medical exam. The next sections will present CT and MRI imaging methods with their advantages and drawbacks [18].

2.1.1 Computerized Tomography Imaging (CT)

CT scan is an imaging modality that uses X-rays to obtain structural and functional information about the human body. The development of the CT scan technique comes mainly from two main limitations of conventional radiography. The first one is that radiography allows having a projective view of the ROI from only one angle. This limits its ability to visualize low-contrast tissue and structures. The second limitation is the poor X-ray absorption efficiency of radiographic screens (about 25%) which induces a lot of background noise [19]. These limitations induced an investigation to improve conventional radiography and it would lead to the development of CT scanners. Figure 2.1 represents the functioning of a CT scanner. The basic principle behind CT was to consider the subject as axial slices. Then the objective is to scan all of these slices and stack them to reconstruct a 3D image of the patient. To do so, a CT scan is composed of an X-rays source and a detector positioned on opposite ends of the patient. These components can rotate around the patient to take an image along one axis, called a view, at very high speeds. By mathematical reconstruction, a cross-sectional image can be created by combining all the views around a patient. Indeed, X-rays are electromagnetic waves that are absorbed differently by the different types of tissue present in the human body. Dense tissue like bones will appear white while soft tissue will appear more gray. Cavities filled with air appear black on CT scan [18–20].

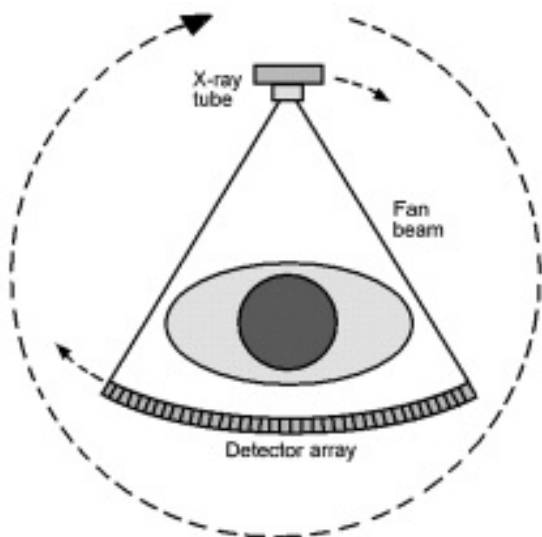


Figure 2.1: Functioning of a CT scan (of the third generation) [21]

Advantages of CT [2]

- Less expensive and wide availability
- High spatial resolution with modern multi-slice scanners
- Short scan time
- Higher sensitivity than MR for sub-arachnoids hemorrhage
- Higher sensitivity in detecting intra-cranial calcifications

Disadvantages of CT [2]

- Inferior soft tissue contrast compared to MRI as it is X-ray-based
- Radiation exposure

Main artifacts [2,22]

- **Partial volume effect:** loss of contrast between two adjacent tissue in an image caused by insufficient resolution so that more than one tissue type occupies the same voxel.
- **Streak artifacts:** occur near high-attenuation materials (bone, metal, etc.) producing dark and white streaks due to beam hardening and scatter.
- **Motion artifacts:** artifacts coming from voluntary or involuntary patient movement during image acquisition.
- **Beam hardening artifacts:** when passing through materials, only high energy photons are left, resulting in an increase mean beam energy.
- **Ring artifacts:** miscalibration or failure of one or more detector elements resulting in visible artifacts on several slices at the same position.
- **Bloom artifacts:** caused by small high density structures which result in these small objects appearing larger than their true size on the image.

CT scan is a prolific diagnostic imaging tool thanks to its cheapness and wide availability. Indeed, there are numerous indications for performing CT-based exams, including evaluation of cerebrovascular disease, intracranial hemorrhage, sinusitis, pulmonary embolism, aortic dissection, fractures, and many tumors [20]. Obviously, CT scans are also widely used in orthopaedic thanks to their great contrast to observe bone, e.g. to observe the symmetricity of the femur [23].

2.1.2 Magnetic Resonance Imaging (MRI)

MRI is the most widely used acquisition technique in the field of radio imaging. It has the big advantage to not put the human body under any radiation in contrary with the majority of the other imaging techniques. MRI benefits from the magnetic characteristic of water molecules, which compose largely the majority of human tissue, to acquire signals to reconstruct images. Figure 2.2 represents in a schematic form the functioning of an MRI scanner. Under a magnetic field, the water molecules behave like little magnets and align along the direction of the magnetic field. Then, by using RadioFrequency (RF) coils to induce some specific pulse sequence, this allows to deviate the water molecules from their initial alignment. The relaxation of the water molecules to their initial state will produce a signal that will generate information about their position in a data space called k-space. Then, a 3D volumetric image can be reconstructed by using some specific reconstruction methods. The image acquisition can be dependent on the

longitudinal (T1) or transverse (T2) relaxation time of the water. The contrast on MR images is a factor dependent on pulse sequence parameters. The most common pulse sequences are T1-weighted and T2-weighted spin-echo sequences. Figure 2.3 represents the difference between a T1 and a T2 weighted MRI. One can observe that one sequence is approximately the negative of the other one. Depending on the application, one sequence will be preferable to the other. MRI is mainly used to acquire images of the brain, but it can also be used to acquire images of the whole body [24–28].

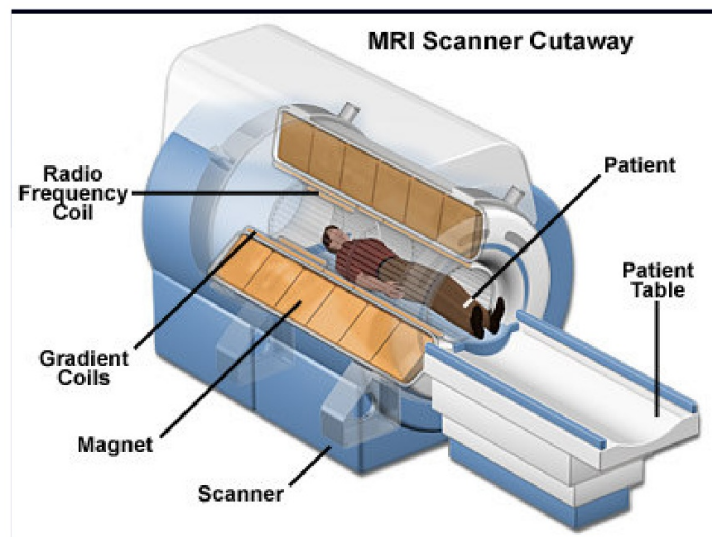


Figure 2.2: Functioning of an MRI scanner [29]

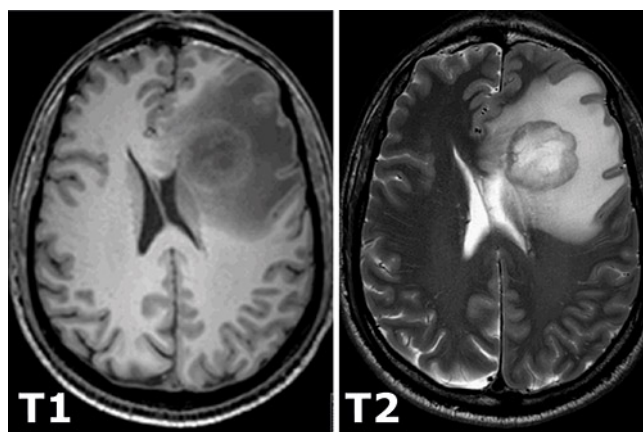


Figure 2.3: Example of a T1 vs a T2 sequence MRI [30]

Advantages of MRI [2]

- It has an excellent capability for soft tissue imaging
- It has very high resolution of the order of 1mm cubic voxels
- It has high signal to noise ratio
- Multi channel images with variable contrast can be achieved by using different pulse sequences; this can be further utilized for segmenting and classifying different structures.

Disadvantages of MRI [2]

- MR acquisition takes considerably longer time as compared to CT
- In case of MR it is more difficult to obtain uniform image quality

Main artifacts [2,31]

- **Partial volume effect:** loss of contrast between two adjacent tissue in an image caused by insufficient resolution so that more than one tissue type occupies the same voxel.
- **RF noise:** result of excessive electromagnetic emissions from other devices present in the room, causing interference.
- **Intensity Inhomogeneity:** spurious smooth varying bias field that causes pixel inconsistency within the same tissue in the MRI image.
- **Motion artifact:** artifacts coming from voluntary or involuntary patient movement during image acquisition.
- **Aliasing or Wrap around artifact:** occurs when the Field of View (FOV) is smaller than the body part being imaged. The part of the body that lies beyond the edge of the FOV is projected onto the other side of the image.
- **Gibbs Ringing:** comes from the inverse fourier transform during the mathematical reconstruction of the image resulting in series of lines in the MR image parallel to abrupt and intense changes in the object.

- **Susceptibility artifacts:** variety of MRI artifacts that share distortions or local signal change due to local magnetic field inhomogeneities from a variety of compounds.

MRI has been broadly developed and improved over these last decades to have a substantial increase in acquisition speed, quality and information extraction [32]. One major benefit in using MRI compared to other imaging techniques is that it offers superior soft-tissue contrast over other cross-sectional imaging techniques [33]. Therefore, this method is widely used for brain imaging which is essentially composed of soft tissue. More especially, MRI segmentation is commonly used for measuring and visualizing different brain structures, for delineating lesions, for analysing brain development, and for image-guided interventions and surgical planning [34]. Finally, functional MRI is another important field of this imaging technique and there has been a lot of development in the research of new pulse sequences over the last generation [35]. However, this field is not really related to the segmentation field which is done on the anatomical image, thus it will not be further explained.

2.2 Basic principles for medical imaging segmentation

2.2.1 Methods based on gray level features

This section will present the basic principles that are used to perform segmentation on medical images nowadays. Bear in mind that complex methods, such as deep learning, that will be presented after, use these basic principles. These principles are, for the most part, fairly intuitive so they will not be that much detailed.

Keep also in mind that the task of medical imaging segmentation is complex due to the difficulty of acquiring these images. Often, the quality and resolution of medical images are rather poor, even more for volume imaging, and they are constrained by partial volume effect, intensity inhomogeneity, presence of artifacts, and the close proximity in gray level of the different soft tissue.

Amplitude segmentation based on histogram features

One has to know that a histogram of gray level can be built from any medical image. This histogram will contain the distribution of the gray level of each pixel or voxel of the image. This kind of histogram can be used for image processing to enhance the contrast of the image, or for segmentation. Using histograms to classify pixels is an intuitive and the simplest idea [36, 37]. The principle is to find peaks in the histogram which will correspond to a cluster of pixels with the same gray level. Then the goal is to choose thresholds in order to separate these peaks as much as possible. This can be observed in the middle figure of Figure 2.4. This is called *thresholding*. Now, based on its position on the histogram, it remains only to assign, for every pixel of the image, the value of the peak that is found between the same threshold as itself. This technique is often used to separate an image into two clusters; the foreground and the background [37]. This process is represented in Figure 2.4. However, it can also be used with several thresholds to separate the image into several ROIs.

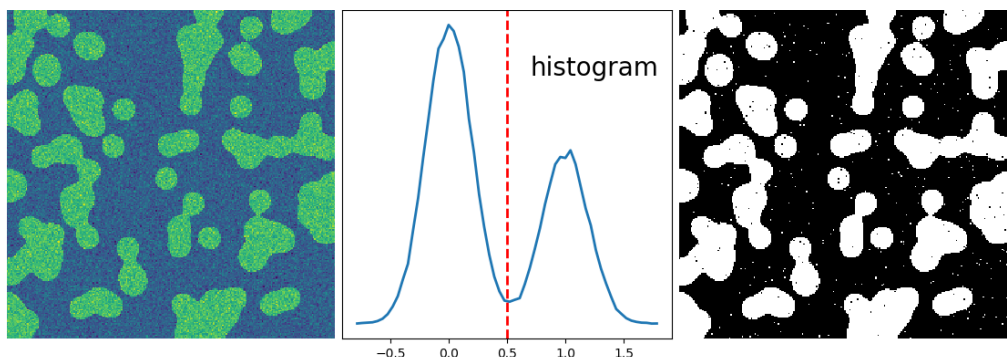


Figure 2.4: Representation of a histogram-based segmentation [38]. The left image corresponds to the original image. The middle one represents the histogram of gray level with a certain value for the threshold (red dotted line). The right figure is the final segmented image

The principle of histogram-based methods has a lot of limitations. First, if the different ROIs and background do not have sufficient different gray levels, this technique will not work. Then, the selection of proper values of thresholds is difficult and the performance is affected by the presence of artifacts or noise [2]. Users can manually choose a threshold value, or a thresholding algorithm, such as Expectation-Minimization, can compute a value automatically [36]. Finally, this technique classifies all pixels into specific regions [2].

Edge-based segmentation

In all images with several ROIs, these distinct regions are separated by edges. An edge is defined by a marking of discontinuities in gray levels in the image. Edges are considered important features of an image containing meaningful pieces of information. The principle for edge-based segmentation is to locate the edges of an image in order to use the information that they contain to separate the image into ROIs [39]. The detection of edges is often based on gradient function (first or second degree) and on the convolution of the image with some specific kernel function. Figure 2.5b) and 2.5c) represents the application of this gradient function on the original image (Figure 2.5a)) in the two principal directions. This allows highlighting the edges of the image which, after some operations that will be explained below, gives the resulting image (Figure 2.5c)). The most commonly used operator is the Sobel operator which has the advantage of providing both a differencing and smoothing effect (it is the one used in Figure 2.5). The smoothing effect is particularly attractive because it counters the fact that derivatives enhance noise [40]. The generalized algorithm for this kind of technique is the following [2]:

1. Apply the derivative operator to detect edges
2. Measure the strength of edges by measuring the amplitude of the gradient
3. Retain all edges having a magnitude superior than a certain threshold T (removal of weak edges)
4. Find the position of crack edges; the crack edge is either retained or rejected based on the confidence it receives from its predecessor and successor edges
5. Step 3 and 4 are repeated with different value for the threshold T until the closed boundaries are found

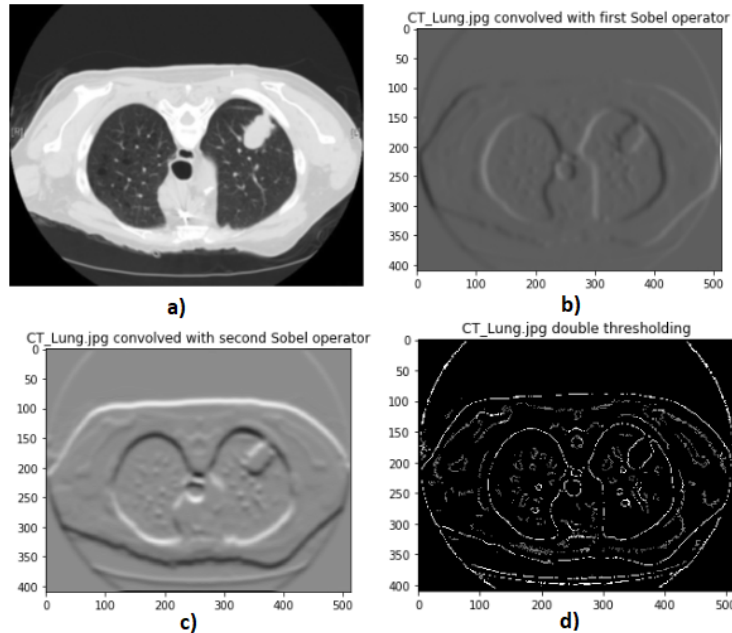


Figure 2.5: The different steps of an edge detection algorithm [41]. a) is the original image, b) and c) corresponds to the results of applying a derivative operator in the two directions, and d) is the final result.

This method can be rather good but still have some important limitations. First of all, it works well with images that present hard edges between the ROIs, which is not always the case for medical images (e.g. the soft tissue on a CT scan). Indeed, with soft edges, the derivative operator will have difficulties detecting the edges if the transition between ROIs is smooth. For the same reason, the performance of this method is greatly affected by the presence of noise in the image. Also, the presence of fake edges and weak edges in the detected image may have a negative impact on segmentation results. Finally, this method detects edges but do not separate the image into ROIs. Thus, in the end, the image is not segmented. Therefore, edge detection techniques are not complete segmentation techniques, they are required to be used with a region-based algorithm to complete the segmentation of the image [2, 42].

Region-based segmentation

The aim of region-based segmentation is to separate the image into different regions of similarities. All the regions are homogeneous if all the pixels of a certain region have similar properties. The difficulty of this method is thus to find a good homogeneity

criterion in order to perform a good segmentation. This homogeneity criterion must follow at least these two conditions [2]:

- $R_1 \cup \dots \cup R_i = I$: the union of all the regions must be equal to the entire image, thus all pixels must be part of a certain region
- $R_1 \cap \dots \cap R_i = \emptyset$: the intersection of all regions must be empty, thus two regions cannot overlap

Now that the homogeneity criterion has been introduced, it can be used with some region-based segmentation algorithms to perform the segmentation of medical images. There are three types of region-based algorithms:

Region merging This method is a semi-automatic method since help from the user is needed at the beginning of the algorithm. Indeed, in this method, some seeding points are required to initialize the process of the segmentation. The result and the performance of this method are dependent on the choice of the seeds. This allows having good interaction with the user. Regions are growing iteratively by merging the neighbouring pixels depending upon the merging criterion. If the neighbouring pixels are considered as part of the region according to the criterion, then they are added to this region. This process is continued until all pixels are assigned to their respective regions as per the merging criterion. This is a simple algorithm but that can work quite well and that is easy to use. The limitation comes from the choice of the homogeneity criterion and the difficulty to choose it [2].

Region splitting The region splitting principle is the opposite of the region merging method. Instead of merging regions, it splits the image into different regions. For more details, the algorithm begins with the whole image, then the image is split continuously until no further splitting is possible, i.e. all the regions are homogeneous upon a certain homogeneity criterion [2].

1. Begin with the whole image as one region
2. For all regions of the image, if the region is not homogeneous, split this region into 4 equal regions.
3. Repeat step 2 until all the regions of the image are considered homogeneous

This is an automatic method and the only human action relies on the choice of the homogeneity criterion which is difficult.

Split and merge method The principle behind this method is to combine the two previous region-based algorithms known as region merging and region splitting to have the best performance possible. The goal is thus to begin by splitting the image into regions and try to merge these new regions if possible according to the homogeneous criterion. Here is the algorithm for the split and merge method [2]:

1. Define a homogeneity criterion
2. Break the image into four square quadrants
3. If any resultant square is not homogeneous:
→ split it further into four quadrants
4. At each level, merge the two or more neighbouring regions if they are satisfying the condition of homogeneity
5. Continue the split and merge until no further split and merge of regions is possible

There are some limitations to this method. First, the homogeneity criterion is quite difficult to choose in order to have some good results. Secondly, there are chances of under and over segmentations in the image. However, there exist some solutions to counter these limitations. To choose the best homogeneity criterion possible, it can be optimally selected via AI technique, which can give some good results for the segmentation but it is more complicated to implement. Also, to enhance the performance of the segmentation, region-based methods can be combined with edge-based ones.

2.2.2 Methods based on textural features

Until now, mainly methods based on gray level features have been seen in order to perform segmentation of medical images. But the problem is that medical images are often a bit noisy and have artifacts that deteriorate greatly these kinds of methods because they are not robust. This is why we are now going to see methods based on textural features, which are more robust. These methods aim to subdivide the image into regions having similar pattern [43].

Let us begin by defining what is a texture. A texture is something consisting of mutually related elements. A texture may be fine coarse, smooth, or grained depending on its tone and structure. The tone of a texture is related to the pixel intensity or its gray level. The structure of a texture considers the spatial relationship between pixels. This kind of texture is called a primitive texture and it represents the simplest or

basic sub pattern [2]. Figure 2.6 represents a segmentation based on the texture where the original image contains a clear difference of textures. Further texture, often more complicated, can be defined as the spatial periodic arrangements of texture primitives or texture elements. There exist three main approaches for texture feature extraction: the statistical, syntactic or structural, and the spectral approach [2, 43].

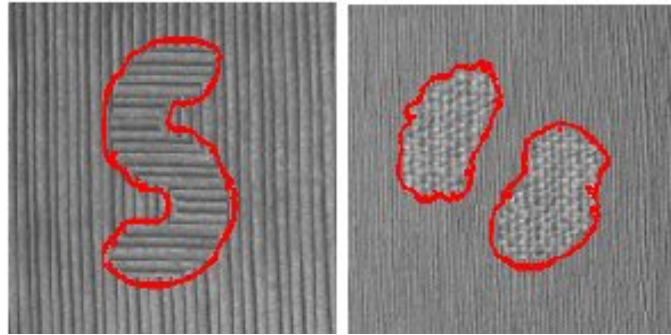


Figure 2.6: Example of an active contour segmentation algorithm based on textural feature [44]

Statistical approach In the case of the statistical approach, the texture is defined as a set of statistically extracted features represented as a vector in a multidimensional feature space. The statistical features can be based on first-order, second-order, or higher-order statistics of gray level of an image. It can use normal, robust, or even Bayesian statistics. The feature vector is assigned to their class by probabilistic or deterministic decision algorithm [2, 43].

Syntactic or structural approach In the case of the syntactic approach, the texture is defined by texture primitives which are spatially organized according to placement rules to generate a complete pattern. In syntactic feature-based pattern recognition, a formal analogy is drawn between the structural pattern and the syntax of a language [2, 43].

Spectral approach In the case of the spectral approach, textures are defined by spatial frequencies and evaluated by the autocorrelation function of a texture. In that case, the extraction of textures is by analysing the spectrum power after having transformed the image into frequency domain [2, 43].

Comparing these three approaches to define texture in the image claims the following observations; spectral frequency-based methods are less efficient while statistical

methods are very useful for random/complex patterns/textures and are very easy to use, even if syntactic or structural methods seem to give better results [2].

2.2.3 Model-based segmentation

The basic idea behind a model-based segmentation is to try to benefit from the fact that the shape, the size, the volume of a certain organ is known. From that, the structure of a certain organ can be modelled probabilistically for variation of shape and geometry. The model can be constructed based on high-resolution images and then used to extract similar anatomy from poor resolution data [45]. Figure 2.7 represents the registration of a 3D model onto a new volumetric image. This method can also be used as a constraint while segmenting other images and it involves:

1. registration of the training data
2. probabilistic representation of variation of registered data
3. statistical influence between model and image

Model-based methods of segmentation involve active shape and appearance models, deformable models and level set based models.

These techniques have two main disadvantages; They require manual interaction to place an initial model and choose appropriate parameters. Also, standard deformable models can exhibit poor convergence to concave boundaries. Finally, this method is very useful when it is required to segment organs, but is very limited for the segmentation of a pathology where the characteristics are not known [2].

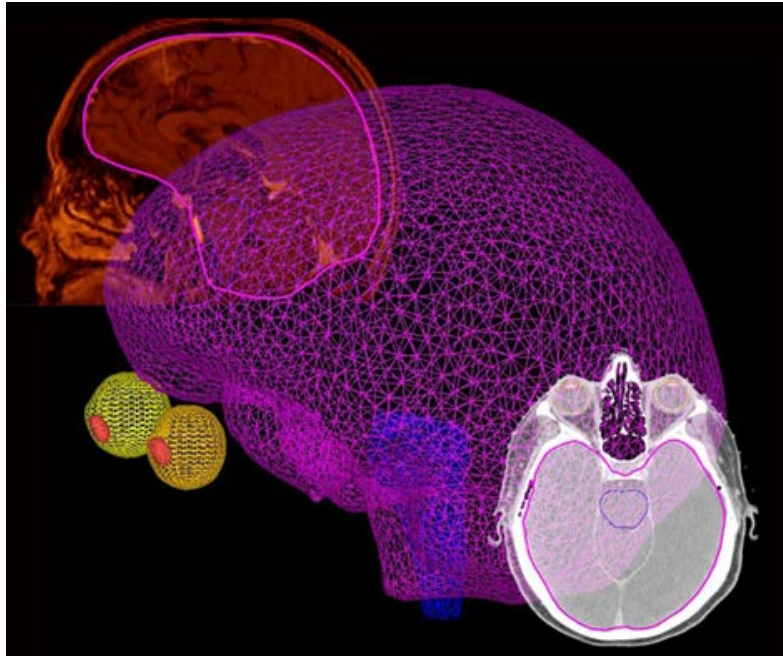


Figure 2.7: Representation of a model based segmentation where a 3D model is registered onto a new image [46].

2.2.4 Atlas-based segmentation

Atlas-based segmentation approaches are the most frequently used and powerful approaches in the field of medical image segmentation. They are among the third generation algorithms. The principle behind this method is to compile in an Atlas or a LookUp Table (LUT) (database) information on anatomy, shape, size, and features of different organs, soft tissue, etc. Then based on the knowledge stored in the Atlas the segmentation is performed on a new medical image. One positive point about the Atlas-based method is that the segmentation and classification are done in one go [2]. One can also use multi-Atlas segmentation methods which use several Atlases that map all of its labelled images onto the target image in order to reduce the segmentation errors [47]. This is represented in Figure 2.8. The performance of this method is dependent on atlas selection, atlas registration procedure, and the manual tracing protocol used in atlas formation. However, they face limitations in segmenting complex structures with variable shapes, sizes, and properties, and expert knowledge is required in building the database.

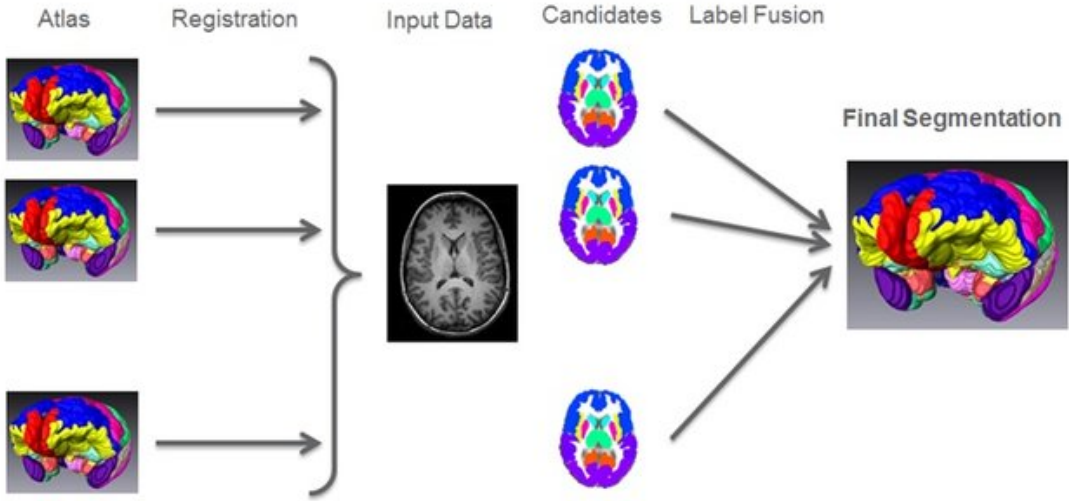


Figure 2.8: Representation of the implementation of a multi-atlases based segmentation algorithm [48].

2.3 AI tools for segmentation and classification

Automatic segmentation methods work with no interaction with a user and are based on Artificial Intelligence (AI). The AI field includes a lot of different subfields such as Machine Learning (ML) and Deep Learning (DL) which are the two main ones when it comes to medical segmentation. These AI methods can be classified as supervised and unsupervised techniques. Supervised algorithms correspond generally to algorithms using Artificial Neural Networks (ANNs) and learning methods based on some training data while unsupervised methods are based on clustering. The next sections will present these two different classes of AI methods [2, 49, 50].

2.3.1 Supervised Methods

Supervised methods correspond generally to algorithms using Artificial Neural Networks (ANNs). An ANN is a network composed of Artificial Neurons (ANs) which corresponds to simple activation units that receive several inputs and produce a simple output according to an activation function, like a biological neuron (this is represented in Figure 2.9).

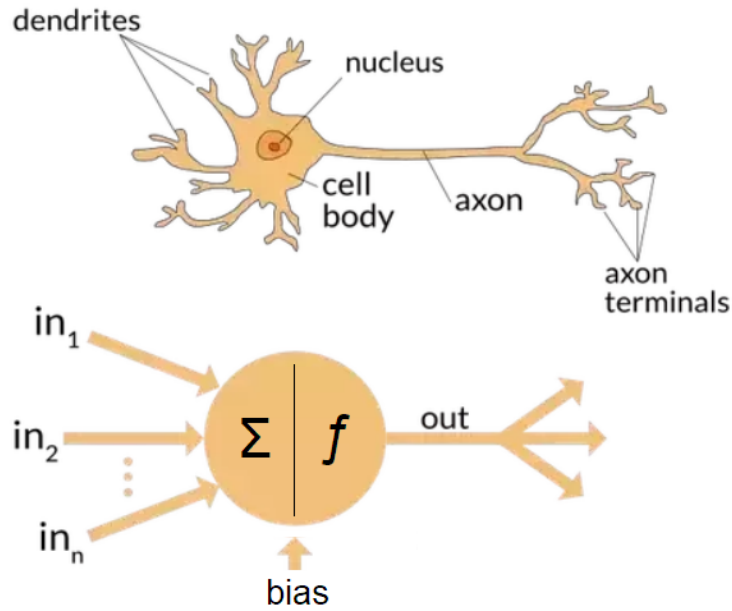


Figure 2.9: Representation of an Artificial Neuron [51].

The main advantages of ANNs are [2]:

- ability to learn adaptively, using training data to solve complex problems.
- capability of self-organization; it can create its own organization depending upon the information it receives during learning time
- capability of performance in real-time because of parallel configuration

The training part of the ANN is the most important and difficult part to create a learning machine. It needs big training datasets to learn accurately. The learning of an ANN will be done by tuning the parameters (weights of the inputs and bias) of the activation function of the ANs composing the network to produce the desired output. A network will typically be composed of layers of parallel ANs, each layer producing the inputs of the next layer (see Figure 2.10). The first one is called the input layer while the last one is called the output layer. When the network is composed of at least two hidden layers (in between the input and the output ones), then the network is said to be deep. Deep Network (DN) will use deep learning techniques. Otherwise, it is called machine learning [2, 49, 50]. For medical segmentation, DL is the most commonly used supervised AI technique. This will thus be further explained below.

Architecture of Artificial Neural Network

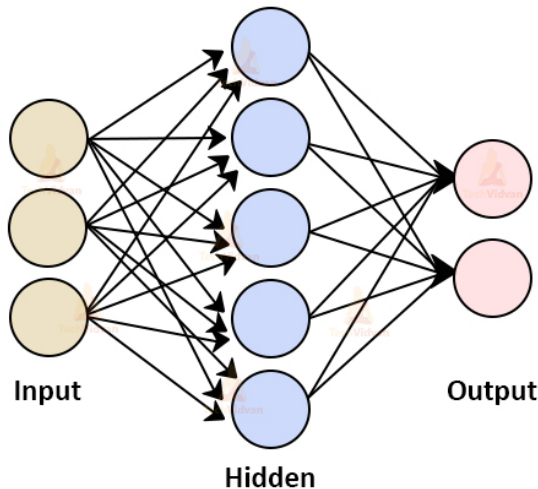


Figure 2.10: Representation of the architecture of a three layers Artificial Neural Network [52].

Deep Learning Methods

In theory, deep learning models can claim excellent performance for the task of segmenting a medical image but, in practice, calculating all the parameters for all the layers has been really difficult when the total number of layers exceeds three. Nowadays, with the advancement of technologies and computing power, ANNs can have hundreds of layers but the computation of all the parameters is still difficult and required some specific techniques and huge training datasets. The basic steps to build a Deep Learning segmentation algorithm will be presented thereafter.

Image preprocessing Image preprocessing is the process of modifying the images before feeding them to the neural networks. This phase is useful in order to accelerate the training phase and to improve the model generalizability. Here are some possible preprocess that can be done [53]:

- Medical images can have a very high resolution, as large as gigapixels. In that case, it is always good to reduce the resolution of the image to reduce the computation

power needed to analyse them. It is still necessary to be careful to not reduce the resolution too much to avoid losing too much information.

- Normalization is a typical preprocessing method that is needed for the training data as well as for the application images. Feature normalization is commonly used in the machine learning field to ensure that different features have a similar effect on the response. Feature normalization generally helps accelerate the convergence of gradient descent algorithms. There are several options to feature normalization, for example, one could rescale the pixel value into [0,1] or [-1,1]. Another option is the standardization of the data, which relate to the fact of transforming the signal from each image channel into a random variable with mean 0 and variance 1. This method is recommended when the images of a dataset are not stationary, i.e. that a certain channel from each image does not follow the same pixel value distribution.
- When using deep networks, huge training datasets are required to tune the huge amount of parameters of the network. Unfortunately, the datasets are often too small. Image data augmentation is a method to create new instances of training images based on the original dataset. This method allows to reduce overfitting, and improve the generalizability of the network.

Model selection and construction This paragraph will present the different and more commonly used network structures for deep learning of medical image segmentation. A network structure consists of a stack of connected layers, each performing a specific operation. The first layer receives the image as input and contains the same number of neurons as the number of pixels of the image so that each neuron receives the value of one pixel of the image. The outputs of each layer become the input of the next one until the last layer is reached and gives the output of the algorithm.

The first architecture that will be detailed is called a Convolutional Neural Network (CNN). Figure 2.11 represents the architecture of this network. The network is composed, in the first place, of convolutional layers and pooling layers, and then it ends up with fully observable layers. The convolutional layers consist of convolving several filters with the input data to extract some features. Each filter, or kernel, is of arbitrary size and responds only to an area of the previous layer, called the receptive field. These convolution layers result in activation maps that can be followed by an activation layer to add non-linearities [54]. They also provide local connectivity between neurons which allows the network to learn features both globally and locally [11]. They are represented by the stack of red layers in Figure 2.11. The next type is the pooling layer that is used to reduce the dimensionality of the output. They are represented by the black arrow in

Figure 2.11. Finally, some fully connected layers are used as a high-level abstraction to give the final output of the network (the purple one in Figure 2.11) [49, 53, 55].

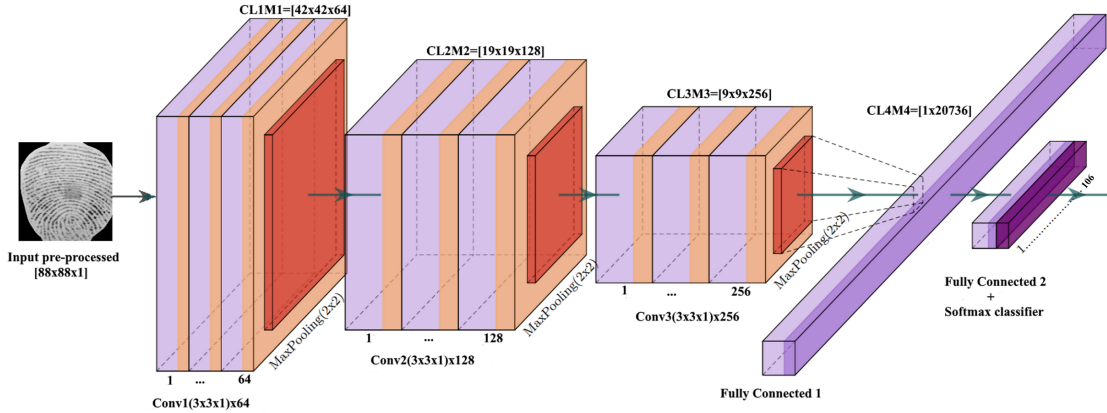


Figure 2.11: Representation of the architecture of a Convolutional Neural Network [56]

The next architecture is the Fully Convolutional Network (FCN). This network is an evolution of CNN. The basic principle behind the FCN is to use the same convolutional and pooling layers as for the CNN (this is the convolution network), but to replace the last fully connected layers by deconvolutional and upsampling layers (deconvolution network). This can be observed in Figure 2.11. This change allows the network to have an output with the same resolution as the input image and therefore pixel-wise instead of patch-wise predictions. The performance of the FCN shows a significant improvement compared the CNN architecture [11, 49, 53–55].

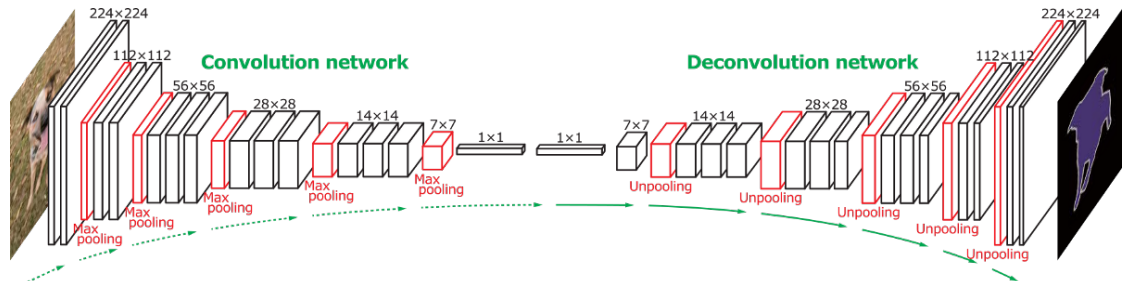


Figure 2.12: Representation of the architecture of a Fully Convolutional Network [57]

Following the same principle of the FCN network, the evolution is the well-known U-Net architecture that is commonly used for medical image segmentation. As for the FCN, U-Net is composing of two parts; the encoding (or analysis) path and the decoding

(or synthesis) path. The major improvement is the presence of shortcut connections between layers of equal resolution in the encoding and decoding phase. This is shown in Figure 2.13. These connections provide the deconvolution layers of the decoding phase with high-resolution features from the encoding phase, thus allowing a better deconvolution. A lot of networks used in medical image segmentation are based on the U-Net architecture [49, 54, 58].

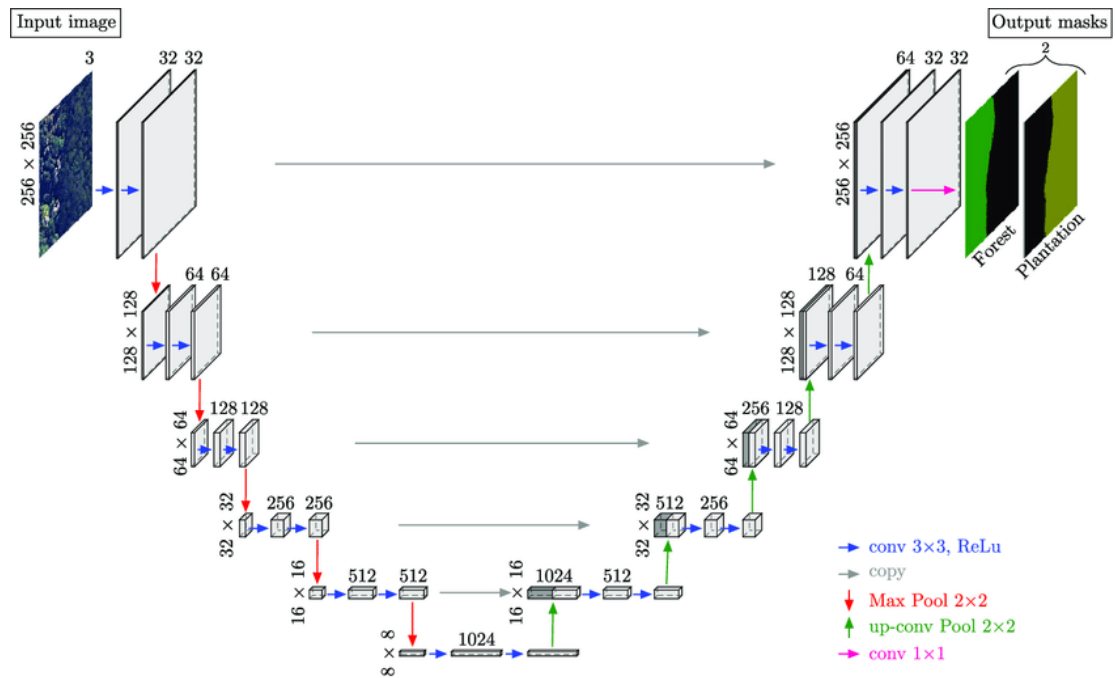


Figure 2.13: Representation of the architecture of a U-Net network [59]

Network Training Techniques This paragraph will present the challenge and the different methods that exist to train a deep network. But first, let us see how a simple network is trained in machine learning. The algorithm used is called *backpropagation* (an illustration is represented in Figure 2.14). It begins at the end of the network and consists of computing the gradient of the loss function (evaluation of the accuracy of the output) with respect to the parameters used in the network. Then, the algorithm tries to minimize the loss at this level by updating the parameters. Once it is done for one layer, it will do the same for the previous one, propagating back iteratively until the first layer is updated [49].

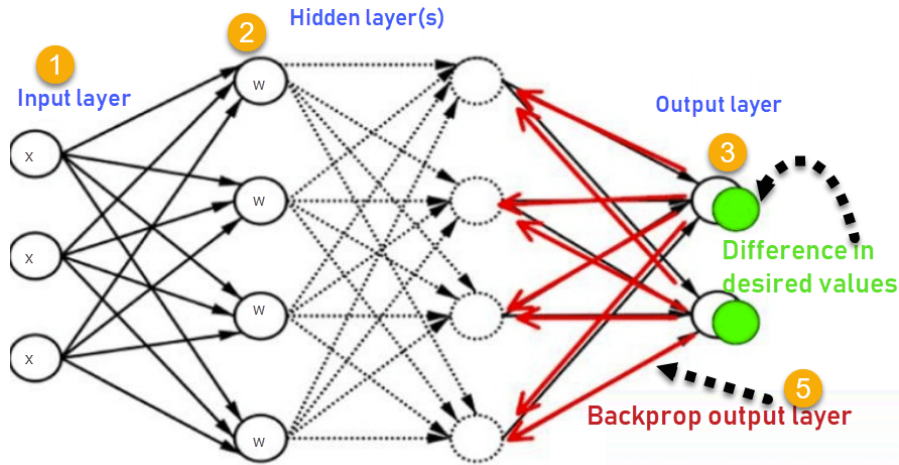


Figure 2.14: Illustration of the backpropagation algorithm [60]

The principal challenge that arises when it comes to training deep networks is the fact that the more layers are composing a network, the more training data is needed. The first training technique for deep networks is called deeply supervised and consists of providing the direct supervision of the hidden layers and propagate it to lower layers, instead of just doing it at the output layer. This method still needs huge training data. The second method is weakly supervised and it tries to train the deep network with not fully annotated images. Instead, the labelled image contains a small zone inside the ROI and a first basic segmentation method, such as region growing, is applied to fully annotate the ROI before feeding it to the deep network to train it. This technique results in less well-trained networks but is easier to implement. The last method uses the principle of transfer learning. The basic idea is to reuse the encoding part of an already trained deep network for a similar application, and to change the decoding part to obtain the results for a new application. Since only the last part needs to be trained, it can use a much smaller training dataset [49, 54].

Post-processing of the results Post-processing of the outputs of a deep network can sometimes be implemented to reduce prediction errors. The basic idea is to use empiric knowledge about the wanted result to remove false positives. This knowledge can be based on the shape, the size, or the area for example [53].

2.3.2 Unsupervised Methods

Most of the unsupervised algorithms are cluster-based and not dependent on training and training data. The basic idea of clustering is to find natural grouping of data in the multidimensional feature space. The two most commonly used algorithms for clustering are K-means and Hard or Fuzzy C-means. Figure 2.15 represents the K-means algorithm (or clustering). The principle of the K-means algorithm is to partition the data into K clusters where each point belongs to the cluster with the nearest centroid (center of a cluster which corresponds to the mean of all the data present in that cluster). Clustering algorithms have some limitations; it is sensitive to the initial partition, to the stopping criterion and it can get stuck in local minima [2, 41, 49].

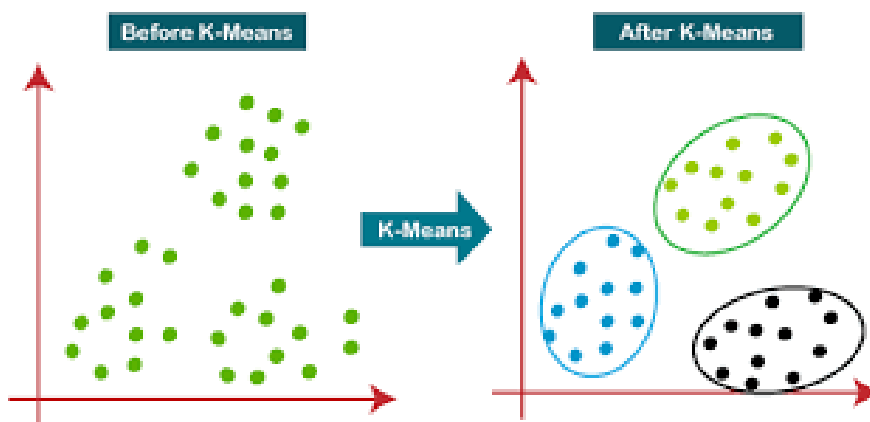


Figure 2.15: Illustration of the K-means algorithm (unsupervised learning method) [61]

2.4 3D Active Contours

The active contour model is a segmentation technique that has been widely used in the first decade of the 2000s because it provides very good frameworks of variational image segmentations [62]. Active contour is a semi-automatic technique to perform segmentation in medical images. The algorithm requires the initial contour to be set by the user and then, it will try to minimize the energy function of the contour. The segmentation depends on the definition of this energy function, so that its minimization will drive the contour towards the boundary of the wanted object [63]. This function is typically composed of a term trying to move the contour towards the objective while another term is used to keep the contour smooth. There exist two main classes of active contour that will be described just below.

2.4.1 Edge-based Active Contour

Edge-based active contour is, as its name suggests, based on the detection of edges in the image. As seen in section 2.2.1, it uses gradients in order to identify the object boundaries. The principal benefit of this method is that it does not take into account the homogeneity of the ROI. Therefore, it can achieve good segmentation even if the region is heterogeneous. The principal limitations are that it is quite sensitive to the image noise and is highly dependent on the initial curve placement. Indeed, the initial curve set by the user is required to be closed to the final boundaries of the object to segment [64, 65].

2.4.2 Region-based Active Contour

Unlike edge-based, region-based active contour uses the description of the regions in the energy function to minimize. This description can be based on several features such as the gray intensity, the texture, the structure tensors, etc. A lot of algorithms use, for the description, some general assumptions or models. Another possibility is to use the information contains in the seeds; all the pixels that are inside the initial contour set by the user. The algorithm will therefore learn the features of the object that it needs to segment, directly from the image [66]. This will be used in the implementation of the algorithm in this work. The principal advantages of region-based approaches include robustness against the initial curve placement and insensitivity to noise. Typically, region-based active contour has better performance than edge-based active contour [65]. However, the big limitation comes when segmenting heterogeneous objects. In that case, the region-based methods will have a lot of trouble producing decent segmentation [64].

Chapter 3

Method

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As a reminder, this thesis aims to investigate the performance of an active contour method to perform the segmentation of VOIs. The use of a semi-automatic method complemented by a Graphical User Interface is interesting because it allows better interaction with non-algorithm expert users such as doctors. The final goal is to see if this technique is interesting enough to be pursued and integrated to an Active Learning scheme.

In this chapter, an intuitive approach will be presented to help understanding the method that will be implemented, then the algorithm in detail and finally, the Graphical User Interface.

3.1 Intuitive approach

In this first section, an intuitive approach to the algorithm will be described to help understanding it. Fortunately, the algorithm is relatively easy to understand and follows the same idea as if a human would have to perform segmentation. The rest of the chapter will present the implementation in more detail, the different choices that have been taken as well as the subtleties and difficulties.

The question is how would a human perform segmentation on a medical image. To answer this question, let us take as an example, the task of segmenting the liver. Firstly, the need for useful empiric information relative to the anatomy of the human body about the object to segment is needed. Indeed, without any prior anatomical knowledge, one would not be able to perform any segmentation. Useful information will be the approximate position of the object, here the liver, to be able to find it on the image or, more especially, to be sure that a certain zone of the image is part of the liver. Other useful information can be the approximate size, the homogeneity of the organ, its approximate shape, etc [66]. All of this information will help to find a beginning zone in the image that corresponds to the structure that we want to segment, here the liver. This is illustrated in Figure 3.1a).

Now, based on that zone, one can learn the characteristics of the liver tissue. These characteristics are the texture, the tone, its average gray level, etc. Based on that, one will be able to encompass the surrounding tissue as liver tissue if the surrounding tissue has roughly the same characteristics as the beginning zone that corresponds to liver tissue. The accent here is put on the surrounding tissue which relates to the spatial relationship. Indeed, all organs are continuous and therefore, the spatial relationship between the pixels composing the same tissue is strong.

By following the same principle just explained above, the segmentation of the organ can be performed. For some object, it can be sufficient if the edges of the organ have enough contrast with the surrounding tissue. Unfortunately, this is not the case for all VOIs in the human body. For the example of the liver, it is known that the right kidney is another organ composed of soft tissue that has roughly the same texture as the soft tissue composing the liver. Moreover, the right kidney is in direct contact with the liver at some point. Therefore, by following the same principle as the one explained above, it is possible that the segmentation does not stop at the edge of the liver but leaks into the kidney as it is shown in Figure 3.1b). The next question is therefore how to prevent this leakage.

To prevent this leakage, one possibility would be to perform the same procedure with the right kidney and try to segment it. By doing so, the region for the right kidney

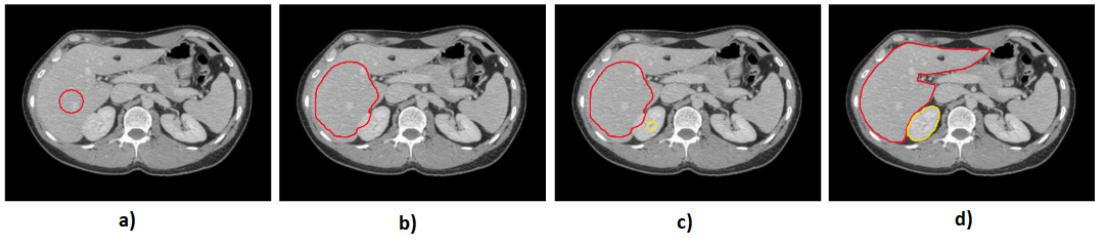


Figure 3.1: Intuitive approach to perform segmentation of the liver. a) Definition of the beginning region for the liver. b) Growing of the region with surrounding tissue until it leaks into the kidney. c) Definition of the beginning region for the kidney. d) Competition until it reaches an equilibrium

would grow until reaching the region of the liver. Then, by following a competition principle, one would take each pixel at the border of the two regions and compare them to see if it is more likely that these pixels belong to the liver or the right kidney. By doing so until an equilibrium is reached, one can hope that at the end, the segmentation of the liver would be performed correctly and that there will not be any leakage into the right kidney anymore [66]. Figure 3.1c) and 3.1d) represents respectively the beginning of the region placed into the kidney and the competition between the segmentation of the liver and the kidney until it reaches an equilibrium.

The here above paragraphs presented an intuitive approach to the problem of segmenting VOIs. The algorithm developed in this thesis is mainly based on the same approach. Here below is a summary of this intuitive approach on which the algorithm is based:

1. Definition of one or more beginning regions with a label map
2. Learning of the textures of the regions
3. Doing the segmentation iteratively:
 - Growing of each region with its surrounding tissue if it has the same texture
 - Competition between the contours if necessary
 - Repeat until equilibrium is reached

This algorithm and of its functionalities will be presented thereafter.

3.2 Definition of the label map

The method selected in this work is a semi-automatic method meaning that the beginning regions are defined by the user. The definition of these regions will be done by the use of a label map. In this work, the label map is defined as an image with the same dimensions as the image to segment and containing only transparent pixels. This allows the label map to be shown on top of the image without masking it. Then the user will be able to paint a region of a certain color on this label map thanks to the Graphical User Interface. This will be presented in the section about the Graphical User Interface (see section 3.5.2). The regions that are painted are not transparent anymore and are seen on top of the image. Once the label map is defined, the algorithm will take over and go through this label map to see where are these beginning regions to perform the segmentation.

3.3 Texture of region

As seen in the section about the intuitive approach, one key element is to be able to decide whether a pixel is part of a region or not based on the characteristics learned about this said region. For this, firstly the computation and the choice of these features, then secondly, the probabilistic model used to decide if a pixel is part of a region will be presented.

3.3.1 Features Selection

The first possibility to choose what will be used to compare if a pixel is likely to be part of a certain region, as explained in section 2.2.1, is to take into account only the gray level of the pixel. The advantage of using only the gray level is that it is very easy, no further computation is needed and it is quite a logical method to use. The problem is that methods using only the gray level of the pixels claims quite poor performance for segmentation, especially for the segmentation of soft tissue with CT scan and MRI where there is some noise and where the contrast between soft tissue is poor. Therefore, this method will not be used in this thesis.

Another way to go when not using gray level is to work with texture. As a reminder, a texture is something consisting of mutually related elements and is composed of a tone and a structure. The tone of a texture is related to the pixel intensity or its

gray level. The structure of a texture considers the spatial relationship between pixels. Segmentation methods based on texture typically claims better results than gray level based methods [2], especially for soft tissue segmentation. **Therefore, a texture-based method is a better choice.**

Before continuing, the need to compile the last sections into a summary of some definitions to better understand the following is required. Here is a recap of all definitions of what has been seen until now. First the image provided by the user to be segmented is denoted as $I : \Omega \rightarrow \mathbb{R}$ where $\Omega \subset \mathbb{R}^d$ with $d \in \{2, 3\}$, the dimension of the image. Likewise, the label image is defined as $L : \Omega \rightarrow \{[0, 0, 0, 0]\} \cup \{[a, b, c, 1]\}$ with $a, b, c \in [0, 1]$. The label map uses an RGBA¹ color model. Therefore, $[0, 0, 0, 0]$ represents the transparent background and $[a, b, c, 1]$ represents any non-transparent color. These non-transparent colors correspond to the target object labels to segment. All pixels having the same non-transparent color is considered to be part of the same object and we assume the distinct label to be consecutively ranging from 1 to N , where N is the number of distinct object to segment in the image. The labelled regions that have been colored by the user, are considered to be the initial "seeds"; which do not have to be close to the desired boundaries of the object. All the voxels that are labelled with non-transparent color are categorized into "seed groups" as $G_i = \{\mathbf{x} \in \Omega : L(\mathbf{x}) = [a_i, b_i, c_i, 1]\}$ with $i \in 1..N$ corresponding to i_{th} color found in the label image. One seed group contains all the voxels of the same non-transparent color in the label image.

The seed groups provided in the label map not only indicate the localization of the target object to be segmented, but also provide some sample voxels contained in the target object region. These sample voxels will be used to try and understand the characteristics of the tissue of the target object instead of making general assumptions on the target characteristics. As explained previously, in many cases, the gray levels of the image are not descriptive enough to have decent performance for the segmentation of soft tissue. Therefore, instead of making the segmentation in the image space, it will be performed in a feature space containing information about the texture of the image. Hence, a feature vector is extracted at each voxel, forming a feature image $\mathbf{f} : \Omega \rightarrow \mathbb{R}^{D_f}$ where D_f corresponds to the dimension of the feature vector. The feature vector contains information about the texture of the image and many choices are possible for this vector. As a reminder, there are three big classes of texture approaches; statistical, syntactic or structural, and spectral approach. Also, comparing these three approaches to define texture in the image claims the following observations; spectral frequency-based methods are less efficient while statistical methods are very useful for random/complex patterns/textures and are very easy to use, even if syntactic or

¹Red, Green, Blue, alpha

structural methods seem to give better results [2]. **In this thesis, the choice of using a statistical approach has been made for the convenience.** Moreover, a statistical method can be computed quickly compared to other approaches.

Now that the choice of a statistical approach has been made, let us focus on which statistics we will be using. Indeed, there is a lot of different statistical functions that exist and that could be used. Firstly, let us see what kind of statistics would be interesting to define a texture. As a reminder, a texture is a combination of mutually related elements composed of a tone, related to the voxel intensity and a structure, related to the spatial relationship of the voxels. The first choice to be made is the choice of the voxels on which the statistics will be computed. The simplest choice for a certain voxel \mathbf{x} is to compute the feature vector $\mathbf{f}(\mathbf{x})$ on the neighborhood $B(\mathbf{x})$ of \mathbf{x} to respect the spatial relationship needed to define a texture. The simplest neighborhood is to take all the voxels surrounding the voxel \mathbf{x} (see Figure 3.2). Therefore, $B(\mathbf{x}) \subset \Omega$ is defined as the $3 \times 3 \times 3$ cube with the voxel \mathbf{x} as the central one. For the voxels that are on the border of the image, even though the probability that they compose a target object is low, **the choice of using a mirroring solution to complete this $3 \times 3 \times 3$ cube has been made.**

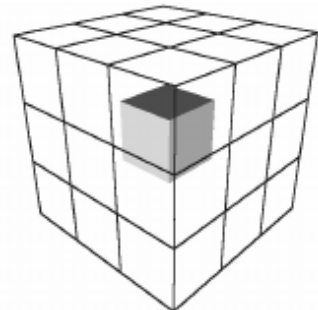


Figure 3.2: Cubic neighborhood $B(\mathbf{x})$ [67]

Now we need to decide which statistics will be used to characterize the texture. To recall, we need some description of the pixel intensity and of the pixel relationship. For example, the choice of using the average of the neighborhood for the tone and the standard deviation and/or the variance for the structure of the texture could seem to be a good idea. This is perfectly admissible to define a texture but there is one major limitation; it is not robust. Indeed, the average, standard deviation and variance are influenced by outliers. In the case of medical imaging, let us recall that the image contains generally a lot of noise and artifacts due to the acquisition techniques. Therefore, the use of these statistics would not claim the best performance. Instead, **the choice of using Robust Statistics, i.e. statistics that are less sensitive to the image noise, is preferable.** An option for the intensity level in Robust Statistics is to use the median (MED) of $B(\mathbf{x})$. Then, two measures of variations inside the neighborhood seem to be a good option to characterize the structure of the texture. To respect the robust statistics constraint, **the choice of using the Inter-Quartile Range (IQR) and Median Absolute Deviation (MAD) has been made.** Consequently, we define the feature vector $\mathbf{f}(\mathbf{x})$ as:

$$\mathbf{f}(\mathbf{x}) = (MED(\mathbf{x}), IQR(\mathbf{x}), MAD(\mathbf{x}))^T \in \mathbb{R}^3$$

with $MED(\mathbf{x}) = median(B(\mathbf{x}))$

$$IQR(\mathbf{x}) = IQR(B(\mathbf{x})) \quad (3.1)$$

$$MAD(\mathbf{x}) = median_{y \in B(\mathbf{x})}(|I(\mathbf{y}) - MED(\mathbf{x})|)$$

3.3.2 Probabilistic model

At this point, we defined a feature space that characterizes the texture of the image. Now, for the segmentation to be performed, we need a probabilistic model to help us decide whether a certain voxel \mathbf{x} , that has a certain feature vector $\mathbf{f}(\mathbf{x})$, is likely to be part of a certain region. The aim of this section is to present this probabilistic model.

The idea behind the construction of this probabilistic model is to use a Kernel Density Estimation (KDE) to build a Probability Density Function (pdf) that characterizes a seed group. In statistics, KDE is a non-parametric way to estimate the pdf of a random variable. Kernel density estimation is a fundamental data smoothing problem where inferences about the population are made, based on a finite data sample. The basic idea is to perform the convolution between the finite sample data and a certain kernel function to build the pdf. In this situation, the feature space has three dimensions, this will influence how to use the kernel density estimation. One possibility is to perform the convolution between the data in 3D and a 3D kernel function to build a pdf in 3D to represent the seed group. The other possibility is to perform a 1D KDE on each dimension of the feature space to build three pdfs, one for each feature, for each seed group and then, to joint these three pdfs afterwards. **The second solution was chosen, easier to implement and allowing a better control.**

The first choice to make for the KDE is to choose the kernel function that will be used to build the pdf. The most common kernel used to create a smooth pdf, which is the best for segmentation, is the gaussian function (this can be observed in Figure 3.3). For this reason, **a gaussian kernel**

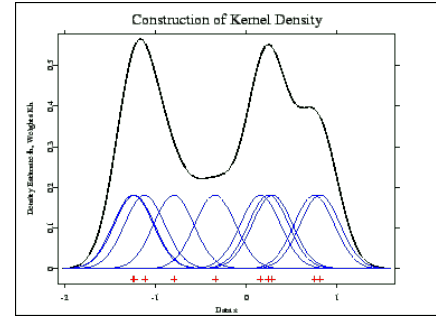


Figure 3.3: Example of a gaussian KDE [68].

function has been selected. The second important feature to choose for the kernel function for a KDE is its bandwidth, or the range for which the kernel function is non-null. For classic KDE applications, there exist methods that can calculate the best bandwidth to have the best pdf, that allow the data to be best represented. In the case of the segmentation, it would be better to have a control on this bandwidth, so that the user can vary the probabilistic model in function of the situation. More especially, greater bandwidth would allow points on the pdf that are further from the peak to still be considered in the group while a narrower bandwidth would have the opposite effect. Therefore, we can take advantage of this property to be able to better control the segmentation.

Let us now define the concept of Intensity Homogeneity (IH). This is defined by the homogeneity of the gray level of a certain region. Indeed, all the tissue in the human body has not the same homogeneity. Some tissue is smooth while others is grained. Thus, one value of bandwidth for the KDE is not optimal for the segmentation of all possible target objects. Thus the user will be able to set a parameter (IH) that will characterize the intensity homogeneity of the region that he ~~want~~ to segment. This parameter will be set between 0 and 1 where zero represents a region heterogeneous while 1 represents a region that is smooth and very homogeneous. Based on this parameter, we need to choose the bandwidth of the KDE to build the pdf for the region. To do that, not only this parameter will be used but also a value of the variation of the gray level intensity of the seed voxels of this region. We decide to use the median absolute deviation feature for that. More specifically, since we want a value of the variation of the gray level intensity of all the region, we will take the median of the third feature of all seed voxels. The choice of the median seemed obvious to keep robust statistics. To compute this bandwidth, a new parameter η will be introduced. This parameter has no signification and it is only used to transform the intensity homogeneity parameter from [0,1] to [0,20] [66]. The bandwidth of the KDE will be computed by dividing the value of the variation of gray levels intensities of a region by η . Here are the following formulas:

$$\eta = IH \times (20 - 0.1) + 0.1 \quad (3.2)$$

$$bandwidth = \frac{MED(MAD(\mathbf{x}_i) \text{ with } \mathbf{x}_i \in G_i)}{\eta} \quad (3.3)$$

Now that the pdf has been computed, there is still one limitation; namely, a pdf integrates to one on all its domain. In this case, the need is to have the probability of a certain value to be part of this region. Therefore, we chose to rescale the pdf to 1

by dividing the pdf by its maximum value. Indeed, it is mostly the shape of the pdf that is interesting. Figure 3.4 represents the evaluated pdf for the first feature of a seed group for different values of the intensity homogeneity parameter. It shows that this parameter influences the bandwidth of the kernel function and therefore the range of voxel values that will be considered as likely to be part of the region.

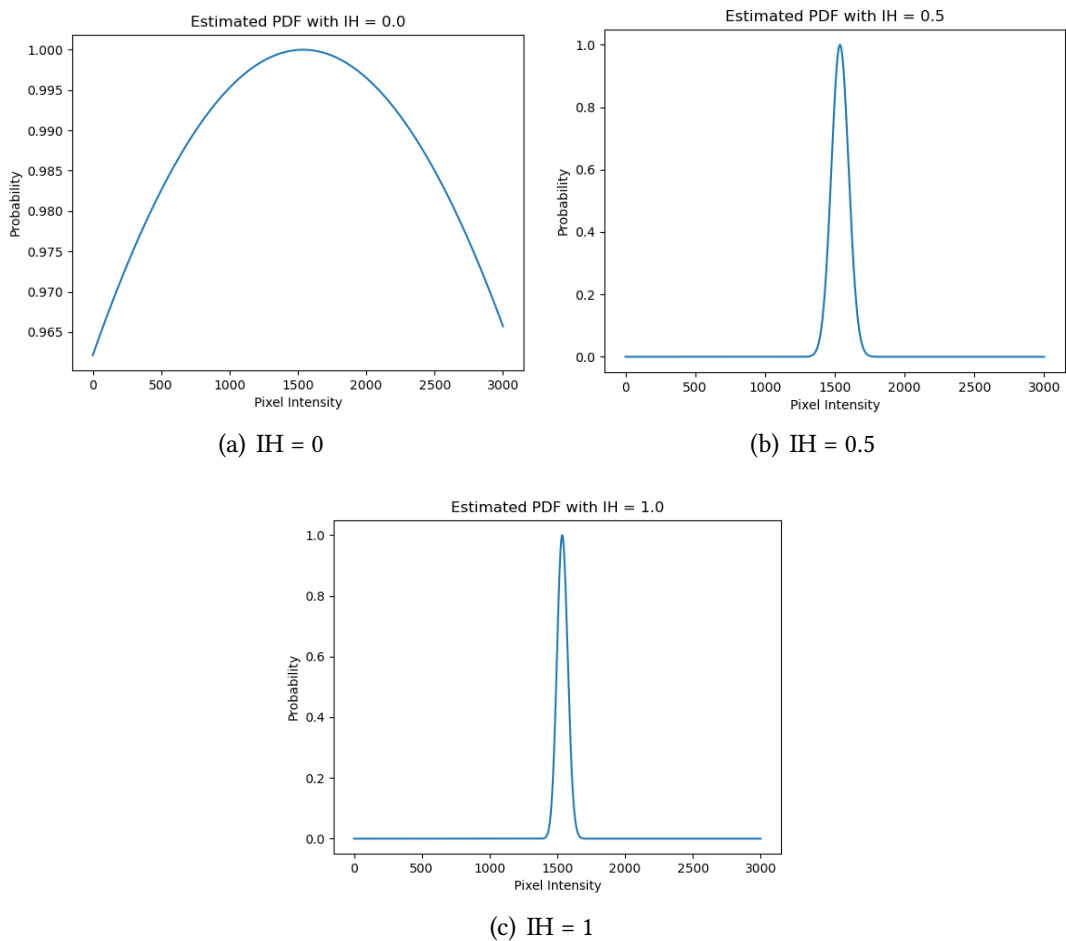


Figure 3.4: Comparison of evaluated PDF for different values for the intensity homogeneity parameter

At this point, the pdfs for the three features of the feature space have been built and in this section, the utility of these pdfs in the algorithm will be presented. The goal is to use them to see if it is likely that a voxel \mathbf{x} is part of a region i . To do so, first the feature vector $\mathbf{f}(\mathbf{x})$ of the voxel must be computed. Then with the three values, the likeliness of that voxel to be part of the region i can be derived thanks to the three pdfs;

$pdf_{i_{med}}$, $pdf_{i_{iqr}}$, $pdf_{i_{mad}}$. From that, the probability $p_{i_{med}}(\mathbf{x})$, $p_{i_{iqr}}(\mathbf{x})$, $p_{i_{mad}}(\mathbf{x})$ following the three features can be computed. Then, the last step is to join the three probabilities together by following this formula:

$$p_i(\mathbf{x}) = p_{i_{med}}(\mathbf{x}) \cdot p_{i_{iqr}}(\mathbf{x}) \cdot p_{i_{mad}}(\mathbf{x}) \quad (3.4)$$

Now, a cut-off probability needs to be chosen to decide whether the voxel is part of a region or not. This will be chosen as a setting parameter for the user so that he can have a little more control over its segmentation. By default, the cut-off probability has been set-up to 0.1, value having demonstrated pretty good results in a majority of cases.

3.4 Active Contours

This section will highlight the active contour part of the algorithm described. First of all, there is a distinction between an image in 2D and 3D. In 2D, an active contour is used while an active surface is used in 3D. Although these two have a lot of similarities, there are still some major differences that need to be pointed out. First, the contour and then the surface will be presented.

Firstly, the contour in this algorithm has been implemented thanks to the use of a double-linked list that contains all the pixels that encompass the region. A double-linked list has the advantage of representing the spatial relation between the different pixels of the contour. This is very useful to compute the mean curvature at a certain position of the contour. Each of the nodes of the contour represents a pixel and contains information about the force that is applied by the algorithm at this position of the contour. Besides, each contour contains information about the region such as the color of the initial seeds and the associated region. The computation of the force at each point gives us information about whether or not the contour needs to expand, but it does not give any information about the direction of the expansion. Initially, the idea was to work with vectors: the computation of the force would give the amplitude of the vector and, the orientation would be given by the perpendicular to the tangent to the curve at this point. This would have allowed leading the expansion of the contour only in a certain direction. Unfortunately, this was not that easy to implement. Indeed, by computing the perpendicular to the tangent to the curve, the orientation of the vector can be found. But, it still misses information about the direction. In one direction, it would lead to the interior of the contour and in the other, this would lead to the contour expanding outside of the region. To counter that problem, the idea was to compute the orientation and then test if it would lead to inside the region or not. If it would lead inside the region, the other direction would be considered. Unfortunately, this method

led to some limit cases where it would not work. Therefore, the idea of using vectors has been aborted. Instead, **the growing of the region is made by expanding the contour with all the pixels that are not already inside the region**. This has been working very well.

Secondly, for the surface, the first idea was to use the same principle as for the contour but just expanded in 3D. Therefore, from a double-linked list for the contour, the aim was to use a graph where each node composing the surface would have a list of all its adjacent nodes in the surface. This would allow keeping the spatial relationship between the voxels. The issue with this method is that using graphs requires the use of graph algorithms to search it and these algorithms are slow. Thus, the use of a graph to represent the surface has been abandoned. Instead, **the spatial relationship between the voxels has not been kept and has been replaced with a simple list of voxels composing the surface**. Each surface is composed of two lists: one list containing the static nodes and one containing the movable nodes. Static nodes designate the ones where the force on them has already been computed, but does not allow the surface to move at this point. On the other hand, the movable voxels are the ones where the force has not been yet computed. This is again a trick to try and optimize the algorithm to increase its speed.

3.4.1 Contour evolution

In this section, the contour evolution scheme will be presented; according to which formula the contour evolution is based on and how it is implemented. First of all, it is based mainly on the definition of the probability that a voxel is part of a region defined earlier. The goal here is to try to maximize the area that is enclosed by a contour. This is achieved by the following variational approach. This approach is valid for both 2D and 3D. First, let us define the family of evolving contours $C_i \subset \Omega$ corresponding to all the contours that encompasses a certain ROI. Without any contour interaction, each of these contours will evolve independently by trying to minimize the following energy function [66]:

$$E_i(C_i) := (1 - \lambda) \int_{\mathbf{x} \in C_i} (p^c - p_i(\mathbf{f}(\mathbf{x}))) d\mathbf{x} + \lambda \int_{C_i} dA, \quad (3.5)$$

where p^c corresponds to the cut-off probability and $\lambda > 0$ is a parameter that weights the importance between the two terms. Moreover, the first term corresponds to a region-growing term that pushes the region to grow while the second term tries to keep the contour from expanding too much. To minimize this formula, we will compute

the first variation of this one to obtain the flow of C_i [66]:

$$\frac{\partial C_i(\mathbf{q}, t)}{\partial t} = [(1 - \lambda)(p^c - p_i(\mathbf{f}(C_i(\mathbf{q}, t)))) + \lambda\kappa_i(\mathbf{q}, t)] \mathbf{N}_i(\mathbf{q}, t), \quad (3.6)$$

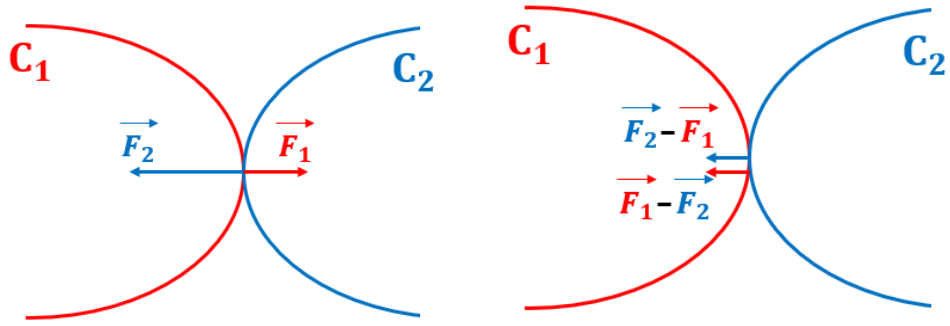
in which \mathbf{q} is the spatial parametrization of the contour, \mathbf{N}_i is the inward unit normal vector field on C_i , and κ_i is the mean curvature of the contour. To explain further, the contour will grow if its flow is negative. In the two terms, the second term is always positive and thus keeps the contour from growing. The first term, concerning the probability of the voxel to be part of a region, can be positive or negative due to the probability cut-off. As explained in a previous section, this probability cut-off is set to 0.1 by default. Therefore, if the probability of a pixel is inferior to p^c , the first term $p^c - p_i(\mathbf{f}(\mathbf{x}))$ will be positive and the contour will not grow. Otherwise, the first term will be negative and the region will grow if the second term is less than the absolute value of the first term [66].

The second term concerns the mean curvature of the contour. The definition of curvature is the inverse of the radius of a circle that follows the contour curve at one point. The mean curvature is thus the mean of the curvature computed at all the points of the contour. Here, it would take an absolutely huge amount of time to compute this curvature at each point and to compute the mean afterwards. The goal is thus to estimate this mean curvature in another way. To do so, we will compute r_p , the radius of the circle based on the length of the surface. We will also compute S_p , the surface of this perfect circle. With that, we can compute c , the ratio $\frac{S_p}{S_a}$ between the perfect surface and the actual surface. This coefficient will be useful to estimate the mean radius of the actual surface thanks to this formula; $r_e = c\sqrt{\frac{S_a}{4\pi}}$. Then, we just have to compute the estimated mean curvature based on this estimated radius;

$$\kappa_i = \frac{1}{r_{ie}} \quad (3.7)$$

3.4.2 Contour interaction

Considering that the algorithm can be used for multiple object segmentation, contour interaction must be implemented. Indeed, two contours may enter in contact and a decision strategy must be implemented to handle these kind of case. Another possibility is to use contour interaction to avoid leakage in the segmentation of a certain VOI, even if the second contour is not use to segment another VOI. In both cases, the strategy will be the same.



(a) Contact between two contours

(b) Result of the competition between the two contours

Figure 3.5: Contour interaction

As explained in the intuitive approach of this algorithm, the strategy that will be used is the competition between the two contours. In more details, when two contours enter in contact, the voxels that are on the edge of the two contours need to be chosen to join one or the other region. For that, the principle used is simple. The computation of the force of one voxel that stand in two contours will be equal to the computed force for one region minus the computed force for the other region. The resultant force will decide whether the voxel should join one region or the other. This will continue until the contours reach an equilibrium. This principle is illustrated in Figure 3.5.

3.5 End user oriented

As a reminder, the objective of this work was to design a software that could be used by doctors and to compare it with currently used software. Therefore, a really important part of this work is the end user orientation. Thus, in this section, the Graphical User Interface (GUI) that we have designed will be presented so that people can easily use this algorithm for any purposes. This section will be divided into different parts, each presenting one tab of the GUI. This GUI has been implemented thanks to the *Tkinter* python library.

3.5.1 Viewer tab

The first tab is called the viewer tab and it is used to load a 3D scan, to view it, and to perform some basic operation on this scan. A snapshot of this tab of the GUI is available in Figure 3.6. Here is a list of the buttons available in this tab with a small description of their action for each of them:

- XY: allows to observe a scan along its XY plan
- XZ: allows to observe a scan along its XZ plan
- YZ: allows to observe a scan along its YZ plan
- Slider: allows to go through the scan along the plan selected
- 90° rotation: allows to rotate the scan of 90° in the plan currently selected
- Reduce Resolution: allows to reduce the resolution of the scan to a maximum of 300 voxel, while keeping the same proportion for the scan

The last part is the rolling menu *File* at the top left of the GUI that contains three actions;

- the *Load* action opens a file browser so that you can choose a file to open
- the *Import Mask* opens a file browser to import a mask for the image (if one already exists) and to observe it in the viewer sub-tab in the segmentation tab (see section 3.5.3)
- the *Save* action allows you to save in a file the result of the segmentation when it is done

A final characteristic is that the XY, XZ, YZ buttons, and the slider are present in all the tab containing a viewer and they are always doing the same actions. Therefore, they will not be repeated in the other tabs.

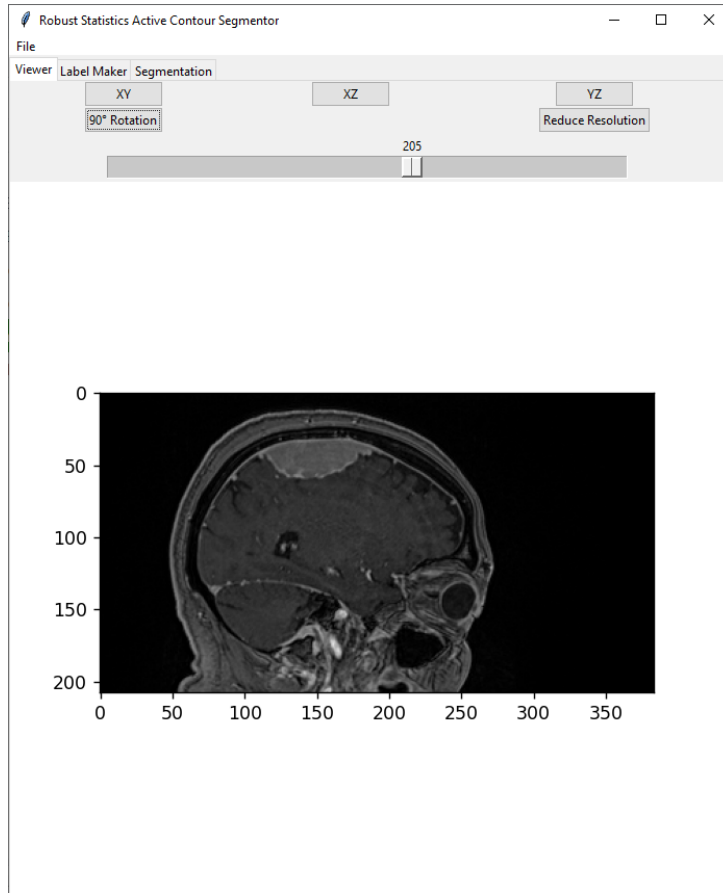


Figure 3.6: Viewer tab of the GUI

3.5.2 Label maker tab

Now, let us present the label maker tab. This tab is represented in Figure 3.7. It is used to draw on the loaded scan to create the label map that will be used by the algorithm for performing the segmentation. In addition to the buttons and sliders that are useful to visualize the scan, there are 5 buttons of color and an eraser that are used to be able to draw onto the 3D scan. Each of these 6 buttons, when pressed, select and keeps in memory the color that you want to draw on the label map. When a color is selected, you can, with your mouse, draw a closed contour and all the voxels included inside this contour will be colored in the selected color. More especially, the coloring is not done by changing the voxel in the 3D scan. A second image, at the beginning transparent, is created to save all the colored drawings which will be used as the label map to the algorithm.. There are two options to use the eraser; either you do a *right*

click on the button and this erases all the label map, or you select it by *left clicking* on the button and by doing the same procedure as for coloring, you can erase part of the label map.

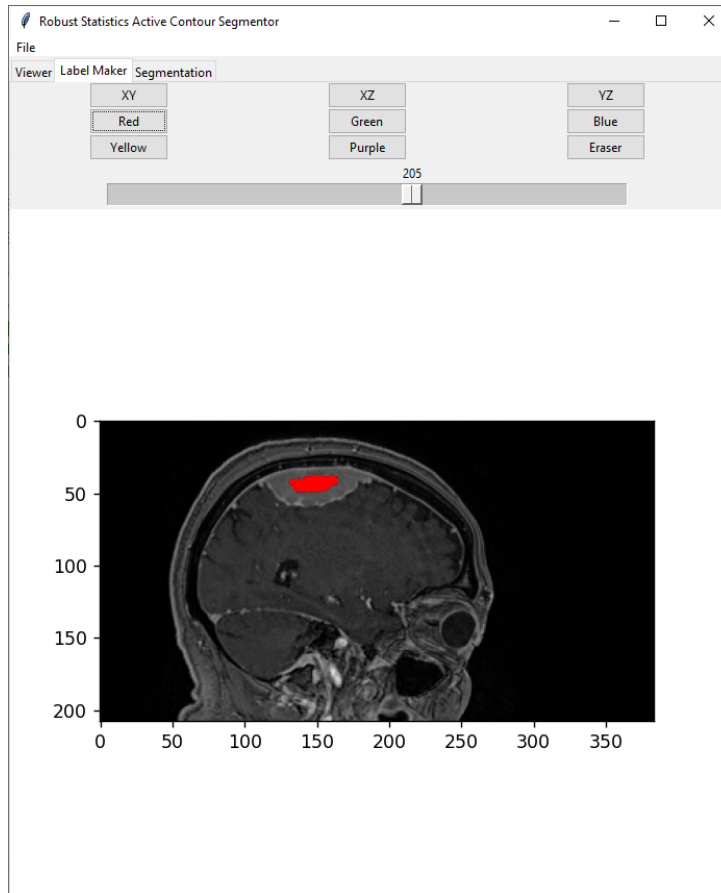


Figure 3.7: Label Maker tab of the GUI

3.5.3 Segmentation tab

The last tab to be presented is the segmentation tab. It is the tab where the segmentation is done, i.e. where the parameters are chosen, where the segmentation is launched, perform and then adjusted if necessary. This tab is separated into two sub-tabs that are each used for a single purpose. These two sub-tabs are presented just below.

Parameters

Figure 3.8 is a screenshot of the parameters sub-tab of the Segmentation tab of the GUI. The purpose of this first sub-tab is to set the parameters that will be used to perform the segmentation. Here is a list of all these parameters with an explanation of their functionality:

- Dimension: this parameter corresponds to the dimension of the image to be segmented; either 2 or 3 (with 3 as the default value)
- Intensity Homogeneity: this parameter is used to define the homogeneity of the intensity of gray level of the VOI. This parameter will be used to choose the bandwidth for the kernel function when computing the pdf function of the texture of the VOI. A value of 0 corresponds to a region with a high variation of gray level while a value of 1 corresponds to a region very homogeneous. The default value is set to 0.5.
- Curvature Weight: this parameter corresponds to the parameter λ in the formula to compute the flow of the contour. More especially, it weights the importance of the smoothness of the curvature compared to the region growing part.
- Max Iteration: this parameter corresponds to the maximum number of iterations that the algorithm will be able to use to perform its segmentation. This can be used to avoid leakage. The default value is set to 1000 but in practice, it can be way less, depending on if you want to use this parameter as a limiting factor.
- Max Running Time: this parameter corresponds to the maximum amount of time allowed to perform the segmentation. The unit is the second and the default value is put to 300.
- Max Volume: this parameter corresponds to the maximum volume that region can grow in. This parameter corresponds to a number of voxels and it is really useful to avoid leakage if the boundary of the VOI is not sharp enough. The default value is set to 25 000.
- Probability Cut-off: this parameter corresponds to the p^c in the formula of the flow of the contour. It is the cut-off value that decides whether a certain pixel is likely to be part of a region or not. The default value is set to 0.1 and this value works well in the majority of the cases. However, we let the possibility to the user to change this value if wanted.

- Inertia: this parameter corresponds to the inertia of the curve, i.e. the importance of the previous force applied on the contour at this point. This can be useful to overcome certain voxels that appear different due to noise but that are still in the region.

Finally, there are still two major buttons yet to be presented, and their actions are important to be understood to be able to fully master the software and exploit it at its full potential. The first button is the *segmentation* button and allows the software to create a new Segmentor object and perform the segmentation with the image, the label image created and the parameters chosen above. One has to know if he wants to segment multiple objects, that each object will use the same parameters, which is not ideal. To counter that among others, the second button *continue segmentation* has been created. This second button allows continuing a segmentation by using the same Segmentor object created with the *segmentation* button. This allows a lot of flexibility as to how to manage a segmentation. For multiple object segmentation, this allows continuing the segmentation for each individual object with a different set of parameters. It can also allow to segment VOI with different texture regions by segmenting one part of the VOI, then segmenting the second part by continuing the segmentation and thus building the segmentation iteratively.

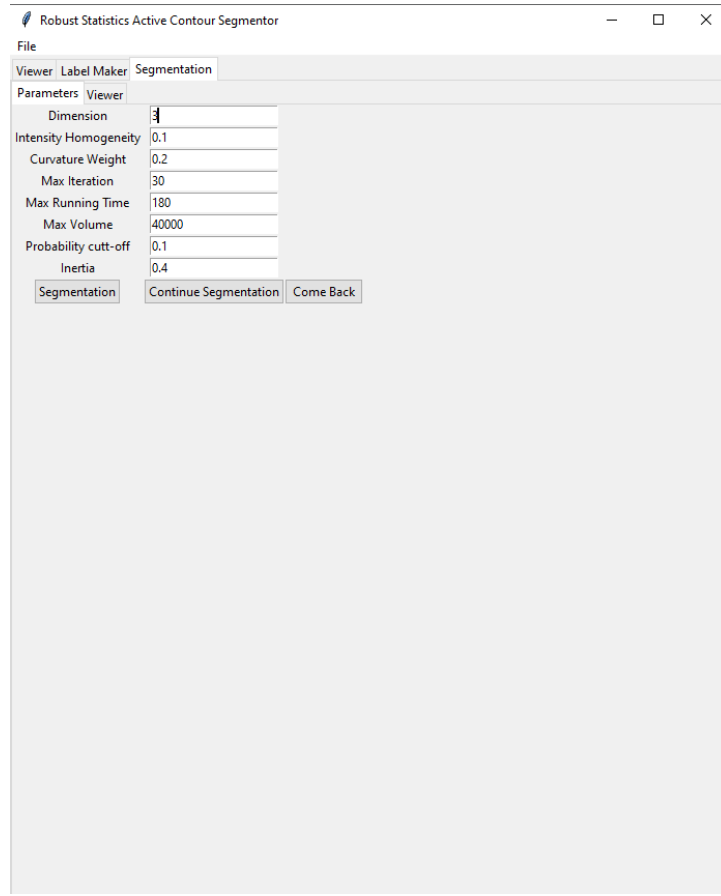


Figure 3.8: Parameters sub-tab of the Segmentation tab of the GUI

Viewer

The viewer sub-tab is exactly the same as the Label Maker Tab (view Figure 3.9). This tab is used by the user to visualize the result of the segmentation algorithm and to modify it if necessary by drawing or erasing parts with the same action as in the Label Maker Tab. Finally, the user can save the result by clicking on the *Save* button in the rolling menu File.

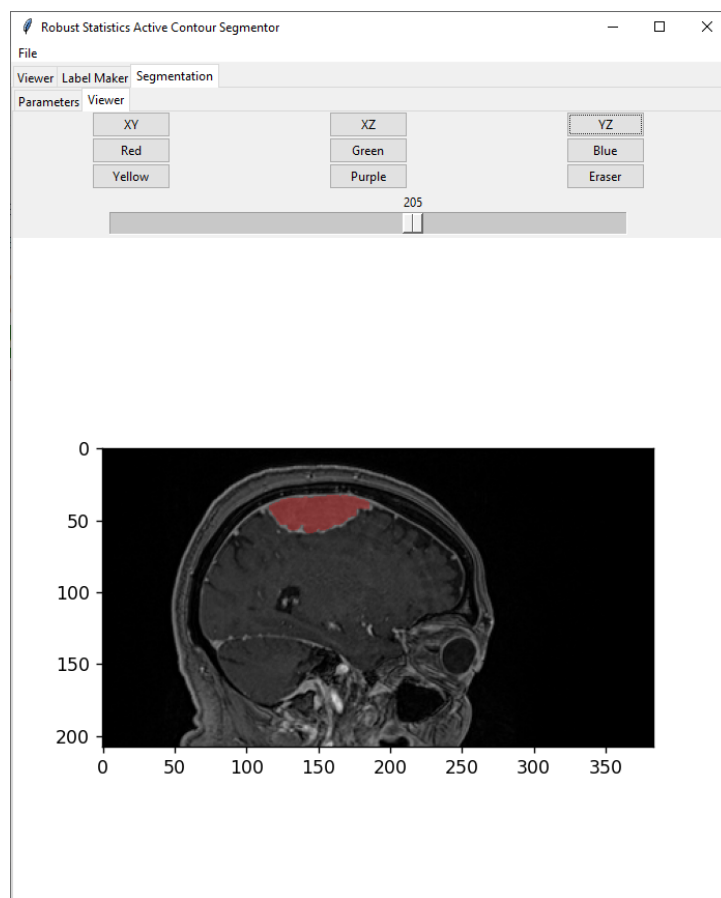


Figure 3.9: Viewer sub-tab of the Segmentation tab of the GUI

3.6 Implementation

Figure 3.10 represents the implementation of the GUI and the algorithm developed in this master thesis.

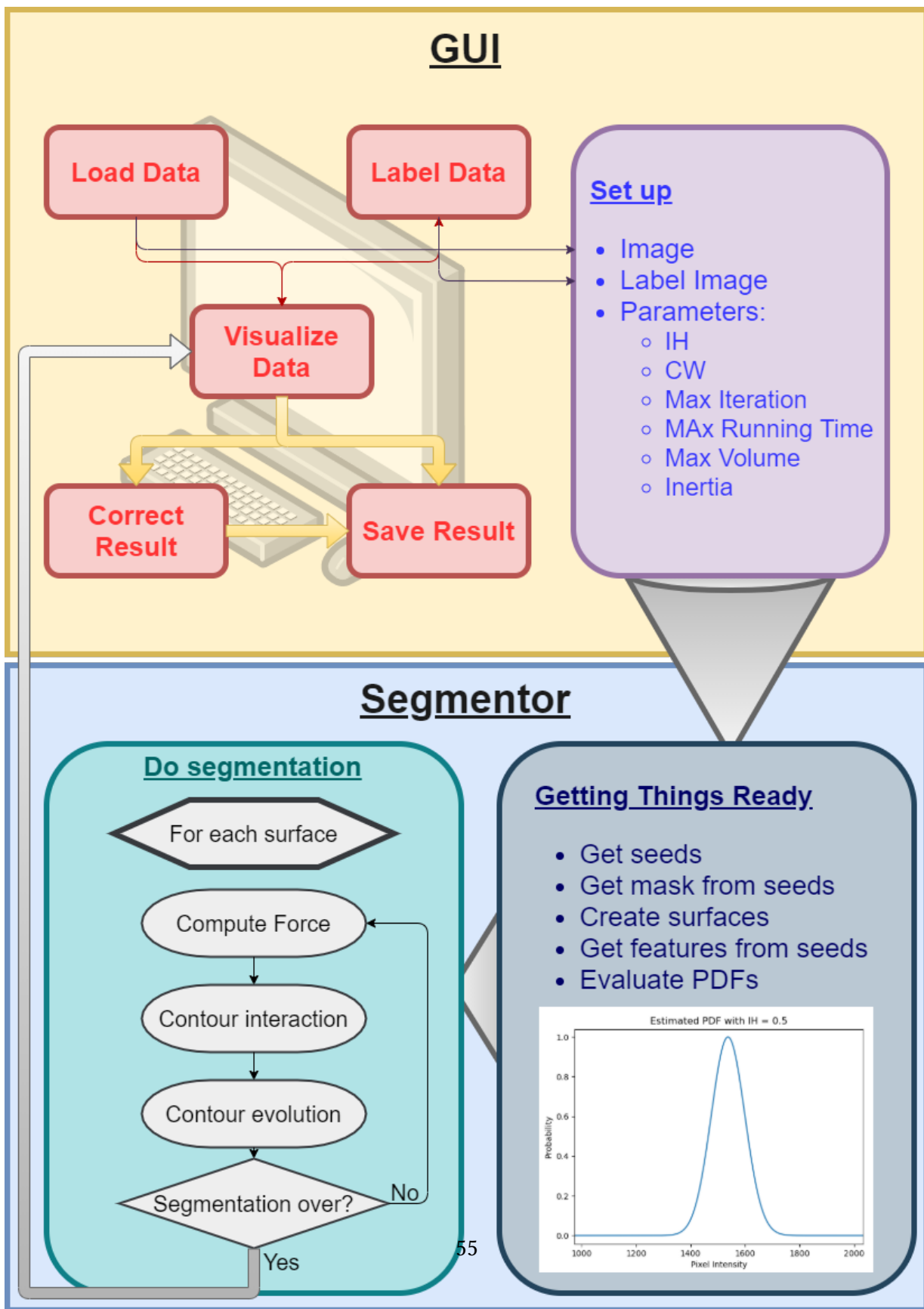


Figure 3.10: Implementation of the algorithm

3.7 Materials used

In this section, the material, i.e. the different 3D images, used in this Master Thesis will be presented. There are related to two different applications; one using 3D CT scans of the pelvis region and the other using different MRI sequences of the brain.

3.7.1 CT dataset

The first dataset corresponds to 3D CT scans of the pelvis region. The objective is to segment the femur, and more specifically the femoral head. Therefore, the use of CT scans is perfect for this kind of application since it allows a really good contrast of the hard tissue like the bone tissue. Figure 3.11 is an example of a CT scan that will be used for the bone application of this work.

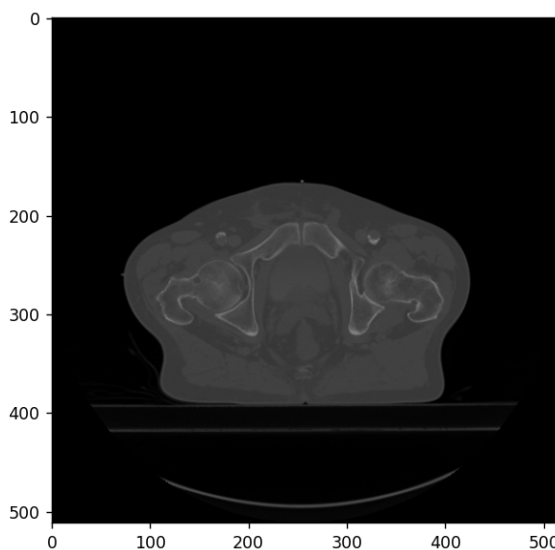


Figure 3.11: CT scan of the hip and femoral joint

3.7.2 MRI dataset

The second dataset corresponds to 3D MRI scans of the brain using different types of sequences. These MRIs are taken from patients that have some kind of brain pathology. MRI is a good choice for brain imaging since it allows to have good contrast between the different soft tissue composing the brain. Figure 3.12 is an MRI of a patient suffering

from lymphoma. This is an example of an ~~MRI that~~ will be used for the application of pathological MRI. This MRI uses a T1 weighted sequence with Gadolinium as a contrast agent.

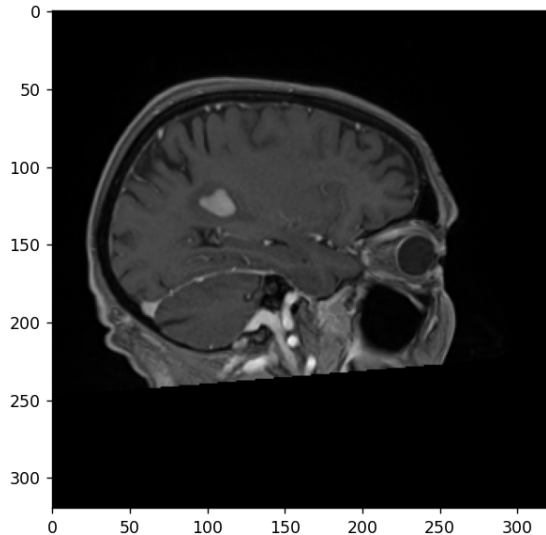


Figure 3.12: MRI of a patient suffering from a lymphoma

3.8 Criteria of comparison

The aim of this section is to describe the different criteria that will be used to compare the performance of a segmentation compared to another one. In this work, we will take into account 3 criteria of comparison; the Dice coefficient, the Jaccard index, and the Hausdorff distance. The two first criteria compare the overlapping of two segmentations in a slightly different way but they are still similar. On the contrary, the Hausdorff distance is based on the distance between the two segmentations to compare them.

3.8.1 Dice coefficient and Jaccard index

The Dice coefficient is a statistic used to gauge the similarity of two samples. In this context, it will be used to compare two segmentations. This coefficient is computed by taking the ratio between two times the overlapping of the two regions and the sum of

these two regions. Therefore, for a region A and a region B:

$$D(A, B) = \frac{2 \cdot |A \cap B|}{|A| + |B|} \quad (3.8)$$

The Jaccard index measures similarity between finite sample sets, and is defined as the size of the intersection divided by the size of the union of the sample sets. Therefore, for a region A and a region B:

$$J(A, B) = \frac{|A \cap B|}{|A \cup B|} = \frac{|A \cap B|}{|A| + |B| - |A \cap B|} \quad (3.9)$$

The Jaccard index is very similar to the Dice coefficient. For both, the result will be a coefficient ranging from [0,1] with 0 corresponding to 0 similarities while 1 characterize perfectly similar samples. The similarities between these two coefficients do not stop here. Another interesting feature is that these two coefficient are positively correlated. That is to say that if classifier A is better than B under one metric, it is also better than classifier B under the other metric. It is tempting then to conclude that the two metrics are functionally equivalent so the choice between them is arbitrary. However, the difference emerges when quantifying how much worse classifier B is than A for any given case. In general, the Jaccard metric tends to penalize single instances of bad result more than the Dice score quantitatively even when they can both agree that this one instance is bad. Therefore, keeping the two metrics can be interesting.

3.8.2 Hausdorff distance

In mathematics, the Hausdorff distance measures how far two subsets of a metric space are from each other. Informally, two sets are close in the Hausdorff distance if every point of either set is close to some point of the other set. The Hausdorff distance is the longest distance you can be forced to travel by an adversary who chooses a point in one of the two sets, from where you then must travel to the other set. In other words, it is the greatest of all the distances from a point in one set to the closest point in the other set. Let X and Y be two non-empty subsets of a metric space (M, d) . We define their Hausdorff distance $d_H(X, Y)$ by

$$d_H(X, Y) = \max\{\sup_{x \in X} d(x, Y), \sup_{y \in Y} d(X, y)\} \quad (3.10)$$

Chapter 4

Applications/Results

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This chapter will present the results obtained for the two applications of this work. It will also give the interpretations of these results to help better understanding them.

4.1 Femur on 3D CT scans

This first section aims to present the results for the first application when the algorithm has been tested with 3D CT scans of the pelvis region. This section begins by detailing the context, the reasons and the objective of this application. Then, we present a detailed explanation of the results for one scan, and finally, the results for all examples are described.

4.1.1 Application

In this application, the goal is to segment the femur in a 3D CT scan of the pelvis region. The femur is a long bone and as for every long bone, it is composed of cortical bone along the length and trabecular bone in the middle and at the extremities of the bone. On a 3D CT scan, the cortical part of the bone has good contrast and is thus easy to identify compared to the surrounding soft tissue. On the other hand, the trabecular bone has a lot of little holes and is, therefore, harder to discriminate with the surrounding soft tissue, especially at the extremities where the edges of the bone are very thin. The real difficulty for this application is to segment the femoral head due to its spongy nature.

Nowadays, the use of automatic algorithms such as Deep Learning methods in the segmentation field is widely recognised as the best method. However, regarding the segmentation of the femur, automatic algorithms are struggling to produce good results due to the lack of contrast between the trabecular bone and the surrounding soft tissue at the extremities. To enhance the quality of these automatic methods, bigger databases containing annotated examples are needed. Unfortunately, it is time-consuming for an expert to segment the scans manually.

This is where our algorithm comes into play. Indeed, the objective of our algorithm is to assist the user by providing a semi-automatic method to segment an image. Therefore, the user does not need to draw on every slice of the scan but can easily correct the segmentation if the algorithm makes some mistakes. The goal is thus to reduce drastically the time that the user spends drawing on the scan while still providing good enough results so that only minor corrections are needed afterwards to have as good a segmentation as if it were done by hand by an expert. Then, these segmented images can be added to the database to provide more examples for automatic methods in order to enhance their performance for this application.

More specifically, two major points will be observed with this application. First of all,

the performance will be assessed to see if the algorithm is able to perform the difficult task of segmenting a femoral head. The second point is to observe the distribution of the parameters' values along all the examples. This aims to see if the algorithm is consistent within a certain application, i.e. if it always uses the same, or a relatively close, set of parameters. This will prove if the algorithm can use knowledge transfer within a certain application.

This section will now explain how the comparison between the segmentation produced by our algorithm and the reference one has been computed to have some quantitative results. First of all, since the reference segmentation only takes into account the femoral head, the lower part of the femur has not been taken into account. Secondly, since the left and right femurs are completely distinct and there is no interaction whatsoever between the two during the segmentation, we decided to separate the two for the comparison. Indeed, if the left femur is badly segmented, it should not have any influence on the computation of the performance of the right one. Now, that we have our cropped images, we can apply the comparison criteria discussed in section 3.8 to the two images. **Unfortunately, the Hausdorff Distance has not been used as a criterion of comparison because its computation was too time-consuming.** Now, the Dice Coefficient and the Jaccard Index have been computed for the segmented image compared to the reference and give us a good quantitative way to evaluate the correctness of our algorithm. But these are not the only results that are interesting for this application. It is also interesting to observe the parameters used to perform the segmentation to see how they vary between different images for the same application.

4.1.2 Detailed results for one scan

As explained just above, the cortical and trabecular parts of the bone are very different on a CT scan and therefore, this algorithm cannot consider the two types of tissue at the same time. Here will be explained the different steps that have been used to segment the whole femur.

- **Step 1: Cortical bone.** The first step to segment the whole femur is to begin by segmenting the cortical bone (see Figure 4.1). This step is rather straightforward given that the cortical bone has good contrast with the rest of the image.

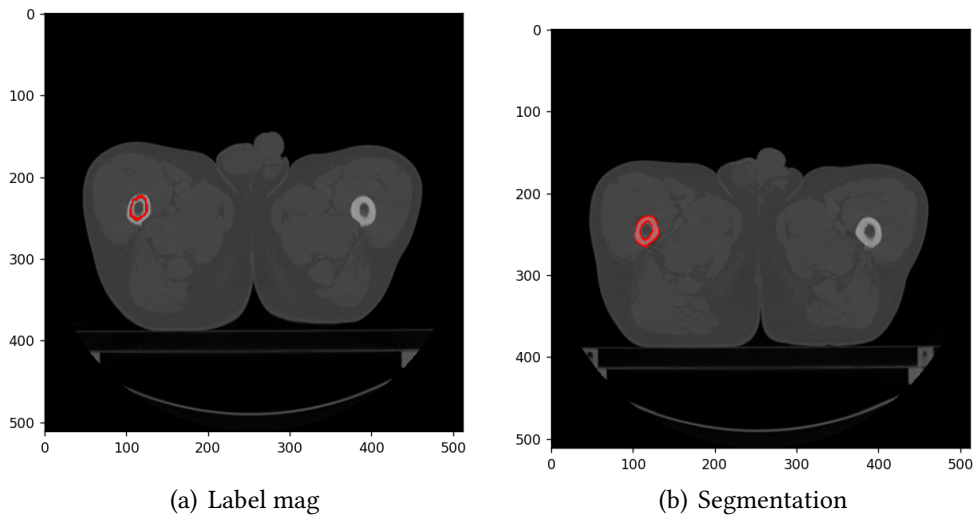


Figure 4.1: Patient 4: Illustration of the first step of the segmentation of the left femur;
(a) is the label map and (b) is the resulting segmentation

- **Step 2: Femoral Head.** The second step consists of segmenting the femoral head, mainly the bulb of the femoral head (see Figure 4.2). This step is more difficult because it is composed of trabecular bone which does not have good contrast with the surrounding soft tissue. Moreover, the edges of the femoral head are pretty thin. To avoid any leaks, it is often good to use the maximum volume parameter or the maximum number of iteration.

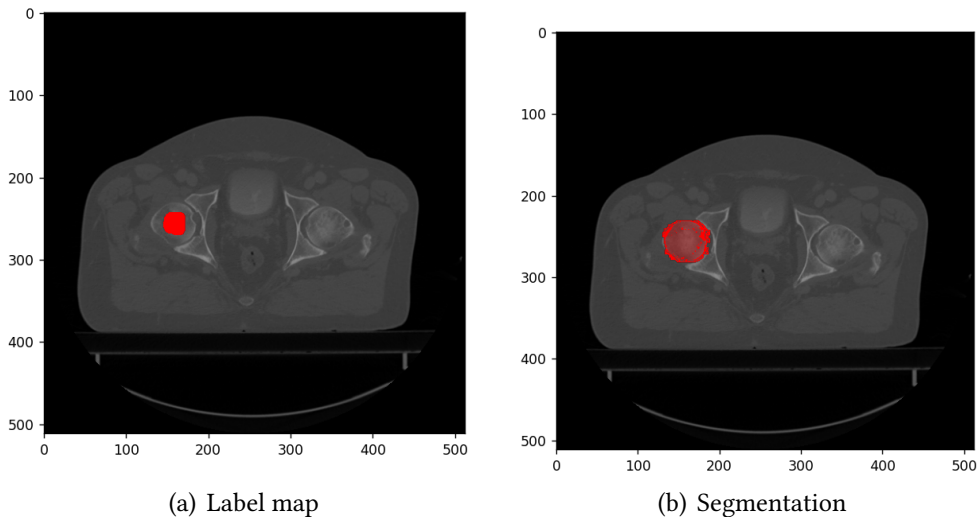


Figure 4.2: Patient 4: Illustration of the second step of the segmentation of the left femur; (a) is the label map and (b) is the resulting segmentation

- **Step 3: Transition.** The third step is to segment the rest of the femoral head, i.e. the transition part between the bulb of the femoral head and the part of the femur composed of cortical bone (see Figure 4.3). This part can also be tricky as the contrast is as low as for the femoral head.

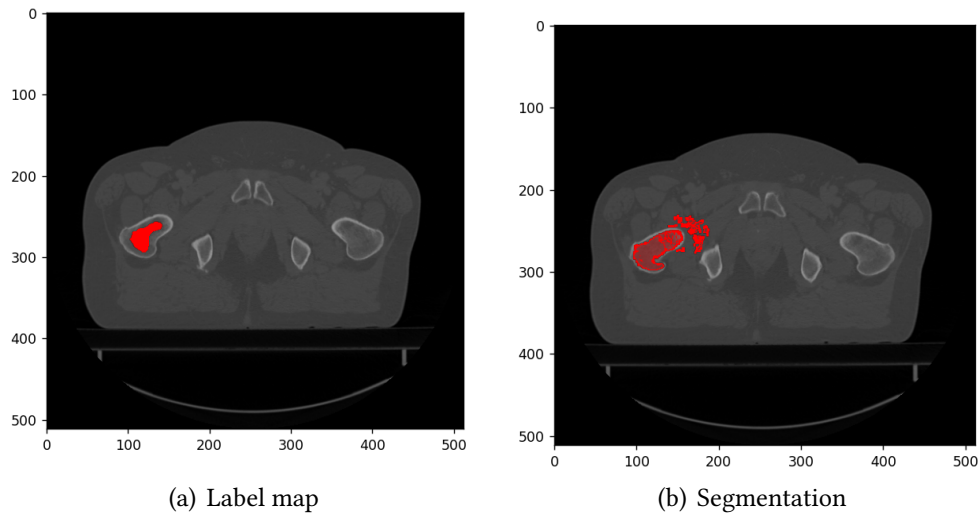


Figure 4.3: Patient 4: Illustration of the third step of the segmentation of the left femur;
(a) is the label map and (b) is the resulting segmentation

- **Step 4: Rest.** The last step consists of segmenting the rest of the femur if needed (see Figure 4.4). Indeed, for some patients, the trabecular bone inside the cortical bone is not always segmented at step 3 (depending on the CT scan). Therefore, it is necessary to add a last step to be sure to segment the whole femur.

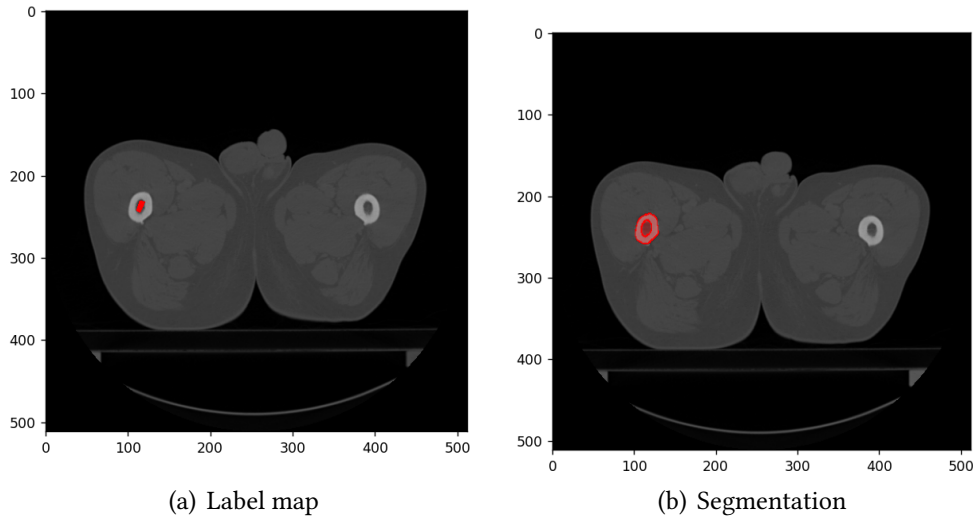


Figure 4.4: Patient 4: Illustration of the fourth step of the segmentation of the left femur; (a) is the label map and (b) is the resulting segmentation

For this application, the main focus is the femoral head as the expert segmentation received was related only to the femoral head and only comparison with this part is possible, which is the most difficult part of the femur to segment.

4.1.3 Results

This section will gather the results for all the examples where this algorithm has been tested for this first application. The interpretations, based on the raw results, will go along to help better understanding them.

Comparison of performance

This section will present the results of the performance for the segmentation of femoral heads on a CT scan of the pelvis region. First of all, you can see in Figure 4.5

the results of the segmentation of both femoral heads of patient 6. The results for the other patients can be found in appendix A.

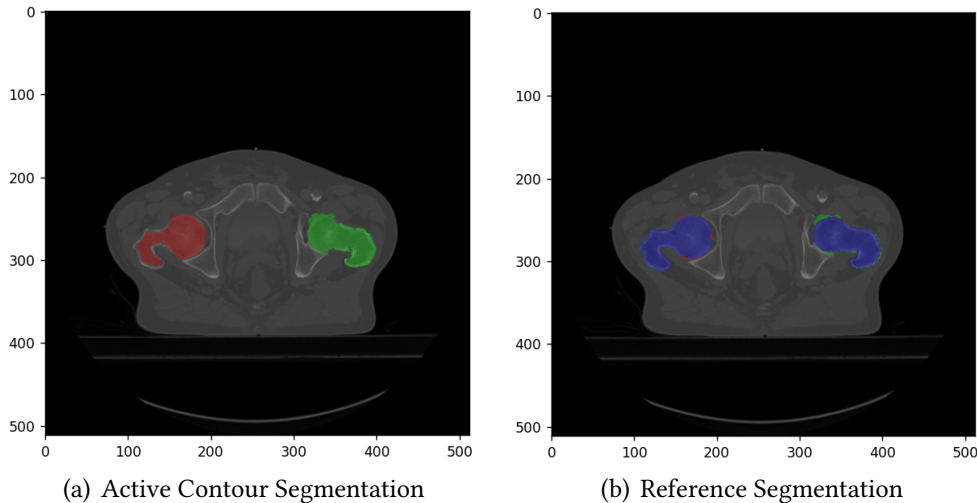


Figure 4.5: Patient 6: comparison of the segmentation performed by the active contour algorithm (a) and with the manual segmentation done by an expert on top (b)

Then, you will find the raw performance results in table 4.1. This table contains the results (Dice Coefficient and Jaccard Index) for the two femoral heads (left (L) and right (R)) of 3 patients (4,6,9).

Patient	Dice Coef	Jac Ind
4-L	0.868	0.767
4-R	0.844	0.731
6-L	0.901	0.82
6-R	0.908	0.832
9-L	0.838	0.721
9-R	0.849	0.738

Table 4.1: Performance for the CT scan application

Since the results in the table may not be the easiest to read, you will also find a boxplot representing the same results in Figure 4.6. This boxplot contains the distribution of the dice coefficient and the jaccard index for the 3 patients.

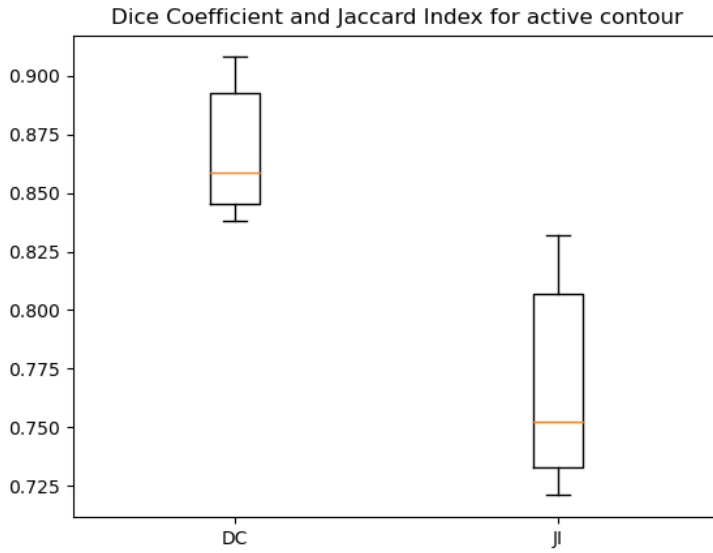


Figure 4.6: Boxplot representing the Dice Coefficient (DC) and the Jaccard Index (JI) of the segmentation of all femoral heads done by the Active Contour algortihm

Now, let us see the interpretation of these results. First of all, the means of dice coefficients and jaccard indexes for the active contour are respectively 0.868 and 0.768. These results are satisfying and prove that the algorithm is able to segment femoral heads from a CT scan of the pelvis region. The JI is lower because it tends to penalize more the errors than the DC. However, the segmentation is not perfect but rather close. This means that only minor manual corrections are needed to have usable segmentation.

Comparison of parameters

In this section, the parameters used to perform the various segmentations for this application will be compared. Table 4.2 represents all the parameters used. The first column *Patient* shows the patient and which femoral head (left or right) is being segmented. The second column *Step* represents which part of the femur the algorithm work on (see section 4.1.2). Then, the last eight columns correspond each to one parameter of the algorithm (see section 3.5.3).

Patient	Step	Dim	IH	CW	Max It	Max Run Time	Max Vol	PC	Inertia
4-L	FC	3	0.5	0	50	300	25000	0.1	0.1
4-L	FH	3	0.35	0.3	30	300	35000	0.1	0.2
4-L	FTrans	3	0.3	0.2	30	300	30000	0.1	0.2
4-L	FT	3	0.5	0.1	30	300	30000	0.1	0.2
4-R	FC	3	0.5	0	50	300	25000	0.1	0.1
4-R	FH	3	0.35	0.3	30	300	30000	0.1	0.2
4-R	FTrans	3	0.25	0.2	25	300	30000	0.1	0.2
4-R	FT	3	0.6	0.1	20	300	30000	0.1	0.1
6-L	FC	3	0.6	0	35	300	25000	0.1	0.1
6-L	FH	3	0.3	0.3	30	300	30000	0.1	0.2
6-L	FTrans	3	0.35	0.2	30	300	35000	0.1	0.2
6-R	FC	3	0.4	0	50	300	35000	0.1	0.1
6-R	FH	3	0.3	0.3	30	300	30000	0.1	0.2
6-R	FTrans	3	0.35	0.2	33	300	35000	0.1	0.2
9-L	FC	3	0.2	0	80	300	60000	0.1	0.1
9-L	FH	3	0.4	0.3	30	300	35000	0.1	0.2
9-L	FTrans	3	0.3	0.2	30	300	35000	0.1	0.3
9-L	FT	3	0.3	0.1	35	300	35000	0.1	0.2
9-R	FC	3	0.2	0	80	300	60000	0.1	0.2
9-R	FH	3	0.3	0.3	30	300	35000	0.1	0.2
9-R	FTrans	3	0.2	0.2	30	300	35000	0.1	0.3
9-R	FT	3	0.3	0.1	35	300	35000	0.1	0.2

Table 4.2: Comparison of the parameters for the CT scan application

As a reminder, the aim is to observe the distribution of the parameters within the same application. Therefore, boxplots of these distributions have been produced. First of all, the Dimension and the PC parameters have not been taken into account since they do not vary between the different segmentations. Then, the Maximum Iteration, Maximum Running Time and Maximum Volume are also not taken into account since they are limiting parameters to stop the algorithm; therefore they have only a marginal implication into the performance of the results. Then, only the Intensity Homogeneity, the Curvature Weight and the Inertia parameter will be further discussed. You will find the boxplots for these parameters respectively in Figures 4.7, 4.8, and 4.9. Each of these boxplots contains the distribution of parameter's value for each possible steps for the segmentation of an entire femur. Narrower distributions would be preferable across the different segmentations as it would prove that the algorithm can use the same set of parameters for the same application and that it does not depend on the patient.

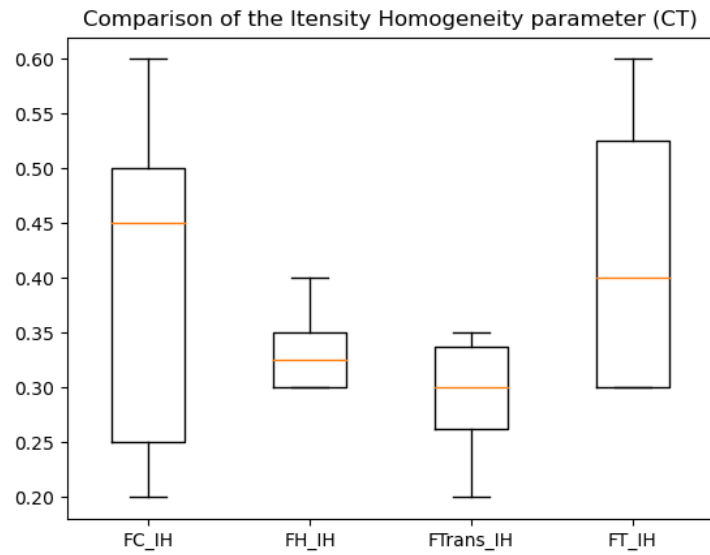


Figure 4.7: Boxplot representing the value of the Intensity Homogeneity (IH) parameter for the different steps of the segmentation of a whole femur

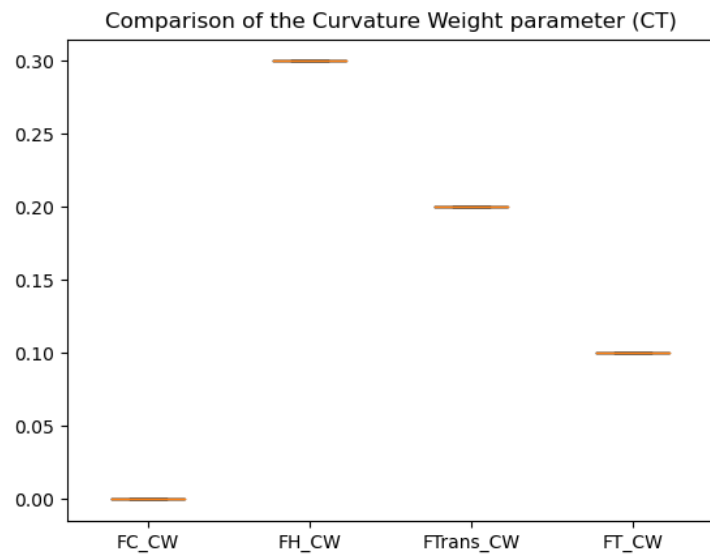


Figure 4.8: Boxplot representing the value of the Curvature Weight (CW) parameter for the different steps of the segmentation of a whole femur

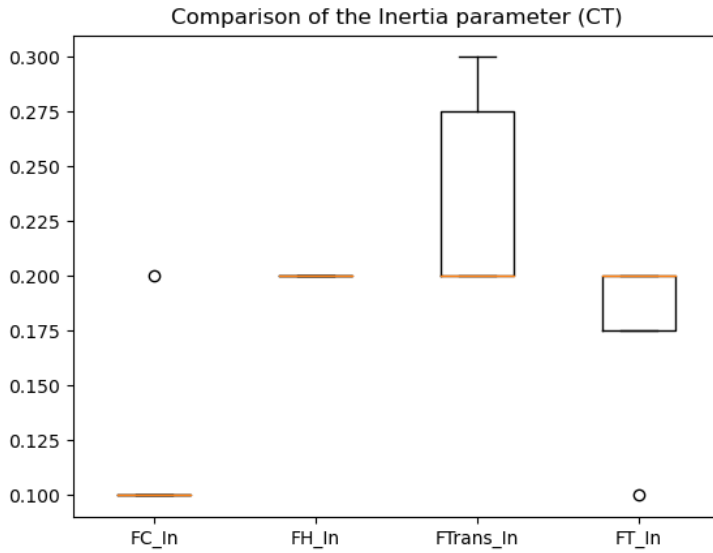


Figure 4.9: Boxplot representing the value of the Inertia (In) parameter for the different steps of the segmentation of a whole femur

Now, interpretations of the different boxplots will be made. For the parameters, unlike for the performance coefficient, it is not the mean but rather the variation of the distribution which is interesting. Therefore, the standard deviation has been computed and will be used to draw interpretations.

For the Intensity Homogeneity parameter, the mean of the standard deviation for each step is 0.093. This deviation, although significant, still stays reasonable. For the Curvature Weight, the mean of the standard deviation is nearly 0 which is perfect, while it corresponds to 0.032 for the Inertia parameter. This is a small value of variation and it allows concluding that the use of the same set of parameters for the same application would claim somewhat good performance.

4.2 Brain pathologies on 3D MRI

The results of the second application about brain pathologies will now be presented. The scans used here are MRI scans, but it is not always the same sequence that is used for every patient. Moreover, some scans use enhancement products. This depends on the type of pathology to be observed. Firstly, the context and the objective of this

application will be detailed. Then, there will be a detailed explanation of the results for one example, and finally, the results for all the patients will be described.

4.2.1 Application

This application takes place in the context of the everyday work of a neurosurgeon. When a patient needs to undergo brain surgery, the neurosurgeon will start by running a scan on the patient (often an MRI) to have a 3D image of the brain and to observe what and where is the pathology. Then, the neurosurgeon will proceed to the segmentation of the VOIs in the brain. This will help him to plan its operation; where to enter, which way to go, which angle, etc, in order to do the least amount of damage possible to the surrounding healthy tissue (this is especially important in the brain). Then the operation can take place. In summary, a big part of the everyday work of a neurosurgeon is to perform segmentation. As seen earlier, nowadays, the segmentation tools used by neurosurgeons are just drawing software with an interpolation function to accelerate the process, but the doctor still needs to draw on nearly every slice where there is the pathology.

The objective of this algorithm is to provide a semi-automatic method for the neurosurgeon to reduce significantly the time that they take to perform segmentation. However, the algorithm needs to produce good enough results so that the time of correction is minor.

In more detail, three main points will be focused on with this application. Firstly, the performance of the segmentation will be assessed to observe if this algorithm produces fine results for brain pathologies. For the comparison of performance, all segmentation performed by our algorithm has been compared to reference images validated by a neurosurgeon. As for the previous application, the Dice Coefficient and the Jaccard Index have been used to assess quantitatively the quality of the segmentation. Then, the influence of the choice of parameters will be detailed, and finally, a comparison of drawing time will be made to observe which of this algorithm or manual segmentation used in hospital is faster.

4.2.2 Detailed results for one scan

For this application, the segmentation is way more straightforward than for the femur. Indeed, in general, the pathological tissue has the same tone and structure. Therefore, the segmentation is built in one step. Unfortunately, depending on the

pathology, the VOI can appear more or less homogeneous in the MRI, which will influence the facility to perform the segmentation. Still, for all patients, the segmentation is always performed by feeding the algorithm with an initial zone representative of the VOI and then choosing a good set of parameters allowing the semi-automatic method to do its best job. This can be observed in Figure 4.10.

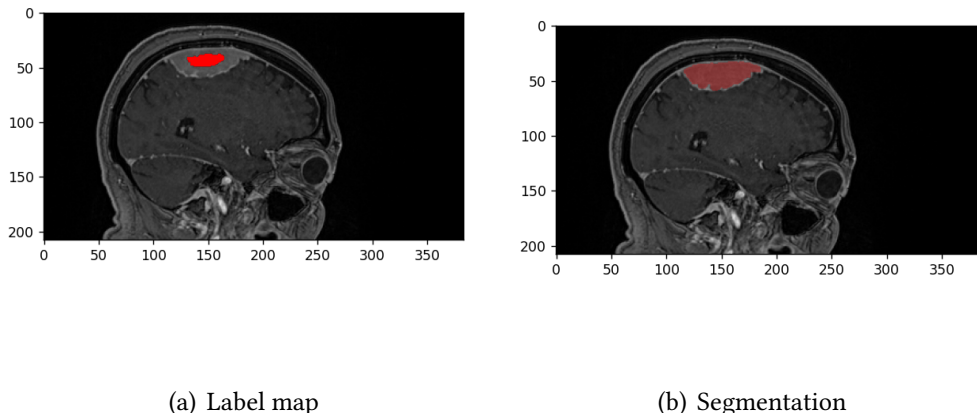


Figure 4.10: Patient 5: Illustration of the segmentation his pathology; (a) is the label map and (b) is the resulting segmentation

4.2.3 Results

This section will gather the results for all the examples where this algorithm has been tested for this application. The interpretations, based on the raw results, will go along to help better understanding them.

Comparison of performance

In this section, the performance of our algorithm will be detailed for the brain pathology application. First of all, you can see in Figure 4.11 the results of the segmentation of the brain pathology for patient 6. The results for the other patients can be found in appendix B.

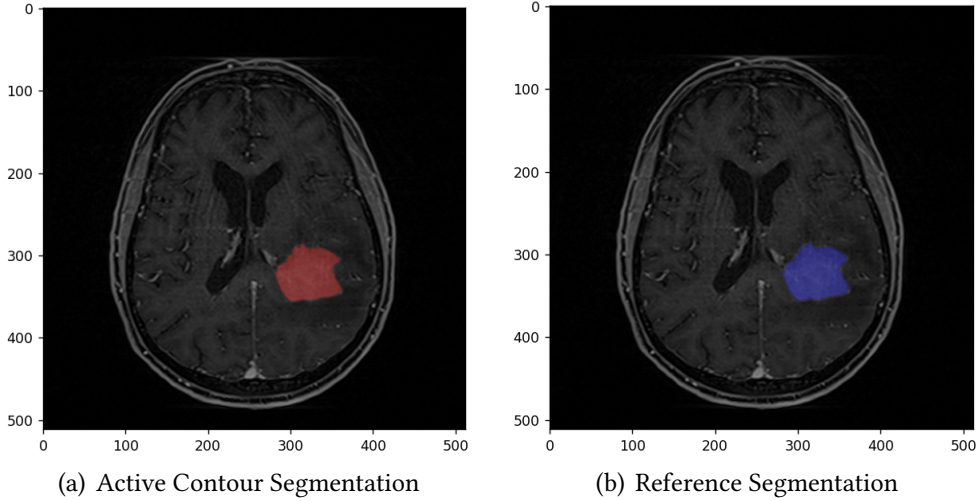


Figure 4.11: Patient 6: comparison of the segmentation performed by the active contour algorithm (a) and with the manual segmentation done by a neurosurgeon on top (b)

Table 4.1 gathers the DC and the JI for all the patients that have been segmented while Figure 4.12 represents via boxplots these values. For more information, the mean DC for this application is 0.834 while the mean JI is 0.747. These values are already convincing but it is not the only observation that can be made. As can be observed on the boxplot, there is the presence of outliers. This can also be seen in the table where the DC and JI are significantly lower for patients 2 and 15. That means that the algorithm does not work for these two pathologies due to their inhomogeneities. If these two outliers are removed, then we obtain a mean DC and JI respectively of 0.9016 and 0.829 which are very promising results for medical image segmentation. This means that the algorithm can be very useful in the segmentation of certain brain pathologies, but not for all of them.

Patient	Dice Coef	Jac Ind	Pathology
1	0.888	0.798	lymphoma
2	0.438	0.28	taxoplasmosis
4	0.933	0.875	radionecrosis
5	0.951	0.906	meningioma
6	0.971	0.943	lymphoma
7	0.802	0.669	lung adenocarcinoma
10 (small lesion)	0.745	0.594	glioblastoma
10 (big lesion)	0.847	0.735	glioblastoma
13 (ventricle)	0.996	0.991	ventricle
13 (lymph)	0.929	0.868	lymphoma
14	0.932	0.872	pilocytic astrocytoma
15	0.559	0.388	lung carcinoma

Table 4.3: Comparison of performance for the brain pathology application

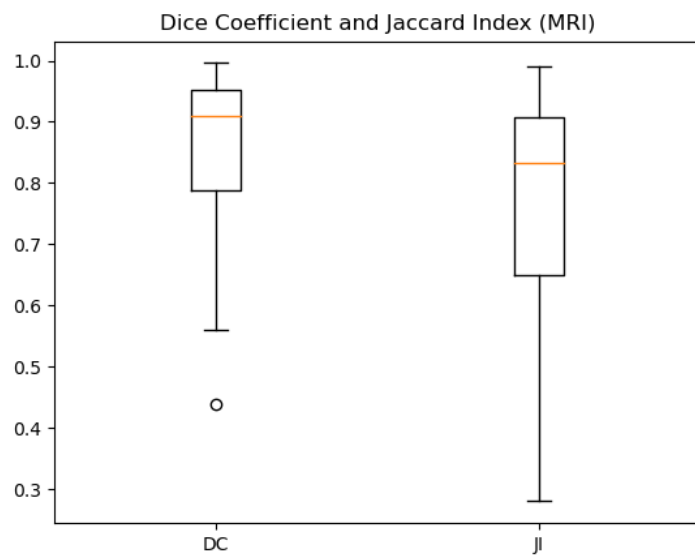


Figure 4.12: Boxplot representing the Dice Coefficient (DC) and the Jaccard Index (JI) of the segmentation of all patients with brain pathologies

Comparison of parameters

Table 4.4 contains the parameters that have been used to perform the segmentation on all the patients.

Patient	Dim	IH	CW	Max It	Max Run Time	Max Vol	PC	Inertia
1	3	0.1	0.1	20	120	5000	0.1	0.1
2	3	0.45	0.2	10	120	3000	0.1	0.2
4	3	0.9	0.1	8	120	2000	0.1	0.1
5	3	0.1	0.2	30	180	40000	0.1	0.4
6	3	0.2	0.2	30	420	80000	0.1	0.2
7	3	0.6	0.33	25	300	40000	0.1	0.3
10 (small lesion)	3	0.8	0.3	10	120	5000	0.1	0.3
10 (big lesion)	3	0.7	0.3	40	300	40000	0.1	0.3
13 (ventricle)	3	0.1	0.1	40	300	65000	0.1	0.3
13 (lymph)	3	0.15	0	40	300	10000	0.1	0.2
14	3	0.2	0.1	40	420	85000	0.1	0.3
15	3	0.675	0.3	15	180	10000	0.1	0.3

Table 4.4: Comparison of parameters for the brain pathology application

Unlike the other application where it is interesting to check the distributions of the parameters' values, this is not the case here since the patients have a variety of different brain pathologies. Instead, another interesting feature to observe is the influence of the parameters on the segmentation. As for the previous application, the focus is put on the Intensity Homogeneity (IH), the Curvature Weight (CW) and the Inertia (In) parameter because they have the most influence on the segmentation. For each patient, segmentations have been performed using different values for these parameters to see their influence.

The first parameter being tested is the IH parameter. As a reminder, the IH parameter is used to choose the bandwidth of the kernel function used for the KDE of the region's features. A high IH means a narrower pdf and thus, the acceptance of further points is lower (see section 3.3.2). Therefore, a high IH value should prevent the region to expand too much. The results for patient 5 and 6 can be observed respectively in Figures 4.13(a) and 4.13(b). The results for the other patients can be seen in appendix B. It can be observed that, for a high value of IH, the region segmented is more constrained than for a small value. This confirms the assumption made on this parameter.

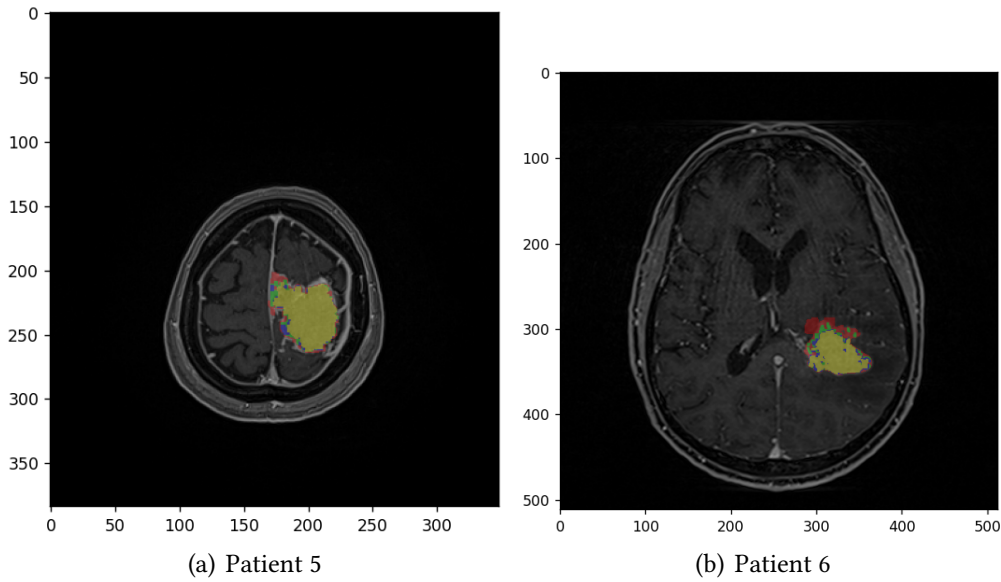


Figure 4.13: Influence of the variation of the IH parameter: $\text{IH}=0.1$, $\text{IH}=0.4$, $\text{IH}=0.6$, $\text{IH}=0.9$

Secondly, the CW parameter is tested. The CW parameter corresponds to the λ coefficient in the formula of the energy of the surface (see equation 3.4.1)¹. It weights the importance of the first term compared to the second term of this formula (see section 3.4.1). The second term of this formula is related to the mean curvature of the surface and it penalizes non smooth surfaces (with a lot of curvature). Therefore, a high CW value should make the surface smoother and with less curvature. The results for patient 4 and 13 (lymph) can be observed respectively in Figures 4.14(a) and 4.14(b). The other results can be seen in appendix B. It can be observed that the regions are less expanded for high values of CW. However, the influence on the smoothness of the contour is not significant.

¹ $E_i(C_i) := (1 - \lambda) \int_{\mathbf{x} \in C_i} (p^c - p_i(\mathbf{f}(\mathbf{x}))) d\mathbf{x} + \lambda \int_{C_i} dA$

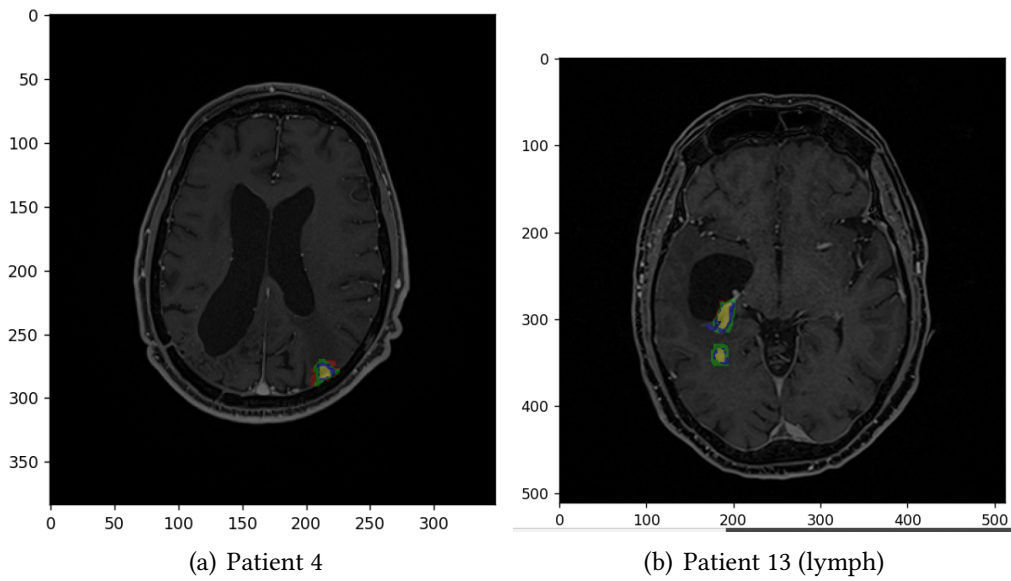


Figure 4.14: Influence of the variation of the CW parameter: $CW=0.1$, $CW=0.2$, $CW=0.3$, $CW=0.4$

The final parameter to be tested is the In parameter. It corresponds to the inertia of the surface, i.e. the influence, for the force at a point on the surface, of the previously computed force (at the previous iteration) at that point. A high value of In should allow the region to grow further. The results for patient 7 and 10 (small) can be observed respectively in Figures 4.15(a) and 4.15(b). The other results can be seen in appendix B. It can be observed that the regions expand more for high values of In, which confirms our assumption.

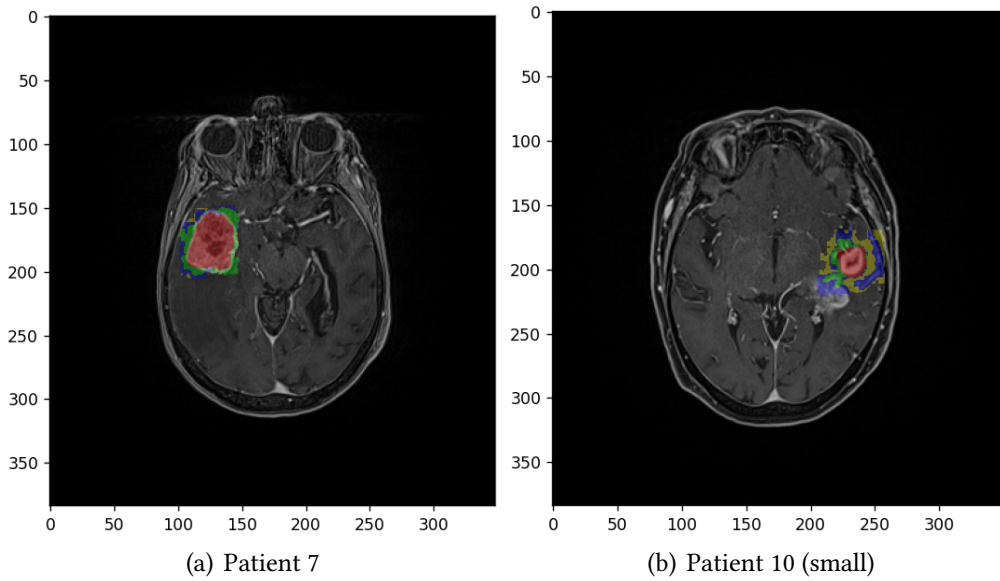


Figure 4.15: Influence of the variation of the In parameter: $In=0.1$, $In=0.4$, $In=0.6$, $In=0.9$

Comparison of time

For this application, the comparison of time has been an important focus. Indeed, the goal was to see if this algorithm allows reducing the drawing time of the user compared to software that is currently used by neurosurgeons in hospitals such as StealthStation by MedTronic. One problem is that the major part of the time to perform the segmentation with active contour is used by the algorithm. Therefore, the user could use this time to do other things. Another issue is that the time taken by the algorithm to perform the segmentation is directly related to the power of the computer on which it runs. To finish, for the MedTronic software, the user nearly only takes time to draw on the scan. For all of these reasons, the comparison will be made only on the drawing times, i.e. the time that the user pass to actively draw on the scan. The table 4.5 shows the time that it took for the same person to perform the segmentation for all patients with the active contour algorithm or the MedTonic software. These values can be better observed in Figure 4.16.

Patient	Active Contours: Time	MedTronic: Time
1	0'44"26	3'01"57
2	0'25"06	3'20"95
4	0'23"13	2'58"23
5	0'38"74	7'23"07
6	0'48"10	8'07"01
7	1'07"77	7'36"26
10 (small lesion)	0'26"47	6'49"61
10 (big lesion)	1'41"79	6'49"61
13 (ventricle)	0'40"98	5'08"00
13 (lymph)	1'03"56	4'42"85
14	0'46"16	7'55"84
15	0'57"92	4'24"04

Table 4.5: Comparison of time for the brain pathology application

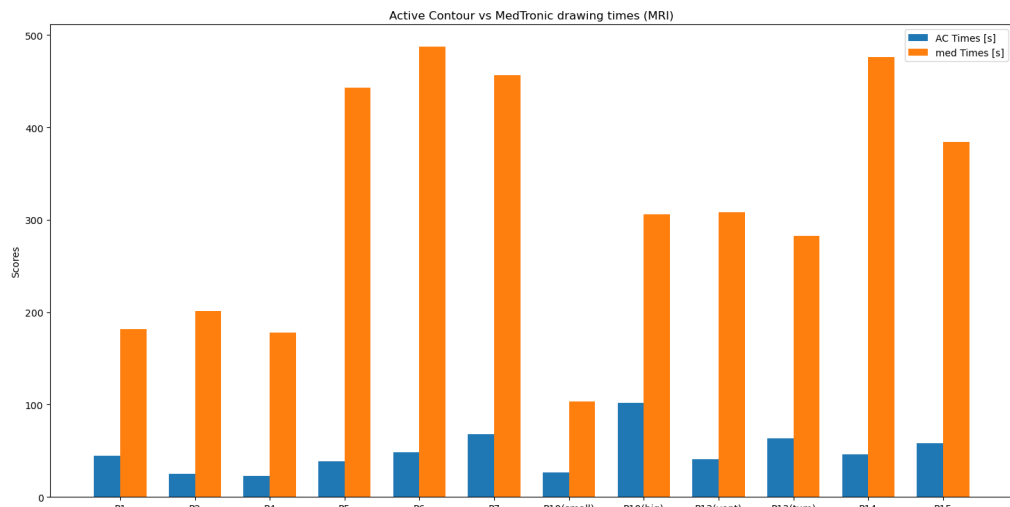


Figure 4.16: Comparison of the drawing times for the Active Contour (AC) algorithm (blue) vs the MedTronic (med) software (orange).

In Figure 4.16, the orange bar represents the time taken by the MedTronic software while the blue bar corresponds to the active contour algorithm. At the first sight, a clear difference can be noticed between the two; the active contour algorithm reduces drastically the drawing time. To have some more quantitative measure, on average, the

drawing time is divided by 6.996 when using the active contour algorithm compared to the MedTronic software, which is a significant improvement.

Chapter 5

Optimization of the parameters

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This chapter will look into one big limitation of the algorithm that arose from the results (see chapter 4). This limitation is the choice of the parameters that is not straightforward and can take a lot of time which is counter productive. This chapter has for objective to overcome this limitation by using a gradient descent method to optimize the choice of parameters.

5.1 Context

Chapter 4 has shown some very promising results for this algorithm. Indeed, the performance of the algorithm is quite satisfying with a mean dice coefficient of 0.868 for the segmentation of the femoral head and of 0.9016 for the segmentation of various brain pathologies. For this last application, it has also been shown that the active contour algorithm helps dividing the drawing time by a mean factor of 6.996, which is a significant improvement. However, there is a big hidden limitation to this reduction of time. Indeed, this only considers the drawing time but it does not consider the time passed by the user to find the best parameters to use. Unfortunately, this step can take several trials and thus a significant amount of time.

This issue is quite problematic since the time taken to find the best parameters can be quite frustrating, which would lead a doctor to prefer doing the segmentation manually. Therefore, the need to find a solution to this problem is important to meet the list of requirements detailed in section 1.3. In order to resolve this issue, there are two major possible solutions;

- Either the user needs to be well trained in order to find the parameters quickly (in a couple of trials)
- Or the parameters need to be found automatically and thus a complementary method needs to be implemented.

The first solution is not a very good solution since it is opposite to the user-friendly requirement. Therefore, **the second solution will be chosen and an automatic method to find the best parameters will be implemented.**

5.2 Possible solutions

Now, the next question to be asked is whether or not it is feasible to find the best parameters to segment an image. This section will be used to discuss this issue and to present possible solutions.

To the question of whether or not it is feasible, the answer is that the optimal parameters for an image cannot be found but an approximation can. Indeed, based only on a new image to segment, the parameters cannot be found. Naturally, another already segmented image is needed as an example. As it has been shown in the results

(see section 4.1.3), the variation of the values of the parameters for the same application is relatively small. Therefore, if the parameters are known for one instance of the application, they can be used for another instance and still claim quite good performance. This proves that it is possible to find automatically the parameters if other examples are available. Below, different possibilities of solutions will be presented with their advantages and limitations.

The first possibility is to use Machine Learning (ML) to optimize the parameters. Since the problem is relatively simple, a simple Artificial Neural Network (ANN) could be used. An ANN is a simple and quite often used solution to optimize parameters and therefore, it could completely suit this application. But there is a major limitation; ML is a method that requires to have large training datasets in order to train the network. Unfortunately, this is not the case here. In this application, the parameters should be found based on one or a couple of examples only. Therefore, this solution has been dropped out because it is not feasible.

Thus, another solution needs to be found to optimize the parameters for one example only. Fortunately, a simple but elegant solution exists, namely a **Gradient Descent**. Indeed, this technique allows finding the best parameters to segment an example image where the solution is known. Now, the parameters found via a Gradient Descent for this example can be reused for another instance of the same application.

5.3 Method

This section will detail the solution that has been chosen to resolve the issue of optimizing the parameters. First, the general principle of Gradient Descent will be explained, and then the implementation realised for this work will be presented.

5.3.1 Gradient Descent

The Gradient Descent is one of the most simple and popular algorithm to perform optimization. It is a first-order iterative optimization algorithm for finding a local minimum of a differentiable function [69]. Gradient Descent is a way to minimize an objective function $J(\Theta)$ parameterized by a model's parameters $\Theta \in \mathbb{R}^d$ by updating the parameters in the opposite direction of the gradient of the objective function $\nabla_{\Theta} J(\Theta)$, with regard to the parameter [70]. This process can be observed in Figure 5.1. From a more intuitive point of view, the algorithm follows the direction of the steepest slope of

the surface created by the objective function downhill until it reaches a local minimum. A final but still important parameter is the learning rate α . This parameter characterizes the size of the steps that the algorithm takes to reach a minimum. If α is small, it will take a lot of steps to reach the minimum. On the other hand, if it is large, the algorithm will lack precision to find the real local minimum of the objective. A last possibility is to have an adaptive parameter α , which would be large at the beginning to quickly descent towards the minimum but would get smaller and smaller towards the end to precisely find the local minimum [49, 71].

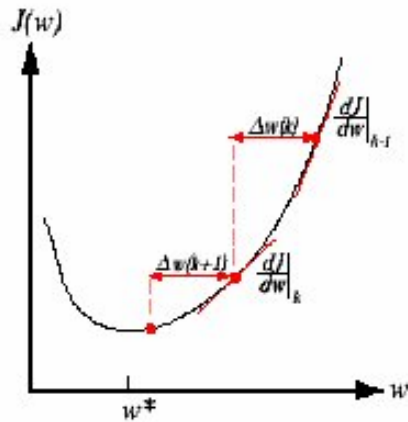


Figure 5.1: Representation of the minimization of an objective function J using a gradient descent algorithm [72].

One big limitation of using Gradient Descent is that it finds a local minimum. But the objective here would be to find the optimal parameters, thus the global minimum. To do so, the initial guess has to be near enough the best solution in order for the gradient descent to follow the path towards the global minimum.

5.3.2 Implementation

The choice of a good objective function is needed to be able to perform a good optimization. Here, the goal is to find the best parameters for the active contour algorithm that will produce the best segmentation possible. As a reminder, there are a lot of parameters that can be tuned (see section 3.5.3). However, a lot of parameters will induce a more difficult and more time-consuming optimization. Fortunately, as seen in section 4.1.3, only three parameters (the Intensity Homogeneity, the Curvature Weight and the Inertia) have a big influence on the segmentation, the other being limiting factors.

Therefore, **the optimization will take place only on these three parameters**. The other parameters will be chosen in order not to influence the segmentation.

Now that the parameters on which the optimization will be performed have been selected, the choice of the objective function is the next step. Fortunately, in our case, a criterion has already been used to quantitatively assess the performance of a segmentation; the Dice Coefficient. Therefore, the algorithm should optimize the parameters in order to produce the biggest DC possible. The only subtlety is that, because it is a Gradient Descent algorithm, it searches for a local minimum. Therefore, the objective function will take the opposite of the DC. Indeed, minimizing the opposite of the DC will maximize the DC.

The optimizer algorithm used is the *brute* function from the *scipy.optimize* python package [73]. This is a largely used optimization package that works really well. The *brute* function minimizes a function over a given range by brute force at first and then switches to a gradient descent to find the minimum. The brute force step computes the objective function at 4 points equally spaced along each dimension (64 points in total). This step is used to find the initial guess for the gradient descent to help it finding the global minimum. This has been working fairly well. Since the optimizer calls iteratively the segmentation algorithm and that this last one takes a bit of time to run, the *RSSegmentor* class has also been a little modified for this application in order to reduce the time taken for this optimization. Still, this optimization is quite time-consuming (couple of hours per example).

5.4 Results

This section aims to present the results of the optimization of the parameters for all the patients of the brain pathologies application. The choice of not working with the femoral head application has been made due to the complexity of doing the segmentation in several steps. This section will also present interpretations of these results. First of all, you will find in Figure 5.2 the representation of the gradient descent on the parameters that has been done for patients 6 and 13 (lymph). The graphs for the other patients can be found in appendix C. These graphs are 3D representations of 4D data (3 variable parameters and the associated DC), thus not straightforward to understand. The x-axis represents the value of the IH parameter and the y-axis the CW parameter. The z-axis represents the opposite of the DC. The In parameter is represented by the color of each point associated with the color bar on the right. Finally, the semi-transparent surface represents the objective function to minimize across the range of all variables. The result of this optimization corresponds to the lowest point

on the graph, from which a set of optimal parameters can be derived.

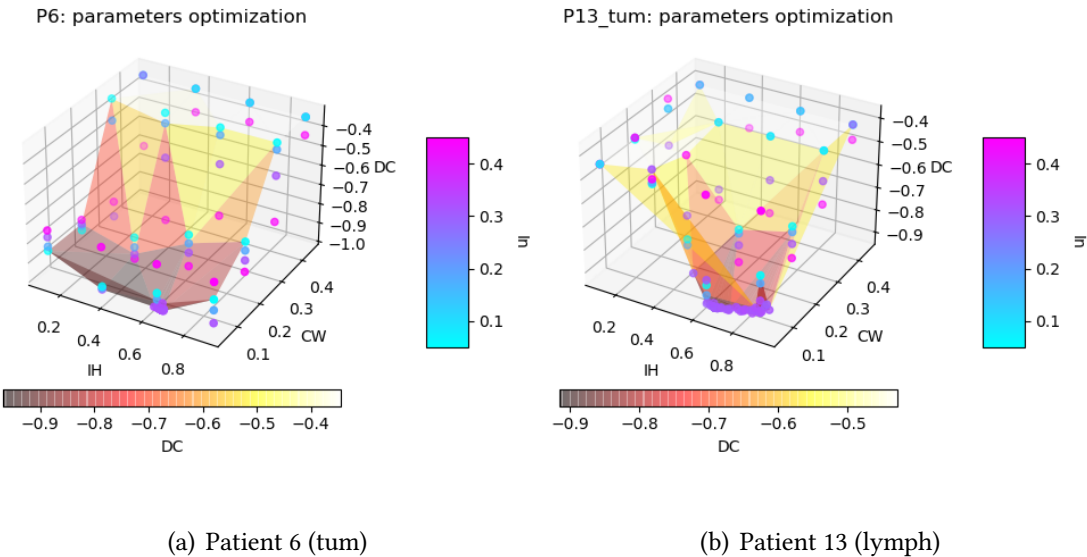


Figure 5.2: Optimization of parameters. Each point corresponds to one evaluation of the objective function for a set of parameters. The color of each point corresponds to the In parameter used. The surface corresponds to the objective function to minimize

This proves that the algorithm used to perform the gradient descent works. The next step is to assess the quality and the utility of this optimization. Table 5.1 represents the DC of the segmentation performed with the parameters found either manually or via optimization. For patient 2, a big improvement can be observed with the optimization as well as for patient 15, while a smaller improvement can be observed for patient 10 (small lesion). However, for patient 4, the optimization has produced a worst result than for what has been found manually. For the rest of the patients, the optimization has produced not significantly different results. The boxplot in Figure 5.3 compares the distribution of the results for the parameters found manually versus via optimization. It can be observed that the distribution is more condensed for the optimization with no outlier. The mean DC is 0.879 compared 0.834 when the parameters are found manually, which shows a little improvement. However, the biggest observation that can be made is that the optimization manages to find parameters for all the pathologies.

Patient	DC	DC (opti)	Pathology
1	0.888	0.8941139831828091	lymphoma
2	0.438	0.7360851699663803	taxoplasmosis
4	0.933	0.8633714152483096	radionecrosis
5	0.951	0.9508582256850155	meningioma
6	0.971	0.9697162494999301	lymphoma
7	0.802	0.8025901755451004	lung adenocarcinosis
10 (small lesion)	0.745	0.8452206651328174	glioblastoma
10 (big lesion)	0.847	0.8706791499479863	glioblastoma
13 (ventricle)	0.996	0.9949653454949143	ventricle
13 (lymph)	0.929	0.9160632481158564	lymphoma
14	0.932	0.9336011472877453	pilocytic astrocytoma
15	0.559	0.7681159420289855	lung carcinosis

Table 5.1: Comparison of performance for the brain pathology application with parameters found either manually or via optimization

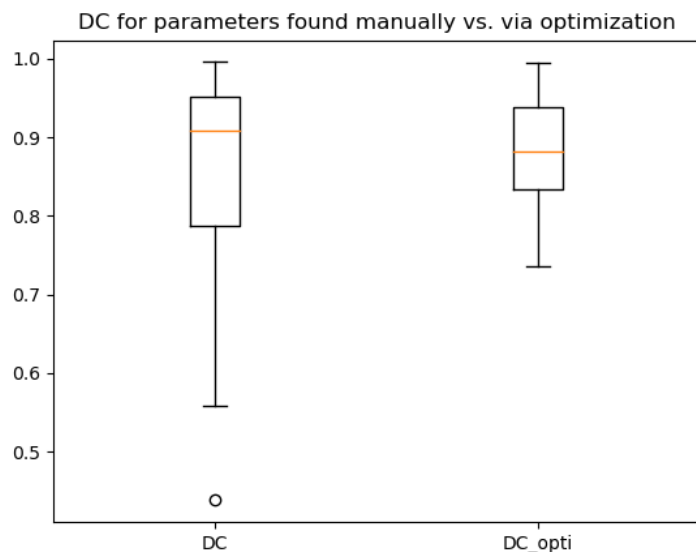


Figure 5.3: Boxplot of the distribution of DC without and with optimization

Finally, table 5.2 represents the comparison between the parameters found by the optimization versus those found manually. It can be observed with this table that not only one set of parameters can be chosen to produce relatively close results. A

second observation can also be made by the known effect of each of the three principal parameters (see section 4.2.3). As a reminder, the IH and CW have a constraining effect on the segmentation when their values increase while it is the opposite effect for the In parameter. This second observation corresponds to the fact that generally, when a parameter is selected to be more constraining, another one will be selected to be less constraining. For patient 5, for example, while the IH parameter is the same (0.1), the CW selected via optimization is more constraining (0.325 vs. 0.2) but is balanced by the selection of the In parameter which is less constraining (0.45 vs. 0.4). The same kind of observation can be made for the majority of the trials.

Patient	Dim	IH	CW	Max It	Max Run Time	Max Vol	PC	Inertia
1	3	0.1	0.1	20	120	5000	0.1	0.1
1 (opti)	3	0.10639411	0.04911527	20	120	5000	0.1	0.31973544
2	3	0.45	0.2	10	120	3000	0.1	0.2
2 (opti)	3	0.9	0.31666667	10	120	3000	0.1	0.05
4	3	0.9	0.1	8	120	2000	0.1	0.1
4 (opti)	3	0.92961263	0.3081632	8	120	2000	0.1	0.45329083
5	3	0.1	0.2	30	180	40000	0.1	0.4
5 (opti)	3	0.10213542	0.32540799	30	180	40000	0.1	0.44929688
6	3	0.2	0.2	30	420	80000	0.1	0.2
6 (opti)	3	0.65875335	0.04713384	30	420	80000	0.1	0.32292469
7	3	0.6	0.33	25	300	40000	0.1	0.3
7 (opti)	3	0.65722673	0.32377921	25	300	40000	0.1	0.17693413
10 (small)	3	0.8	0.3	10	120	5000	0.1	0.3
10 (small) (opti)	3	0.6175733	0.31157215	10	120	5000	0.1	0.05227431
10 (big)	3	0.7	0.3	40	300	40000	0.1	0.3
10 (big) (opti)	3	0.95214759	0.31856174	40	300	40000	0.1	0.31295273
13 (vent)	3	0.1	0.1	40	300	65000	0.1	0.3
13 (vent) (opti)	3	0.09752232	0.04923357	40	300	65000	0.1	0.19515924
13 (lymph)	3	0.15	0	40	300	10000	0.1	0.2
13 (lymph) (opti)	3	0.67011592	0.05541995	40	300	10000	0.1	0.30623441
14	3	0.2	0.1	40	420	85000	0.1	0.3
14 (opti)	3	0.41811658	0.04808355	40	420	85000	0.1	0.31831428
15	3	0.675	0.3	15	180	10000	0.1	0.3
15 (opti)	3	0.63333333	0.05	15	180	10000	0.1	0.31666667

Table 5.2: Comparison of parameters for the brain pathology application found either manually or via optimization

Now that the optimization has been done and that it is known to work, we need

to assess if the knowledge of these optimal parameters is useful and can be transferred. To do so, two experiments have been conducted with this dataset. First, with patient 1, 6 and 13 who have all a lymphoma and then with patient 7 and 15 who have metastases. The goal of this experiment is to perform the segmentation for each patient with the parameters of the other patients with the same pathology and compare their obtained DC. The results for the lymphoma are listed in table 5.3. It can be observed that the transfer of parameters from patient 13 to patient 6 MRI as been really successful. However the rest of the results are not very convincing. Let us try to explain these poor results. First of all, although patients 6 and 13 are using the same MRI sequence (t1_vibe_fs_tra_1mm_iso), patient 1 is using another sequence (t1_vibe_fs_tra_0_8_mm_CAIPI). Basically, it means that the resolution used is not the same between patient 1 and patients 6 and 13. This can explain the variations of DC for patient 1 and when using patient 1's parameters. Still, the DC for patient 13 using patient 6's parameters is quite poor. This can be explained by the fact that the lymphoma of Patient 13 is small and very irregular while the lymphoma for patient 6 is big and regular.

		Patients' Parameters		
		1	6	13
Patients	1	0.8941139831828091	0.7995802728226653	0.7788649706457925
	6	0.7898693427572006	0.9697162494999301	0.9603777096455054
	13	0.44430408301376045	0.7138800734925672	0.9160632481158564

Table 5.3: Comparison of the DC when using optimal parameters vs. the parameters of other patients with the same pathology (lymphoma)

The results for the second experiment about metastases are listed in table 5.4. Both MRIs are using the same sequence (t1_vibe_fs_tra_0_8_mm_CAIPI). The results for this experiment are not really good either. Indeed, a difference of 0.07267 and of 0.07126 between the use of the optimal parameters and the parameters for another instance can be observed. This can be explained by the inhomogeneity of a metastasis on an MRI and by the difficulty of this algorithm to segment this pathology. However, it can be noted that, although the results of this transfer of parameters are usually significantly lower, it can still be a good starting point to find the best parameters for a new instance.

		Patients' Parameters	
		7	15
Patients	7	0.8025901755451004	0.7299174494733845
	15	0.6968532557898016	0.7681159420289855

Table 5.4: Comparison of the DC when using optimal parameters vs. the parameters of other patients with the same pathology (metastasis)

Chapter 6

Discussion

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This chapter aims to discuss the results obtained in chapter 4 and 5. Firstly, the performance, parameters and optimization, and the time improvement will be discussed. Then, the limitations of the algorithm developed in this thesis will be presented. Finally, some future perspectives will be proposed.

This section aims to discuss the overall work of this thesis. As a reminder, the objective of this work was to develop and implement a user-friendly semi-automatic tool to perform segmentation on medical images that would reduce the task time taken by doctors. Firstly, the discussion will focus directly and precisely on the results obtained (see chapter 4 and section 5.4). Secondly, the limitations, followed by a general discussion of this thesis' work will be presented. Finally, a set of future perspectives about my program will be proposed.

6.1 Performance

This section aims to discuss the results of the performance of the algorithm. These results are described in sections 4.1.3, 4.2.3, 5.4. For the first application about the segmentation of the femoral bone, the algorithm claims a mean DC of 0.868 and a mean JI of 0.768. Typically, in the literature, DC value in the 0.8 corresponds to good segmentation, while 0.9 is very good and it is considered excellent above 0.95 [74–76]. Since the JI is more punitive, values of JI starting from 0.75 are already considered as good [77, 78]. Therefore, for this application, the algorithm produces some promising results, although not excellent. This being said, it is important to remember that the segmentation of the femoral head is a challenging application. Fully automated methods, which generally produce better results, obtain DC values of 0.92 to 0.94 for the same application [79].

For the second application about brain pathologies, the algorithm claims results with a mean DC and a mean JI of respectively 0.834 and 0.747. However, there were some outliers and by removing them, the mean DC and JI become 0.9016 and 0.826. These results show that it can produce very good results for the majorities of pathology tested, but it does not work for every pathology. It depends mainly on the homogeneity of the region to be segmented, which is one of the biggest limitations of this algorithm. That was for the segmentation with the parameters found manually. The optimization of the parameters showed a little improvement throughout this second application with a mean DC of 0.879. However, when looking at the performance for all of the patients individually, the optimization considerably improved the results for the patient with initially bad performance. For the rest of the patients, there were no significant difference between the segmentation with the parameters found manually or via optimization.

6.2 Parameters and optimization

Now, the implementation of the different parameters and their effect will be discussed, followed by their possible optimization. The results can be found in sections 4.1.3, 4.2.3 and 5.4. There are three main parameters that control the behaviour of the algorithm during a segmentation; the Intensity Homogeneity, the Curvature Weight and the Inertia parameter. The other parameters have a limiting effect but do not influence the behaviour of the segmentation. It has been seen that the IH and In parameters have been well implemented and that their influence is the one expected. However, the CW parameter, which is supposed to control the smoothness of the contour, is not totally convincing. It has a constraining effect as expected but its influence on the smoothness of the contour is marginal. This is due to the fact that it considers the mean curvature over the whole contour. Moreover, the computation of this mean curvature is an approximation. It would be better if the curvature of the contour was computed on several precise points (the more, the better) along the contour, so that the constraining effect would be different at different points. It should have a more constraining effect at the places that have the biggest curvature. However, this would take a lot more computation power and time.

Regarding the optimization of the parameters, a simple solution is to perform a gradient descent to find the optimal parameters. This solution is simple, works well and has the advantage of only needing one example. Although the overall performance improved with the optimization with a mean DC of 0.879 versus 0.834 without, it has been shown that it only improved significantly the bad performance. The second observation is that there is more than one set of parameters that can produce very similar result, due to the effect of the three main parameters. Finally, the transfer of parameters between one instance to another is not very convincing. However, the examples used for this experiment were not optimal; although the patients had the same pathologies, the images were quite different in shape, size, resolution, etc. Still, it worked well for one example and for the other, it has been shown that it could be a good starting point to try to find the best parameters.

6.3 Time improvement

One objective of this thesis was to develop a segmentation tool that allows reducing the time taken by doctors in their everyday work. Section 4.2.3 of the results chapter has shown that the algorithm divided the drawing time by a mean factor of 6.996 for the brain application. This is a major improvement and it could reduce drastically the time

taken for segmentation by neurosurgeons. However, this result, although promising, is to be taken with caution. Indeed, it only relates to the drawing time. Yet, this is not the only action needed to perform a complete segmentation. First, there is the running time of the algorithm. Although it is quite long, it can be neglected since the user can do something else during this process. Moreover, this time is dependent on the machine on which it is running. But it would still be interesting to optimize the code to accelerate this process. Secondly, the time passed to choose the parameters is significant and needs to be taken into account. The optimization of the parameters for another example can give a pretty good initial guess but this optimization is quite time-consuming too (a couple of hours). However, once it is done for one example, the set of parameters can be used for all the future instances of this application. Finally, the most important one is the correction time. Indeed, for all segmentations of all applications, the user will still need to assess manually the quality of the segmentation and correct the few mistakes if needed. Unlike manual segmentation where the segmentation and validation are done at the same time.

6.4 Limitations

This section will present the three main limitations of this thesis work. The first one is about the algorithm itself and its performance. This algorithm was supposed to be as general as possible. However, it does not work for every application. The struggle concerns principally the homogeneity of the region to be segmented. This has always been a struggle for region-based active contour algorithms and this one does not overcome this issue. Also, the performance, although pretty good, does not compete with well-trained automated techniques. However, these automated techniques are used only for specific applications while this algorithm is more flexible.

The second principal limitation is about the time. Although the algorithm reduces significantly the drawing time, the rest still needs to be improved. Firstly, the running time of the algorithm is very long and needs to be optimized. The principal reason is that it is implemented in *python*. This language is very easy to use but is not at all optimized since it is an interpreted and dynamically typed language. The use of another language like *C* or *C++* should reduce drastically the running time. Secondly, choosing the parameters is also time-consuming. Choosing them manually requires the user to be trained with this method and requires several trials. The second option is to choose the parameters from another instance of the same application, after having optimized the parameters for this instance. Unfortunately, this optimization takes a lot of time (matters of hours). Finally, once the segmentation has been made, the user still needs to assess the results and perform manual corrections since the results produced are

generally good but not excellent.

The final big limitation is the GUI. While one of the objectives was to develop a well designed GUI to allow new users to easily take in hand this algorithm, the GUI is not very well implemented. This is principally due to a lack of time, but also to a lack of competence (this was the first time that I was implementing a GUI). After a neurosurgeon tried it, its feedback is that the GUI is not very intuitive and that the drawing tool is not pleasant to use [6].

6.5 Perspectives

The algorithm developed in this thesis shows promising results to be used as a segmentation tool for a wide range of applications. Moreover, it is directly destined to medical professionals and their daily work in hospitals. This is not the case for a lot of research in the segmentation field that stays as research projects. However, this work still needs further improvement in order to fulfill all the requirements. Key elements of potential improvements will be highlighted in the section below.

First of all, the algorithm can be improved even if it already claims very promising results. The first improvement would be to optimize fully the algorithm so that it takes less time. Reimplementing it in *C* or *C++* is a good starting point. The CW parameter can be reworked to have a more significant effect on the smoothness of the contour. That would really help to segment spherical versus irregular objects. Then, some new features could be added to the algorithm like combining this region-based active contour to an edge-based detection algorithm to help the segmentation to stop precisely at the edges. Another perspective would be to add some other medical imaging post-processing techniques. For example, adding a dilation and erosion process can be very useful. In image processing, dilation is the process of adding pixels to the boundaries of an object while erosion is removing pixels. These two techniques are useful to increase or decrease the boundaries of an object. Moreover, the combination of the two is also very interesting. Opening is the process of applying an erosion followed by a dilation. This eliminates the thin protrusions of the obtained image. Closing is the inverse process and eliminates the small holes from the obtained image [41]. These little post-processing techniques can be very useful for the user to easily and rapidly correct the segmented image.

Secondly, more experiments should be conducted on the transfer of parameters with the optimization to see if it is promising. It should be using a better dataset of images, to overcome the issues found in this thesis.

Last but not least, the GUI also needs to be improved to enhance the user-friendly interface. It should be composed of more intuitive buttons (with small icons for example) and allow for a better display of the image. Secondly, the drawing tool needs to be reimplemented to be more pleasant to use. Finally, it should also allow the possibility to use the post-processing techniques discussed just above.

The perspective of this work is to insert it in the workflow of an Active Learning segmentation scheme. This would involve a dynamic deep learning algorithm to segment the images with a certainty coefficient. If this coefficient is low, it means that the DL algorithm has difficulty to segment the image. Thus, it would ask the user to segment manually the image and it would learn new features from this trial. The objective is thus to offer to the user the semi-automatic tool developed in this work to help him accelerate this process.

Conclusion

Nowadays, segmentation is an important tool in the diagnosis and prognosis as well as for the treatment planning of medical disorders. With the increase in the use of volumetric medical imaging, especially CT and MRI, rapid advancements in the segmentation field has been observed over the past decade. In hospitals, the two principal departments that need segmentation are the neurosurgery and the radiotherapy departments. The first one uses currently mainly manual segmentation tools while the second one uses also AI-based and atlas-based segmentation for the easiest applications.

The literature review showed, in a first time, the different medical image acquisition methods that are the most commonly used. Then, some basic principles of segmentation have been explained such as the textural-based methods that are used in this thesis. Then, some precision about more advanced segmentation methods such as Deep Learning and Active Contour have been presented, with their advantages and limitations.

The choice of using a semi-automatic method has been made because it allows good interactions with the user. In this work, we developed a region-based active contour algorithm. The description of the regions is made by using a textural approach where the texture is described by robust statistics. A probabilistic model over the texture of the region built via a KDE is used to decide if a new area is part of the region. The contour evolution is described by a formula composed of an expanding term and a second term that assures the smoothness of the contour. The interactions between different contours are managed via a competition principle. Secondly, a GUI has been implemented to facilitate the interactions between the users and the algorithm. This GUI allows to load medical images, to draw on them, to initiate the segmentation algorithm and to save the results. Finally, the algorithm has been tested for two different applications; the segmentation of femur on CT scan and the segmentation of several brain pathologies on MRI.

The results have shown that the algorithm produces promising but not yet excellent results. However, the positive point of this algorithm is that it can be used for a variety

of applications over two different image acquisition methods compared to existing tools usually restricted to a limited range of applications. The parameters used to perform the segmentation behaves as expected, even if the CW parameter could be better implemented. The optimization allows to find optimal parameters to segment an image. However, the transfer of these parameters of another instance of the same application needs to be further investigated. This algorithm allows reducing the segmentation task time.

Finally, some perspectives for further development were also given. Firstly, the algorithm can be improved to increase the performance and optimized to reduce its running time. Then, some new post-processing methods, such as dilation and erosion, can be added to help the correction of the segmented image. Finally the GUI needs to be completely updated in order to be more user-friendly and pleasant to use.

In conclusion, a region-based active contour algorithm is a flexible segmentation tool that produces promising results. It allows to have good interactions with the user and, with a good GUI, can be used by doctors in their everyday work for a variety of applications. The tool can also be inserted in the workflow of an active learning segmentation scheme to further improve the segmentation process and provide the medical staff with a new efficient and robust way to label images.

Appendices

Appendix A

All results from the bone CT application

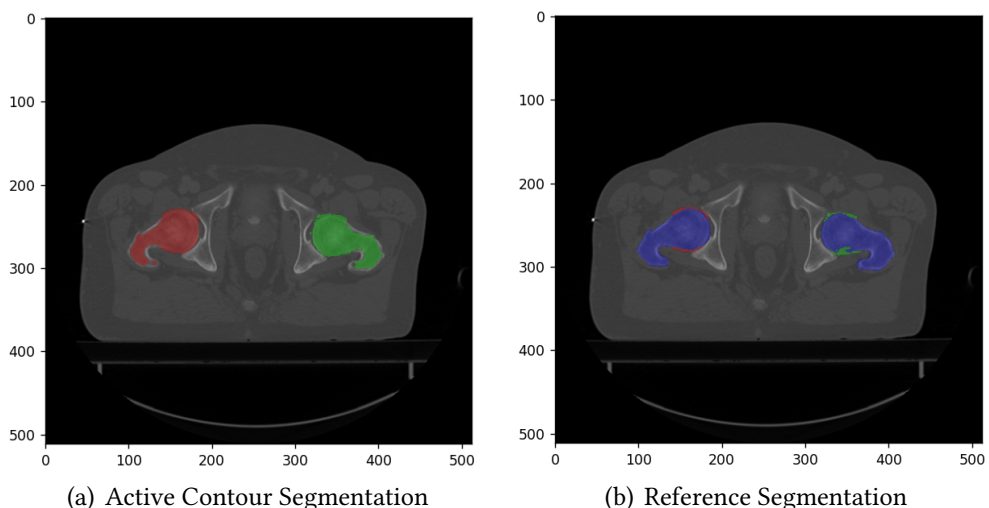


Figure A.1: Patient 4: comparison of the segmentation performed by the active contour algorithm (a) and with the manual segmentation done by an expert on top (b)

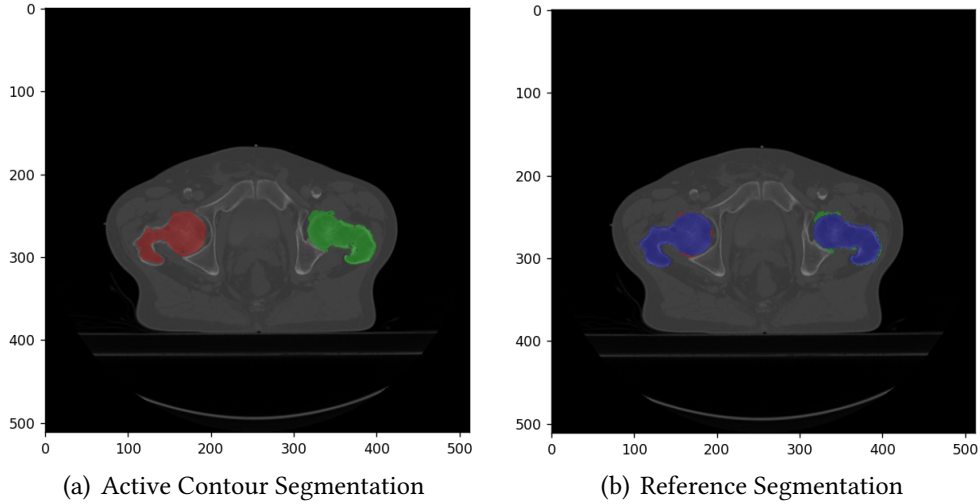


Figure A.2: Patient 6: comparison of the segmentation performed by the active contour algorithm (a) and with the manual segmentation done by an expert on top (b)

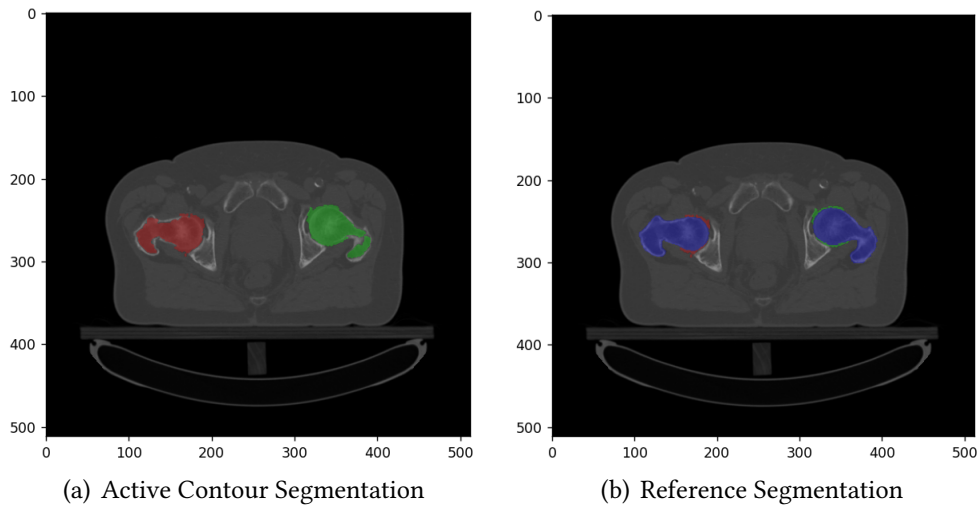


Figure A.3: Patient 9: comparison of the segmentation performed by the active contour algorithm (a) and with the manual segmentation done by an expert on top (b)

Appendix B

All the results from the MRI application

B.1 Performances

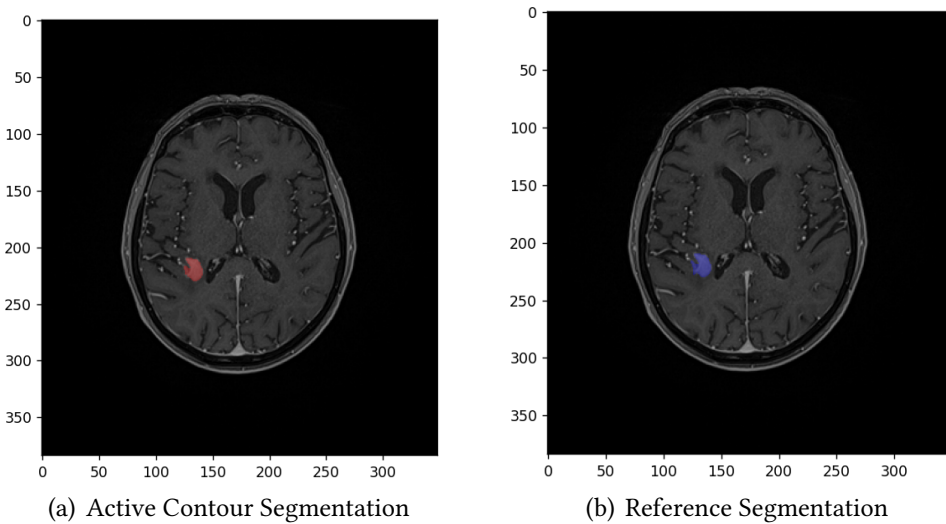


Figure B.1: Patient 1: comparison of the segmentation performed by the active contour algorithm (a) and with the manual segmentation done by a neurosurgeon on top (b)

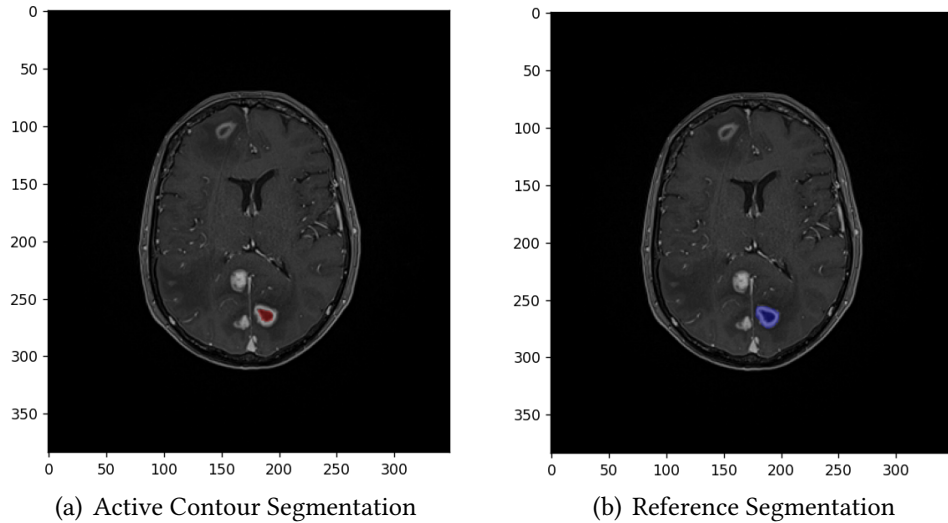


Figure B.2: Patient 2: comparison of the segmentation performed by the active contour algorithm (a) and with the manual segmentation done by a neurosurgeon on top (b)

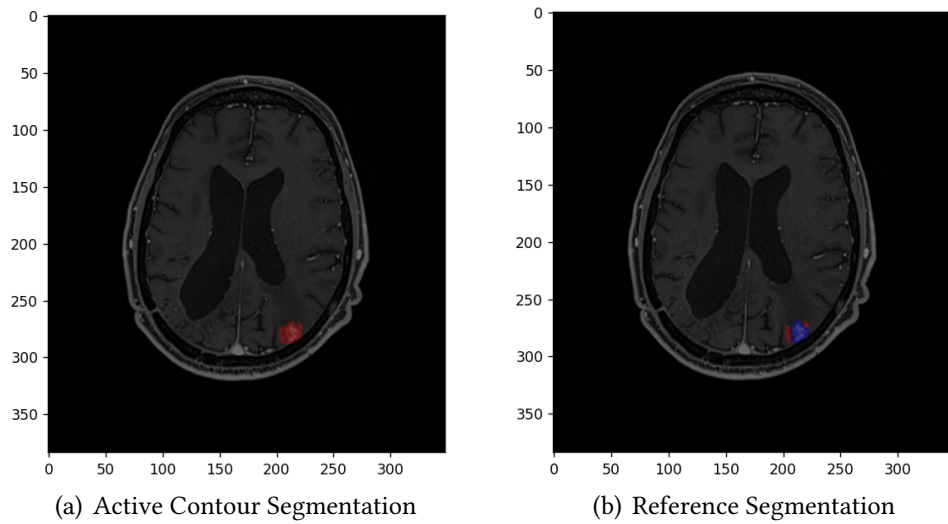


Figure B.3: Patient 4: comparison of the segmentation performed by the active contour algorithm (a) and with the manual segmentation done by a neurosurgeon on top (b)

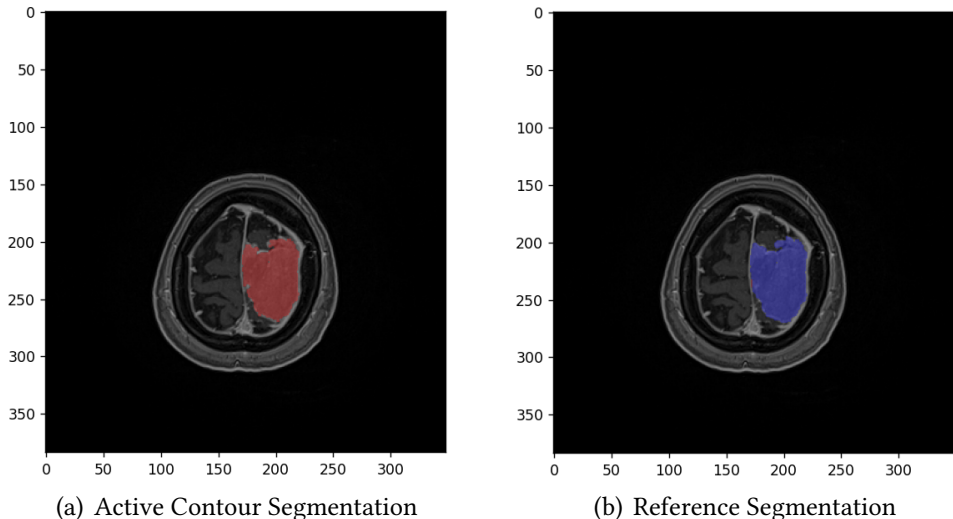


Figure B.4: Patient 5: comparison of the segmentation performed by the active contour algorithm (a) and with the manual segmentation done by a neurosurgeon on top (b)

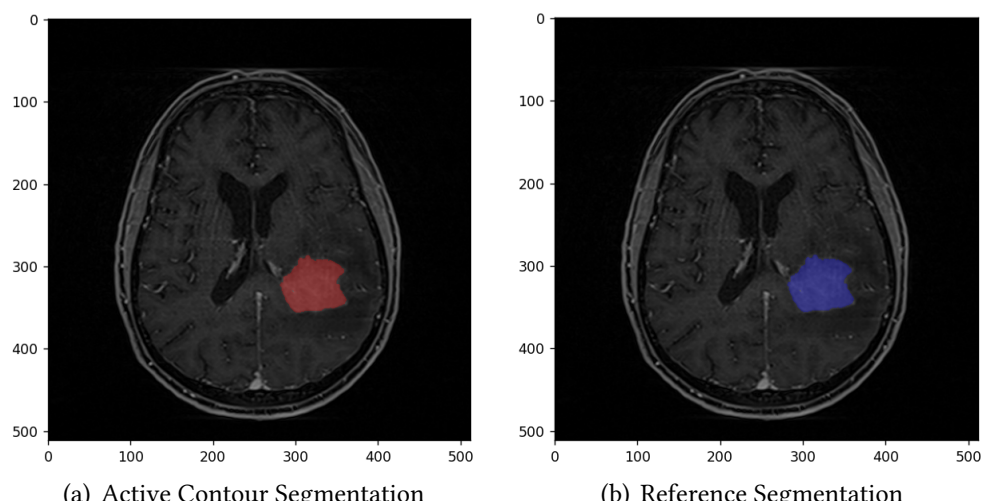


Figure B.5: Patient 6: comparison of the segmentation performed by the active contour algorithm (a) and with the manual segmentation done by a neurosurgeon on top (b)

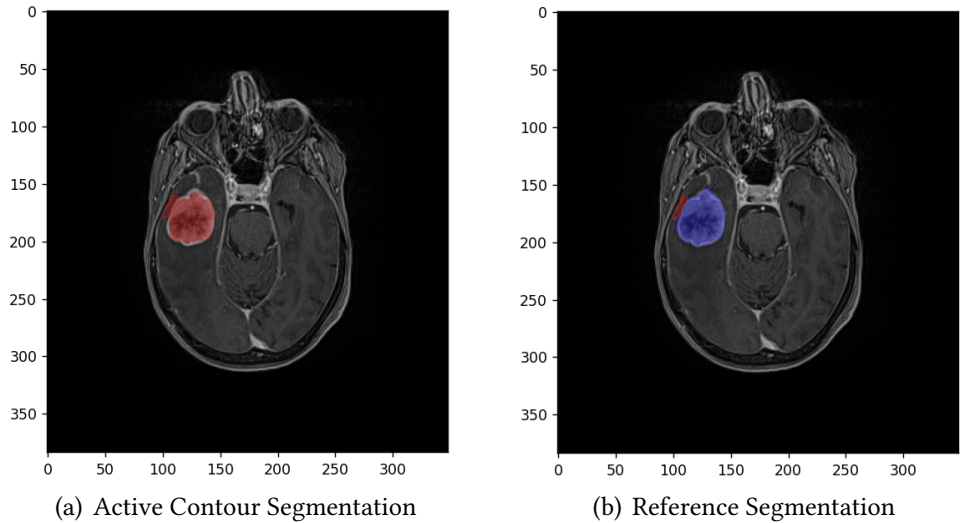


Figure B.6: Patient 7: comparison of the segmentation performed by the active contour algorithm (a) and with the manual segmentation done by a neurosurgeon on top (b)

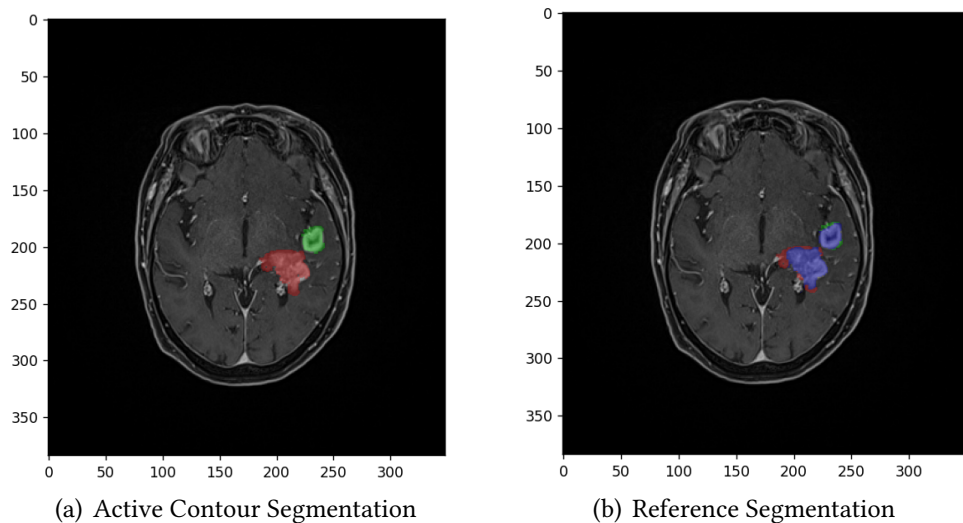


Figure B.7: Patient 10: comparison of the segmentation performed by the active contour algorithm (a) and with the manual segmentation done by a neurosurgeon on top (b)

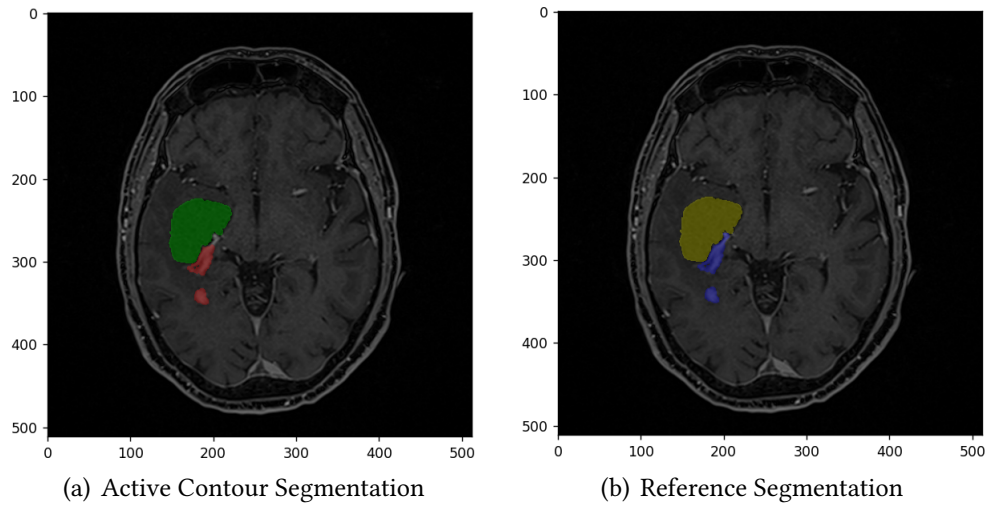


Figure B.8: Patient 13: comparison of the segmentation performed by the active contour algorithm (a) and with the manual segmentation done by a neurosurgeon on top (b)

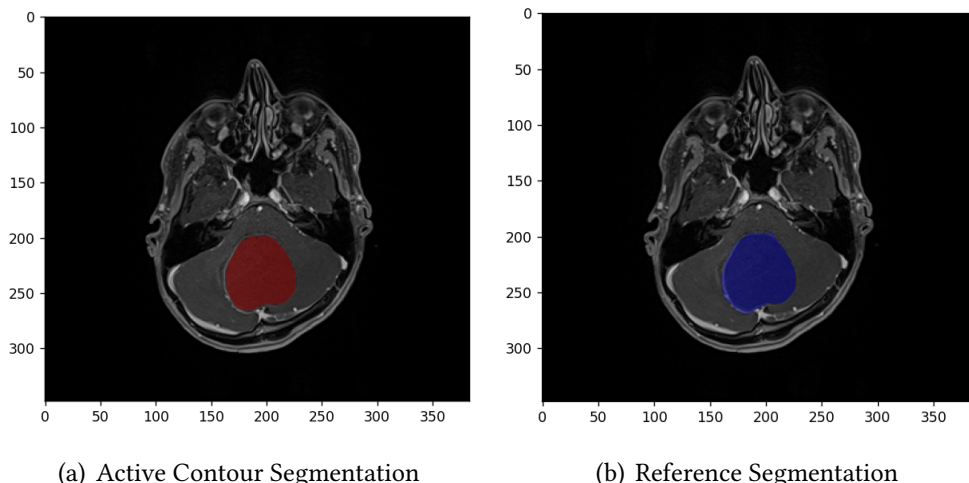


Figure B.9: Patient 14: comparison of the segmentation performed by the active contour algorithm (a) and with the manual segmentation done by a neurosurgeon on top (b)

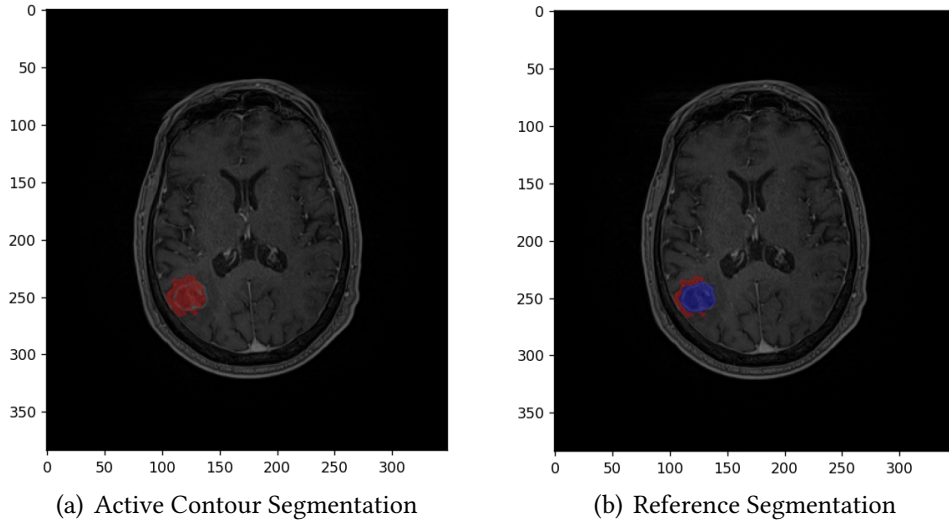


Figure B.10: Patient 15: comparison of the segmentation performed by the active contour algorithm (a) and with the manual segmentation done by a neurosurgeon on top (b)

B.2 Comparison of parameters

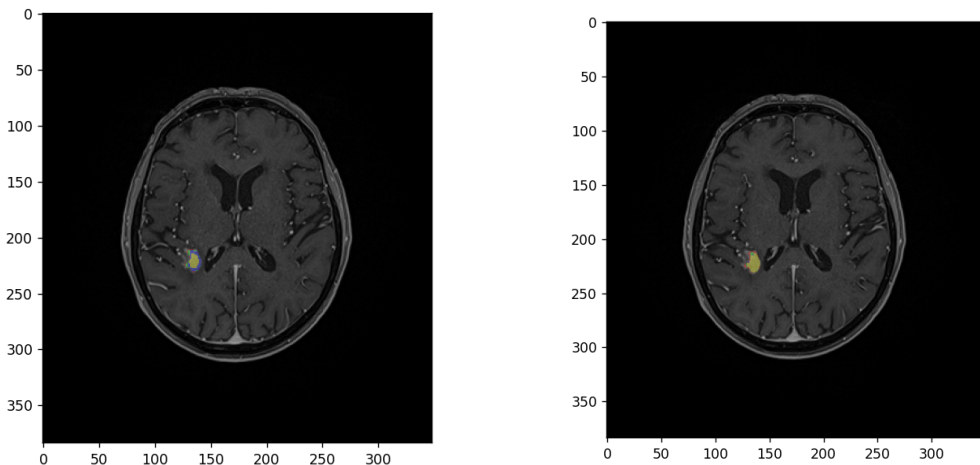


Figure B.11: Patient 1: Comparison of the Curvature Weight parameter; $CW=0.1$, $CW=0.2$, $CW=0.3$, $CW=0.4$

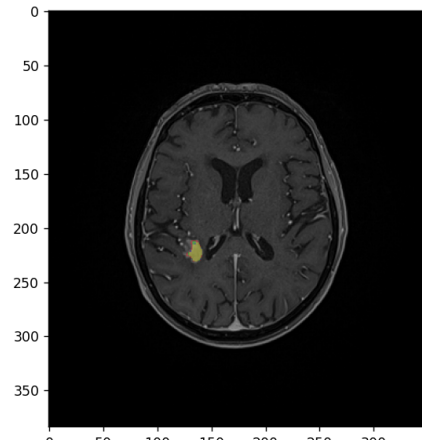


Figure B.12: Patient 1: Comparison of the Intensity Homogeneity parameter; $IH=0.1$, $IH=0.4$, $IH=0.6$, $IH=0.9$

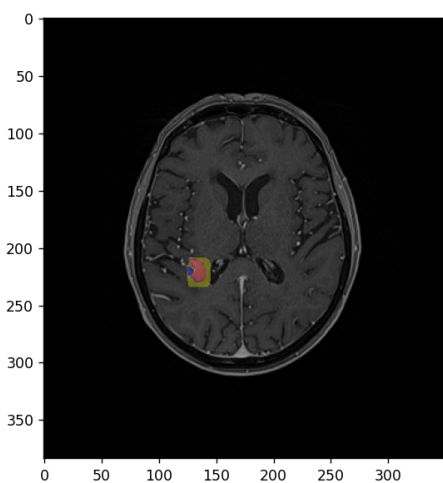


Figure B.13: Patient 1: Comparison of the Inertia parameter; $In=0.1$, $In=0.2$, $In=0.3$, $In=0.4$

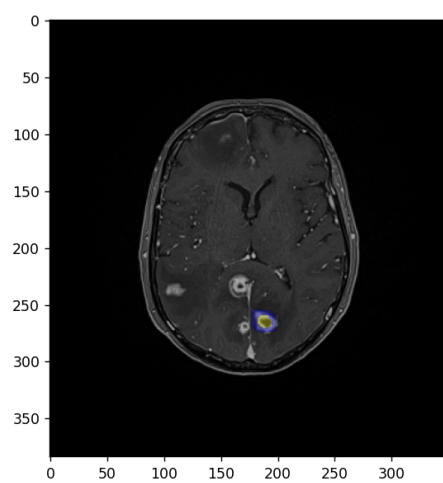


Figure B.14: Patient 2: Comparison of the Curvature Weight parameter; $CW=0.1$, $CW=0.2$, $CW=0.3$, $CW=0.4$

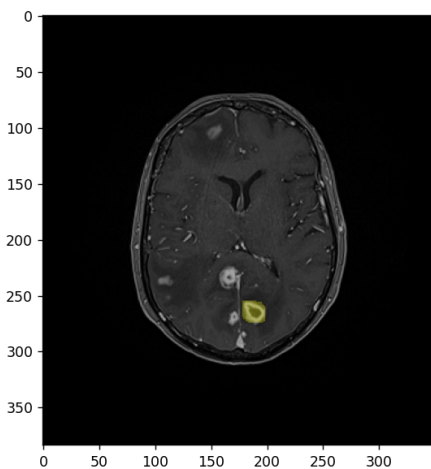


Figure B.15: Patient 2: Comparison of the Intensity Homogeneity parameter; $IH=0.1$, $IH=0.4$, $IH=0.6$, $IH=0.9$

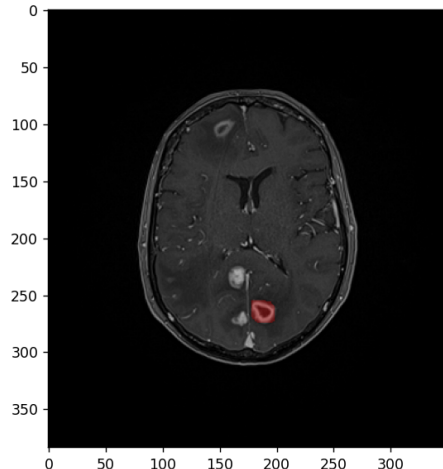


Figure B.16: Patient 2: Comparison of the Inertia parameter; $In=0.1$, $In=0.2$, $In=0.3$, $In=0.4$

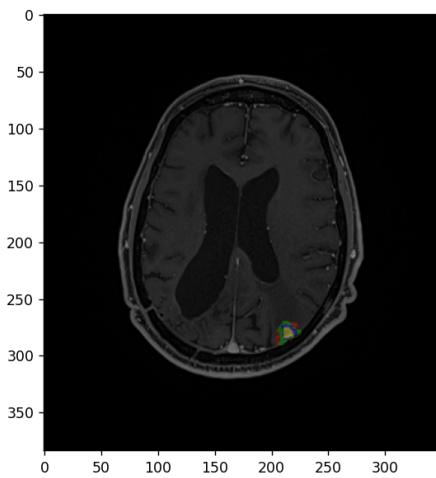


Figure B.17: Patient 4: Comparison of the Curvature Weight parameter; $CW=0.1$, $CW=0.2$, $CW=0.3$, $CW=0.4$

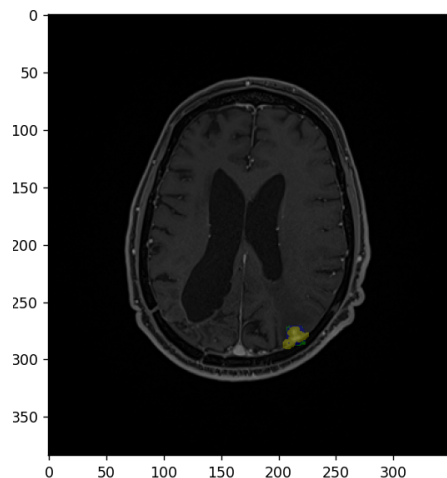


Figure B.18: Patient 4: Comparison of the Intensity Homogeneity parameter; $IH=0.1$, $IH=0.4$, $IH=0.6$, $IH=0.9$

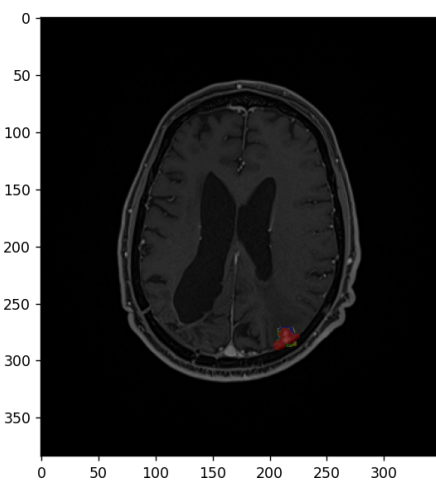


Figure B.19: Patient 4: Comparison of the Inertia parameter; $In=0.1$, $In=0.2$, $In=0.3$, $In=0.4$

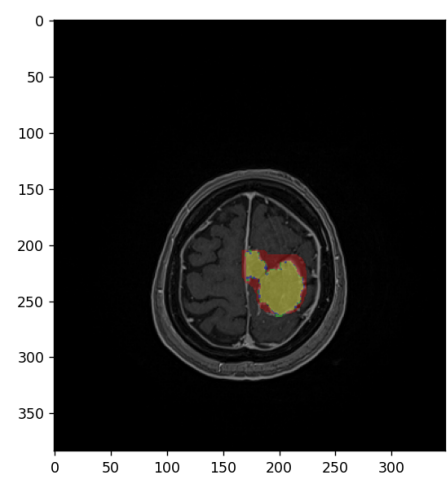


Figure B.20: Patient 5: Comparison of the Curvature Weight parameter; $CW=0.1$, $CW=0.2$, $CW=0.3$, $CW=0.4$

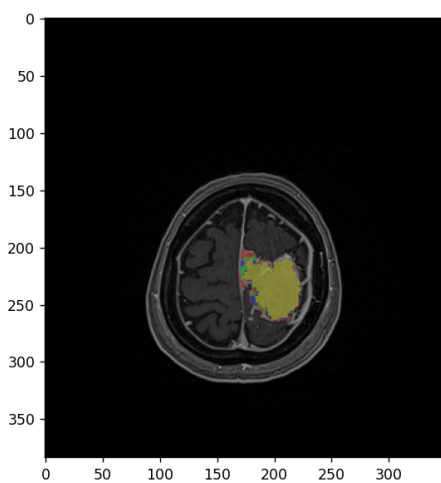


Figure B.21: Patient 5: Comparison of the Intensity Homogeneity parameter; $\text{IH}=0.1$, $\text{IH}=0.4$, $\text{IH}=0.6$, $\text{IH}=0.9$

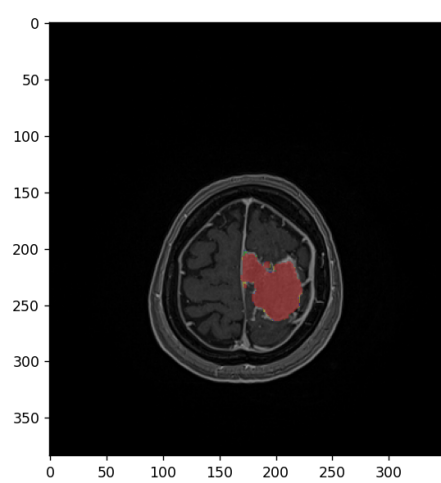


Figure B.22: Patient 5: Comparison of the Inertia parameter; $\text{In}=0.1$, $\text{In}=0.2$, $\text{In}=0.3$, $\text{In}=0.4$

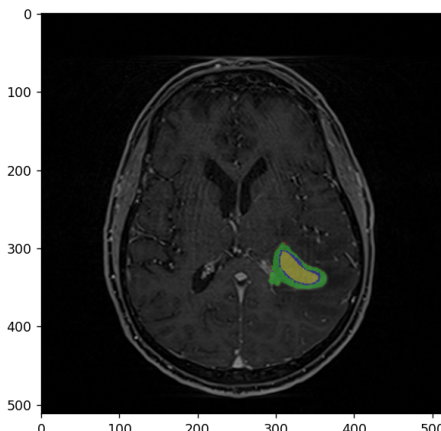


Figure B.23: Patient 6: Comparison of the Curvature Weight parameter; $\text{CW}=0.1$, $\text{CW}=0.2$, $\text{CW}=0.3$, $\text{CW}=0.4$

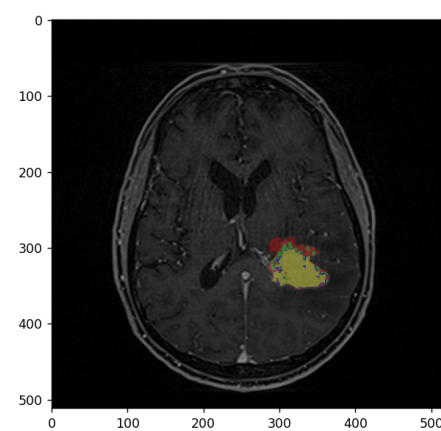


Figure B.24: Patient 6: Comparison of the Intensity Homogeneity parameter; $\text{IH}=0.1$, $\text{IH}=0.4$, $\text{IH}=0.6$, $\text{IH}=0.9$

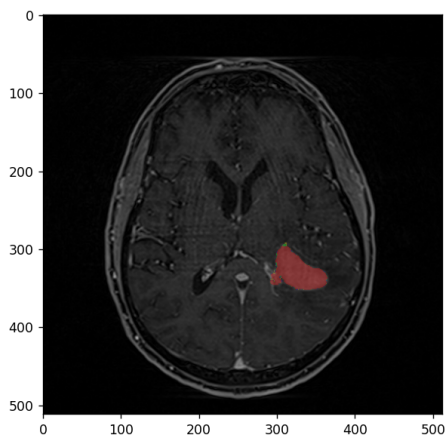


Figure B.25: Patient 6: Comparison of the Inertia parameter; $In=0.1$, $In=0.2$, $In=0.3$, $In=0.4$

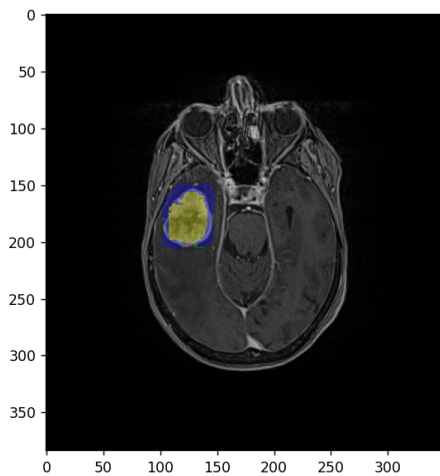


Figure B.26: Patient 7: Comparison of the Curvature Weight parameter; $CW=0.1$, $CW=0.2$, $CW=0.3$, $CW=0.4$

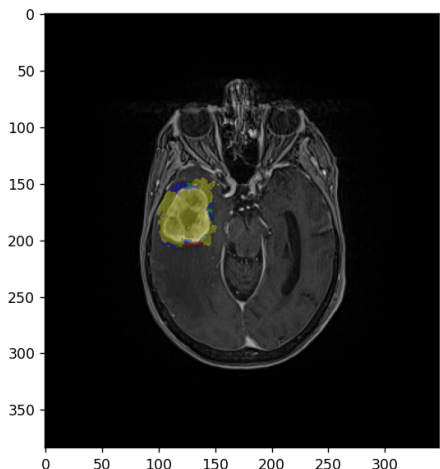


Figure B.27: Patient 7: Comparison of the Intensity Homogeneity parameter; $IH=0.1$, $IH=0.4$, $IH=0.6$, $IH=0.9$

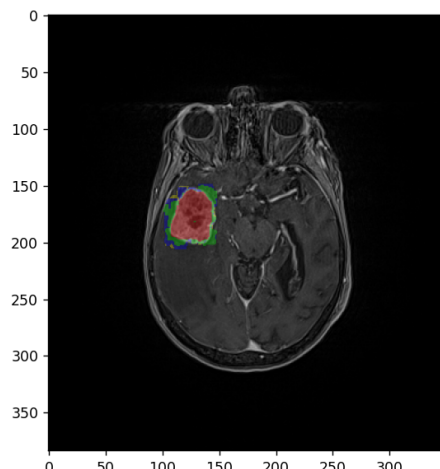


Figure B.28: Patient 7: Comparison of the Inertia parameter; $In=0.1$, $In=0.2$, $In=0.3$, $In=0.4$

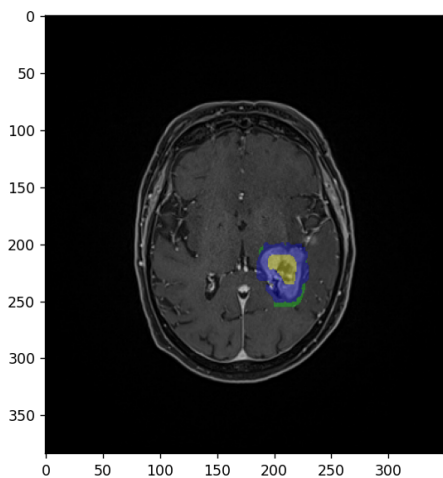


Figure B.29: Patient 10 (big): Comparison of the Curvature Weight parameter; CW=0.1, CW=0.2, CW=0.3, CW=0.4

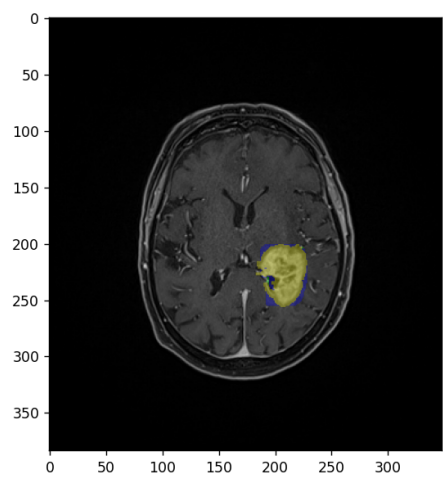


Figure B.30: Patient 10 (big): Comparison of the Intensity Homogeneity parameter; IH=0.1, IH=0.4, IH=0.6, IH=0.9

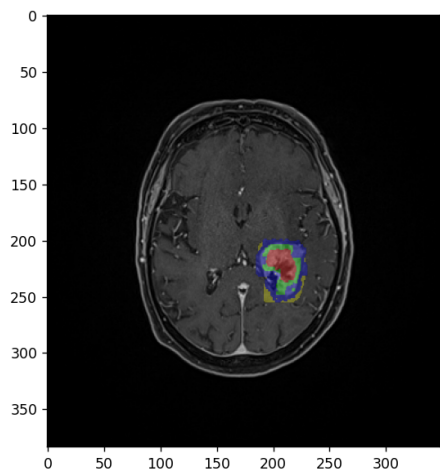


Figure B.31: Patient 10 (big): Comparison of the Inertia parameter; In=0.1, In=0.2, In=0.3, In=0.4

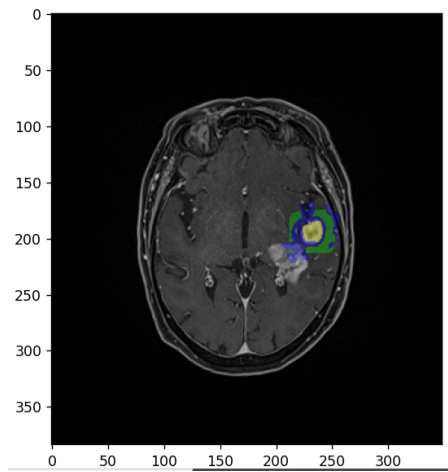


Figure B.32: Patient 10 (small): Comparison of the Curvature Weight parameter; CW=0.1, CW=0.2, CW=0.3, CW=0.4

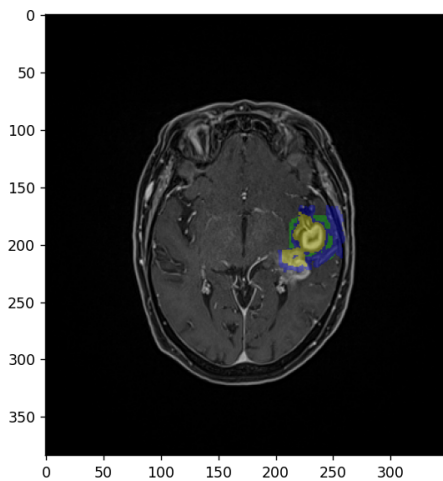


Figure B.33: Patient 10 (small): Comparison of the Intensity Homogeneity parameter; $\text{IH}=0.1, \text{IH}=0.4, \text{IH}=0.6, \text{IH}=0.9$

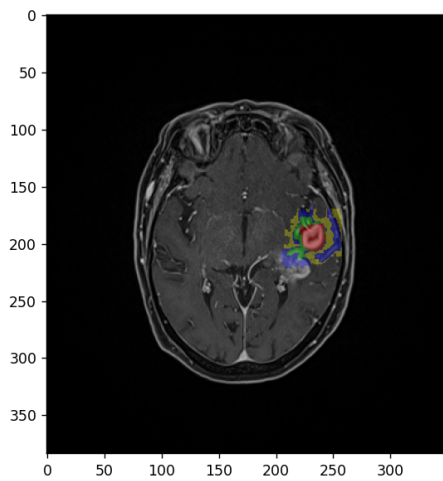


Figure B.34: Patient 10 (small): Comparison of the Inertia parameter; $\text{In}=0.1, \text{In}=0.2, \text{In}=0.3, \text{In}=0.4$

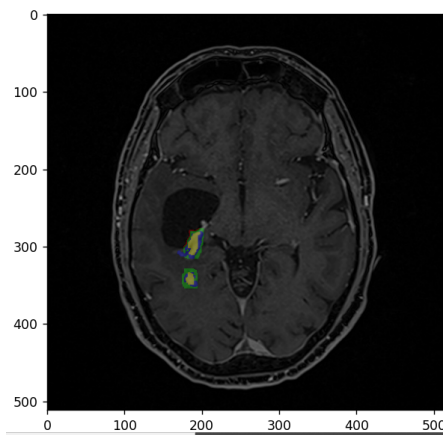


Figure B.35: Patient 13 (lymph): Comparison of the Curvature Weight parameter; $\text{CW}=0.1, \text{CW}=0.2, \text{CW}=0.3, \text{CW}=0.4$

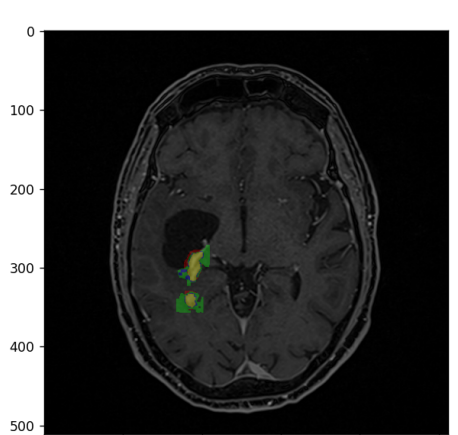


Figure B.36: Patient 13 (lymph): Comparison of the Intensity Homogeneity parameter; $\text{IH}=0.1, \text{IH}=0.4, \text{IH}=0.6, \text{IH}=0.9$

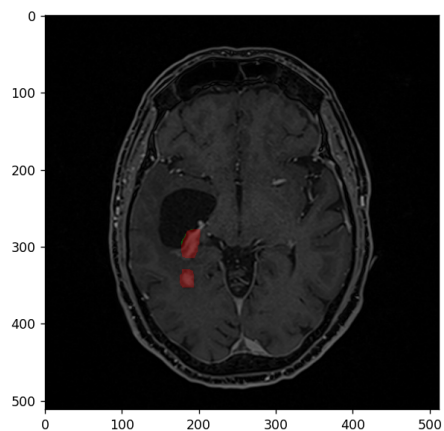


Figure B.37: Patient 13 (lymph): Comparison of the Inertia parameter; $In=0.1$, $In=0.2$, $In=0.3$, $In=0.4$

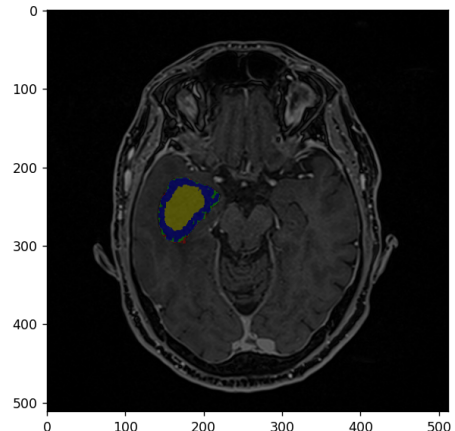


Figure B.38: Patient 13 (vent): Comparison of the Curvature Weight parameter; $CW=0.1$, $CW=0.2$, $CW=0.3$, $CW=0.4$

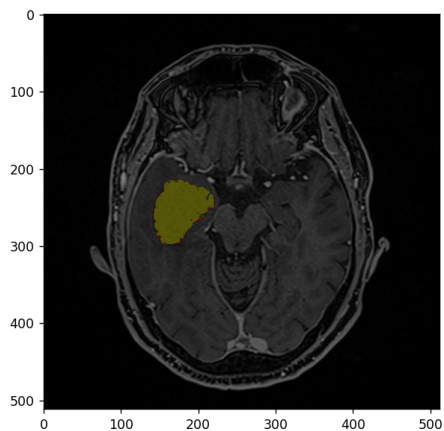


Figure B.39: Patient 13 (vent): Comparison of the Intensity Homogeneity parameter; $IH=0.1$, $IH=0.4$, $IH=0.6$, $IH=0.9$

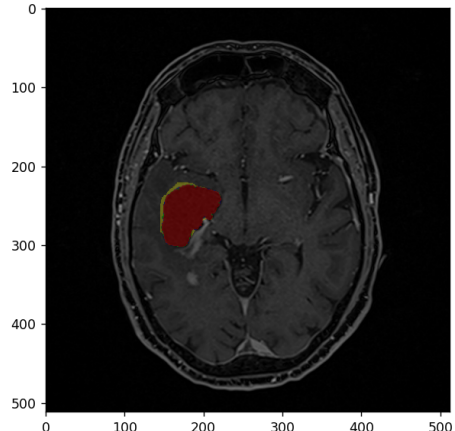


Figure B.40: Patient 13 (vent): Comparison of the Inertia parameter; $In=0.1$, $In=0.2$, $In=0.3$, $In=0.4$

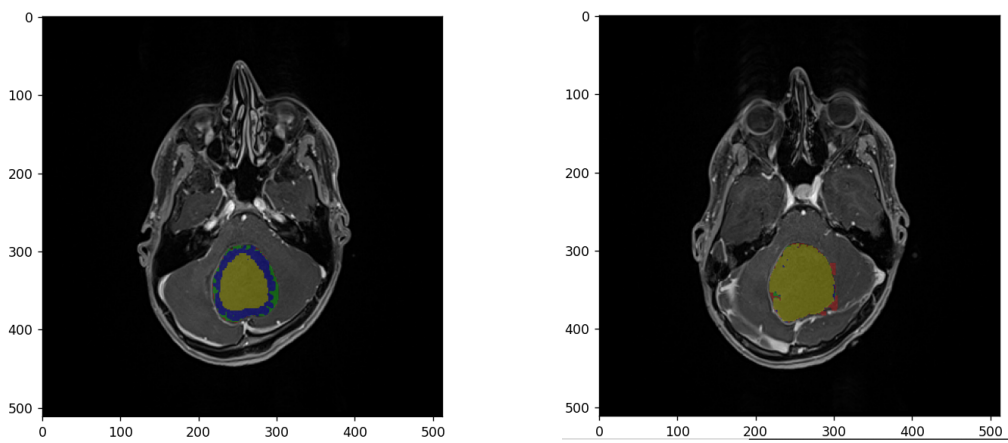


Figure B.41: Patient 14: Comparison of the Curvature Weight parameter; $CW=0.1$, $CW=0.2$, $CW=0.3$, $CW=0.4$

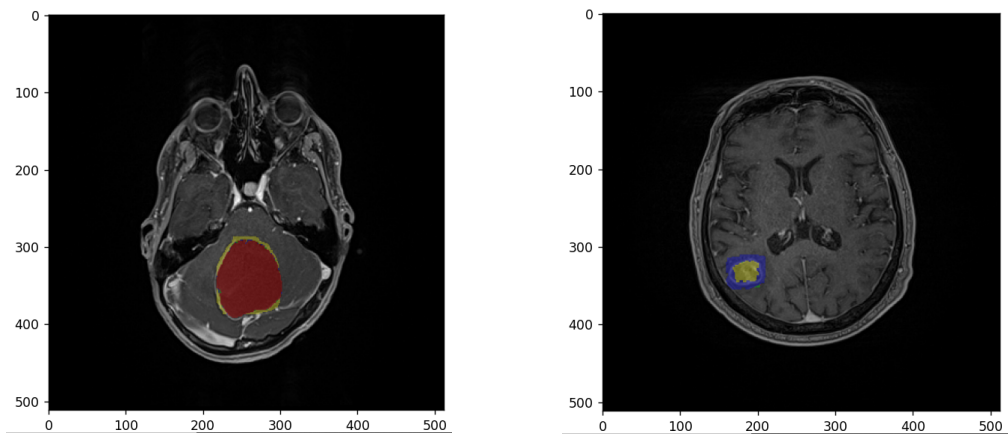


Figure B.42: Patient 14: Comparison of the Intensity Homogeneity parameter; $IH=0.1$, $IH=0.4$, $IH=0.6$, $IH=0.9$



Figure B.43: Patient 14: Comparison of the Inertia parameter; $In=0.1$, $In=0.2$, $In=0.3$, $In=0.4$



Figure B.44: Patient 15: Comparison of the Curvature Weight parameter; $CW=0.1$, $CW=0.2$, $CW=0.3$, $CW=0.4$

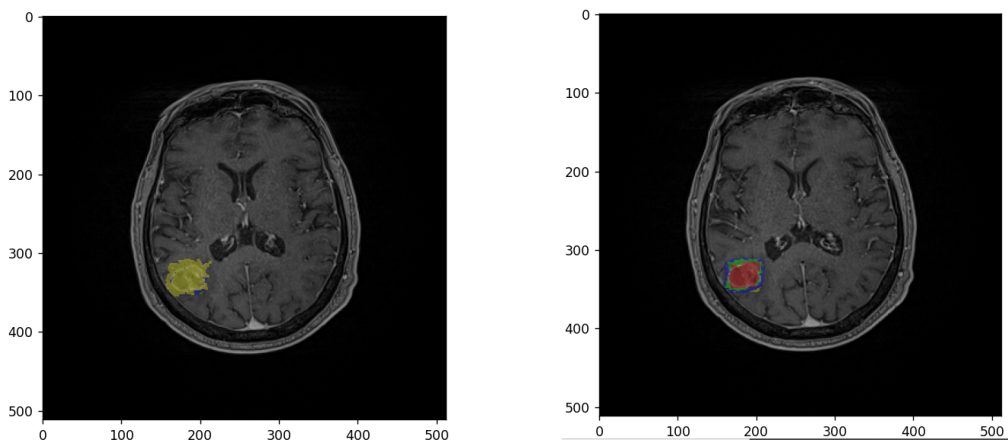


Figure B.45: Patient 15: Comparison of the Intensity Homogeneity parameter; $IH=0.1$, $IH=0.4$, $IH=0.6$, $IH=0.9$

Figure B.46: Patient 15: Comparison of the Inertia parameter; $In=0.1$, $In=0.2$, $In=0.3$, $In=0.4$

Appendix C

All results for the optimization of parameters

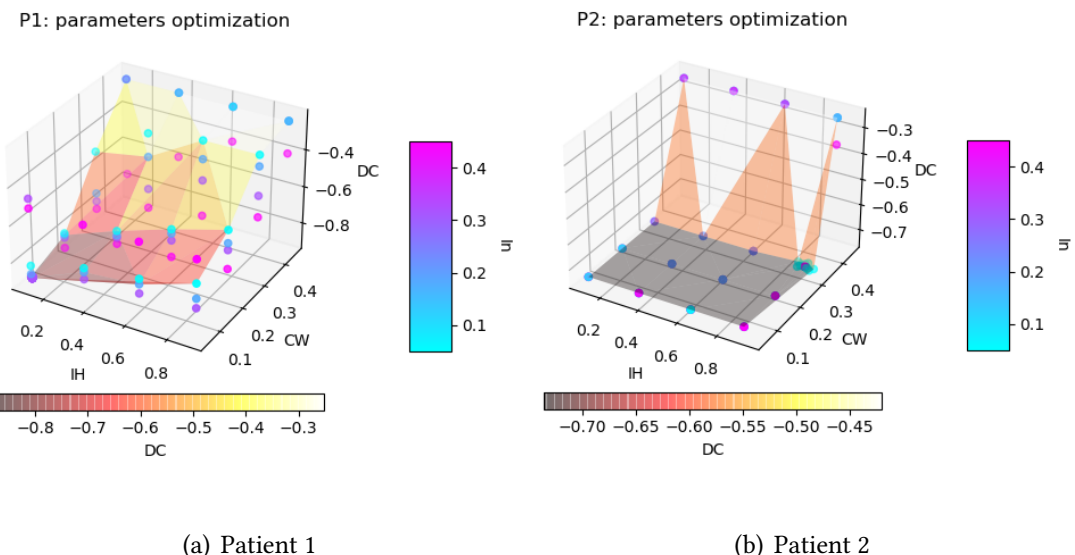
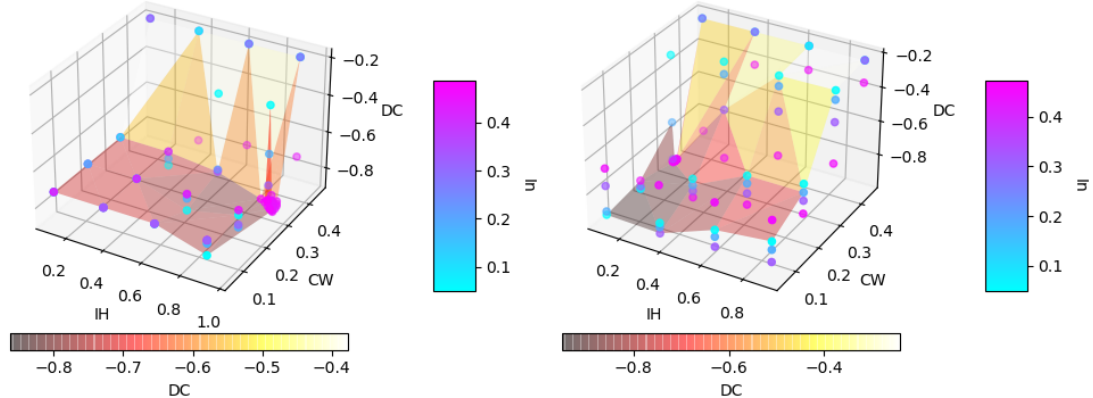


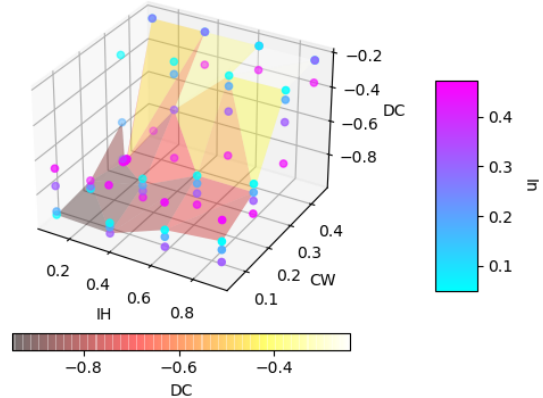
Figure C.1: Optimization of parameters. Each point corresponds to one evaluation of the objective function for a set of parameters. The color of each point corresponds to the I_H parameter used. The surface corresponds to the objective function to minimize

P4: parameters optimization



(a) Patient 4

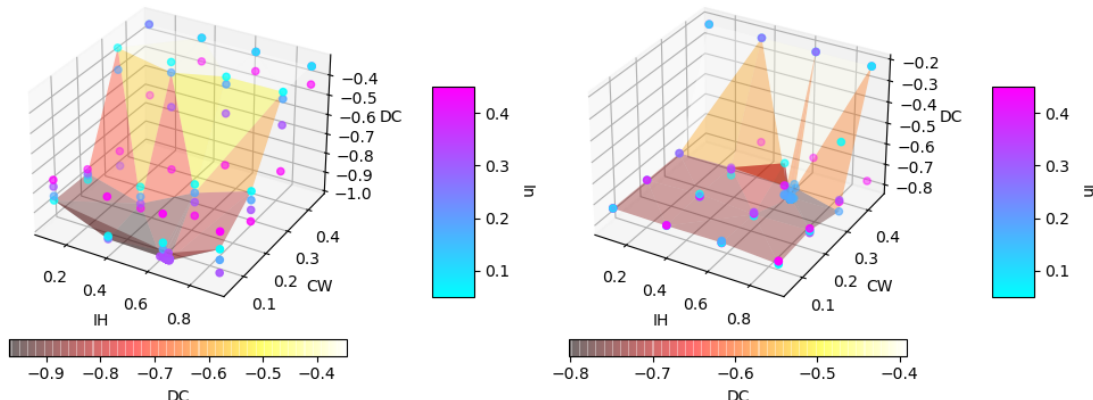
P5: parameters optimization



(b) Patient 5

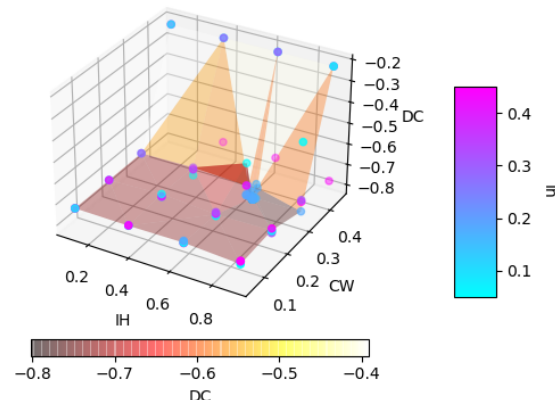
Figure C.2: Optimization of parameters. Each point corresponds to one evaluation of the objective function for a set of parameters. The color of each point corresponds to the In parameter used. The surface corresponds to the objective function to minimize

P6: parameters optimization



(a) Patient 6

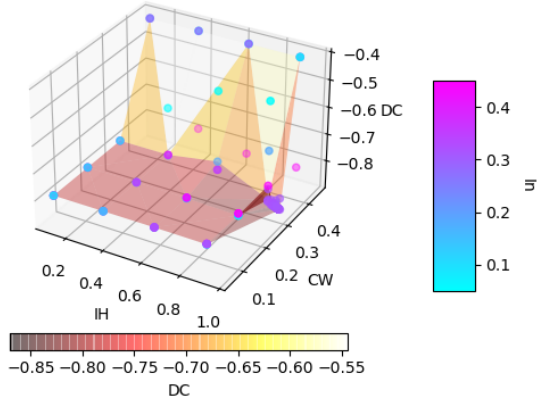
P7: parameters optimization



(b) Patient 7

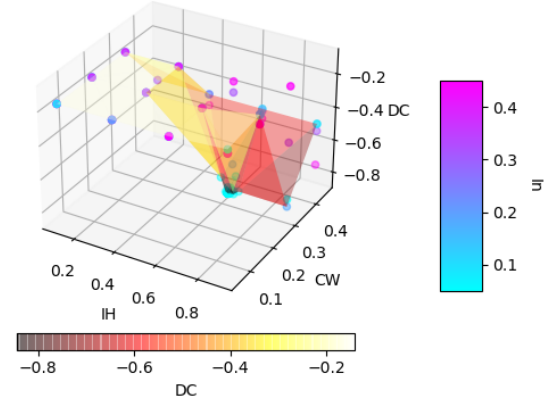
Figure C.3: Optimization of parameters. Each point corresponds to one evaluation of the objective function for a set of parameters. The color of each point corresponds to the In parameter used. The surface corresponds to the objective function to minimize

P10_big: parameters optimization



(a) Patient 10 (big)

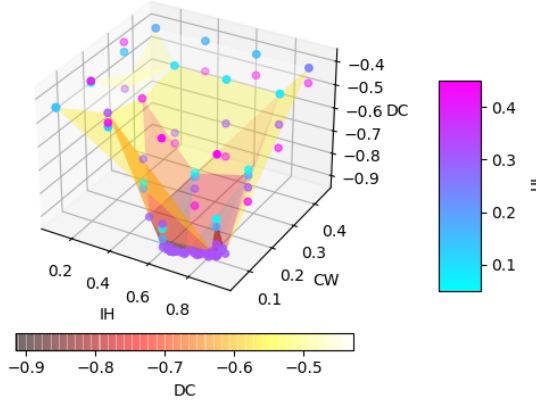
P10_small: parameters optimization



(b) Patient 10 (small)

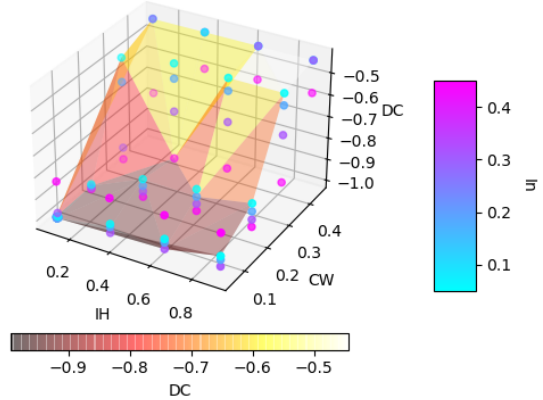
Figure C.4: Optimization of parameters. Each point corresponds to one evaluation of the objective function for a set of parameters. The color of each point corresponds to the In parameter used. The surface corresponds to the objective function to minimize

P13_tum: parameters optimization



(a) Patient 13 (lymph)

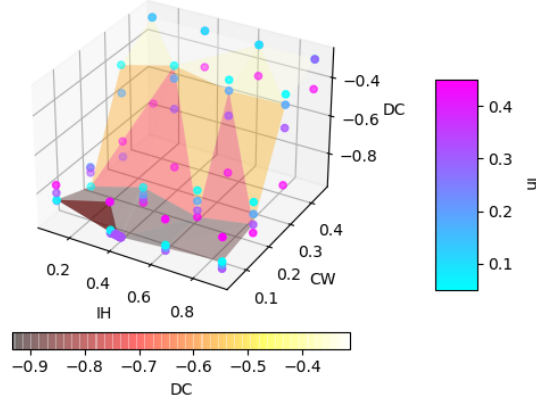
P13_vent: parameters optimization



(b) Patient 13 (vent)

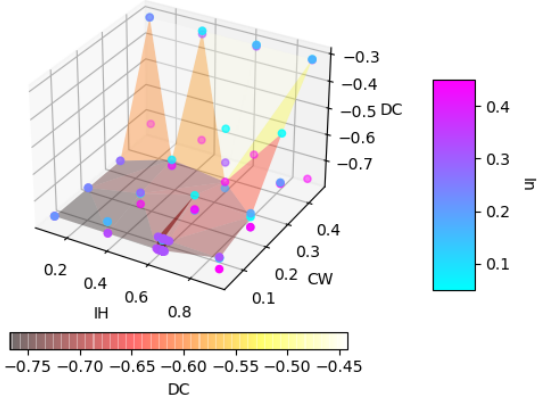
Figure C.5: Optimization of parameters. Each point corresponds to one evaluation of the objective function for a set of parameters. The color of each point corresponds to the In parameter used. The surface corresponds to the objective function to minimize

P14: parameters optimization



(a) Patient 14 (tum)

P15: parameters optimization



(b) Patient 15 (vent)

Figure C.6: Optimization of parameters. Each point corresponds to one evaluation of the objective function for a set of parameters. The color of each point corresponds to the In parameter used. The surface corresponds to the objective function to minimize

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