Do Neural Networks for Segmentation Understand Insideness?

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Abstract

The insideness problem is an image segmentation modality that consists of determining which pixels are inside and outside a region. Deep Neural Networks (DNNs) excel in segmentation benchmarks, but it is unclear that they have the ability to solve the insideness problem as it requires evaluating long-range spatial dependencies. In this paper, the insideness problem is analysed in isolation, without texture or semantic cues, such that other aspects of segmentation do not interfere in the analysis. We demonstrate that DNNs for segmentation with few units have sufficient complexity to solve insideness for any curve. Yet, such DNNs have severe problems to learn general solutions. Only recurrent networks trained with small images learn solutions that generalize well to almost any curve. Recurrent networks can decompose the evaluation of long-range dependencies into a sequence of local operations, and learning with small images alleviates the common difficulties of training recurrent networks with a large number of unrolling steps.

This material is based upon work supported by the Center for Brains, Minds and Machines (CBMM), funded by NSF STC award CCF-1231216.
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The insideness problem is an image segmentation modality that consists of determining which pixels are inside and outside a region. Deep Neural Networks (DNNs) excel in segmentation benchmarks, but it is unclear that they have the ability to solve the insideness problem as it requires evaluating long-range spatial dependencies. In this paper, the insideness problem is analysed in isolation, without texture or semantic cues, such that other aspects of segmentation do not interfere in the analysis. We demonstrate that DNNs for segmentation with few units have sufficient complexity to solve insideness for any curve. Yet, such DNNs have severe problems to learn general solutions. Only recurrent networks trained with small images learn solutions that generalize well to almost any curve. Recurrent networks can decompose the evaluation of long-range dependencies into a sequence of local operations, and learning with small images alleviates the common difficulties of training recurrent networks with a large number of unrolling steps.

1 Introduction

A key component of image segmentation is to determine whether a pixel is inside or outside a region, i.e. the “insideness” problem [39, 40]. This problem involves evaluating long-range spatial dependencies. Capturing such long-range dependencies may be challenging for artificial neural networks as pointed out in Minsky & Papert’s historic book Perceptrons [31] and recent works on capturing other spatial relationships such as containment [20] and connectedness [26].

Deep Neural Networks (DNNs) have been tremendously successful in image segmentation benchmarks, but it is not well understood whether DNNs represent insideness or how. Insideness has been overlooked in DNNs for segmentation since they have been mainly applied to the modality of “semantic segmentation”, i.e. labelling each pixel with its object category [33, 44, 41, 3, 7, 29, 22]. The same could be said for DNNs for more advanced segmentation modalities and applications that have been recently introduced, e.g. segmentation of individual object instances rather than object categories [23, 25, 37, 6, 16, 30, 28, 14] and generating realistic images [46]. In such cases, insideness has not been considered because solutions rely on texture, shape and other visual cues. Yet, investigating whether DNNs understand insideness could reveal new insights about their ability to capture long-range spatial relationships, which is key for a full image understanding.

In this paper, we investigate analytically-derived and learned representations in DNNs for insideness. We take the reductionist approach by isolating insideness such that other components of image segmentation do not provide additional cues and ensure that our analysis focuses on the long-
We now introduce the paradigm that will serve to analyze the ability of DNNs to solve insideness. Thus, we analyze the segmentation of Jordan curves, i.e. closed curves synthetically generated without texture nor object category.

We analytically demonstrate that state-of-the-art network architectures, i.e. DNNs with dilated convolutions [44,7] and convolutional LSTMs (ConvLSTMs) [43], among other networks, can exactly solve the insideness problem for any curve with network sizes that are easily implemented in practice. The proofs draw on algorithmic ideas from classical work on visual routines [39,40], namely, the ray-intersection method and the coloring method, to derive equivalent neural networks that implement these algorithms. In a series of experiments with synthetically generated closed curves, we evaluate the capabilities of these DNNs to learn to solve the insideness problem. The experiments show the DNNs do not learn general solutions for insideness, even though the architectures are complex enough to capture the long-range relationships. This is because the DNNs learn to recognize specific features of the family of curves of the training set that do not generalize to new families of curves. Only the recurrent networks, such as the ConvLSTM, when trained on small images generalize to almost any given curve of any size. This is because training on small images alleviates the well-known difficulties of training recurrent networks with a large number of unrolling steps. Also, it facilitates learning a strategy that deals with long-range dependencies by breaking them into local operations that are reusable for any curve, even curves in larger images that are not seen during training.

These results add to the growing body of work that shows that DNNs have problems in learning to solve some elemental visual tasks [26,27,42,36]. Shalev-Shwartz et al. [36] introduced several tasks that DNNs can in theory learn, as demonstrated mathematically, but were unable to solve in practice, even for the training dataset, due to difficulties in the optimization with gradient descent. In contrast, the challenges we report for insideness are related to poor generalization rather than optimization, as our experiments show the networks succeed in solving insideness for the family of curves seen during training. Linsley et al. [26] and Kim et al. [19] introduced new architectures that better capture the long-range dependencies in images. Here, we show that the training strategy has a significant impact on capturing the long-range dependencies, as even DNNs with the capacity to capture such dependencies do not learn a general solution with standard training strategies. Our results highlight the need to decompose the long-range dependencies in a sequence of local operations, that can be learned with recurrent networks by controlling the number of unrolling steps with the image size.

2 The Reductionist Approach to Insideness

We now introduce the paradigm that will serve to analyze the ability of DNNs to solve insideness. Rather than using natural images, we use synthetic stimuli that solely contains a closed curve. In this way, we do not mix the insideness problem with other components of image segmentation found in natural images, e.g. discontinuity of segments, representation of the hierarchy of segments, etc. This reductionist methodology that we apply here has the advantage of minimizing the interference of these other factors in the analysis of the abilities of DNNs to capture long-range spatial dependencies. Note that if we would include other factors, it will obfuscate if the behaviour of the network is caused by some of the individual factors or by the combination of them.

Let \( X \in \{0,1\}^{N \times N} \) be an image or a matrix of size \( N \times N \) pixels. We use \( X_{i,j} \) and \( (X)_{i,j} \), indistinguishably, to denote the value of the image in position \((i,j)\). We use this notation for indexing elements in any of the images and matrices that appear in the rest of the paper. Also, in the figures we use white and black to represent 0 and 1, respectively.

Insideness refers to finding which pixels are in the inside and which ones in the outside of a closed curve. We assume without loss of generality that there is only one closed curve in the image and that it is a digital version of a Jordan curve [21], i.e. a closed curve without self-crosses nor self-touches and containing only horizontal and vertical turns, as shown in Fig. [1]. We further assume that the curve does not contain the border of the image. The curve is the set of pixels equal to 1 and is denoted by \( \mathcal{F}_X = \{(i,j)|X_{i,j} = 1\} \).

The pixels in \( X \) that are not in \( \mathcal{F}_X \) can be classified into two categories: the inside and the outside of the curve [21]. We define the segmentation of \( X \) as \( S(X) \in \{0,1\}^{N \times N} \), where

\[
S(X)_{i,j} = \begin{cases} 
0 & \text{if } X_{i,j} \text{ is inside} \\
1 & \text{if } X_{i,j} \text{ is outside} 
\end{cases}
\]  

(1)
Figure 1: Intersections of the Ray and the Curve. (a) Example of ray going from one region to the opposite one when crossing the curve. (b) Example of ray staying in the same region after intersecting the curve. (c) All cases in which a ray could intersect a curve. In the three cases above the ray travels from one region to the opposite one, while in the two cases below the ray does not change regions.

and for the pixels in $\mathcal{F}_X$, the value of $S(X)_{i,j}$ can be either 0 or 1. Note that unlike object recognition, the definition of insideness is rigorously and uniquely determined by the input image itself.

The number of all digital Jordan curves is enormous even if the image size is relatively small, e.g. it is more than $10^{47}$ for the size $32 \times 32$ (App. A). In addition, insideness is a global problem; whether a pixel is inside or outside depends on the entire image, and not just on some local area around the pixel. Therefore, simple pattern matching, i.e., memorization, is impossible in practice.

3 Can DNNs for Segmentation Solve Insideness?

The universal approximation theorem [8] tells us that even a shallow neural network is able to solve the insideness problem. Yet, it could be that the amount of units is too large to be implementable in practice. In this Section, we introduce two DNN architectures that are able to solve the insideness problem at perfection and they are easily implementable in practice. One architecture is feed-forward with dilated convolutions [44, 7] and the other is recurrent, a ConvLSTM [43].

3.1 Feed-Forward Architecture with Dilated Convolutions

Dilated convolutions facilitate capturing long-range dependencies which are key for segmentation [44, 7]. To demonstrate that there are architectures with dilated convolutions that can solve the insideness problem, we borrow insights from the ray-intersection method. The ray-intersection method [39, 40], also known as the crossings test or the even-odd test [12], is built on the following fact: Any ray that goes from a pixel to the border of the image alternates between inside and outside regions every time it crosses the curve. Therefore, the parity of the total number of such crossings determines the region to which the pixel belongs. If the parity is odd then the pixel is inside, otherwise it is outside (see Fig. 1a).

The definition of a crossing should take into account cases like the one depicted in Fig. 1b, in which the ray intersects the curve, but does not change region after the intersection. To address these cases, we enumerate all possible intersections of a ray and a curve, and analyze which cases should count as crossings and which ones should not. Without loss of generality, we consider only horizontal rays. As we can see in Fig. 1c, there are only five cases for how a horizontal ray can intersect the curve. The three cases at the top of Fig. 1c, are crosses because the ray goes from one region to the opposite one, while the two cases at the bottom (like in Fig. 1b) are not considered crosses because the ray remains in the same region.

Let $\vec{X}(i, j) \in \{0, 1\}^{1 \times N}$ be a horizontal ray starting from pixel $(i, j)$, which we define as

$$\vec{X}(i, j) = [X_{i,j}, X_{i,j+1}, X_{i,j+2}, \ldots, X_{i,N}, 0, \ldots, 0],$$

where zeros are padded to the vector if the ray goes outside the image, such that $\vec{X}(i, j)$ is always of dimension $N$. Let $\vec{X}(i, j) \cdot \vec{X}(i + 1, j)$ be the inner product of the ray starting from $(i, j)$ and the ray starting from the pixel below, $(i + 1, j)$. Note that the contribution to this inner product from the three cases at the top of Fig. 1c (the crossings) is odd, whereas the contribution from the other two intersections is even. Thus, the parity of $\vec{X}(i, j) \cdot \vec{X}(i + 1, j)$ is the same as the parity of the total number of crosses and determines the insideness of the pixel $(i, j)$, i.e.

$$S(X)_{i,j} = \text{parity} \left( \vec{X}(i, j) \cdot \vec{X}(i + 1, j) \right).$$
Dilated convolutions, also called atrous convolutions, are convolutions with upsampled kernels, which enlarge the receptive fields of the units but preserve the number of parameters of the kernel [44, 7]. In App. [3] we prove that equation 3 can be easily implemented with a neural network with dilated convolutions. The demonstration is based on implementing the dot product in equation 3 with multiple layers of dilated convolutions, as they enable capturing the information across the ray. The number of dilated convolutional layers is equal to \( \log_2(N) \) (recall \( N \) is the image size). The dot product can also be implemented with two convolutional layers, but with the drawback of using a long kernel of size \( 1 \times N \). The multiple dilated convolutions using \( 3 \times 3 \) kernels are equivalent to the long kernel of \( 1 \times N \). Finally, the parity function in equation 3 is implemented by adapting the network introduced by [56], which yields a two-layer convolutional network with a maximum of \( 3N/2 \) kernels of \( 1 \times 1 \) size (see App. [C]). Thus, DNNs with dilated convolutions can solve the insideness problem and are implementable in practice, since the number of layers and the number of kernels grow logarithmically and linearly with the image size, respectively. In Fig. 3.4 we depict the neural network architecture and the process to derive the equivalence with dilated convolutions.

Note that the proof introduces the smallest network we could find that solves the insideness problem with dilated convolutions. Larger networks than the one we introduced can also solve the insideness problem, as the network size can be reduced by setting kernels to zero and layers to implement the identity operation.

3.2 Recurrent Architecture: Convolutional LSTMs

Convolutional LSTM (ConvLSTM) [43] is another architecture designed to handle long-range dependencies and is used in state-of-the-art segmentation architectures [41, 24, 2]. We now show that a ConvLSTM with just one kernel of size \( 3 \times 3 \) is sufficient to solve the insideness problem. This is achieved by exploiting its internal back-projection of the LSTM, i.e., the flow of information from a posterior layer to an anterior layer.

Our demonstration is inspired by the coloring method [39, 40], which is another algorithm for the insideness problem. This algorithm is based on the fact that neighboring pixels not separated by the curve are in the same region. We present a version of this method that will allow us to introduce the network with an LSTM. This method consists of multiple iterations of two steps: (i) expand the outside region from the borders of the image (which by assumption are in the outside region) and (ii) block the expansion when the curve is reached. The blocking operation prevents the outside region from expanding to the inside of the curve, yielding the solution of the insideness problem, as depicted in Fig. 3.2. We call one iteration of expanding and blocking coloring routine.

We use \( E^t \in \{0, 1\}^{N \times N} \) (expansion) and \( B^t \in \{0, 1\}^{N \times N} \) (blocking) to represent the result of the two operations after iteration \( t \). The coloring routine can then be written as (i) \( E^t = \text{Expand}(B^{t-1}) \) and (ii) \( B^t = \text{Block}(E^t, \mathcal{F}_X) \). Let \( B^{t-1} \) maintain a value of 1 for all pixels that are known to be outside and 0 for all pixels whose region is not yet determined or belong to the curve. Thus, we initialize \( B^{0} \) to have value 1 (outside) for all border pixels of the image and 0 for the rest. In step (i), the outside region of \( B^{t-1} \) is expanded by setting also to 1 \((\text{outside})\) its neighboring pixels, and the result is assigned to \( E^t \). Next, in step (ii), the pixels in \( E^t \) that were labeled with 1 \( (\text{outside}) \) and belong to the curve, \( \mathcal{F}_X \), are reverted to 0 \( (\text{inside}) \), and the result is assigned to \( B^t \). This algorithm ends when the outside region can no longer expand, which is at most after \( N^2 \) iterations (worst case where each iteration expands the outside region by only one pixel). Therefore, we have \( E^{N^2} = S(X) \).

In App. [D] we demonstrate that a ConvLSTM with one kernel applied on an image \( X \) can implement the coloring algorithm. In the following we provide a summary of the proof. Let \( I^t, F^t, O^t, C^t, \) and \( H^t \in \mathbb{R}^{N \times N} \) be the activations of the input, forget, and output gates, and cell and hidden states of a ConvLSTM at step \( t \), respectively. By analyzing the equations of the ConvLSTM (equation 11 and equation 12 in App. [D]) we can see that the output layer, \( O^t \), back-projects to the hidden layer, \( H^t \). In the coloring algorithm, \( E^t \) and \( B^t \) are related in a similar manner. Thus, we define \( O^t = E^t \) (expansion) and \( H^t = \frac{1}{2} B^t \) (blocking). The \( \frac{1}{2} \) factor is a technicality due to non-linearities, which is compensated in the output gate and has no relevance in this discussion.

We initialize \( H^0 = \frac{1}{2} B^0 \) (recall \( B^0 \) is 1 for all pixels in the border of the image and 0 for the rest). The output gate expands the hidden representations using one \( 3 \times 3 \) kernel. To stop the outside region from expanding to the inside of the curve, \( H^t \) takes the expansion output \( O^t \) and sets the pixels
Figure 2: The Coloring Method with ConvLSTM. (a) The coloring method consists of several iterations of the coloring routine, i.e. expanding the outside region and blocking it on the curve. (b) Diagram of the ConvLSTM implementing the coloring method, we highlight the connections between layers that are used for insideness. \( \neg X \) denotes the element-wise “Boolean not” of \( X \).

at the curve’s location to 0 (inside). This is the same as the element-wise product of \( O \) and the “Boolean not” of \( X \), which is denoted as \( \neg X \). Thus, the blocking operation can be implemented as \( H' = \frac{1}{2}(O \odot \neg X) \), and can be achieved if \( C' \) is equal to \( \neg X \). In Fig. 2b, we depict these computations.

In App. D we show that the weights of a ConvLSTM with just one kernel of size \( 3 \times 3 \) can be configured to reproduce these computations. A key component is that many of the weights use a value that tends to infinity. This value is denoted as \( q \) and it is used to saturate the non-linearities of the ConvLSTM, which are hyperbolic tangents and sigmoids. Note that it is common in practice to have weights that asymptotically tend to infinity, e.g. when using the cross-entropy loss to train a network [38]. In practice, we found that saturating non-linear units using \( q = 100 \) is enough to solve the insideness problem for all curves in our datasets. Note that only one kernel is sufficient for ConvLSTM to solve the insideness problem, regardless of image size. Furthermore, networks with multiple stacked ConvLSTM and more than one kernel can implement the coloring method by setting unnecessary ConvLSTMs to implement the identity operation (App. D) and the unnecessary kernels to 0.

Finally, we point out that there are networks with a much lower complexity than LSTMs that can solve the insideness problem, although these networks rarely find applications in practice. In App. E we show that a convolutional recurrent network as small as having one sigmoidal hidden unit per pixel, with a \( 3 \times 3 \) kernel, can also solve the insideness problem for any given curve.

4 Can DNNs for Segmentation Learn Insideness?

After having identified DNNs that with few units have sufficient complexity to solve the insideness problem, we focus on analyzing whether these solutions can be learnt from examples. In the following, we first describe the experimental setup and then analyze the generalization capabilities of the DNNs trained in standard manner.

4.1 Experimental Setup

Recall that the goal of the network is to predict for each pixel in the image whether it is inside or outside of the curve.

Datasets. Given that the number of Jordan curves explodes exponentially with the image size, a procedure that could provide curves without introducing a bias for learning is unknown. We introduce three algorithms to generate different types of Jordan curves. For each dataset, we generate \( 95K \) images for training, \( 5K \) for validation and \( 10K \) for testing. All the datasets are constructed to fulfill the constraints introduced in Sec. 2. In addition, for testing and validation sets, we only use images that are dissimilar to all images from the training set. Two images are considered dissimilar if at least 25% of the pixels of the curve are in different locations. In the following, we briefly introduce each dataset (see App. F for details). Fig. 3, shows examples of curves for each dataset.

- Polar Dataset (92 \( \times \) 92 pixels): We use polar coordinates to generate this dataset. We randomly
select the center of the figure and a random number of vertices that are connected with straight lines. The vertices are determined by their angles and distance with respect to the center of the figure. We generate 5 datasets with different maximum amount of vertices, namely, 4, 9, 14, 19 and 24, and refer to each dataset by this number, e.g. 24-Polar.
- Spiral Dataset (42 × 42 pixels): The curves are generated by growing intervals of a spiral in random directions from a random starting point. The spiral has a random thickness at the different intervals.
- Digs Dataset (42 × 42 pixels): We generate a rectangle of random size and then, we create “digs” of random thicknesses in the rectangle. The digs are created sequentially a random number of times.

Evaluation metrics. From the definition of the problem in Sec. 2, the pixels in the Jordan curve \( \mathcal{F}_X \) are not evaluated. For the rest of the pixels, we use the following metrics:
- Per pixel accuracy (%): It is the average of the accuracy for inside and outside, evaluated separately. In this way, the metric weights the two categories equally, as there is an imbalance of inside and outside pixels.
- Per image accuracy (%): We use a second metric which is more stringent. Each image is considered correctly classified if all the pixels in the image are correctly classified.

Architectures. We evaluate the network architectures that we analyzed theoretically and also other relevant baselines:
- Feed-forward Architectures: We use the dilated convolutional DNN (Dilated) introduced in Sec. 3.1. We also evaluate two variants of Dilated, which are the Ray-intersection network (Ray-int.), which uses a receptive field of \( 1 \times N \) instead of the dilated convolutions, and a convolutional network (CNN), which has all the dilation factors set to \( d = 1 \). Finally, we also evaluate U/Net, which is a popular architecture with skip connections and de-convolutions.
- Recurrent Architectures. We test the ConvLSTM (1-LSTM) corresponding to the architecture introduced in Sec. 3.2. We initialize the hidden and cell states to 0 (inside) everywhere except the border of the image which is initialized to 1 (outside), such that the network can learn to color by expanding the outside region. We also evaluate a 2-layers ConvLSTM (2-LSTM) by stacking one 1-LSTM after another, both with the initialization of the hidden and cell states of the 1-LSTM. Finally, to evaluate the effect of such initialization, we test the 2-LSTM without it (2-LSTM w/o init.), i.e. with the hidden and cell states initialized all to 0. We use backpropagation through time by unrolling 60, 30 or 10 time steps for training (we select the best performing one). For testing, we unroll until there is no change in the output labeling.

Learning. The parameters are initialized using Xavier initialization. The derived parameters we obtained in the theoretical demonstrations obtain 100% accuracy but we do not use them in this analysis as they are not learned from examples. The ground-truth consists on the insideness for each pixel in the image, as in equation. For all experiments, we use the cross-entropy with softmax as
the loss function averaged across pixels. Thus, the networks have two outputs per pixel (note that this does not affect the result that the networks are sufficiently complex to solve insideness, as the second output can be set to a constant threshold of 0.5). We found that the cross-entropy loss leads to better accuracy than other losses. Moreover, we found that using a weighted loss improves the accuracy of the networks. The weight, which we denote as $\alpha$, multiplies the loss relative to inside, and $(1 - \alpha)$ multiplies the loss relative to outside. This $\alpha$ is a hyperparamater that we tune and can be equal to 0.1, 0.2, and 0.4. We try batch sizes of 32, 256 and 2048 when they fit in the GPUs’ memory (12GB), and we try learning rates from 1 to $10^{-5}$ (dividing by 10). We train the networks for all the hyperparameters for at least 50 epochs, and until there is no more improvement of the validation set loss. In the following, we report the testing accuracy for the hyperparameters that achieved the highest per image accuracy at the validation set. We test a large set of hyperparameters (we trained several thousands of networks per dataset), which we report in detail in App. G.

4.2 Results

We evaluate the DNNs trained in the standard manner, *ie.* using backpropagation with the labeled datasets as described previously.

**Intra-dataset Evaluation.** In Fig[3] and c we show per pixel and per image accuracy for the networks trained on the same Polar dataset that are being tested. Dilated, 2-LSTM and UNet achieve a testing accuracy very close to 100\%, but Ray-int. and 1-LSTM perform much worse. Training accuracy of Ray-int. and 1-LSTM is the same as their testing accuracy (Fig[1]a and b). This indicates an optimization problem similar to the cases reported by [36], as both Ray-int. and 1-LSTM are complex enough to generalize. It is an open question to understand why backpropagation performs so differently in each of these architectures that can all generalize in theory. Finally, note that the per pixel accuracy is in most cases very high, and from now on, we only report the per image accuracy.

**Cross-dataset Evaluation.** We evaluate if the networks that have achieved very high accuracies (Dilated, 2-LSTM and UNet), have learnt the general solution of insideness that we introduced in Sec. 3. To do so, we train on one dataset and test on the different one. In Fig[3]d and e, we observe that Dilated and 2-LSTM do not generalize to Polar datasets with larger amount of vertices than the
Figure 6: Visualization of the Units Learnt by Dilation. Each block are the 9 images that produce the maximum activation of a units in a convolutional layer across the test set. The gray dot indicates the location of the unit in the feature map. Fig. 7 shows more examples.

Figure 7: Activation Maps of the Learnt Representations by 2-LSTM. Each row corresponds to a different layer and each column to a different time step. For the first stacked LSTM, we show two different features maps (ex.1 and ex.0).

Polar dataset on which they were trained. Only if the networks are trained in 24-Polar, the networks generalize in all the Polar datasets. The same conclusions can be extracted for UNet (Fig. 6c).

We further test generalization capabilities of these networks beyond the Polar dataset. In this more broad analysis, we also include the CNN and 2-LSTM w/o init, by training them on 24-Polar, Spiral and both 24-Polar and Spiral, and test them on 24-Polar, Spiral and Digs separately. We can see in Fig. 4 that all tested networks generalize to new curves of the same family as the training set. Yet, the networks do not generalize to curves of other families. In Fig. 5, we show qualitative examples of failed segmentations produced by networks trained on 24-Polar and Spiral and tested on the Digs dataset.

Furthermore, note that using a more varied training set (“Both”) does not necessarily lead to better cross-dataset accuracy in all cases. For example, for UNet and 2-LSTM w/o init., training on Polar achieves better accuracy in Digs than when training on “Both”. Also, for Dilated, training on “Both” harms its accuracy: the accuracy drops more than 6% in 24-Polar and Spiral. In this case, the training accuracy is close to 100%, which indicates a problem of overfitting. We tried to address this problem by regularizing using weight decay, but it did not improve the accuracy (App. H).

Understanding the Lack of Generalization. We visualize the networks to understand why the representations learnt do not generalize. In Fig. 6, we analyze different units of Dilated trained on 24-Polar and Spiral. We display two units of the same kernel from the second and sixth layers, by showing the nine images in the testing set that produce the unit to be most active across all images. For each image, we indicate the unit location in the feature map by a gray dot. The visualizations suggest that units of the second layer are tuned to local features (e.g. Unit 19 is tuned to close parallel lines), while in layer 6 they are tuned to more global ones (e.g. Unit 27 captures...
the space left in the center of a spiral). Thus, the units are tuned to characteristics of the curves in the training set rather than to features that could capture the the long-range dependencies to solve insideness, such as the ones we derived theoretically.

In Fig. 7 we display the feature maps of 2-LSTM trained on 24-Polar and Spiral. The figure shows the feature maps of the layers at different time steps. We can see that the network expands the borders of the image, which have been initialized to outside. Yet, it also expands the curve and it is impossible to know the direction of expansion with only local operations, as in one step of the convLSTM. Note that our analytical solution only expands the borders of the image (see Fig. 9 for a visualization of the analytical solution).

Training in Natural Images. Finally, we evaluate if training with the large-scale datasets in natural images solves the generalization issues that we reported before. We use the off-the-shelf state-of-the-art architectures, namely DEXTR [30] for instance segmentation and DeepLabv3+ [5] for semantic segmentation, which have been trained on PASCAL VOC 2012 [9] and ImageNet [35]. These methods fail to determine the insideness for a vast majority of curves, even after fine-tuning in the Both dataset (Deeplabv3+ achieved 36.58% per image accuracy in Both dataset and 2.18% in Digs, see implementation details and qualitative examples in App. K).

5 Learning the Coloring Routine with Small Images

Training recurrent networks with backpropagation through time consists of applying backpropagation to the unrolled version of the network. Recall that for the insideness problem the prediction error is evaluated after the last unrolling step. Thus, depending on the image size and the shape of the curve a large number of unrolling steps may be required to train the network (e.g. the number of unrolling steps for the networks shown in Section 3.2 is bounded by $N^2$). It is well known that for a large number of unrolling steps backpropagation through time has difficulties capturing long-range relationships [4]. Moreover, the memory requirements are proportional to the number of unrolling steps, making it difficult to scale in practice.

A plethora of attempts have been made to alleviate the problems of learning with a large number of unrolling steps, e.g. [15, 32, 11]. Here we introduce a simple yet effective strategy to reduce the number of unrolling steps to learn insideness. Our strategy consists of training the network with small images as they require a smaller number of unrolling steps. To test the network with bigger image sizes, we just unroll it a larger number of steps. Since the recurrent network can learn the coloring routine (Section 3.2), it is possible to generalize to larger images and more complex curves. As we have shown before, this routine does not contain long-range operations because it just takes into account $3 \times 3$ neighbourhoods; the long-range relationships are captured by applying the coloring routine multiple times.

Note that by using small images there is the risk that the dataset lacks enough variability and the network does not generalize beyond the few cases seen during training. Thus, there is a trade-off between the variability in the dataset and the number of unrolling steps required. In the following, we explore this trade-off and show that adjusting the image size and the number of unrolling steps leads to large gains of generalization accuracy.

5.1 Experimental Setup

The Random Walk Dataset. Polar, Spiral and Digs datasets are hard to adapt to small images without constraining the dataset variability. We introduce a new family of curves which we call the Random Walk dataset. The algorithm to generate the curves is based on a random walk following the connectivity conditions of the Jordan curve we previously introduced. It starts by selecting a random location in the image and at every iteration chooses a valid direction with uniform probability among all possible directions. If there are no available directions to expand in, the algorithm backtracks until there are, and continues until the starting point is reached. In Fig. 8, we show several examples of the curves in images of different sizes.

Architecture. In order to show the generality of our results, we test the 2-LSTM and another recurrent neural network which we denote as RNN. We conducted an architecture search to find a recurrent network that succeeded: a convolutional recurrent neural network with a sigmoidal hidden layer and an output layer that is backwardly connected to the hidden layer. The kernel sizes are $3 \times 3$ and $1 \times 1$.
LSTM Tested on 42x42pix size

RNN Tested on 42x42pix size

Examples of Curves of the Random Walk Dataset

Figure 8: Datasets and Results of Controlling the Number of Unrolling Steps. (a) Images of the curves of the Random Walk dataset used to train the recurrent networks with different image sizes. Each row correspond to a different image size. Cross-dataset evaluation (per image accuracy) using the Polar, Spiral and Digs testing sets for (b) 2-LSTM and (c) RNN networks. Evaluation of generalization accuracy based on increasing numbers of unrolling steps for (d) 2-LSTM and (e) RNN network.

for the hidden and output layers, respectively, and we use 5 kernels. Observe that this network is sufficiently complex to solve the insideness problem, because it is the network introduced in App. E with an additional layer and connections.

Learning. We train both the 2-LSTM and the RNN on the Random Walk dataset on 7 different image sizes (from $8 \times 8$ to $42 \times 42$ pixels) and test it with the original images of the 24-Polar, Spiral and Digs datasets ($42 \times 42$ pixels). To explore the effect of the number of unrolling steps, for each training image size we train the networks using 5, 10, 20, 30 and 60 unrolling steps.

5.2 Results

In Fig. 8a and c, we show the cross-dataset accuracy for the 2-LSTM and RNN, respectively, for different training image sizes. The optimal number of unrolling steps is optimized for each training image size. To evaluate the cross-dataset generalization accuracy we apply the trained models to the full size images of the Polar, Spiral, and Digs datasets, using 60 unrolling steps (larger number of unrolling steps does not improve the accuracy). Note that when the networks are trained with small images ($14 \times 14$ is best for RNN, $18 \times 18$ is best for 2-LSTM), at least 80% of the images in all the datasets are correctly segmented. This is a massive improvement of the cross-dataset accuracy compared with the networks trained with large image sizes, as shown in previous section (Fig. 4) and confirmed here again with the Random Walk dataset (less than 40% of the images are correctly segmented in Spiral and Digs datasets).

In Fig. 8b and e, we show the performance of 2-LSTM and RNN, trained with their optimal training image size, when varying the number of unrolling steps. We can observe that both networks generalize with a small number of unrolling steps and fail to generalize as the number of unrolling steps is increased. Also, note that as expected, 2-LSTM is more robust than the RNN to large numbers of unrolling steps during training. These positive results demonstrate that training with smaller images and number of unrolling steps is extremely useful in enabling different types of recurrent networks to learn a general solution of insideness across different families of curves not seen during training.

Furthermore, the feature maps of both 2-LSTM and RNN show that the solution implemented by the network is the Coloring Routine, as the features maps are a non-binary version of the theoretical solution in Fig. 9, rather than the learned solution with big images (Fig. 7), which does not generalize because it expands the border of the curve. Note that the Coloring Routine emerges in the network and we do not enforce it in any way besides training the network with small images and small number of unrolling steps. Note that we could enforce the network to learn the Coloring Routine by providing the ground-truth produced by the routine at each step rather than waiting until the last step. In App. F we show that this per-step strategy also leads to successful cross-dataset generalization accuracy. This
Figure 9: Visualization of Convolutional LSTM with the Mathematically Derived Parameters. We can see that only the border of the image (outside) is propagated, and not the curve, as in the learnt solution.

result is however less interesting as it requires the per-step ground-truth derived from the analytical solution.

6 Conclusions and Future Work

We have shown that DNNs with dilated convolutions and convolutional LSTM networks with few units are sufficiently complex to solve the insideness problem for any given curve. Yet, when using the standard training strategies, the units in these networks become specialized to detect characteristics of the curves in the training set and only generalize to curves of the same family as the training. We introduced a strategy to alleviate this limitation by showing that recurrent networks learn the coloring routine. The coloring routine breaks the evaluation of long-range relationships into local operations and when trained with small images, it generalizes substantially better to new families of curves as it alleviates the well-known difficulties of training recurrent networks with a large number of unrolling steps.

We also hope that this research will help establish the reductionist approach to understand DNNs, such that specific problem components are understood by minimizing the interference of other factors. In future work, we plan to use the reductionist approach to tackle other important aspects of segmentation beyond insideness (e.g. the discontinuity of segments and the hierarchical structure of segments). Also, we hope that this research can lead to improvements of state-of-the-art in image segmentation and in other applications that require understanding of complex spatial relationships. Insideness can be especially helpful when there are few cues available besides the boundaries of the objects, i.e. when there is lack of texture and color, such as in cartoons or sketches, or when objects have different textures and shapes from those seen during training.

Acknowledgements. We are grateful to Tomaso Poggio for his insightful advice and warm encouragement. We also thank Shimon Ullman and Pawan Sinha for helpful feedback and discussions. This work is supported by the Center for Brains, Minds and Machines (funded by NSF STC award CCF-1231216), Fujitsu Laboratories Ltd. (Contract No.40008401) and the MIT-Sensetime Alliance on Artificial Intelligence.

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A Number of Digital Jordan Curves

We now introduce a procedure to derive a lower bound of the number of Jordan curves in an image. We represent an image of size $N \times N$ pixels by using a grid graph (square lattice) with $N \times N$ vertices. We employ 4-adjacency for black pixels and corresponding grid points, and 8-adjacency for white pixels and their counterpart. Then, a curve (the set of black pixels) corresponds to a subgraph of the base grid graph (Fig. A.1a).

In this representation, a digital Jordan curve is defined as a subgraph specified by a sequence of vertices $(v_0, v_1, \ldots, v_L)$ satisfying the following conditions [34, 21]:

1. $L \geq 4$,
2. $v_r = v_s$ if and only if $r = s$, and
3. $v_r$ is 4-adjacent to $v_s$ if and only if $r \equiv s \pm 1 \pmod{L + 1}$.

Note that conditions 1 and 2 defines a cycle [13] in a grid graph. Therefore, any digital Jordan curve is a cycle but not vice versa. Figure A.1b shows examples of cycles that are not digital Jordan curves.

The numbers of all cycles in grid graphs of different sizes were computed up to $27 \times 27$ vertices [17, 1], and we utilize this result to get lower bounds for the number of digital Jordan curves with the following considerations.

Although a cycle in a grid graph is not necessarily a digital Jordan curve as shown above, we can obtain a digital Jordan curve in a larger image from any cycle by “upsampling” as shown in Fig. A.2a. Note that there are other digital Jordan curves than the ones obtained in this manner (therefore we get a lower bound with this technique). See Fig. A.2b for examples.

We also consider “padding” shown in Fig. A.3 to assure that a digital Jordan curve does not contain the border of an image (this is what we assume in the main body of the paper).

Taking everything into consideration, we can obtain a lower bound of the number of digital Jordan curves in an $N \times N$ image that does not contain border pixels utilizing the above-mentioned result [17, 1], upsampling and padding. Table 1 shows lower bounds obtained in this way. For example, starting with the row 2 of [18] in [1] (this represents the number of all cycles in the grid graph with $3 \times 3$ vertices), we get a lower bound 13 for the number of digital Jordan curves in $5 \times 5$ images by considering the upsampling and get the same number as a lower bound for the number of digital Jordan curves that do not contain border pixels in $7 \times 7$ images by considering the padding.
Figure A.2: “Upsampling” Operation and Its Limitations. (a) Depiction of “upsampling” operation. (b) Digital Jordan curves that cannot be obtained by the upsampling shown in (a). (Left) The issue is the place of the digital Jordan curve. We can get the same curve on the upper-left, upper-right, lower-left and lower-right corners but cannot get the one in the center. (Right) The issue is the length of the side. We cannot get a side with 4 vertices (4 pixels) nor with even number vertices (pixels) in general.

Figure A.3: Depiction of “Padding”.

B Insideness with Dilated Convolutional Networks

We first introduce a feed-forward convolutional DNN for which there exist parameters that reproduce equation 3. Then, we show that one of the layers in this network can be better expressed with multiple dilated convolutions.

B.1 Convolutional DNN to Implement the Ray-Intersection Method

The smallest CNN that we found that implements the ray-intersection method has 4-layers. As we show in the following, the first two layers compute \( \vec{X}(i,j) \cdot \vec{X}(i+1,j) \), and the last two layers compute the parity. We use \( H^{(k)} \in \mathbb{R}^{N \times N} \) and \([\cdot]_+\) to denote the activations of the units at the \( k \)-th layer, and the ReLU activation function, respectively. Fig. B.4 depicts the architecture.

First and Second Layer: Inner product. For the sake of simplicity, we only use horizontal rays, but the network that we introduce can be easily adapted to any ray direction. The first layer implements all products needed for the inner products across all rays in the image, ie. \( X_{i,j} \cdot X_{i+1,j}, \forall (i,j) \). Note that there is exactly one product per pixel, and each product can be reused for multiple rays. For convenience, \( H_{i,j}^{(1)} \) represents the product in pixel \( (i,j) \), ie. \( H_{i,j}^{(1)} = X_{i,j} \cdot X_{i+1,j} \). Since the input consists of binary images, each product can be reformulated as

\[
H_{i,j}^{(1)} = \begin{cases} 
1 & \text{if } X_{i,j} = X_{i+1,j} = 1 \\
0 & \text{otherwise}
\end{cases}.
\]

This equality can be implemented with a ReLU: \( H_{i,j}^{(1)} = [1 \cdot X_{i,j} + 1 \cdot X_{i+1,j} - 1]_+ \). Thus, \( H^{(1)} \) is a convolutional layer with a \( 2 \times 1 \) kernel that detects the intersections shown in Fig. 1. This layer can also be implemented with a standard convolutional layer with a \( 3 \times 3 \) kernel, by setting the unnecessary elements of the kernel to 0.

The second layer sums over the products of each ray. To do so, we use a kernel of dimension \( 1 \times N \) with weights equal to 1 and bias equal to 0, ie. \( H_{i,j}^{(2)} = I_{1 \times N} \cdot H_{i,j}^{(1)} = \vec{X}(i,j) \cdot \vec{X}(i+1,j) \), in which \( I_{1 \times J} \) denotes the
We use this parity network to all units in $H^{(3)}$ via convolutions, reproducing the network for each unit. Since a ray through a closed curve in an $N \times N$ image can not have more than $N$ crossings, $C$ is upper bounded by $N$. Thus, the third layer has $3N/2$ kernels, and both the third and output layer are convolutions with a $1 \times 1$ kernel. At this point we have shown that the DNN explained above is feasible in practice, as the number of kernels is $O(N)$, and it requires no more than 4 convolutional layers with ReLUs. The network has a layer with a kernel of size $1 \times N$, and next we show that this layer is equivalent to several layers of dilated convolutions of kernel size $3 \times 3$.

### B.2 Dilated Convolutions to Implement the $1 \times N$ kernel

We use $*_{d}$ to denote a dilated convolution, in which $d$ is the dilation factor. Let $H \in \mathbb{R}^{N \times N}$ be the units of a layer and let $K \in \mathbb{R}^{k \times k}$ be a kernel of size $k \times k$. A dilated convolution is defined as follows: $(H \ast_{d} K)_{i,j} = \sum_{x \in \mathbb{Z}} H_{i,dv,j+dw} \cdot K_{v,w}$, in which $H_{i,dv,j+dw}$ is 0 if $i + dv$ or $j + dw$ are smaller than 0 or larger than $N$, i.e., we abuse notation for the zero-padding. Note that in the dilated convolution the kernel is applied in a sparse manner, every $d$ units, rather than in consecutive units. See [44][7] for more details on dilated convolutions.

Recall the kernel of size $1 \times N$ is set to $I_{1 \times N}$ so as to perform the sum of the units corresponding to the ray in the first layer, i.e., $\sum_{0 \leq v < N} H_{i,j,v}^{(1)}$. We can obtain this long-range sum with a series of dilated convolutions using the following $3 \times 3$ kernel:

$$K = \begin{bmatrix} 0 & 0 & 0 \\ 0 & 1 & 1 \\ 0 & 0 & 0 \end{bmatrix}.$$  (5)

First, we apply this $K$ to the $H^{(1)}$ through $*_{1}$ in order to accumulate the first two entries in the ray, which yields: $(H^{(2)})_{i,j} = (H^{(1)} \ast_{1} K)_{i,j} = \sum_{0 \leq v \leq 1} H_{i,j,v}^{(1)}$. As shown in Fig. B.4a, to accumulate the next entries of the ray, we can apply $K$ with a dilated convolution of dilation factor $d = 2$, which leads to $(H^{(3)})_{i,j} = \sum_{0 \leq v < 4} H_{i,j,v}^{(1)}$. To further accumulate more entries of the ray, we need larger dilation factors. It can be seen in Fig. B.4b that these dilation factors are powers of 2, which yield the following expression:

$$(H^{(l)})_{i,j} = (H^{(l-1)} \ast_{2^{l-2}} K)_{i,j} = \sum_{0 \leq v < 2^{l-2}} H_{i,j,v}^{(1)}.$$  (6)
Observe that when we reach layer $l = \log_2(N) + 2$, the units accumulate the entire ray of length $N$, i.e. $\sum_{0 \leq v < N} \mathcal{H}^{(l)}_{i,v}$. Networks with dilation factors $d = 2^l$ are common in practice, e.g. [44] uses these exact dilation factors.

In summary, DNNs with dilated convolutions can solve the insideness problem and are implementable in practice, since the number of layers and the number of kernels grow logarithmically and linearly with the image size, respectively.

C Parity Network by [36]

To calculate the parity of each unit’s value in $H^{(2)}$, we borrow the DNN introduced by Shalev-Shwartz et al. (namely, Lemma 3 in the supplemental material of [36]). This network obtains the parity of any integer bounded by a constant $C$. The network has $\frac{3C}{2}$ hidden units with ReLUs and one output unit, which is 1 if the input is even, 0 otherwise. Since such a network requires an upper bound on the number whose parity is being found, we define $C$ as the maximum number of times of a horizontal ray can cross $\mathcal{F}_X$. This number can be regarded as an index to express the complexity of the shape.

There is a subtle difference between the network introduced by [36] and the network we use in the paper. In [36], the input of the network is a string of bits, but in our case, the sum is done in the previous layer, through the dilated convolutions. Thus, we use the network in [36] after the sum of bits is done, i.e. after the first dot product in the first layer in [36].

To calculate the parity, for each even number between 0 and $C$ (0 included), $\{2i \mid 0 \leq i \leq \lfloor C/2 \rfloor\}$, the network has three hidden units that threshold at $(2i - \frac{1}{2})$, $2i$ and $(2i + \frac{1}{2})$, i.e. $-0.5, 0, 0.5, 1.5, 2, 2.5, 3.5, 4, 4.5, \ldots$. The output layer linearly combines all the hidden units and weights each triplet of units by $2, -4$ and $2$. Observe that when the input is an odd number, the three units in the triplet are either all below or all above the threshold. The triplets that are all below the threshold contribute 0 to the output because the units are inactive, and the triplets that are all above the threshold also contribute 0 because the linear combination is $2(2i - \frac{1}{2}) - 4(2i) + 2(2i + \frac{1}{2}) = 0$. For even numbers, the triplet corresponding to that even number has one unit below, equal and above the threshold. The unit that is above the threshold contributes 1 to the output, yielding the parity function.

D Implementing the Coloring Routine with a Convolutional LSTM

Here we prove that a ConvLSTM can implement the coloring routine, namely, the iteration of the expansion and the blocking operations. A ConvLSTM applied on an image $X$ is defined as the following set of layers (see [43] for a comprehensive introduction to the ConvLSTM):

$$
I^t = \sigma \left( W^{zi} \ast X + W^{hi} \ast H^{t-1} + b^i \right),
$$

$$
F^t = \sigma \left( W^{zf} \ast X + W^{hf} \ast H^{t-1} + b^f \right),
$$

$$
C^t = \tanh \left( W^{zc} \ast X + W^{hc} \ast H^{t-1} + b^c \right),
$$

$$
O^t = \sigma \left( W^{zo} \ast X + W^{ho} \ast H^{t-1} + b^o \right),
$$

$$
H^t = O^t \odot C^t + I^t \odot \tilde{C}^t,
$$

where $I^t, F^t, C^t, O^t$ and $H^t \in \mathbb{R}^{N \times N}$ are the activation of the units of the input, forget, cell state, output and hidden layers at $t$, respectively. Note that $C^t$ has been decomposed with the help of the auxiliary equation defining $\tilde{C}^t$. Note also that each of these layers use a different set of weights that are applied to $X$ and to $H^t$ denoted as $W \in \mathbb{R}^{N \times N}$ with superindices that indicate the connections between layers, e.g. $W^{zi}$ are the weights that connect $X$ to $I$. Similarly, the biases are denoted as $b \in \mathbb{R}^{N \times N}$ with the superindices indicating the layers. The symbols $\ast$ and $\odot$ denote the (usual, not dilated) convolution and the element-wise product, respectively. Finally, $\sigma$ and $\tanh$ are the sigmoid and the hyperbolic tangent, which are used as non-linearities.

We can see by analyzing equation [11] and equation [12] that the output layer, $O^t$, back-projects to the hidden layer, $H^t$. In the coloring algorithm, $E^t$ and $B^t$ are related in a similar manner. Thus, we define $O^t = E^t$ (expansion) and $H^t = \frac{1}{2}B^t$ (blocking), as depicted in Fig. [3]. The $\frac{1}{2}$ factor will become clear below, and it does not affect the correctness. We initialize $H^0 = \frac{1}{2}B^0$ (recall $B^0$ is 1 for all pixels in the border of the image and 0 for the rest). We now show how to implement the iteration of the expansion and the blocking operations with the ConvLSTM:
We introduce here a convolutional recurrent network that uses sigmoids as non-linearities. Since it is a ConvLSTM, because \( \tanh(0) = 0 \), we can obtain \( C_t = \lim_{q \to \infty} \sigma(q) = 1 \). Also, when all neighboring elements of the unit are 0, then no expansion occurs because \( O_{t,j} = \lim_{q \to \infty} \sigma(-\frac{q}{2}) = 0 \).

**(ii) Blocking, \( H^t \):** To stop the outside region from expanding to the inside of the curve, \( H^t \) takes the expansion output \( O^t \) and sets the pixels at the curve’s location to 0 (inside). This is the same as the element-wise product between \( O^t \) and the element-wise “Boolean not” of \( X \), which is denoted as \( \neg X \). Thus, the blocking operation can be implemented as \( H^t = \frac{1}{2}(O^t \odot \neg X) \). Observe that if \( C^t = X \), this is equal to equation 12 of the LSTM, because tanh(0) = 0 and tanh(1) = 1/2, i.e.,

\[
H^t = O^t \odot \tanh(C^t) = \frac{1}{2}O^t \odot \neg X. \tag{14}
\]

We can obtain \( C_t = \neg X \), by imposing \( I^t = \neg X \) and \( C^t = I^t \), as shown in Fig. 2b. To do so, let \( W^{si} = -qI_{1 \times 1}, W^{bi} = 0_{N \times N} \), and \( b^i = \frac{2}{q}I_{N \times N} \), and equation 13 becomes the following expression:

\[
I^t = \lim_{q \to \infty} \sigma \left( -qI_{1 \times 1} * X + \frac{q}{2}I_{N \times N} \right). \tag{15}
\]

Observe that when \( q \) tends to infinity, we have \( I_{t,j} = \lim_{q \to \infty} \sigma(\frac{q}{2}) = 1 \) when \( X_{i,j} = 0 \) and \( I_{t,j} = \lim_{q \to \infty} \sigma(-\frac{q}{2}) = 0 \) when \( X_{i,j} = 1 \), which means \( I^t = \neg X \). Next, to obtain \( C^t = I^t \), we set \( W^{si} = W^{bi} = W^{bc} = 0_{N \times N}, b^i = -qI_{1 \times N} \) and \( b^c = qI_{N \times N} \). This leads to the desired result:

\[
F^t = \lim_{q \to \infty} \sigma(-qI_{N \times N}) = 0_{N \times N}, \tag{16}
\]

\[
\tilde{C}^t = \lim_{q \to \infty} \tanh(qI_{N \times N}) = I_{N \times N}, \tag{17}
\]

Thus, the coloring method can be implemented with a network as small as one ConvLSTM with one kernel. A network with more than one kernel and multiple stacked ConvLSTM can also solve the insideness problem for any given curve. The kernels that are not needed to implement the coloring method can be just set to 0 and the ConvLSTM that are not needed should implement the identity operation, i.e. the output layer is equal to the input. To implement the identity operator, equation 11 can be rewritten in the following way:

\[
O^t = \lim_{q \to \infty} \sigma \left( qI_{1 \times 1} * X - \frac{q}{2}I_{N \times N} \right) \tag{18}
\]

where \( W^{ho} = 0_{1 \times 1} \) is to remove the connections with the hidden units, and \( q \) is the constant that tends to infinity. Observe that if \( X_{i,j} = 1 \), then \( O^t = \lim_{q \to \infty} \sigma(q/2) = 1 \). If \( X_{i,j} = 0 \), then \( O^t \) is asymptotically close to 0 or 1, otherwise the coloring routine would fade after many steps. It is easy to check that \( h_{t+1,j} = \sigma \left( q \left( \sum_{k \in N_{ij}} h_k - 5X_{i,j} - 1/2 \right) \right) \) implements the coloring routine, where \( q \) is the factor that ensures saturation of the sigmoid.

**E Coloring Routine with a Sigmoidal Convolutional RNN**

There are other recurrent networks simpler than a ConvLSTM that can also implement the coloring algorithm. We introduce here a convolutional recurrent network that uses sigmoids as non-linearities. Since it is a convolutional network, for the sake of simplicity we just describe the operations done to obtain an output pixel in a step. The network has only one hidden layer, which also corresponds to the output of the network. Let \( \{h_{k}^t\}_{k \in N_{ij}} \) be the hidden state of the output pixel indexed by \( i,j \) and its 4-neighbourhood, at step \( t \). Let \( X_{i,j} \) be the only relevant input image pixel. A necessary condition is that the outputs of the sigmoid should asymptotically be close to 0 or 1, otherwise the coloring routine would fade after many steps. It is easy to check that \( h_{t+1,j} = \sigma \left( q \left( \sum_{k \in N_{ij}} h_k - 5X_{i,j} - 1/2 \right) \right) \) implements the coloring routine, where \( q \) is the factor that ensures saturation of the sigmoid.

**F Dataset Generation**

In Fig. 2, we show more examples of curves in the datasets. In the following we provide a more detailed description of the algorithms to generate the curves:
Examples of Jordan Curves of Each Dataset

Figure F.5: Datasets. Images of the curves used to train and test the DNNs. Each row correspond to a different dataset.

- **Polar Dataset** (32 × 32 pixels): We use polar coordinates to generate this dataset. We randomly select the center of the figure and a random number of vertices that are connected with straight lines. These lines are constrained to follow the definition of digital Jordan curve in Sec. 2 in the main paper (and App. A in this supplementary material). The vertices are determined by their angles, which are randomly generated. The distance with respect to the center of the figure are also randomly generated to be between 3 to 14 pixels away from the center. We generate 5 datasets with different maximum amount of vertices, namely, 4, 9, 14, 19 and 24. We refer to each of these datasets as Polar with a prefix with the amount of vertices.

- **Spiral Dataset** (42 × 42 pixels): The curves in these data set are generated from a random walk. First, a starting position is chosen uniformly at random from \([10, 20] \times [10, 20]\). Then, a segment of the spiral is built in the following way: a random direction (up, down, left, right) and a random length \(r\) from 3 to 10 are chosen so that the walk is extended by turning \(r\) pixels in the given direction. However, such extension can only happen if adding a random thickness \(t \in \{1, 2, 3, 4\}\) to both sides of this segment does not cause self intersections. These segments are added repeatedly until there is no space to add a new segment without violating the definition of a Jordan curve.

- **Digs Dataset** (42 × 42 pixels): We generate a rectangle of random size and then, we create “digs” of random thicknesses in the rectangle. The number of “digs” is a random number between 1 to 10. The digs are created sequentially and they are of random depth (between 1 pixel to the length of the rectangle minus 2 pixels). For each new “dig”, we made sure to not cross previous digs by adjusting the depth of the “dig”.

### G Hyperparameters

In this Section we report all the tried hyperparameters for all architectures. In all cases, the convolutional layers use zero-padding.

- **Dilated Convolution DNN (Dilated)**: This network was introduced in Sec. 3.1. We use the same hyperparameters as in [44]: 3 × 3 kernels, a number of kernels equal to \(2^l \times \{2, 4, 8\}\), where \(l\) is the number of layers and ranges between 8 to 11, with \(d = 2^l\) (the first layer and the last two layers \(d = 1\)). The number of kernels in the layer that calculates the parity can be \{5, 10, 20, 40, 80\}.

- **Ray-intersection network (Ray-int.):** This is the architecture introduced in Sec. 3.1 which uses a receptive field of \(1 \times N\) instead of the dilated convolutions. The rest of the hyperparameters are as in Dilated.

- **Convolutional DNN (CNN):** To analyze the usefulness of the dilated convolutions, we use the Dilated architecture with all dilation factors \(d = 1\). Also, we try adding more layers than in Dilated, up to 25.

- **UNet:** This is a popular architecture with skip connections and de-convolutions. We use similar hyperparameters as in [33]: starting with 64 kernels (3 × 3) at the first layer and doubling this number after each max-pooling; a total of 1 to 3 max-pooling layers in all the network, that are placed after sequences of 1 or 2 convolutional layers.
- **Convolutional LSTM (1-LSTM):** This is the architecture with just one ConvLSTM, introduced in Sec. 3.2. We use backpropagation through time by unrolling 60, 30 or 10 time steps for training (we select the best performing one). For testing, we unroll until there is no change in the output labeling. We initialize the hidden and cell states to 0 (inside) everywhere except the border of the image which is initialized to 1 (outside).

- **2-layers Convolutional LSTM (2-LSTM):** We stack one convolutional LSTM after another. The first LSTM has 64 kernels, and the hidden and cell states are initialized as in the 1-LSTM.

- **2-layers Convolutional LSTM without initialization (2-LSTM w/o init.):** this is the same as the 2-LSTM architecture the hidden and cell states are initialized to 0 (outside).

**H Additional Experiments of Feed-Forward Networks**

In Fig. 4, we have observed that Dilated trained on both 24-Polar and Spiral datasets, obtains a test accuracy of less than 95% on these datasets while the accuracy in the training set is very close to 100%. We added weight decay in all the layers in order to regularize the network. We tried values between $10^{-5}$ to 1, scaling by a factor of 10. In all these experiments we have observed overfitting except for a weight decay of 1, in which the training never converged.

Also, note that the CNN does not have this overfitting problem. Yet, the number of layers needed is 25, which is more than the double than for Dilated, which is 9 layers. We added more layers to Dilated but the accuracy did not improve.

**I Additional Figures and Visualizations**

Figure I.6: *Training Accuracy in the Polar Dataset.* Intra-dataset evaluation using (a) per pixel accuracy and (b) per image accuracy on the training set, which are very similar to the test accuracy reported in Fig. 3b and c. (c) Intra-dataset evaluation of Unet.
Figure I.7: More examples of Visualization of the Units Learnt by Dilation.
J  Per-step Learning of the Coloring Routine

The Coloring Routine can be learned by enforcing to each step the ground-truth produced by the routine, rather than waiting until the last step. The inputs of a step are the image and the hidden state of the previous step. Recall that the Coloring Routine determines that a pixel is outside if there is at least one neighbor assigned to outside that is not at the curve border. All input cases (64) are depicted in Fig. J.8, leaving the irrelevant inputs for the Coloring Routine at 0. During learning, such irrelevant pixels are assigned randomly a value of 0 or 1.

We could not make any of the previously introduced LSTM networks fit a step of the coloring routine due to optimization problems. Yet, we found that the RNN network was able to learn with the per-step training. The RNN reached 0 training error about 40% of the times after randomly initializing the parameters. After training the RNN in one step, we unroll it and apply it to images of Jordan curves. We can see that with less than 1000 examples the RNN is able to generalize to any of the datasets for more than 99% of the images. This demonstrates the great potential of decomposing the learning to facilitate the emergence of the routine.

K  Networks Pre-trained on Natural Images

We chose two state-of-the-art networks on Instance Segmentation, DEXTR [30] and DeepLabv3+ [5], to investigate their ability in solving the insideness problem.

DEXTR. Deep Extreme Cut (DEXTR) is a neural network used for interactive instance segmentation. We use the pre-trained model on PASCAL VOC 2012 [9] and show some of the qualitative results in Fig. K.9.

DeepLabv3+. This architecture extends DeepLabv3 [7] by utilizing it as an encoder network and adding a decoder network to refine segmentation boundaries. The encoder employs dilated convolution and Atrous Spatial Pyramid Pooling module for feature extraction. We use DeepLabv3+ with Xception backbone pretrained on PASCAL VOC 2012, and fine-tune its last layer with Polar and Spiral datasets for training. The ratio of input image spatial resolution to encoder output image is referred to as output stride and varies according to dilation rates. We use output strides of 8 and 16 as suggested in the paper; loss weight ($\alpha$) of 0.1, 0.2 and 0.4; and initial learning rates from 0.1 to $10^{-5}$ (dividing by 10). We train the network on Polar and Spiral datasets until there is no improvement of the accuracy at the validations set, and we then reduce the learning rate by a ratio of 10 and stop at the next plateau of the validation set accuracy.
Figure K.9: Qualitative Results with DEXTR on the Polar Dataset. We use the publicly available pre-trained DEXTR model \cite{30}. DEXTR uses 4 points marked by the user (indicated with crosses). We report the best found points, two examples of them per image.
Figure K.10: Results of DeepLabv3+ on Polar, Spiral and Digs Datasets. The network is fine-tuned on Polar and Spiral. The results show that the network predicts well most of the pixels except in the borders. For the cross-dataset evaluations in the Digs dataset, the network is not able to generalize.