

LEARNING TO SOLVE $\mathcal{NP}\text{-}\mathsf{COMPLETE}$ PROBLEMS

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OUTLINE

1. Introduction

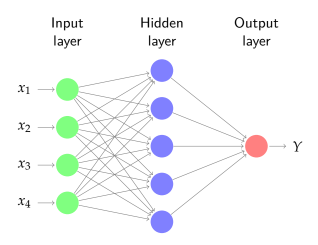
2. Graph Neural Networks

3. A GNN for the Decision TSP

NEURAL SYMBOLIC LEARNING

- Deep learning has been successfully applied to a wide range of domains (images, audio, NLP, RL)
 [6, 14, 7, 5, 4, 1, 8, 9, 12, 13]
- Until now, however, DL has mostly succeeded in applications where the inputs are numerical signals (i.e. pixels)
- How do you design a DL model to learn on a molecule / social network / relational database / multiparticle system / symbolic expressions?
- Classical DL does not understand relational input!
- Combining DL with combinatorial generalization is seen as a key step forward for AI [2]

 A typical ANN is composed of multiple neurons connected according to a given topology



 A recurrent neural network unit is just a neural unit with a self-loop: it is fed with its own output

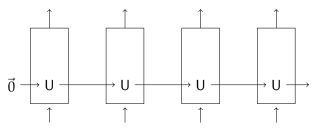


Figure 1: Pictorial representation of the unrolling of a recurrent unit "U" into four iterations. Because the parameters of all four blocks are shared, the resulting network can be thought of as iterating the same operation that many times.



- Parameter sharing is powerful because it allows us to exploit redundancy in the problem domain
- Instead of a RNN, we could just instantiate 4 unrelated neural units and connect them. The model could learn the correct association for 4 fixed iterations, but would not learn the "for loop"
- This would be equivalent to writing a C program that repeats the same subroutine 4 times. It works, but does not generalize for $\neq 4$
- Key insight: if your problem has redundancy, you can use parameter sharing to do the same work while spending much less parameters (easier training)

CONVOLUTIONAL NEURAL NETWORKS

 Convolutional neural networks employ redundancy in space (i.e. learn a number of kernels which will be repeatedly applied throughout the image)



Figure 2: Google's DeepDream [3]

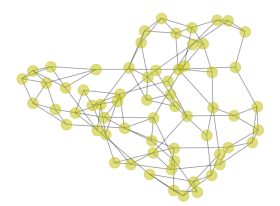
CONVOLUTIONAL NEURAL NETWORKS

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- CNNs assume a grid structure (matrix of pixels). What if this topology changed with the input?



Figure 3: Google's DeepDream [3]

- Suppose you want to perform some computation on a graph $\mathcal{G} = (\mathcal{V}, \mathcal{E})$
- Envision a distributed algorithm in which nodes communicate with one another to learn about their neighborhood



GRAPH NEURAL NETWORKS MESSAGE PASSING IN GRAPHS

- To archive this knowledge, we must allocate a "memory" x_i for each node $v_i \in \mathcal{V}$ (which can be initialized randomly or with zeros)
- Then each node v_i can compute a "message" $msg(x_i)$ to send to each of its neighbors
- Upon receiving a set of messages $X = \{msg(x_j) \mid v_j \in \mathcal{N}(v_i)\}$ from its neighbors v_j , each node v_i can update its memory through some "update" function $x_i' \leftarrow update(x_i, X)$

- Conceivably, if the functions msg() and update() are well-designed, upon many iterations each node could be able to "enrich" its memory with valuable information about its neighborhood
- So what if msg() and update() are learned?
- Concretely: what if msg() is modelled as a MLP and update() as a RNN (specifically LSTM)?

- If we instantiate $msg(): \mathbb{R}^d \to \mathbb{R}^d$ as a MLP and $update(): \mathbb{R}^d, \mathbb{R}^d \to \mathbb{R}^d$ as a LSTM, we can use them as "neural modules" to assemble different neural architectures
- For a given graph $\mathcal{G} = (\mathcal{V}, \mathcal{E})$:
 - 1. Assign a multidimensional vector $x_i \in \mathbb{R}^d$ to each vertex $v_i \in \mathcal{V}$ and collect all vectors into a matrix $\mathbf{X} \in \mathbb{R}^{|\mathcal{V}| \times d}$
 - 2. Compute a $(|\mathcal{V}| \times d)$ matrix of messages $MSGS \leftarrow msg(\mathbf{X})$
 - 3. Multiply by \mathcal{G} 's adjacency matrix: $\mathbf{A} \times MSGS$
 - This yields a $|\mathcal{V}| \times d$ matrix where the i-th line is the sum of all the messages received by vertex v_i
 - 4. Now pass through the update function: $update(\mathbf{X}, \mathbf{A} \times MSGS)$
 - This yields a $|\mathcal{V}| \times d$ matrix where the i-th line is the updated vector of vertex v_i

- Note that this process is completely differentiable and can be accomplished just by function composition and matrix multiplication
- Also note that in principle we can iterate this process by "unrolling" (i.e. composing the function with itself n times)
- What we obtain is a end-to-end differentiable message-passing algorithm on graphs, which outputs a set of refined vertex embeddings at the end [10]

GRAPH NEURAL NETWORKS GRAPH NEURAL NETWORKS

- If we want to learn to compute for example a decision problem on graphs, we can just "reduce" the matrix X to a scalar. For example by appending a mean operator at the end of the pipeline, like mean(X)
- Finally: if we perform gradient descent on $loss = (Y mean(\mathbf{X}))^2$, we can learn to solve a decision problem on graphs, given enough examples
- Note that the input for each problem is just a adjacency matrix $\mathbf{A} \in \mathbb{R}^{|\mathcal{V}| \times |\mathcal{V}|}$, and the output is just a boolean $Y \in \{0,1\}$

- In summary: the core of a GNN is just a RNN applied over a matrix multiplication between an adjacency matrix and a function applied linewise on a matrix of embeddings. Iterated many times
- Or, in math talk:

$$\mathbf{X}^{(t+1)} \leftarrow update(\mathbf{X}^{(t)}, \mathbf{A} \times msg(\mathbf{X}^{(t)}))$$
 (1)

- Selsam et al: You can learn to solve boolean satisfiability (SAT) with the following GNN [11]:
- $\mathbf{A} \in \{0,1\}^{|\mathcal{C}| \times |\mathcal{L}|}$ is an adjacency matrix between clauses and literals

$$\mathbf{C}^{(t+1)} \leftarrow update(\mathbf{C}^{(t)}, \mathbf{A} \times msg_{L \to C}(\mathbf{L}^{(t)}))$$

$$\mathbf{L}^{(t+1)} \leftarrow update(\mathbf{L}^{(t)}, \mathbf{A}^T \times msg_{C \to L}(\mathbf{L}^{(t)}), F(\mathbf{L}^{(t)}))$$
 (2)

• Clauses send messages to literals, literals send messages to clauses and literals send messages to their negated counterparts (i.e. x_1 and $\neg x_1$)

- Given a weighted graph $\mathcal{G} = (\mathcal{V}, \mathcal{E})$ and a number $\mathcal{C} \in \mathbb{R}$:
- Train a GNN to decide whether ${\cal G}$ admits a Hamiltonian route with cost no larger than ${\cal C}$

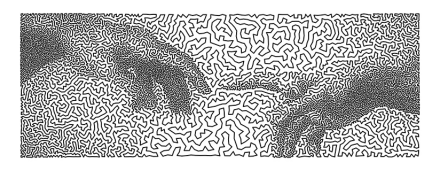


Figure 4: Recreation of Michelangelo's "The Creation of Adam" by Robert Bosch as a TSP solution

- Two problems to address:
 - 1. How to feed the model with numerical information (edge weights) in addition to relational (graph) data?
 - 2. How to tell the model the weight of each edge if only vertices have embeddings?
- Maybe assign embeddings to edges!
- Also initialize each edge embedding with its corresponding weight

- Vertices send messages to edges of which they are endpoints (i.e. vertex v sends messages to $\forall (v_1, v_2) \mid v = v_1 \lor v = v_2$)
- Each edge (v_1, v_2) sends a message to v_1 and a message to v_2
- Over t_{max} message-passing iterations, each edge will become enriched with relational & numerical information relevant to the TSP problem
- This information is destilled into a scalar which is interpreted as that edge's "vote" (i.e. the probability with which it thinks that a route exists)
- Finally, all votes are averaged into the final prediction

Algorithm 1 Graph Neural Network TSP Solver

1: **procedure** GNN-TSP(
$$\mathcal{G} = (\mathcal{V}, \mathcal{E}), \mathcal{C}$$
)
2: $\mathbf{A}[i,j] \leftarrow 1$ iff $(\exists v' | e_i = (v_j, v', w) \lor e_i = (v', v_j, w)) | \forall e_i \in \mathcal{E}, \forall v_j \in \mathcal{V}$
3: $\mathbf{E}[i] \leftarrow \underset{init}{E}(w, \mathcal{C}) | \forall e_i = (s, t, w) \in \mathcal{E}$
4: **for** $t = 1 \dots t_{max}$ **do**

$$(t+1) \quad (t+1) \quad (t) \quad (t) \quad \mathbf{X} \times \underset{msg}{E}(\mathbf{E})$$
6: \mathbf{V}_h , $\mathbf{V} \leftarrow V_u(\mathbf{V}_h, \mathbf{A} \times \underset{msg}{E}(\mathbf{E}))$
6: \mathbf{E}_h , $\mathbf{E} \leftarrow E_u(\mathbf{E}_h, \mathbf{A}^T \times \underset{msg}{V}(\mathbf{V}))$
7: $\mathbf{E}_{\mathbf{logits}} \leftarrow \underset{vote}{E} \begin{pmatrix} t_{max} \\ \mathbf{E} \end{pmatrix}$
8: $\mathbf{P}_{\mathbf{r}}$ prediction \leftarrow sigmoid $(\langle \mathbf{E}_{\mathbf{logits}} \rangle)$

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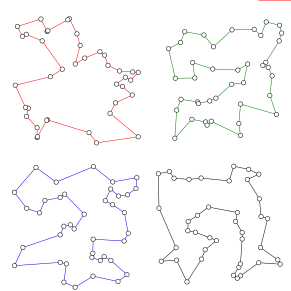
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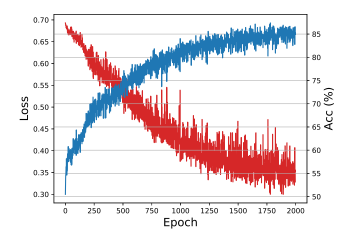
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 - 7. A function $E_{vote}: \mathbb{R}^d \to \mathbb{R}$ to compute a logit probability given an edge embedding (MLP)

- OK, so how do we train it?
- Idea: show the model very similar instances with opposite answers
- Given a graph G with optimal TSP cost C*:
- The model should answer **NO** for inputs $X^- = (\mathcal{G}, 0.98C^*)$ and **YES** for inputs $X^+ = (\mathcal{G}, 1.02C^*)$
- So: create random graphs and feed each graph to the model two times; one with -2% and the other with +2% deviation from the optimal cost
- Advantage: we can create our own training instances!

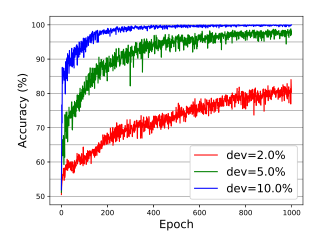
- Batches of $2 \times 8 = 16$ instances
- 128 batches per training epoch
- 2^{20} training instances, to drastically reduce the possibility of overfitting (we're expected to cycle through the entire dataset only after $2^{20-7-4} = 512$ epochs)
- Each instance is a complete, euclidean graph obtained from n points uniformly distributed in the $\frac{\sqrt{2}}{2} \times \frac{\sqrt{2}}{2}$ square
- The size of each graph is chosen uniformly at random between 20 and 40 vertices: $n \sim \mathcal{U}(20,40)$



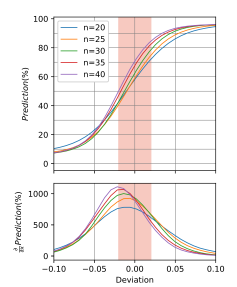
• With -2%, +2% deviations we can achieve 85% training accuracy (80% test) in 2000 epochs



• The greater the deviation, the easier it is to train

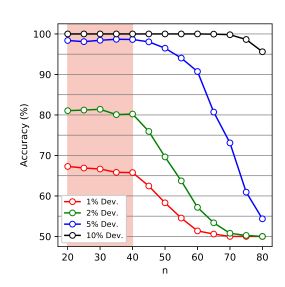


- The probability with which the model thinks that there is a route undergoes something reminiscent of a phase transition as a function of the deviation from the optimal cost
- Curves for big n are higher because the larger the size the larger the probability of (proportionally) cheap routes

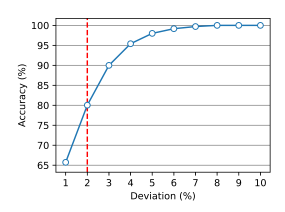


- We can exploit these "acceptance curves" to approximate the TSP optimal cost:
- Intuitively, the closer the model is to absolute uncertainty (50%), the closer we are to the optimal cost
- So: compute lower and upper bounds to the optimal cost and perform a binary search
- We can approximate costs within 1.5% of the optimal cost through on average 8.9 iterations of binary search

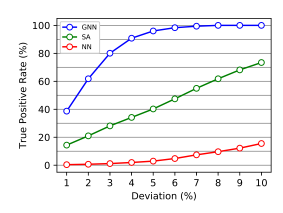
- The model is able to generalize to some extent to larger instances
- Accuracy decreases progressively until the baseline of 50% with increasing n
- Larger deviations yield higher accuracy



 Accuracy increases with deviation, approaching 100% for dev > 8%



We compare the trained model with a predictor for the decision TSP obtained from the solutions yielded by a 1) Nearest Neighbor strategy and 2) a Simulated Annealing strategy (2-exchange)



 The model (which is trained with euclidean instances) can generalize somewhat to random edge weights iff the instances satisfy the triangle inequality

Deviation	Accuracy (%)			
	Euc. 2D	Rand. Metric	Rand.	
1	66	57	50	
2	80	64	50	
5	98	82	50	
10	100	96	50	

Table 1: Test accuracy averaged over 1024 n-city instances with $n \sim \mathcal{U}(20,40)$ for varying percentual deviations from the optimal route cost for differing random graph distributions: bidimensional euclidean distances, "random metric" distances and random distances.

Instance	Size	Relative D GNN	eviation (%) SA
ulysses161 ulysses221	16 22	-22.80 -27.20	+1.94 +1.91
eil51	51	-18.37	+18.07
berlin52	52	-8.73	+21.45
st70	70 70	-11.87	+14.47
eil76 kroA100	76 100	-13.91 -2.00	+19.24
eil101	100	-2.00 -9.93	+30.73 +20.46
lin105	105	+6.37	+17.77

1 These instances had their distance matrix computed according to Haversine formula (great-circle distance).

- Github repository:
 https://github.com/
 machine-reasoning-ufrgs/
 TSP-GNN
- Our library eases the prototyping of GNNs, which can be described briefly

```
= GraphNN(
    'V': d,
    'E': d
},
{
    'EV': ('E','V')
},
    'V_msg_E': ('V', 'E'),
    'E msg V': ('E','V')
},
{
    'V': [
             'mat': 'EV'
             'msg': 'E_msg_V',
             'transpose?': True.
             'var': 'E'
    ],
    1E1 : 1
             'mat': 'EV'.
             'msg': 'V_msg_E',
             'var': 'V'
})
```

- Ideally we want to obtain Hamiltonian routes from the model (even though it was not explicitly trained to do so)
- [11] were able to extract satisfying assignments from the literal embeddings by performing 2-clustering, but we were not successful. Hypothesis: euclidean graphs may be too easy
- Train with other graph distributions (variable connectivity, random weights etc.)
- In principle we can train the model to compute (approximated) optimal costs directly

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