Chapter 11
Automatic Multiorgan Segmentation Using Hierarchically Registered Probabilistic Atlases

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Abstract We propose a generic method for the automatic multiple-organ segmentation of 3D images based on a multilabel graph cut optimization approach which uses location likelihood of organs and prior information of spatial relationships between them. The latter is derived from shortest-path constraints defined on the adjacency graph of structures and the former is defined by probabilistic atlases learned from a training dataset. Organ atlases are mapped to the image by a fast (2+1)D hierarchical registration method based on SURF keypoints. Registered atlases are also used to derive organ intensity likelihoods. Prior and likelihood models are then introduced in a joint centroidal Voronoi image clustering and graph cut multiobject segmentation framework. Qualitative and quantitative evaluation has been performed on contrast-enhanced CT and MR images from the VISCERAL dataset.

11.1 Introduction and Related Work

Clinical practice today, especially whole-body CT and MR imaging, often generates large numbers of high-resolution images, which makes tasks of efficient data access, transfer, analysis and visualization challenging. This is especially true in distributed...
computing environments which have seen a growing use of hand-held terminals for interactive data access and visualization of anatomy. Therefore, there is great interest in efficient and robust medical image segmentation algorithms for the purposes of creating patient-specific anatomical models, clinical applications, medical research and education, and visualization and semantic navigation of full-body anatomy [3, 26].

Traditionally, single-object- or pathology-oriented, recent image processing methods [9, 12, 14, 15, 19, 23, 25, 27] have made the analysis and the segmentation of multiple anatomical structures increasingly possible. However, CT and MR images have intrinsic characteristics that render their automatic segmentation challenging. They are commonly degraded by various noise sources and artefacts due to limited acquisition time and resolution, and patient motion which all reduce the prominence of intensity edges in images. In addition, MR images suffer from spatial distortion of tissue intensity due to main magnetic field inhomogeneity. Regardless of the imaging modality and related artefacts, many anatomically and functionally distinct structures, especially those corresponding to soft tissues, have similar intensity levels in images and, furthermore, blend into surrounding tissues which have intensities close to their own. It is impossible to identify and segment such structures automatically on the basis of intensity information only. Hence, most advanced segmentation methods exploit some form of prior information on structure location [12, 19, 27] or interrelations [9, 14, 23, 25] to achieve greater robustness and precision. Hierarchical approaches to segmentation [23, 25, 32] rely on hierarchical organizations of prior information and algorithms that proceed in a coarse-to-fine manner according to anatomical level of detail.

Graph cut methods, which have been widely applied to single-object segmentation problems [4], rely on a maximum-flow binary optimization scheme of a discrete cost function on the image graph. For a particular class of cost functions which frequently arises in segmentation applications [16], these methods produce provably good approximate solutions in multiobject [5] and global optima in single-object segmentation. In addition, simultaneous multiobject segmentation approaches are superior to their sequential counterparts in that they raise questions neither on the best segmentation sequence to follow nor on how to avoid the propagation of errors of individual segmentations [9].

While widely used by the computer vision community, keypoint-based image description and matching methods, such as SIFT [20] and SURF [2], have found relatively few application proposals in medical image processing. These methods proceed by first detecting some points of interest (edges, ridges, blobs, etc.) within the image, then compute vectors describing local neighbourhoods around these points and use them as content descriptors. The approach has been successfully applied to image indexing, content-based image retrieval, object detection and recognition, and image registration [30]. In medical imaging, 3D versions of SIFT have been used in brain MR image matching [6], linear registration of radiation therapy data [1], and nonlinear (deformable) registration of thoracic CT [31] and brain MR [22] images. A SURF-based method [10] has also been successfully applied to the intermodality
registration of 2D brain images. A review of keypoint-based medical image registration can be found in [28].

We propose a generic method for the automatic multiple-organ segmentation of 3D images based on multilabel graph cut optimization which uses location and intensity likelihoods of organs and prior information of their spatial configuration. The spatial prior is derived from shortest-path pairwise constraints defined on the adjacency graph of structures [14], and the organ location likelihood is defined by probabilistic atlases [24] learned from the VISCERAL training dataset [11]. We register organ atlases to the image prior to segmentation using a fast (2+1)D registration method based on SURF keypoints. Registered atlases are also used to derive organ intensity likelihoods. Prior and likelihood models are then introduced in a joint centroidal Voronoi image clustering and graph cut multiobject segmentation framework. We present the results of qualitative and quantitative evaluation of our method on contrast-enhanced CT and MR images from the VISCERAL dataset.

### 11.2 Methods

In the following, we present the different elements of our approach in detail, namely the keypoint-based image registration method and its use in organ atlas construction as well as its hierarchical application in segmentation. Image clustering and segmentation methods are detailed next, followed by a presentation of evaluation results in the subsequent section.

#### 11.2.1 SURF Keypoint-Based Image Registration

We first outline our fast (2+1)D algorithm for the rigid registration of 3D medical images using content features. Our method has the following properties:

- Features are extracted in 2D volume slices. This has the advantage of being fast and easily parallelizable. Another advantage is that medical data are usually stored in a picture archiving and communication system (PACS) in the form of volume slices as opposed to full 3D volumes. Our method easily fits into such medical environments. Note that while feature extraction is done in 2D images, registration is still performed in 3D, hence the (2+1)D definition.
- Partial matching is well handled, thus making our algorithm suitable for general medical data.
- Total processing time is on the order of seconds.
- The (2+1)D paradigm currently restricts our method to image volumes with consistent orientations. A pair of images featuring patients with orthogonal orientations cannot be registered for now.
11.2.1.1 2D Feature Extraction and Matching

As previously mentioned, we extract features from 2D slices of the image volume. We currently use the SURF image descriptor [21]; however, our method is generic and would work with other descriptors as well. To reduce computation time, we first downsample the input volume to a user-specified size. As a rule of thumb, we isotropically resample each volume so that its second longest dimension is equal to the desired resolution $R$. For example, with $R = 100$, the VISCERAL training dataset image 10000108_1_CTce_ThAb of dimensions $512 \times 512 \times 468$ and spacing $0.7 \text{mm} \times 0.7 \text{mm} \times 1.5 \text{mm}$ is resampled to a $100 \times 100 \times 198$ volume with an isotropic spacing of $3.54 \text{mm}$.

Next, we extract 2D SURF features from each slice. As these operations are completely independent, this step is carried out in a parallel manner. Figure 11.1 shows feature extraction results on a pair of axial slices from VISCERAL training dataset images 10000108_1_CTce_ThAb (left) and 10000109_1_CTce_ThAb (right). The top row shows all features extracted from these slices. The number of extracted features is 11500 and 9400, respectively.

Once all features are extracted, they are matched using the widely used second closest ratio criterion [20]. If the first volume $V_1$ contains $n_1$ slices and the second volume $V_2$ contains $n_2$ slices, we have to compute $n_1 \times n_2$ image matches, which again is easily carried out in a parallel fashion. The bottom row of Fig. 11.1 shows the nine matching couples (pairs of keypoints) found in both slices.

The output of the matching step is a similarity matrix $S$ of $n_1 \times n_2$ of, possibly empty, matching couple sets $H_{i,j}$. Figure 11.2 illustrates the 2D matching procedure on test images 10000108_1_CTce_ThAb and 10000109_1_CTce_ThAb. Figure 11.3 shows similarity matrices reflecting the number of matching couples between any pair of slices for three settings of downsampling resolution $R$ (grey level is inversely proportional to the number of matching couples). Matrix diagonals are clearly visible, confirming the fact that input volumes contain similar structures. In total, 2561 matching couples were found between this pair of volumes with $R = 100$.

11.2.1.2 (2+1)D Registration

Once 2D matches are found, we are able to proceed with volume registration. For robustness purposes, we use a simple “scale + translation” transformation model:

\[
\begin{bmatrix}
x' \\
y' \\
z'
\end{bmatrix} = s \begin{bmatrix} x \\ y \\ z \end{bmatrix} + \begin{bmatrix} t_x \\ t_y \\ t_z \end{bmatrix}.
\] (11.1)

We estimate the four parameters $s$, $t_x$, $t_y$ and $t_z$ in similar spirit to the RANSAC method [8], using the set of matching couples between the slices of the pair of volumes, computed as indicated in the previous subsection. RANSAC is an iterative
parametric model estimation method known to be very efficient in the presence of outliers. One RANSAC iteration usually consists in randomly picking a small number of samples to estimate the model parameters, then counting the number of data samples consistent with the model, rejecting outliers. After performing all iterations, the model providing the highest number of consistent data samples is kept as the solution. In our case, we carry out parameter estimation in a two-stage fashion; first, we fix the parameters $s$ and $t_z$ which allow us to work only on a subset of $S$, then we estimate the remaining parameters $t_x$ and $t_y$. More specifically, we carry out the $n^{th}$ RANSAC iteration as follows:
Fig. 11.2  Feature matching. The two input volumes (a, b) are sliced (c, d), and each slice from the first volume is compared against every slice from the second volume.

Fig. 11.3  Similarity matrices of test volumes 10000108_1_CTce_ThAb and 10000109_1_CTce_ThAb for different values of downsampling resolution $R$.

- Randomly pick a line $L_n$ crossing $S$. This fixes half of the transform parameters, that is, the parameters $s$ and $t_z$. The top row of Fig. 11.4 shows three different randomly picked lines on $S$.
- Build the couple set $M_n$ as the union of all couple sets $H_{i,j}$ in $S$ within a distance $d_L$ to $L_n$. In our experiments, we set $d_L$ to 2.5. For the three cases illustrated in the top row of Fig. 11.4, couple sets $M_n$ correspond to image pixels covered by the red lines.
- Randomly pick one couple from $M_n$, which allows to estimate the remaining transform parameters $t_x$ and $t_y$.
- Count the number of couples $N_n$ in $M_n$ which are consistent with the transform, excluding outliers and forbidding any keypoints to appear in multiple matching couples. If $f_1$ and $f_2$ are the coordinates of a couple, consistency checking is done by transforming the coordinates of $f_1$ into $f'_1$ using Eq. 11.1, and verifying that $f'_1$ is within a fixed distance $d_c$ from $f_2$. In practice, we set $d_c$ to 20 mm.
In all experiments, we perform \(2 \times 10^5\) iterations to register a pair of volumes. The bottom row of Fig. 11.4 illustrates the final registration of test images 10000108_1_CTce_ThAb and 10000109_1_CTce_ThAb showing only respective patient body envelopes.

### 11.2.2 Organ Atlas Construction

Using contrast-enhanced CT and MR images and available ground-truth annotations from the VISCERAL training dataset, we construct modality-specific probabilistic atlases for the following 20 structures: thyroid; trachea; sternum; liver; spleen; pancreas; gall bladder; first lumbar vertebra; aorta; urinary bladder; right and left
lungs, kidneys, adrenal glands, psoas major and rectus abdominis muscle bodies. In addition, we create atlases for three additional image and body regions: background (BKG), thorax and abdomen (THAB) and a body envelope (ENV) from annotations generated automatically as follows. BKG is created by thresholding the image followed by morphological processing in order to isolate the background from the body region. THAB is created as the dilated union of the aforementioned 20 structures and their bounding 3D ellipse, from which the structures are subtracted after dilation. Finally, ENV is defined as the image minus BKG and THAB. Note that ENV is a crude body envelope that comprises skin, fat, muscle and bone structures. Figure 11.8c, f illustrate the additional annotations.

To create probabilistic atlases, we choose a representative image per modality from the dataset and use it as a reference onto which we register all remaining images in the modality via the method described in Sect. 11.2.1. We register each structure separately in a bounding box of a given margin in the intensity image, defined according to the corresponding annotation image, and apply the obtained transform subsequently to the annotation image. We accumulate annotations thus registered in a 3D histogram of reference image dimensions which is normalized to produce the corresponding probability map. Refer to Fig. 11.6a for an illustration of probabilistic atlases.

11.2.3 Image Clustering

The full-resolution voxel representation is often redundant because objects usually comprise many similar voxels that could be grouped. Therefore, we simplify the image prior to segmentation by an image-adaptive centroidal Voronoi tessellation (CVT), which strikes a good balance between cluster compactness and object boundary adherence and helps to place subsequent segmentation boundaries precisely. We have shown that the clustering step improves the overall run-time and memory footprint of the segmentation process up to an order of magnitude without compromising the quality of the result [14].

Let us define a greyscale image as a set of voxels $I = \{v \mid v = (x, y, z)\}$ and associate with each voxel $v \in I$ a grey level $I_v$ from some range $I \subset \mathbb{R}$. Given a greyscale image $I$ and $n$ sites $c_i \in I$, a CVT partitions $I$ into $n$ disjoint clusters $C_i$ associated with each centroid $c_i$ and minimizes the following energy:

$$F(v; c_i) = \sum_{i=1}^{n} \left( \sum_{v \in C_i} \rho(v) \left( \|v - c_i\|^2 + \alpha \|I_v - I_i\|^2 \right) \right).$$  \hspace{1cm} (11.2)

In Eq. 11.2, $\rho(v)$ is a density function defined according to the intensity gradient magnitude at voxel $v$, $\rho(v) = |\nabla I_v|$, $\alpha$ is a positive scalar and $I_i$ is the grey level of the cluster $C_i$ defined as the mean intensity of its voxels. Intuitively, minimizing Eq. 11.2 corresponds to maximizing cluster compactness in terms of both geometry
11.2.4 Multiorgan Image Segmentation

We formulate image segmentation as a labelling problem, defined as the assignment of a label from a set of labels \( L \) representing the structures to be segmented to each of the variables in a set of \( n \) variables, indexed by \( \mathcal{I} \), corresponding to the clusters of a CVT-clustered image. Assume that each variable \( i \in \mathcal{I} \) is associated with the corresponding node in the graph \( \mathcal{G} \) of the CVT defined in Sect. 11.2.3. An assignment of labels to all variables is called a configuration and is denoted by \( \ell \in \mathcal{L} \). An assignment of a label to a single variable is denoted by \( \ell_i \). We cast the labelling problem in a maximum a posteriori estimation framework and solve it by minimizing the following energy function of label configurations via the expansion moves multilabel graph cut algorithm [5], which has been shown to outperform popular multilabel optimization algorithms in terms of both speed and quality of obtained solutions [29]:

Fig. 11.5 An image-adaptive CVT clustering and its dual graph for a circle image

and grey level. Refer to Fig. 11.5 for an illustration in 2D. To minimize Eq. 11.2, we apply a variant of the clustering algorithm in [7], which approximates a CVT in a computationally efficient manner, involving only local queries on voxels located on boundaries of pairs of clusters.

For referral in later sections, we shall define the graph of a CVT, illustrated in Fig. 11.5b. Denote the surface of a cluster \( C_i \) by \( \partial C_i \). Given a CVT clustering \( \mathcal{C} \), let the set \( \mathcal{I} \) index its clusters, and let \( \mathcal{G} = (\mathcal{I}, \mathcal{E}) \) be an undirected graph on cluster centroids where pairs of clusters having nonzero area common surface define the set of edges \( \mathcal{E} = \{ \{i, j\} \mid i, j \in \mathcal{I}, |\partial C_i \cap \partial C_j| \neq 0 \} \). Consequently, the neighbourhood of a node \( i \in \mathcal{I} \) is defined as \( \mathcal{N}_i = \{ j \mid j \in \mathcal{I}, \exists \{i, j\} \in \mathcal{E} \} \).
\[ E(\ell) = t_1 \sum_{i \in \mathcal{I}} D_i(\ell_i) + t_2 \sum_{i \in \mathcal{I}} P_i(\ell_i) + \frac{1}{2} \sum_{i \in \mathcal{I}} \sum_{j \in \mathcal{N}_i} V_{i,j}(\ell_i, \ell_j). \] (11.3)

In Eq. 11.3, \( t_1 \) and \( t_2 \) are temperature hyperparameters, and \( \mathcal{N}_i \) is the neighbourhood of the variable \( i \in \mathcal{I} \). The first and second sums in Eq. 11.3 correspond, respectively, to organ intensity and location (atlas) likelihood energies, and the third is the energy of a prior distribution of label configurations expressed as a Markov random field \[18\] with respect to the graph \( \mathcal{I} \). We shall define these terms in detail.

### 11.2.4.1 Spatial Configuration Prior

Pairwise terms of Eq. 11.3 encode prior information on interactions between labels assigned to pairs of neighbouring variables encouraging the spatial consistency of labelling with respect to a reference model. We define these terms according to the piecewise-constant vicinity prior model proposed in \[14\], which, unlike the standard Potts model, incurs multiple levels of penalization capturing the spatial configuration of structures in multiobject segmentation. It is defined as follows.

Let \( \mathcal{R} \) be the set of symmetric adjacency relations on pairs of distinct labels (corresponding to image structures), \( \mathcal{R} = \{ r \mid a r b, a, b \in L, a \neq b \} \). \( \mathcal{R} \) can be represented by a weighted undirected graph on \( L, \mathcal{A} = (L, W) \), with the set of edges \( W = \{ \{ a, b \} \mid \exists r \in \mathcal{R}, a r b, a \neq b \} \), where edge weights are defined by \( w(\{ a, b \}) = 1 \), such that \( w(\{ a, b \}) = \infty \) if \( \not\exists r \in \mathcal{R}, a r b \).

Given the graph \( \mathcal{A} \), we define the pairwise term in Eq. 11.3 as follows:

\[ V_{i,j}(\ell_i, \ell_j) = |\partial C_i \cap \partial C_j| \omega(a, b), \quad \ell_i = a, \ell_j = b. \] (11.4)

where \( \omega(a, b) \) is the shortest-path weight from \( a \) to \( b \) in \( \mathcal{A} \). The adjacency graph of structures according to which we define the spatial prior in our experiments is given in Fig. 11.6b. In Eq. 11.4, the area of the common surface of adjacent clusters \( |\partial C_i \cap \partial C_j| \) is introduced, so that for \( a, b \in L \) the sum of pairwise energies in (11.3) is equal to the area of the common surface between the corresponding pair of structures multiplied by the shortest-path weight. This definition ensures that the segmentation energy is independent of the CVT clustering resolution \[13\].

### 11.2.4.2 Intensity and Location Likelihoods

Unary terms of Eq. 11.3 measure the cost of assigning labels to variables. They are defined as negative log-likelihood functions derived from organ observed intensity and location probabilities:
In Eq. 11.5b, $X_v$ denotes the object-space coordinates of the voxel $v$. Conditional probabilities in Eq. 11.5a and 11.5b correspond, respectively, to those of voxel intensity and location given the structure $\ell_i$. To estimate the conditional probability distribution $Pr(I | \ell)$ for a given label $\ell \in L$, we first register the corresponding organ atlas to the image, then estimate the conditional probability as a Gauss-smoothed and normalized intensity histogram derived from voxels in high-probability regions of the registered atlas according to a threshold value. Conditional probability distributions $Pr(X | \ell)$ are defined directly from registered atlases. The next section outlines our hierarchical registration method which maps organ atlases to an image prior to its segmentation.

### 11.2.4.3 Hierarchical Registration of Organ Atlases

We register probabilistic organ atlases, constructed as described in Sect. 11.2.2, to an image in a three-step hierarchical fashion starting at the full image scale, then on an intermediate level corresponding to the THAB region and finally on individual organs. After performing registration on each scale, we apply the obtained transform to the corresponding atlas as well as to those of organs contained in the registered region. As in Sect. 11.2.2, we register each structure separately in a bounding box of a given margin in the intensity image, defined according to the corresponding atlas.

$$D_i(\ell_i) = -\ln \prod_{v \in C_i} Pr(I_v | \ell_i) , \quad \text{(11.5a)}$$

$$P_i(\ell_i) = -\ln \prod_{v \in C_i} Pr(X_v | \ell_i) . \quad \text{(11.5b)}$$
Figure 11.7 illustrates the hierarchical registration procedure, and Fig. 11.6 gives an example of registered organ atlases overlaid on the CT image to which they have been registered.

### 11.3 Evaluation Results and Discussion

We have carried out qualitative evaluation on several contrast-enhanced CT and MR images from the VISCERAL training dataset. Figure 11.8 shows a pair of segmentations on images 10000109_1_CTce_ThAb and 10000324_4_MRT1cefs_Ab. Their dimensions, respectively, are $512 \times 512 \times 428$ and $312 \times 72 \times 384$. For this pair of images, the number of CVT clusters is set, respectively, to 3 and 20% of image voxel count. In all experiments, we set temperature parameters $t_1$ and $t_2$ so that intensity and location likelihood-based unary terms have the same magnitude in Eq. 11.3. Likewise, for intensity likelihood estimation in all experiments, we fix the probability threshold mentioned in Sect. 11.2.4.2 to 0.9 times that of the maximum probability of the registered probabilistic atlas. The spatial configuration prior is defined according to the adjacency graph given in Fig. 11.6b. We note that, due to a smaller field of view, VISCERAL dataset contrast-enhanced MR images exclude thoracic organs, namely the pair of lungs, the trachea, the sternum and the thyroid. Naturally, we do not construct probabilistic atlases for these structures nor do we take them into account for the segmentation of MR images.

Table 11.1 presents the results of quantitative evaluation of our segmentation method on contrast-enhanced CT images during the VISCERAL Anatomy 2 Benchmark and those of its more recent evaluation on contrast-enhanced MR images. We report results corresponding to the best setting of temperature parameters out of the allowed five. For CT images, the settings for $t_1$ are as follows: 0.15, 0.20, 0.25, 0.30 and 0.40. For MR images, tested settings of this parameter are as follows: 0.6, 0.8, 1.0, 1.2 and 1.4. The parameter $t_2$ was set to 0.2 $t_1$ in both cases. These ranges of parameter values were experimentally found to give the best results in offline evaluations on the VISCERAL training dataset. For each structure, the Table 11.1 gives the
number of produced segmentations out of an attempted 10, mean Dice and average distance (in millimeters) measurements. “N/A” indicates an absent structure, while a dash “–” indicates one for which the segmentation was missed or was not attempted.

Mean run-time and memory footprint figures of our algorithm are given in Table 11.2. These measurements are taken on the 20 contrast-enhanced CT images in the VISCERAL training dataset, the average dimension of which is $512 \times 512 \times 426$. The number of CVT clusters is set to 5% of image voxel count. The algorithm is run on a cluster computer of heterogeneous nodes with an average CPU speed of 2.1 GHz, an average number of cores of 20 and an average memory size of 87 GB.
Table 11.1 Quantitative evaluation results of the proposed method on contrast-enhanced CT and MR images

| Structures          | CT                    | MR                    |
|--------------------|-----------------------|-----------------------|
|                    | # | Dice | Avg. Dist. | # | Dice | Avg. Dist. |
| Trachea            | 9 | 0.62 | 18.56      | N/A | N/A | N/A        |
| Lung (R)           | 10| 0.95 | 0.30       | N/A | N/A | N/A        |
| Lung (L)           | 10| 0.96 | 0.20       | N/A | N/A | N/A        |
| Pancreas           | 7 | 0.35 | 11.45      | 6  | 0.37 | 11.99      |
| Gall bladder       | 2 | 0.14 | 21.82      | 1  | 0.30 | 1.90       |
| Urinary bladder    | 10| 0.77 | 1.08       | 10 | 0.40 | 3.67       |
| Sternum            | 10| 0.63 | 6.59       | N/A | N/A | N/A        |
| Lumbar vertebra    | 10| 0.49 | 9.74       | 7  | 0.26 | 6.65       |
| Kidney (R)         | 10| 0.81 | 1.81       | 10 | 0.80 | 3.90       |
| Kidney (L)         | 10| 0.86 | 0.89       | 8  | 0.74 | 1.69       |
| Adrenal gland (R)  | – | –    | –          | –  | –    | –          |
| Adrenal gland (L)  | – | –    | –          | –  | –    | –          |
| Psoas major muscle (R) | 10| 0.71 | 2.70       | 10 | 0.69 | 1.73       |
| Psoas major muscle (L) | 10| 0.79 | 1.22       | 10 | 0.66 | 2.28       |
| Rectus abdominis muscle (R) | 9 | 0.26 | 30.25      | –  | –    | –          |
| Rectus abdominis muscle (L) | 10| 0.13 | 24.43      | –  | –    | –          |
| Aorta              | 10| 0.58 | 5.43       | 3  | 0.27 | 17.40      |
| Liver              | 10| 0.93 | 0.34       | 10 | 0.77 | 1.91       |
| Thyroid            | 3 | 0.04 | 13.77      | N/A | N/A | N/A        |
| Spleen             | 10| 0.84 | 1.29       | 9  | 0.53 | 3.31       |

Table 11.2 Mean memory footprint and run-time figures of proposed algorithms measured on 20 contrast-enhanced CT images from the VISCERAL training dataset

| Memory (MB) | Registration (s) | Clustering (s) | Segmentation (s) | Total run-time (m) |
|-------------|------------------|----------------|------------------|---------------------|
| 10520.87    | 4294.60          | 8995.20        | 2598.48          | 264.80              |

From these results, we can readily see that our method performs better on CT than on MR images. This is due to the fact that tissues in CT images have consistent appearances, whereas in MR images, they suffer intensity inhomogeneity. In addition, MR images in the VISCERAL dataset have lower resolution compared to CT images. We can observe furthermore that our method performs better on larger, well-contrasted structures than on smaller, low-contrasted ones such as the gall bladder, the thyroid and the adrenal glands. This is mainly due to the inaccurate localization of these structures by our registration method and the subsequent flawed estimation of the structure intensity likelihood. For most structures however, even though our hierarchical approach of mapping atlases to the image relies on a rigid registration method, unlike many hierarchical methods which use non-rigid...
deformable registration [17], it helps localizing structure boundaries in segmentation quite well. This is because location information roughly registered atlases provide is complemented by intensity similarity and spatial consistency criteria. Furthermore, full-body modelling by the introduction of BKG, ENV and THAB annotations not only complements location information and allows for hierarchical registration, but also increases the discriminative power of the spatial prior by a higher penalization of inconsistent configurations.

11.4 Concluding Remarks and Future Work

It should not go without notice that without the VISCERAL platform and the dataset, we would not have been able to test and understand the limits and the properties of our algorithms, to improve them and to develop new ones. We hope that our active participation in benchmarks and our regular feedback on software and data have been valuable for the VISCERAL project.

We are currently scrutinizing our hierarchical registration method in view of multiresolution extensions, possibly bypassing anatomical hierarchy, which would help better localize structures, especially small, low-contrasted ones. We are also investigating the introduction of a better, more robust intensity likelihood estimation method. If an inaccurate registration could be detected and quantified, then it may be possible to “correct” it. Otherwise, with a large training dataset to draw upon, techniques from machine learning could easily be used. Another interesting venue for future research is the extension of the spatial prior model to express other types of relations, such as spatial directionality, and the possibility of taking into account the uncertainty of relations.

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