Beyond pixel-wise supervision for segmentation: A few global shape descriptors might be surprisingly good!

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Abstract

Standard losses for training deep segmentation networks could be seen as individual classifications of pixels, instead of supervising the global shape of the predicted segmentations. While effective, they require exact knowledge of the label of each pixel in an image.

This study investigates how effective global geometric shape descriptors could be, when used on their own as segmentation losses for training deep networks. Not only interesting theoretically, there exist deeper motivations to posing segmentation problems as a reconstruction of shape descriptors: First, annotations to obtain approximations of low-order shape moments could be much less cumbersome than their full-mask counterparts, and anatomical priors could be readily encoded into invariant shape descriptions, which might alleviate the annotation burden. Also, some shape descriptors could be readily used to "encode" biomarkers, leading to better interpretability. Finally, and most importantly, we hypothesize that, given a task, certain shape descriptions might be invariant across image acquisition protocols/modalities and subject populations, which might open interesting research avenues for generalization in medical image segmentation.

We introduce and formulate a few shape descriptors in the context of deep segmentation, and evaluate their potential as stand-alone losses on two different, challenging tasks. Inspired by recent works in constrained optimization for deep networks, we propose a way to use those descriptors to supervise segmentation, without any pixel-level label. Very surprisingly, as little as 4 descriptors values per class can approach the performance of a segmentation mask with 65k individual discrete labels. We also found that shape descriptors can be a valid way to encode anatomical priors about the task, enabling to leverage expert knowledge without requiring additional annotations. Our implementation is publicly available and can be easily extended to other tasks and descriptors: https://github.com/hkervadec/shape_descriptors.

Keywords: Semantic segmentation, constraints, weak supervision, shape moments.

1. Introduction

In the recent years, image semantic segmentation has received considerable attention in the medical imaging research community, and almost all contemporary methods rely on deep learning and fully convolutional neural networks (Ronneberger et al., 2015; Litjens et al., 2017). While network architectures have been extensively studied (Ronneberger et al., 2015; Milletari et al., 2016; Chen et al., 2017), the loss functions used to train them have received relatively less attention, and most of the existing methods rely on variants of either the cross-entropy (Ronneberger et al., 2015) or dice loss (Sudre et al., 2017; Milletari et al., 2016). Ultimately, all those losses are actually performing "pixel-wise classification", and do not account for the image spatial domain—for instance, standard implementations in popular frameworks completely discard the image dimension¹. Other methods that take into account the distances to the segmentation boundary (Kervadec et al., 2019a), or weakly supervised methods that have access to only partial/uncertain annotations (Qu et al., 2019; Rajchl et al., 2016; Papandreou et al., 2015; Bearman et al., 2016; Lin et al., 2016)—in a weakly supervised segmentation setting—eventually supervise a subset of pixels individually. Informally, we could say that the existing segmentation methods are "micro-managing" pixels,treating each one as a separate classification problem, instead of supervising the global shape information of segmentation prediction. (Some additional related methods are discussed in Appendix C.)

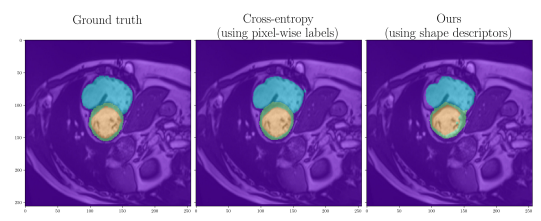
The traditional computer vision literature abounds of global mathematical descriptions that characterize the shapes of objects (Nayak and Stojmenovic, 2008), for instance, shape moments, length, total variation, Fourier transforms, etc. It has also been shown that descriptions based on a few geometric shape moments could be enough to re-construct complex shapes (Milanfar et al., 2000), via solving an inverse problem. Furthermore, such geometric shape moments could be made invariant with respect to geometric transformation (e.g., rotation, translation and scaling) by pure mathematical manipulations, which is convenient for segmentation (Klodt and Cremers, 2011). This includes the well-known Hu's invariant moments (Hu, 1962). While less popular today in computer vision than they used to be, those remain powerful regularization and shape-description tools for segmentation methods. So powerful, perhaps, that they could be fully used to characterize the objects that we want to segment, while providing intrinsic invariance; in short, supervising the overall shape prediction of a segmentation networks, not through individual pixels but rather global shape descriptions.

This paper studies how effective global geometric shape descriptors can be, when used on their own as segmentation losses for training deep neural networks. Not only interesting theoretically, there exist deeper motivations to posing segmentation problems as a reconstruction of shape descriptors: First, annotations to obtain approximations of low-order shape moments could be much less cumbersome than their full-mask counterparts (e.g., from a few mouse clicks by the user). Furthermore, anatomical priors can also be readily translated into shape descriptions, which is not feasible when using dense label masks. This might alleviate the annotation burden for training deep segmentation networks. Also, some shape descriptors could be readily used to "encode" biomarkers, leading to better interpretability. Finally, and most importantly, we hypothesize that, given a task, certain shape descriptions might be invariant across image acquisition protocols/modalities and subject populations, which might open interesting research avenues for generalization in segmentation.

Our contributions can be summarized as follow:

• we re-introduce and reformulate different shape descriptors, in the context of deep semantic segmentation;

^{1.} https://pytorch.org/docs/stable/generated/torch.nn.CrossEntropyLoss.html



(a) A visual comparison of the different supervision methods on the ACDC dataset.

Pixel	Label	Shape descriptor	Class			
0	RV	(in pixels)	RV	Муо	LV	
1 2	Background LV	$\begin{array}{c} \text{Object volume } \mathfrak{V} \\ \text{Centroid location } \mathfrak{C} \end{array}$	3100 (125, 80)	800 (125,	1600 (125)	
65536	: Background	Avg. dist. to centroid \mathfrak{D} Object length \mathfrak{L}	(20, 15) 750	(15, 20) 1000	(10, 10) 500	
(b) Pixel-wise labels (65k discrete values)		(c) Shape descriptors (16 continuous values)				

Figure 1: RV, MYO and LV stands for "right-ventricle", "myocardium" and "left-ventricle", respectively. The shape descriptors are detailed in Subsection 2.2.

- inspired by recent works in inequality constraints, we propose a way to use those descriptors to supervise deep neural networks;
- as such, we benchmark a combination of those descriptors and show that—surprisingly using only a few shape descriptors can go a long way, even in more complex settings (Figure 1). In fact, we found that as little as 4 descriptors values per class could approach the performance of a segmentation mask with 65k individual discrete labels.;
- we discuss future research directions that could benefit from those surprising findings.

2. Formulation

2.1. Notation and background

Let $\Omega \subset \mathbb{R}^2$ denotes the image spatial domain² and $I : \Omega \to \mathbb{R}$ an input image, with $G : \Omega \to \hat{\Delta}^K$ its associated ground truth. Here, Δ^K refers to the K-simplex, and $\hat{\Delta}^K$ to its

^{2.} For readability and simplicity, we detail here only the case of 2D-images, but the method could be extended to N dimensions in a straightforward way.

vertices, i.e., a one-hot encoding for K classes. Our goal is to train a network $\mathcal{N}_{\theta}: I \mapsto s_{\theta}$, with parameters θ predicting a dense probability map $s_{\theta}: \Omega \to \Delta^{K}; s_{\theta}^{(i,k)}$ denotes the predicted softmax probability for class $k \in \{0, ..., K\}$ at pixel $i \in \Omega$. For each pixel i, its coordinates in the 2D space are represented by the tuple $(x_{(i)}, y_{(i)}) \in \mathbb{R}^{2}$.

Shape and central moments have been widely studied in traditionnal computer vision (Nayak and Stojmenovic, 2008; Milanfar et al., 2000), where they can be used to characterize a shape. Each moment is parametrized by its orders $p, q \in \mathbb{N}$, and each order represents a different characteristic of the shape.

Shape moment Shape moments can be defined, in their general form, as functions of a deep-network softmax predictions for a given class k as follows:

$$\mu_{p,q}^{(k)}(s_{\boldsymbol{\theta}}) := \sum_{i \in \Omega} s_{\boldsymbol{\theta}}^{(i,k)} x_{(i)}^p y_{(i)}^q,$$

where $p, q \in \mathbb{N}$ are the moment orders.

Central moment The central moment is closely related to the shape moment, with the difference being that coordinates $x_{(i)}$ and $y_{(i)}$ are shifted by their respective centroids, for translation invariance (more details in the next sub-section). It is given by:

$$\bar{\mu}_{p,q}^{(k)} := \sum_{i \in \Omega} s_{\theta}^{(i,k)} \left(x_{(i)} - \frac{\mu_{1,0}^{(k)}}{\mu_{0,0}^{(k)}} \right)^p \left(y_{(i)} - \frac{\mu_{0,1}^{(k)}}{\mu_{0,0}^{(k)}} \right)^q$$

Image Laplacian The Laplacian of an image is defined by the underlying graph structure \mathcal{G}_{Ω} , which describes the connectivity between each pair of pixels. A sparse graph (i.e., each pixel is connected only to its 4 or 8 direct neighbors) is often used and can be encoded with a sparse adjacency matrix $A_{\Omega} \in \{0, 1\}^{|\Omega| \times |\Omega|}$, where $A_{\Omega,i,j} = 1$ means that i, j are neighbors, and $A_{\Omega,i,j} = 0$ means that they are not. The Laplacian is constructed directly from A_{Ω} :

$$L_{\Omega} := A_{\Omega} - \operatorname{diag}(\mathbf{1}^{\top} A_{\Omega}),$$

where diag : $\mathbb{R}^a \to \mathbb{R}^{a \times a}$ builds a diagonal matrix and $\mathbf{1}^{\top} A_{\Omega} \in \mathbb{R}^{|\Omega|}$ encodes the number of neighbors for each pixel *i*. For a 8-neighbors connectivity, this number will be the same for all pixels, except the image edges which will have less. Notice that L_{Ω} depends only on the image spatial domain, but not the image values. As such, it can be efficiently pre-computed and cached for all the samples in a dataset—assuming they share the same resolution. There exists some edge-sensitive variants, which define A_{Ω} so that it accounts for pixel similarities (e.g., intensity differences), but this is beyond the scope of this study.

2.2. Shape descriptors

With those different standard building blocks, it is possible to define shape descriptors, measuring actual properties of the object, rather than listing a list of pixels that should belong to it. All following descriptors hold for some input image I and some class k:

Volume The volume of the predicted segmentation is simply a summation of the predicted probabilities—which is a special case of shape moments. As such:

$$\mathfrak{V}^{(k)}(s_{\boldsymbol{\theta}}) := \mu_{0,0}^{(k)}(s_{\boldsymbol{\theta}}).$$

Centroid The centroid of a class can be computed by dividing the first shape moment by the volume. We can see it as the average of the pixel-coordinates for class k:

$$\mathfrak{C}^{(k)}(s_{\theta}) := \left(\frac{\mu_{1,0}^{(k)}(s_{\theta})}{\mu_{0,0}^{(k)}(s_{\theta})}, \frac{\mu_{0,1}^{(k)}(s_{\theta})}{\mu_{0,0}^{(k)}(s_{\theta})}\right).$$

Average distance to the centroid It measures how far the object should spread around its centroid, on average. It is the standard deviation of pixel-coordinates for class k:

$$\mathfrak{D}^{(k)}(s_{\theta}) := \left(\sqrt[2]{\frac{\bar{\mu}_{2,0}^{(k)}(s_{\theta})}{\mu_{0,0}^{(k)}(s_{\theta})}}, \sqrt[2]{\frac{\bar{\mu}_{0,2}^{(k)}(s_{\theta})}{\mu_{0,0}^{(k)}(s_{\theta})}} \right).$$

Length The length of a segmentation, or rather, the length of its boundary, can be efficiently computed by re-using the pre-computed image Laplacian. To summarily describe it, each difference of classification between two neighbors will be counted as 1, while neighbors with the same predicted class will count as 0; which is a standard Potts model. It is trivial to relax this definition to plug the predicted (continuous) probabilities:

$$\mathfrak{L}^{(k)}(s_{\theta}) := \sum_{i,j \in \mathcal{G}_{\Omega}} |s_{\theta}^{(i,k)} - s_{\theta}^{(j,k)}| L_{\Omega,i,j}.$$

Ratio of descriptors In the multi-class setting, some relationships between different classes might be known in advance, using anatomical priors, for instance. While exact values are not necessarily required, inequalities could provide useful information. As such, we can define an additional descriptors for pairs of classes k and l, for a specific descriptor $\mathfrak{f} \in {\mathfrak{V}, \mathfrak{C}, \mathfrak{D}, \mathfrak{L}}$:

$$\mathfrak{R}_{\mathfrak{f}}^{(k,l)}(s_{\boldsymbol{\theta}}) := \frac{\mathfrak{f}^{(k)}(s_{\boldsymbol{\theta}})}{\mathfrak{f}^{(l)}(s_{\boldsymbol{\theta}})}.$$

2.3. Supervision with constraints

Instead of optimizing a pixelwise loss, we design loss functions, which penalize the deviations between the global shape descriptors computed from the predicted segmentation and those corresponding to the ground truth, e.g., $\mathfrak{C}^{(k)}(s_{\theta}) = \mathfrak{C}^{(k)}(G)$. This could be formulated as a hard equality-constrained optimization problem. Here, we propose to relax the constraint to add a lower and upper bound centered around the ground truth value (this may mimic imprecise information about shape descriptors when these are derived, for instance, from anatomical prior knowledge and not from ground truth):

$$\begin{array}{ll} \underset{\boldsymbol{\theta}}{\operatorname{arg\,min}} & \mathcal{L}_{\boldsymbol{\theta}} & (1) \\ \text{subject to} & 0.9\tau_{\mathfrak{f}}^{(k)} \leq \mathfrak{f}^{(k)}(s_{\boldsymbol{\theta}}) \leq 1.1\tau_{\mathfrak{f}}^{(k)} & \forall k, \; \forall \mathfrak{f} \in \{\mathfrak{V}, \mathfrak{C}, \mathfrak{D}, \mathfrak{L}\} \\ & a \leq \mathfrak{R}_{\mathfrak{f}}^{(k,l)} \leq b & \text{for some } \mathfrak{f}, a, b, k, l, \end{array}$$

where $\tau_{\mathfrak{f}}^{(k)} = \mathfrak{f}^{(k)}(G)$.

In the context of deep neural networks, standard constrained optimization techniques (such as Lagrangian or interior-point methods) are not directly applicable for tractability reasons. The inequality constraints can be tackled directly as a loss function using a *logbarrier-extension* penalty $\tilde{\psi}_t(\cdot)$ (details can be found in Section B), controlled by a parameter t that is increased over time to make the bounds tighter and tighter. Such log-barrier penalties were introduced recently in (Kervadec et al., 2019c) in the general optimization context for constrained deep networks. As, for the sake of the study, we want to completely forego pixel-wise supervision, we set $\mathcal{L}_{\boldsymbol{\theta}} := 0$. As such, our final model is:

$$\underset{\boldsymbol{\theta}}{\operatorname{arg\,min}} \quad \sum_{\mathfrak{f}} \sum_{k} \left[\widetilde{\psi}_{t} \left(0.9\tau_{\mathfrak{f}}^{(k)} - \mathfrak{f}^{(k)}(s_{\boldsymbol{\theta}}) \right) + \widetilde{\psi}_{t} \left(\mathfrak{f}^{(k)}(s_{\boldsymbol{\theta}}) - 1.1\tau_{\mathfrak{f}}^{(k)} \right) \right]. \tag{2}$$

Bounds for $\mathfrak{R}_{\mathfrak{f}}^{(k,l)}$ can be included in the same fashion, if available and relevant—depending on the task at hand. The bound values for \mathfrak{R} do not rely on G, but rather on expert knowledge about the task. We will give some examples in the next section.

3. Experiments

3.1. Datasets

Heart segmentation on cine-MRI The main dataset that we use in our experiments is the publicly available 2017 ACDC Challenge (Bernard et al., 2018), which contains 4 classes to segment: left and right ventricles, myocardium, and background. The dataset consists of 100 cine magnetic resonance (MR) exams covering well defined pathologies: dilated cardiomyopathy, hypertrophic cardiomyopathy, myocardial infarction with altered left ventricular ejection fraction and abnormal right ventricle. It also included normal subjects. We chose this dataset because it is a good benchmark for shape descriptors. Not only a multi-class setting, the myocardium and left-ventricle share a common centroid, and the myocardium completely surround the left-ventricle—which is more challenging to describe. We constraint $\mathfrak{S}, \mathfrak{C}, \mathfrak{D}, \mathfrak{L}$, and $\tau_{\mathfrak{C}}^{(MYO)} = \tau_{\mathfrak{C}}^{(LV)}$. Moreover, the relationship between myocardium and left-ventricle can be formulated with the following bounds on their relative length: $2 \leq \mathfrak{R}_{\mathfrak{L}}^{(MYO,LV)} \leq 3$. We retained 70 exams for training, 10 for validation and 20 for testing.

Prostate segmentation on MR-T2 The second dataset that we use is the PROMISE12 challenge (Litjens et al., 2014). It contains the transversal T2-weighted MR images of 50 patients acquired at different centers, with multiple MRI vendors and different scanning protocols. The images include patients with benign diseases, as well as with prostate cancer. We employed 35 patients for training, 5 for validation, and 10 for testing. The difficulty of this dataset lies in its low-contrast, and the very variable shape of the prostate. We supervise $\mathfrak{S}, \mathfrak{C}, \mathfrak{D}, \mathfrak{L}$.

3.2. Implementation details

We use the ENet architecture (Paszke et al., 2016) for experiences on ACDC, and a modified fully residual UNet for the experience on PROMISE12—the prostate is a harder task that requires a more powerful network, and it also enables us to validate the supervision method

on different network architectures. We perform blurring, shifting, and scaling as online data augmentations, and we use the same network initialization for all settings, with the same scheduler and hyper-parameters (ADAM scheduler (Kingma and Ba, 2014), with learning rate of 5e-4 and $\beta = (0.9, 0.99)$). The shape descriptors are computed from the annotated mask, and we use a relaxed value by $\pm 10\%$ as bounds. Most of the implementation was done in the PyTorch framework, and experiments were run on a NVIDIA Titan RTX. All descriptors can be efficiently vectorized, resulting in minimal slowdown during training (less than 10% compared to a cross-entropy loss). The computation of the Laplacian \mathcal{L}_{Ω} is done once per image shape (usually a single one per dataset after pre-processing), and cached using standard Python utilities (lru_cache from functools). Our code is publicly available at https://github.com/hkervadec/shape_descriptors, and can easily be extended to other shape descriptors.

4. Results

Surprisingly, using only a few shape descriptors in place of dense pixel-wise supervision is capable to segment the objects of interest, as we can see in Figure 2. On ACDC, what remains the most difficult to learn is the hierarchy between the left-ventricle and its surrounding myocardium: some noisy myocardium pixels can sometime remain inside the predicted leftventricle. Nonetheless, we can consider that the network has properly learned the overall structure of the heart. On PROMISE12, the task is difficult even for cross-entropy with full annotations. Despite the more powerful network used, the low-contrast can still trick both methods. Nevertheless, supervision with shape descriptors is capable to predict a rough location and shape of the prostate, which is much more than we initially expected. Actual testing DSC values can be found in Table 1, and the plots of training and validation metrics over time can be found in Appendix D.

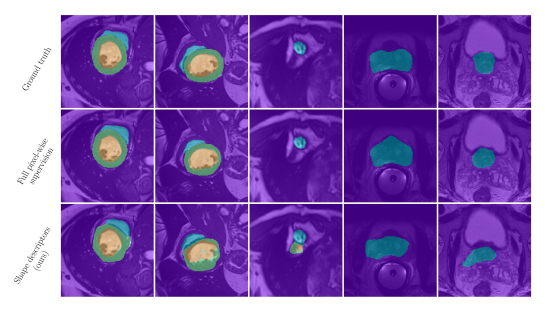


Figure 2: Visual comparison for both ACDC and PROMISE12 on the testing set, including some failure cases.

	ACDC				Promise12
Method	RV	Муо	LV	Overall	Prostate
Cross-entropy (pixel-wise) Ours (shape descriptors)	· · · ·	· · · · · ·	$\begin{array}{c} 0.919 \ (0.059) \\ 0.819 \ (0.086) \end{array}$		· · · ·

Table 1: Average and standard deviation of DSC on the testing set, for both datasets.

5. Discussion and conclusion

We have shown that simple and light shape descriptors can be effective supervision tools for semantic segmentation, allowing us to avoid completely pixel-wise supervision; and proving how powerful shape descriptors can be. In a multi-class setting, the neural network is able to learn the inherent relationship between classes and the anatomical structure of the heart.

While not needed on the two datasets that we benchmarked on, it is very easy to compute the orientation and elongation of an object (Nayak and Stojmenovic, 2008), which would be very useful for certain tasks (for instance, esophagus segmentation). Spatial relationship between classes, that would be translation invariant, could be very beneficial in some settings, such as the co-segmentation of esophagus and trachea—both long objects next to each others.

We found empirically that using only shape descriptors without online data augmentation was more sensitive to network initialization than its pixel-wise counterpart. It is entirely plausible that the random networks' initializations, designed and tuned with crossentropy in mind (Sutskever et al., 2013), are not optimal for shape descriptors. As such, future works could investigate other network initialization strategies.

A main limitation of the method is its inability to be sub-patched and processed in different batches (any loss requiring a sum over an area bigger than the current patch shares this limitation, including the very popular Dice loss and its derivatives.) Recently, for a similar ill-suited problem (enforcing a prior of the distribution of the classes, over the whole training set), (Zhou et al., 2019) showed that a primal-dual approach can be a promising avenue.

We believe that we barely scratched the surface for the potential of invariant shape descriptors: shape and central moments orders can go much higher than two. Depending on the task, some invariant and higher-order descriptors could be common to all the samples and would not require additional annotations, but rather exploit existing anatomical knowledge. Time-series could also benefit, as some descriptors might not vary across time, reducing the annotation burden and enabling the re-use of previously computed statistics. We describe such a setting in Appendix A. All in all, this might open interesting avenues for generalization across subject populations and acquisition protocols.

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Appendix A. Time independent 3D shape descriptors

As mentioned in Section 2.1, all shape descriptors can be extended quite easily to 3D. In the case of the ACDC dataset, this could be very powerful: as the training data comes from actual 4D Cine-MRI scans (3D images over time), with annotations for two time points at the beginning of the systole and diastole: when the heart is at its biggest and smallest, respectively.

By computing the descriptors at those two extremes, one could get *patient-wise* (and not *image-wise*) upper and lower bounds for our descriptors, valid at any time-point. Figure 3 shows one ellipsoid per class k (based on the *average distance to the centroid*, $\mathfrak{D}^{(k)}$, and centered around its centroid $\mathfrak{C}^{(k)}$). The thick lines represent the shift of the centroid between the two phases. We can clearly see the "shrink" between the two phases, and the slight shift of the centroids toward the center of the scan. The bounds given by those two annotations would allows to constrain the remaining unannotated 3D volumes at train time; increasing the training set size by one order of magnitude. The benefits are clear, especially in low-data settings.

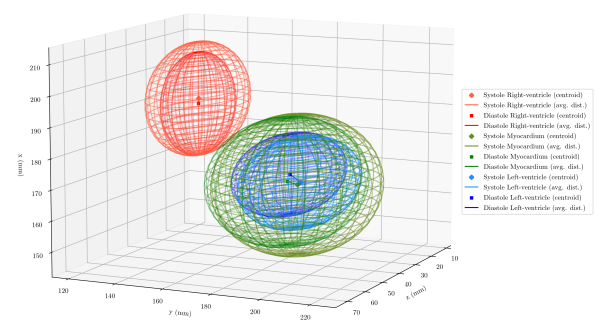


Figure 3: 3D-mesh representation of the ellipsoids of the *average distance to the centroid* at systole and diastole, for patient001 (ACDC dataset).

Appendix B. Extended log-barrier

The extended log-barrier was introduced in (Kervadec et al., 2019c), as standard Lagrangian or interior-point methods are not directly applicable to deep learnign settings. If we take a simple constrained optimization setting:

$$\begin{array}{ll} \underset{\boldsymbol{\theta}}{\operatorname{arg\,min}} & \mathcal{L}_{\boldsymbol{\theta}} \\ \text{subject to} & z \leq 0, \end{array}$$

then its extended log-barrier equivalent is:

$$\underset{\boldsymbol{\theta}}{\operatorname{arg\,min}} \quad \mathcal{L}_{\boldsymbol{\theta}} + \widetilde{\psi}_t(z) \\ \widetilde{\psi}_t(z) = \begin{cases} -\frac{1}{t}\log(-z) & \text{if } z \leq -\frac{1}{t^2} \\ tz - \frac{1}{t}\log(\frac{1}{t^2}) + \frac{1}{t} & \text{otherwise,} \end{cases}$$

where t is the slope parameter of the log-barrier that is increased over time, eventually "closing" the barrier when $t \to \infty$. This is illustrated in Figure 4

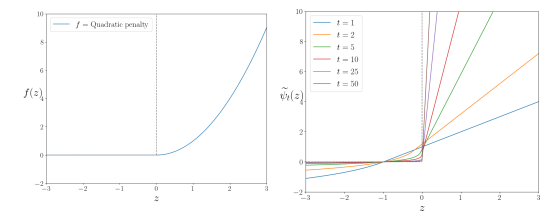


Figure 4: Illustration of the extended log-barrier (Kervadec et al., 2019c), for increasing t values, compared to a fixed quadratic penalty (Kervadec et al., 2019b).

The advantage of the log-barrier are two-fold:

- it allows to gradually increase the tightness of the constraints that we want to satisfy;
- once satisfied, it gently pushes back the constrained function toward the feasible set, preventing it to go out of bounds.

Appendix C. Related works

While there is, to the best of our knowledge, no other work attempting to supervise the shape of the predicted segmentation in the way we do—with direct losses that can be plugged on top of any existing network—there exist a few works that created custom architectures to regularize the shape of the predicted segmentation. They all require full-mask annotations, use a base cross-entropy loss, and have been evaluated only on simpler binary settings.

ACNN (Oktay et al., 2017) trains an auto-encoder with example annotations, to generate a *shape embedding* of the task at hand. The encoder is then used when training the main segmentation network, by minimizing (on top of the base cross-entropy loss) the euclidean distance between the encoding of the predicted segmentation and the encoding of the (fully labeled) ground truth.

(Milletari et al., 2017) integrate PCA values (computed from an annotated dataset) into a dedicated "PCA-aware" CNN architecture, which is used to regularize the shape of the predicted segmentation. However, the heavily-customized layers make it incompatible with existing FCN architectures, and the PCA pre-computed values are hardly interpretable.

(Nguyen and Ray, 2020) integrates a "star-shape" prior into their training on top of the base cross-entropy loss. This was used to regularize the contour of ventricle segmentation in 2D slices, given a user-provided centroid. The original prior (Ray et al., 2012) was optimized using dynamic programming, which is not usable for deep neural networks. In that paper, the authors proved that it was possible to "learn" that dynamic programming part using an additional, trainable network module. However, to be useful, the method requires user-provided centroid *even at inference*.

Appendix D. Training curves

The training curves shows that the training is fairly stable over time, though in the case of PROMISE12 it takes a few epochs for the network to start producing meaningful predictions. This is related, we think, to the random initialization procedure used in standard deep learning settings, which might not be the most optimal method when using different forms of supervision.

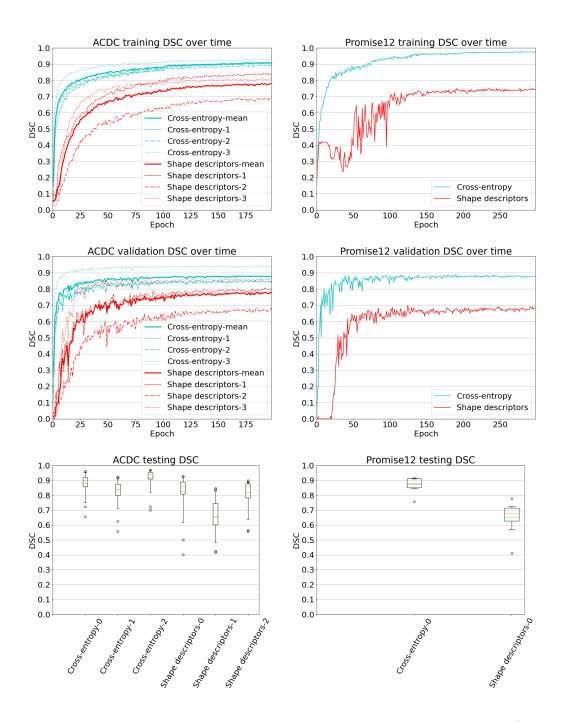


Figure 5: Comparison of the training and validation dice, and the distribution of (patientwise) testing dice for each individual class, for both datasets. In the case of ACDC, we plot one curve per class (RV=1, MYO=2, LV=3), as well as an average of the three.