### DeepPSL: End-to-end perception and reasoning

Sridhar Dasaratha<sup>1</sup>, Sai Akhil Puranam<sup>1</sup>, Karmvir Singh Phogat<sup>1</sup>, Sunil Reddy Tiyyagura<sup>1</sup>, Nigel Duffy<sup>2</sup>

<sup>1</sup> EY Global Delivery Services India LLP <sup>2</sup> Ernst & Young (EY) LLP USA

{Sridhar.Dasaratha,Sai.Puranam,Karmvir.Phogat}@gds.ey.com, Sunil.Tiyyagura@gds.ey.com,Nigel.P.Duffy@ey.com

#### Abstract

We introduce DeepPSL a variant of probabilistic soft logic (PSL) to produce an end-to-end trainable system that integrates reasoning and perception. PSL represents first-order logic in terms of a convex graphical model - hinge-loss Markov random fields (HL-MRFs). PSL stands out among probabilistic logic frameworks due to its tractability having been applied to systems of more than 1 billion ground rules. The key to our approach is to represent predicates in first-order logic using deep neural networks and then to approximately back-propagate through the HL-MRF and thus train every aspect of the first-order system being represented. We believe that this approach represents an interesting direction for the integration of deep learning and reasoning techniques with applications to knowledge base learning, multi-task learning, and explainability. Evaluation on three different tasks demonstrates that DeepPSL significantly outperforms state-of-the-art neuro-symbolic methods on scalability while achieving comparable or better accuracy.

#### **1** Introduction

Many machine learning problems involve rich and structured domains with numerous dependenbetween its elements. Statistical relational cies [Richardson and Domingos, 2006; learning (SRL) Koller et al., 2007] methods seek to represent these dependencies and create graphical models using rule based representations. A fundamental challenge faced by SRL approaches is balancing scalability with the expressivity of the dependency structure.

[Bach *et al.*, 2017] introduced HL-MRFs a class of probabilistic graphical models that are both tractable and expressive enabling scalable modelling of rich structured data. In addition they provide a powerful formalism, probabilistic soft logic (PSL) that can define the HL-MRF using first order logic and introduce a scalable inference algorithm. The continuous nature of HL-MRFs enable PSL to scale beyond what was previously feasible for SRL frameworks [Augustine and Getoor, 2018]. Using PSL, problems with tens of millions of ground rules have been solved in minutes [Kouki *et al.*, 2017]. Recent advances using tandem inference make inference tractable even for extremely large systems (billions of random variables) [Srinivasan *et al.*, 2020]. The PSL technique has been successfully applied to problems from various domains ranging from knowledge extraction [Rospocher, 2018], cyberattack prediction [Perera *et al.*, 2018], relational fairness [Farnadi *et al.*, 2020] to hybrid recommender systems [Rodden *et al.*, 2020].

While PSL has significantly advanced SRL methods, there have also been remarkable advances in the field of perception driven by deep learning methods. It would be highly desirable to integrate these perception capabilities into the PSL framework: however currently there is no mechanism to provide this integration. We tackle this challenge with DeepPSL, an end-to-end integration of PSL with deep learning thus achieving a major enhancement of PSL capabilities. DeepPSL fully inherits the scalability of PSL both during inference and training.

The first order expressions in PSL are built from predicates that capture the truth of an assertion with soft truth values in [0,1]. For instance, "HasClaws" and "HasStripes" could be predicates that represent whether claws and stripes are detected in an image. Given some mechanism to compute these truth values and combined with knowledge of animal attributes, PSL can infer whether a specific animal is present in the image. On the other hand neural nets (NN) can learn to identify animals directly from an image, typically from large quantities of training data.

With DeepPSL we integrate these two approaches: some of the predicates are replaced by neural nets and the input data (for e.g., text or image) is processed through a NN to generate the predicate values which are then used by PSL for inference. End-to-end training of this architecture on training data for a given task then permits the NN to learn concepts without any data on the concept itself. In contrast to PSL where one must define the predicate, the training in DeepPSL directly learns predicates that are optimized for the task at hand. Further, these concepts can now be utilized in other tasks where we may have only limited or no task specific data.

Training of this architecture poses significant challenges as it requires back-propagation through an HL-MRF that does not have continuous derivatives. This optimization problem cannot be solved by adapting existing convex optimization methods but solving it is critical to integrating deep learning and PSL. The key contribution of our work is a novel and nonobvious approach to back-propagate through the HL-MRF to learn the parameters of these deep networks. The proposed algorithm enables end-to-end training of DeepPSL which in turn helps fully realize the benefits of the proposed architecture.

We evaluate the efficacy and performance of DeepPSL on three different tasks: digit addition, semi-supervised classification and entity resolution. Experiment results demonstrate the superior scalability of DeepPSL over other state-of-the-art neuro-symbolic approaches while achieving similar or better accuracy.

#### 2 Related Work

Relational Neural machine (RNM) [Marra *et al.*, 2020], an extension of Deep logic models (DLM) [Marra *et al.*, 2019], model reasoning using a Markov random field and backpropagate through that field to learn underlying neural models. Unlike DeepPSL, RNMs do not require backpropagation through an arg min and do not allow any learned values to be used directly in logical rules. Rather they add potentials that couple the learned values to output variables which must be either observed or latent in the Markov random field potentially resulting in a large increase in the number of latent variables. DLM and RNM are related to semantic-based regularization [Diligenti *et al.*, 2017], logic tensor networks [Donadello *et al.*, 2017] and neural logic machines [Dong *et al.*, 2019] which allow logical constraints to constrain the learning of deep networks.

Neural theorem prover [Rocktäschel and Riedel, 2016] is an end- to-end differentiable prover. TensorLog [Cohen *et al.*, 2020] is a recent framework to reuse the deep learning infrastructure of TensorFlow to perform probabilistic logical reasoning. Neither of these methods model predicates using deep learning. [Hu *et al.*, 2016] presents an iterative distillation method that transfers structured information of first-order logic rules into the weights of the NNs. [Gridach, 2020] generalized the approach to include rules built using PSL.

DeepProbLog [Manhaeve et al., 2018] augments the probabilistic logic programming language ProbLog [Raedt et al., 2007] by incorporating neural predicates, and makes predictions by employing marginal inference and sampling. DeepProbLog and other methods such as NeurASP [Yang et al., 2020] do not scale well as they rely on the computationally expensive possible world semantics. DeepStochLog [Winters et al., 2022] achieves better scalability as compared to DeepProbLog and NeurASP using stochastic definite clause grammars. In contrast, DeepPSL scales to significantly larger systems with millions of ground rules by leveraging the continuous nature of HL-MRFs that cast MAP inference as a convex optimization [Pryor et al., 2022] propose a neuro-symbolic problem. approach based on an energy based modeling extension of the PSL framework. Their work uses a novel "energy loss" that doesn't require back propagating through a convex optimization problem. Our approach allows for arbitrary convex differentiable loss functions including all of the most commonly used losses.

End-to-end training of a DeepPSL model requires solving a bi-level optimization problem. The techniques discussed in [Sinha et al., 2017; Ghadimi and Wang, 2018; Dempe, 2018] for solving a bi-level optimization computes gradient of the loss function which needs computation of inverse of the Hessian that is expensive to compute at each iteration. Other methods consider neural network layers consisting of a variety of optimization problems: arg min and arg max problems [Gould et al., 2016], quadratic programming problems [Amos and Kolter, 2017; Lee et al., 2019], convex problems [Agrawal et al., 2019], cone programs [Agrawal et al., 2020]. All of these methods require that the optimization functions have continuous derivatives and make use of second derivatives to allow backpropagation through the optimization problems. HL-MRFs do not have continuous derivatives and therefore are not amenable to these approaches.

#### 3 Background

#### 3.1 HL-MRFs: Hinge Loss Markov Random Fields

HL-MRFs are defined with k continuous potentials  $\phi = \{\phi_1, \dots, \phi_k\}$  of the form:

$$\phi_j(\boldsymbol{x}, \boldsymbol{y}) = (max\{l_j(\boldsymbol{x}, \boldsymbol{y}), 0\})^{p_j}$$
(1)

where  $\phi_j$  is a potential function of n free random variables  $\boldsymbol{y} = \{y_1, \ldots, y_n\}$  conditioned on n' observed random variables  $\boldsymbol{x} = \{x_1, \ldots, x_{n'}\}$ , each random variable can take soft values between [0, 1]. The function  $l_j$  is linear in  $\boldsymbol{y}$  and  $\boldsymbol{x}$  and  $p_j \in \{1, 2\}$ .<sup>1</sup> Collecting the definitions from above, a hinge-loss energy function f is defined as

$$f_{\boldsymbol{\theta}}(\boldsymbol{x}, \boldsymbol{y}) = \sum_{j=1}^{k} \theta_j \phi_j(\boldsymbol{x}, \boldsymbol{y})$$
(2)

where  $\boldsymbol{\theta} = (\theta_1, \dots, \theta_k)$  and  $\theta_j$  is a positive weight corresponding to the potential function  $\phi_j$ . A HL-MRF over random variables  $\boldsymbol{y}$  and conditioned on random variables  $\boldsymbol{x}$  is a probability density defined as

$$P(\boldsymbol{y}|\boldsymbol{x}) = \frac{1}{Z(\boldsymbol{x})} \exp\left(-f_{\boldsymbol{\theta}}(\boldsymbol{x}, \boldsymbol{y})\right)$$
(3)

where  $Z(\boldsymbol{x})$  is the partition co-efficient. Maximum a posteriori (MAP) inference finds the most probable assignment to the free variables  $\boldsymbol{y}$  given the observed variables  $\boldsymbol{x}$ . MAP inference is done by maximizing the probability density  $P(\boldsymbol{y}|\boldsymbol{x})$  while satisfying the constraint that the random variable  $\boldsymbol{y} \in [0,1]^n$ . Since the normalizing function Z in (3) is not a function of  $\boldsymbol{y}$ , maximizing  $P(\boldsymbol{y}|\boldsymbol{x})$  is equivalent to minimizing the energy function f, i.e.,

$$\arg\max_{\boldsymbol{y}\in[0,1]^n} P(\boldsymbol{y}|\boldsymbol{x}) \equiv \argmin_{\boldsymbol{y}\in[0,1]^n} f_{\boldsymbol{\theta}}(\boldsymbol{x},\boldsymbol{y})$$
(4)

<sup>1</sup>In this work,  $p_j = 2$  is used for quadratic hinge-loss.

Critically the function f is convex in y, for a given x, allowing for tractable inference even for very large HL-MRFs. In this study, the inference problem is solved by employing stochastic gradient descent<sup>2</sup> (SGD) algorithm. However, one may employ other alternative algorithms such as distributed optimization algorithm, alternating direction method of multipliers (ADMM), as discussed in [Bach *et al.*, 2017].

#### 3.2 PSL rules

PSL uses first order logic as a template language for HL-MRFs. A PSL program defines a set of rules in first order logic. These rules in the PSL program are grounded over the base of ground atoms each of which represents an observation or unknown of interest. These ground atoms are associated with random variables (x, y) and can take any value in [0, 1]. Each ground rule is then translated into a weighted hinge-loss potential. The sum of these potentials defines a HL-MRF, and minimizing the HL-MRF conditioned over x gives values for the inferred predicates y.

It is beyond the scope of this paper to provide a detailed description of how first order logic rules are used as a template language for HL-MRFs, see [Bach *et al.*, 2017] for further details.

#### 4 DeepPSL

#### 4.1 Deep Learning based Predicates

The key difference between PSL and DeepPSL is that some of the predicates are modeled with deep neural networks (DNNs). In PSL, the observed predicates x are available through a knowledge base, while in DeepPSL a feature vector u is processed through DNNs to compute some of the predicates. Typically, there is no data available on these predicates to learn their corresponding DNNs. Hence, end-to-end training of DeepPSL system is needed to learn these predicates.

#### 4.2 Learning

The key problem that needs to be solved is to determine how to train this system end-to-end. In the proposed DeepPSL framework, the features u are first processed through a neural net with tunable weights  $\omega$  to generate estimates of xwhich are predicates for the PSL. The estimates of predicates in the DeepPSL are modeled by a deep neural network  $p(u; \omega)$ . These predicates then go through PSL inference to produce the final values of the random variables y. For endto-end training, we need to enable backpropagation through the PSL inference. Since PSL inference is a convex optimization problem, there is no direct way to backpropagate and update the weights of the predicate network. We now describe our solution to address this problem.

#### **Optimization objective**

The prime objective of training this end-to-end learning model is to determine weights  $w = (\omega, \theta)$  such that the HL-MRFs inference yields variables y which are close to their true values  $\hat{y}$  in the training data. These free variables y represent the outputs of DeepPSL. For example, a y might represent a belief that a given image contains a zebra.

We want to obtain good outputs or predictions where "good" is measured by a loss relative to their true values  $\hat{y}$ . We restrict our analysis here to HL-MRFs in which all y correspond to outputs.

In order to measure if the inferred values y are close enough to the true values  $\hat{y}$ ; let us consider a differentiable convex loss function:

$$\mathbb{R}^n \times \mathbb{R}^n \ni (\hat{\boldsymbol{y}}, \boldsymbol{y}) \mapsto L(\hat{\boldsymbol{y}}, \boldsymbol{y}) \in \mathbb{R}$$
(5)

The DeepPSL inference problem (4) is approximated with soft constraints <sup>3</sup> and  $w = (\omega, \theta)$  as

$$\boldsymbol{y}^* = \operatorname*{arg\,min}_{\boldsymbol{y}} \quad \tilde{f}(\boldsymbol{u}, \boldsymbol{w}, \boldsymbol{y})$$
 (6)

where

$$\tilde{f}(\boldsymbol{u}, \boldsymbol{w}, \boldsymbol{y}) = f_{\boldsymbol{\theta}}(\boldsymbol{p}(\boldsymbol{u}; \boldsymbol{\omega}), \boldsymbol{y}) + \sum_{i=1}^{n} \underline{\gamma_i} (\max\{0, -y_i\})^2 + \sum_{i=1}^{n} \overline{\gamma_i} (\max\{0, y_i - 1\})^2$$

with fixed  $\overline{\gamma_i}, \underline{\gamma_i} > 0$ . Therefore, the weight training problem is set up as a nonlinear optimization

$$\min_{\boldsymbol{w},\boldsymbol{y}} L(\hat{\boldsymbol{y}},\boldsymbol{y})$$
subject to  $\boldsymbol{y} = \operatorname*{arg\,min}_{\bar{\boldsymbol{y}}} \tilde{f}(\boldsymbol{u},\boldsymbol{w},\bar{\boldsymbol{y}})$ 
(7)

#### **Gradient Following Algorithm**

We develop a gradient descent procedure for solving the nonlinear optimization (7). This task is challenging because we need to back-propagate through the arg min. The most direct approach involves inverting the Hessian  $\nabla_{yy} \tilde{f}(u, w, y)$ which is not well-defined for HL-MRFs as they do not have continuous derivatives.

We take an alternative approach which avoids this pitfall. We assume that the functions L and f are differentiable. Furthermore, we assume that the gradient  $\nabla_y \tilde{f}$  and the neural network p are locally Lipschitz continuous. In general, DNNs are designed to be trained using gradient based techniques and that requires DNNs to be locally Lipschitz continuous, see [Rockafellar, 1981; Scaman and Virmaux, 2018; Jordan and Dimakis, 2020]. Therefore, these assumptions are general enough and the proposed technique is applicable to a wide class of problems.

Consider the neural network weights  $w_{t-1}$  such that  $y_t = \arg \min_{y} \tilde{f}(u, w_{t-1}, y)$ . The objective is to find a  $w_t$  such that

$$L(\hat{\boldsymbol{y}}, \boldsymbol{y}_{t+1}) < L(\hat{\boldsymbol{y}}, \boldsymbol{y}_t)$$
 where  $\boldsymbol{y}_{t+1} = \operatorname*{arg\,min}_{\boldsymbol{y}} \tilde{f}(\boldsymbol{u}, \boldsymbol{w}_t, \boldsymbol{y})$ 

To this end, we first linearly approximate the constraint  $\tilde{y}_{t+1} = \arg \min_{\boldsymbol{u}} \tilde{f}(\boldsymbol{u}, \boldsymbol{w}, \boldsymbol{y})$  in the neighborhood of  $\boldsymbol{y}_t$  by

<sup>&</sup>lt;sup>2</sup>The SGD optimization in PSL suffers from a drawback that hard constraints cannot be strictly enforced. However, this limitation can be circumvented by employing ADMM in place of SGD.

<sup>&</sup>lt;sup>3</sup>The constraints in (4) are incorporated using Lagrange multipliers in a fairly standard way.

using continuous dependence<sup>4</sup> of  $\tilde{y}_{t+1}$  on w and that is given by

$$\tilde{\boldsymbol{y}}_{t+1} = \boldsymbol{y}_t - \delta \nabla_{\boldsymbol{y}} \tilde{f}(\boldsymbol{u}, \boldsymbol{w}, \boldsymbol{y}_t)$$
(8)

for sufficiently small  $\delta > 0$ . Therefore, using the approximation (8), the optimization (7), in the neighborhood of  $y_t$ , translates to

$$\underset{\boldsymbol{w}}{\arg\min} L(\hat{\boldsymbol{y}}, \boldsymbol{y}_t - \delta \nabla_{\boldsymbol{y}} \tilde{f}(\boldsymbol{u}, \boldsymbol{w}, \boldsymbol{y}_t))$$
(9)

and that is linearly approximated, in the neighborhood of  $y_t$ , to

$$\boldsymbol{w}_{t} = \operatorname*{arg\,max}_{\boldsymbol{w}} \nabla_{\boldsymbol{y}} \tilde{f}(\boldsymbol{u}, \boldsymbol{w}, \boldsymbol{y}_{t}) \cdot \nabla_{\boldsymbol{y}} L(\hat{\boldsymbol{y}}, \boldsymbol{y}_{t})$$
(10)

It is worth noting that if  $\nabla_{\boldsymbol{y}} \tilde{f}(\boldsymbol{u}, \boldsymbol{w}_t, \boldsymbol{y}_t) \cdot \nabla_{\boldsymbol{y}} L(\hat{\boldsymbol{y}}, \boldsymbol{y}_t) > 0$  then  $L(\hat{\boldsymbol{y}}, \tilde{\boldsymbol{y}}_{t+1}) < L(\hat{\boldsymbol{y}}, \boldsymbol{y}_t)$ . On the other hand if  $\nabla_{\boldsymbol{y}} \tilde{f}(\boldsymbol{u}, \boldsymbol{w}_t, \boldsymbol{y}_t) \cdot \nabla_{\boldsymbol{y}} L(\hat{\boldsymbol{y}}, \boldsymbol{y}_t) \leq 0$  then local optimality is attained at  $\boldsymbol{w}_{t-1}$ . Furthermore, to ensure constraint satisfaction at each iteration, the local linear approximation  $\tilde{\boldsymbol{y}}_{t+1}$  is replaced with the inference optimization

$$\boldsymbol{y}_{t+1} = \operatorname*{arg\,min}_{\boldsymbol{y}} \tilde{f}(\boldsymbol{u}, \boldsymbol{w}_t, \boldsymbol{y}) \tag{11}$$

Recall that  $\lim_{h\to 0} \frac{g(v+hz)-g(v)}{h} = \nabla g(v) \cdot z$  and therefore, (10) can be rewritten as

$$\boldsymbol{w}_{t} = \underset{\boldsymbol{w}}{\operatorname{arg\,min}} \quad \tilde{f}(\boldsymbol{u}, \boldsymbol{w}, \boldsymbol{y}_{t} - \alpha \nabla_{\boldsymbol{y}} L(\hat{\boldsymbol{y}}, \boldsymbol{y}_{t})) - \tilde{f}(\boldsymbol{u}, \boldsymbol{w}, \boldsymbol{y}_{t})$$
(12)

for sufficiently small  $\alpha > 0$ .

#### Regularization

A PSL potential  $\phi_j$  corresponding to a rule j, defined in (1), translates in DeepPSL setup to

$$\phi_j(\boldsymbol{u}, \boldsymbol{\omega}, \boldsymbol{y}) = \max \left( l_j(\boldsymbol{p}(\boldsymbol{\omega}; \boldsymbol{u}), \boldsymbol{y}), 0 \right)^{p_j}$$

where  $l_j$  is a linear function,  $p_j \in \{1, 2\}$ ,  $p(\omega; u)$  is a neural network with weights  $\omega$  and input u. Note that, for a certain  $\tilde{\omega}$ , the potential  $\phi_i$  does not trigger, i.e.,

$$\phi_j(\bar{\boldsymbol{u}}, \tilde{\boldsymbol{\omega}}, \bar{\boldsymbol{y}}) = 0$$
 for any  $\bar{\boldsymbol{u}} \in \mathcal{D}$  and  $\bar{\boldsymbol{y}} \in [0, 1]^n$ 

where  $\mathcal{D}$  is the dataset, and therefore, there will be no updates to the neural network weights  $\tilde{\omega}$  from the potential  $\phi_j$ . This loss of weight updates might lead to locally optimal solutions to the training optimization (7), and that may be avoided by adding a penalty term to the optimization (12) for each potential  $\phi_j$  as

$$\psi_j(\boldsymbol{u}, \boldsymbol{\omega}, \boldsymbol{y}) = \mu \left( l_j(\boldsymbol{p}(\boldsymbol{u}; \boldsymbol{\omega}), \boldsymbol{y}) \right)^2 \quad \text{with} \quad \mu > 0 \quad (13)$$

The penalty term  $\psi_j$  penalizes the objective function, in case, the hinge loss potential  $\phi_j$  does not trigger. The optimization (12) with the regularizer is given by

$$\boldsymbol{w}_{t} = \underset{\boldsymbol{w}}{\operatorname{arg\,min}} \begin{array}{l} f(\boldsymbol{u}, \boldsymbol{w}, \boldsymbol{y}_{t} - \alpha \nabla_{\boldsymbol{y}} L(\hat{\boldsymbol{y}}, \boldsymbol{y}_{t})) \\ \\ - \tilde{f}(\boldsymbol{u}, \boldsymbol{w}, \boldsymbol{y}_{t}) + \mu \Omega(\boldsymbol{u}, \boldsymbol{w}, \boldsymbol{y}_{t}) \end{array}$$
(14)

where

$$\Omega(\boldsymbol{u}, \boldsymbol{w}, \boldsymbol{y}_t) = \sum_{j=1}^k \psi_j(\boldsymbol{u}, \boldsymbol{\omega}, \boldsymbol{y}_t - \alpha \nabla_{\boldsymbol{y}} L(\hat{\boldsymbol{y}}, \boldsymbol{y}_t)) - \psi_j(\boldsymbol{u}, \boldsymbol{\omega}, \boldsymbol{y}_t)$$

for k potentials, and  $\mu > 0$ .

These two optimizations (11) and (14) are executed alternatively until convergence in Algorithm 1.

Algorithm 1 Joint optimization: backpropagating loss to the neural network

Initialization:  $t = 1; \alpha, \eta, \mu > 0; N, T \ge 1$ .

Neural network weights  $w_0$  are initialized using standard techniques.

while 
$$t \leq T$$
 do  
 $y_t = \arg \min_y \tilde{f}(u, w_{t-1}, y)$  {MAP inference}  
 $w_t = w_{t-1}$   
for  $i = 1, ..., N$  do  
 $w_t = \eta \nabla_{w_t} [\tilde{f}(u, w_t, y_t - \alpha \nabla_y L(\hat{y}, y_t)) - \tilde{f}(u, w_t, y_t) + \mu \Omega(u, w_t, y_t)]$   
end for  
 $t = t + 1$   
end while

## 4.3 Scalability

DeepPSL fully inherits the scalability of PSL, that is, they have the same Big O behavior. The first step in DeepPSL inference consists of a forward pass through the NNs to compute the groundings, only adding time linear in number of ground terms. The second step is standard PSL inference, that is extremely efficient and scales linearly in number of potentials, see Section 3.1.

DeepPSL training requires alternate execution of gradient descent step for minimizing loss (14) and an inference step. The gradient descent step can become memory intensive. As the loss function can be rewritten as summation of losses for each of the grounding, we address the problem using gradient accumulation. Weight updates are made after accumulating gradients from all the grounded potentials. Preserving the inherent scalability of PSL inference and addressing the challenges during training makes DeepPSL highly scalable.

#### **5** Experimental Evaluation

We evaluate DeepPSL on three tasks – T1: a digit addition task that shows that DeepPSL can learn predicates that are hard to specify in PSL, T2: a document classification relational problem that is of moderate scale and T3: a challenging large scale entity resolution problem. We compare our method with state-of-the-art neuro-symbolic methods and in some cases, with other methods that are more specific to the task. Each of these tasks contains train, validation and test splits in the corresponding dataset(s). We select the model and hyper-parameters, see Table 1, that give the highest performance on the validation set. The selected model is then

<sup>&</sup>lt;sup>4</sup>Please note that, in general, convexity of a differentiable function f is not sufficient to ensure continuous dependence of  $\arg\min_{\boldsymbol{y}} \tilde{f}(\boldsymbol{u}, \boldsymbol{w}, \boldsymbol{y})$  on  $\boldsymbol{w}$ . However,  $\arg\min_{\boldsymbol{y}} \tilde{f}(\boldsymbol{u}, \boldsymbol{w}, \boldsymbol{y}) + \nu ||\boldsymbol{y} - \boldsymbol{y}_t||^2$  for any  $\nu > 0$  is augmented to ensure uniqueness of the solution and so continuous dependence on  $\boldsymbol{w}$ .

evaluated on the test set to report the performance. The performance metric is reported with a 95% confidence interval, calculated by repeating each experiment multiple times. The experiments are performed on a MacBook Pro with 2.6GHz Intel i7 processor having 6 cores.

Table 1: Hyperparameters used for DeepPSL

Inference parameters	T1	T2	T3
Optimizer	SGD	SGD	SGD
Learning rate	5e-3	5e-3	1e-2
Loss change threshold	1e-6	5e-3	1e-1
Max iterations	5000	1000	5000
$\underline{\gamma_i}, \overline{\gamma_i}$	20	20	20
Training Parameters *			
Optimizer	Adagrad	Adam	Adagrad
$\alpha$	5e-5	1	5e-3
$\mu$	1e-3	12e-2	0
Update steps $(N)$	2	10	2
Epochs $(T)$	10	75	100

\* Rule weights are initialized randomly by drawing samples from a normal distribution  $\mathcal{N}(1.0, 0.1)$ .

#### 5.1 T1: Addition of handwritten digits

The goal of this task is to predict the sum of digits present in two MNIST images<sup>5</sup> [Manhaeve *et al.*, 2018], a relatively simpler problem that is already well addressed by several neuro-symbolic methods. Note that the training data provides only the sum of the digits, and the digit labels are not provided. Hence, it is not possible to directly learn or specify a predicate to classify individual images, making it difficult for PSL to solve this problem. We investigate whether DeepPSL can learn the image predicate and achieve performance comparable to other neuro-symbolic approaches.

The datasets for this problem are generated following the procedure described in [Winters *et al.*, 2022]. Rules provided in Figure 1 are used to add two digits in DeepPSL system. We use the predicate DIGIT, a convolutional neural network (CNN), to recognize the digits present in the input images (Img1, Img2). DeepPSL is trained end-to-end to learn the rule weights as well as the CNN parameters, by minimizing the cross-entropy loss between the inferred sums (S) and the ground truth.

The CNN consists of two convolution (CONV) layers with 32 and 64 filters of size 3 and stride 1 with ELU activation. Max pool layer with size 2 is applied on the output of last CONV layer. This is followed by two fully connected layers with 128 and 10 nodes on which ELU and softmax activations are applied respectively. A batch size of 16 is used for training. The learning rate for rule weights and CNN are decayed for the 10 epochs according to  $\eta = \eta_0/(E+1)$  where,  $\eta_0$  (5e–4 for rule weights and 1e–3 for CNN parameters) is the initial learning rate and *E* is the epoch number (starting

with 0). A weight decay of 1e-7 is used for CNN parameters from the second epoch of training. Weight decay is not used for rule weights. In the ruleset, the summation constraint is implemented as a soft constraint with a fixed weight 10.

# Addition of handwritten digits rules
$\theta_1: \mathrm{DIGIT}(\mathrm{Img1}, \mathrm{D1}) \land \mathrm{DIGIT}(\mathrm{Img2}, \mathrm{D2}) \land (\mathrm{Img1} \neg = \mathrm{Img2}) \rightarrow \mathrm{SUM}(\mathrm{S})^2$
$\theta_2: DIGIT(\texttt{Img1},\texttt{D1}) \land DIGIT(\texttt{Img2},\texttt{D2}) \land (\texttt{Img1} \neg = \texttt{Img2}) \rightarrow \neg SUM(\texttt{S})^2$
# Summation Constraint SUM $(+S) = 1$

Figure 1: DeepPSL rule set for addition of handwritten digits

The average classification accuracy of DeepPSL over 10 runs is shown in Table 2. The performance numbers for other methods are obtained from [Winters *et al.*, 2022]. We also evaluate the CNN corresponding to DIGIT predicate on the test split of MNIST data set. The CNN achieves an accuracy of 98.2% demonstrating that DeepPSL could successfully learn the predicate even without any explicit data and thus achieves performance comparable to other neuro-symbolic methods.

Table 2: Test accuracy on addition of handwritten digits

NeurASP	DeepProbLog	DeepStochLog	DeepPSL
$97.3\pm0.3$	$97.2\pm0.5$	$97.9\pm0.1$	$96.2\pm0.2$

#### 5.2 T2: Semi-supervised Classification

The goal is to classify unlabeled documents in citation networks given some documents that are labeled. The problem is more challenging than task T1 as there is significantly more relational information available, which in turn poses a challenge to the scalability of neuro-symbolic methods. We use data from the Cora and Citeseer scientific datasets [Yang *et al.*, 2016]. The Cora dataset contains 2708 documents in 7 categories, with 5429 citation links, and each document is represented by indicating the absence or presence of the corresponding word from a dictionary of 1433 unique words. Similarly, the Citeseer dataset contains 3327 documents in 6 categories, with 4591 citation links, and each document is represented by indicating the absence or presence of the corresponding word from a dictionary of 3703 unique words.

Table 3: Dataset splits for semi-supervised classification task

Dataset	Nodes	Split 1 (Train/ Val/ Test)	Split 2 (Train/ Val/ Test)
Cora	2708	140/ 500/ 1000	1708/ 500/ 500
Citeseer	3327	120/ 500/ 1000	2327/ 500/ 500

The predicate CITE(A, B) defines a citation from node A to node B, and the number of neighbors of A are  $n_{\rm A}$  =

<sup>&</sup>lt;sup>5</sup>A detailed explanation on digit addition problem may be found in [Manhaeve *et al.*, 2018].

 $|\{B | CITE(B, A)\}|$ . Furthermore, we define 2-hop neighbors and 3-hop neighbors as, for any A! = B,

 $CITEP(A, B) = CITE(A, C) \land CITE(C, B) \text{ for any node } C,$  $CITEQ(A, B) = CITEP(A, C) \land CITE(C, B) \text{ for any node } C.$ 

In the rule set shown in Figure 2, the predicates CITE, CITEP and CITEQ are directly observed from the data. The inferred predicate LABEL identifies the labels of the documents. The deep learning predicates NEURAL and SIMI-LAR represent the neural net classifier output and the similarity between documents, respectively.

$$\begin{split} & \text{\# Semi-supervised classification rules} \\ & \theta_1 : \text{NEURAL}(\mathbb{A}, \mathbb{Y}) \rightarrow \text{LABEL}(\mathbb{A}, \mathbb{Y}) \\ & \frac{\theta_2}{n_\mathbb{A}} : \text{LABEL}(\mathbb{B}, \mathbb{Y}) \wedge \text{SIMILAR}(\mathbb{B}, \mathbb{A}) \wedge \text{CITE}(\mathbb{B}, \mathbb{A}) \rightarrow \text{LABEL}(\mathbb{A}, \mathbb{Y}) \\ & \frac{\theta_3}{n_\mathbb{A}} : \text{LABEL}(\mathbb{B}, \mathbb{Y}) \wedge \neg \text{SIMILAR}(\mathbb{B}, \mathbb{A}) \wedge \text{CITE}(\mathbb{B}, \mathbb{A}) \rightarrow \neg \text{LABEL}(\mathbb{A}, \mathbb{Y}) \\ & \theta_4 : \frac{1}{|\mathbb{B}|} \text{NEURAL}(+\mathbb{B}, \mathbb{Y}) \rightarrow \text{LABEL}(\mathbb{A}, \mathbb{Y}) \{\mathbb{B} : \text{CITE}(\mathbb{A}, \mathbb{B})\} \\ & \theta_5 : \frac{1}{|\mathbb{B}|} \text{NEURAL}(+\mathbb{B}, \mathbb{Y}) \rightarrow \text{LABEL}(\mathbb{A}, \mathbb{Y}) \{\mathbb{B} : \text{CITEP}(\mathbb{A}, \mathbb{B})\} \\ & \theta_6 : \frac{1}{|\mathbb{B}|} \text{NEURAL}(+\mathbb{B}, \mathbb{Y}) \rightarrow \text{LABEL}(\mathbb{A}, \mathbb{Y}) \{\mathbb{B} : \text{CITEQ}(\mathbb{A}, \mathbb{B})\} \\ & \text{\# Constraints} \\ & \text{LABEL}(\mathbb{A}, +\mathbb{Y}) = 1 \end{split}$$

Figure 2: DeepPSL rule set for semi-supervised classification

The predicate NEURAL is represented by a feedforward neural network with an input layer that takes the document features, followed by a hidden layer with 16 nodes (RELU activation), a drop out layer with a rate of 0.2, and a final softmax layer with 7 nodes corresponding to the output classes. The predicate LABEL is represented by a Siamese network composed of two identical networks that share the first layer of the feedforward network, and a distance layer with 16 nodes. We minimize cross entropy loss and use learning rate 2e-3 for rule weights and 2e-2 for NN parameters. The weight decay parameter of Adam optimizer is set to 3e-4 (Split 1) and 3e-5 (Split 2), and is only used for NN parameters. The summation constraint is implemented as a soft constraint with a fixed weight 20.

We compare the performance of DeepPSL against various baselines on the Cora and Citeseer datasets, using randomly generated data splits described in Table 3. The results, presented in Table 4, show classification class averaged accuracy on the test sets. The results for GCN<sup>6</sup> [Kipf and Welling, 2017], PSL and DeepPSL are generated by running each method 100 times, while the results for ManiReg [Belkin *et al.*, 2006], SemiEmb [Weston *et al.*, 2012], LP [Zhu *et al.*, 2003], DeepWalk [Perozzi *et al.*, 2014], ICA [Lu and Getoor, 2003], Planetoid [Yang *et al.*, 2016] are taken from [Kipf and Welling, 2017]. As DeepStochLog training was slow, we ran it only 5 times and don't report error bars. Both splits are evaluated for

GCN, PSL and DeepPSL while other results are reported only for Split 1.

DeepPSL achieves the highest accuracy for both data sets and both splits, outperforming DeepStochLog by a large margin and surpassing even methods that are specialized for semi-supervised learning. The expressivity of PSL for relational systems enables DeepPSL to leverage the labels and features of its neighbors, while the end-to-end training helps optimize the weights of the neural predicates to maximize classification accuracy. When compared to other neurosymbolic methods, DeepPSL demonstrates excellent scalability. DeepProbLog , timed out for both networks while Deep-StochLog training time was  $\sim 50x$  that of DeepPSL.

Table 4: Classification accuracy on test nodes for Task T2

Algorithm	Cora	Citeseer
ManiReg	59.5	60.1
SemiEmb	59.0	59.6
LP	68.0	45.3
DeepWalk	67.2	43.2
ICA	75.1	69.1
Planetoid	75.7	64.7
DeepStochLog	69.4	65
DeepProbLog	timeout	timeout
GCN	$80.08 \pm 0.34$	$67.96 \pm 0.32$
PSL	$62.97 \pm 0.52$	$64.88 \pm 0.38$
DeepPSL	$81.31 \pm 0.28$	$69.11 \pm 0.27$
GCN(Split 2)	$87.46 \pm 0.30$	$75.96 \pm 0.34$
PSL(Split 2)	$85.94 \pm 0.28$	$75.66 \pm 0.32$
DeepPSL(Split 2)	$88.94 \pm 0.25$	$76.01 \pm 0.31$

#### 5.3 T3: Entity Resolution

We perform entity resolution on CiteSeer database [Bhattacharya and Getoor, 2007] using DeepPSL to identify duplicate references to authors and published papers. As the task requires predicting the edges between every pair of author nodes and paper nodes in a large network, the problem results in millions of ground rules posing a major challenge for existing neuro-symbolic approaches.

The data consists of author names, paper titles and relational information such as authorship of papers. There are around 3000 author references and 1500 paper references. We use the train and test splits provided in PSL entity resolution example.<sup>7</sup> We extract a third of data in the provided train set to create a validation set.

For this problem, pair-wise similarity of author names and pair-wise similarity of paper titles provide key information to identify duplicates. Traditionally, this similarity is computed with hand crafted metrics and is used for inference. It has been shown that the performance of different standard string similarity metrics varies greatly based on the application domain [Cheatham and Hitzler, 2013]. With DeepPSL, we do not rely on pre-determined string similarity metrics.

<sup>&</sup>lt;sup>6</sup>https://github.com/tkipf/gcn

<sup>&</sup>lt;sup>7</sup>https://github.com/linqs/psl-examples/tree/master/entity-resolution

Rather, the system learns the optimal way to compute similarities from the provided application specific data.

# Entity resolution rules
$\theta_1$ : AuthorName(A1, N1) $\land$ AuthorName(A2, N2) $\land$ SimName(N1, N2)
$\rightarrow$ SameAuthor (A1, A2) <sup>2</sup>
$\theta_2$ : PaperTitle (P1, T1) $\land$ PaperTitle (P2, T2) $\land$ SimTitle (T1, T2)
$\rightarrow$ SamePaper (P1, P2) <sup>2</sup>
$\theta_3$ : AuthorBlk (A1, B1) $\land$ AuthorBlk (A2, B1) $\land$ AuthorOf (A1, P1) $\land$
$AuthorOf(\texttt{A2},\texttt{P2}) \land SamePaper(\texttt{P1},\texttt{P2}) \rightarrow SameAuthor(\texttt{A1},\texttt{A2})^2$
$\theta_4$ : AuthorBlk (A1, B) $\land$ AuthorBlk (A2, B) $\land$ AuthorBlk (A3, B) $\land$
SameAuthor (A1, A2) $\land$ SameAuthor (A2, A3) $\land$ (A1 $\neg$ = A3) $\land$
$(A2 \neg = A3) \land (A1 \neg = A2) \rightarrow SAMEAUTHOR(A1, A3)^2$
$ heta_5: \neg S$ AMEAUTHOR (A1, A2) $^2$
$\theta_6: \neg SAMEPAPER(P1, P2)^2$
# Identity Constraints
SameAuthor $(A, A) = 1$
SamePaper $(P, P) = 1$

Figure 3: DeepPSL rule set for entity resolution

Rules<sup>7</sup> in Figure 3 incorporate the author name and paper title similarity values in conjunction with other relational information to identify duplicates. The inferred predicates SAMEAUTHOR and SAMEPAPER identify if authors and papers are same respectively. AUTHORNAME, PAPERTITLE, AUTHOROF and AUTHORBLOCK are directly observed from the data. In the DeepPSL system, we associate author name similarity predicate SIMNAME and title similarity predicate SIMTITLE with siamese networks[Koch et al., 2015]. The NN to compute author name similarity takes 56 dimensional character ([A-Za-z .,']) based one hot encoding of the names as input. NN for title similarity operates on mean of 300 dimensional vectors derived from GloVe embeddings (pretrained on Wikipedia 2014 and Gigaword5 corpus.<sup>8</sup>) of the words in the title. Each twin in the architecture has a hidden layer of 50 nodes and distance layer of 50 nodes for both NNs. We minimize cross entropy loss to learn rule weights and weights of NNs for author name and paper title similarities. During training of the DeepPSL system, we do not perform any sampling over the graph. We use learning rate of 5e-2 for NN parameters and 25e-3 for rule weights. Weight decay is not used for rule weights but for NN parameters, 1e-3 is used. The identity constraints are implemented with a fixed weight 10. The model which gives the highest average of F1-scores for same author and same paper identification is selected based on validation set.

After trivial potential removal, there are 6.5 million and 3.1 million ground rules during train and test respectively. The scale of this relational problem is out of scope for neural PLP approaches such as DeepProbLog. We attempted a comparison with DeepStochLog9. However, Deep-StochLog could not scale to the size of this problem, encountering memory errors and timing out even when using only a subset of the rules. We report performance of DeepPSL and PSL over 10 runs in Table 5. PSL setup is built with same ruleset as in DeepPSL but the author name and paper title similarities are computed using metrics mentioned in [Bhattacharya and Getoor, 2007]. PSL-A uses Soft TFIDF with Jaro-Winkler [Cohen et al., 2003] for both author name similarity and paper title similarity. PSL-B uses Soft TFIDF for author names and cosine similarity of GloVe based vectors for paper titles. We also experiment with PSL setups using other standard similarity metrics (as mentioned in [Cheatham and Hitzler, 2013; Gali et al., 2016; Yih and Meek, 2007]): Cosine, Jaccard, Monge Elkan (with Levenshtein). The PSL systems corresponding to these metrics are designated as PSL-C, PSL-J and PSL-M respectively. In each of these setups, we use the same similarity metric for computing author name and paper title similarities. Strings are not preprocessed apart from tokenization when necessary. Error bars are not reported for PSL as there is insignificant variation across runs. DeepPSL outperforms all the PSL se-

Table 5: Performance on Task T3

Algorithm	Author F1 score	Paper F1 score
PSL-A	0.9271	0.8276
PSL-B	0.8958	0.8501
PSL-C	0.7852	0.7656
PSL-J	0.8073	0.7631
PSL-M	0.8008	0.8884
DeepStochLog	timeout	timeout
DeepPSL	$0.9468 \pm 0.0061$	$0.9228 \pm 0.0036$

tups for both author and paper entity resolution, see Table 5. The PSL performance depends on the pre-determined similarity metric used and it is hard to find a similarity metric that yields the highest accuracy. DeepPSL achieves excellent performance by directly learning optimal similarity functions for the entity resolution task. Moreover, these results highlight that DeepPSL scales to a difficult problem which proves to be out of reach for competing methods.

#### 6 **Conclusions and Future Work**

We introduced DeepPSL an end-to-end trainable system that integrates reasoning and perception. We proposed a novel algorithm to enable end-to-end training. Experimental results on three different tasks demonstrated the broad applicability of the method. DeepPSL scales to relational problems that prove to be challenging for competing methods. DeepPSL has some limitations. Firstly, as DeepPSL uses deep networks to model predicates, learning is not convex, and therefore, the proposed approach may suffer from local minima. While we did not observe any significant sensitivity to local minima in our experiments, further research is needed to understand this better. Secondly, the work described in this article does not address latent variables. Future work will report on extensions of DeepPSL to the latent variable case.

#### Disclaimer

<sup>&</sup>lt;sup>8</sup>https://nlp.stanford.edu/projects/glove

The views reflected in this article are the views of the authors <sup>9</sup>https://github.com/MLKULeuven/deepstochlog/tree/main/examples.and do not necessarily reflect the views of the global EY or-

ganization or its member firms.

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#### A DeepPSL Scalability

We demonstrate experimentally that DeepPSL fully inherits the scalability of PSL, that is, they have the same Big O behavior. To this end, we perform run time and scalability analysis for semi-supervised classification task (**T2**) and entity resolution task (**T3**). We report average training and inference time (per epochs) of DeepPSL over 5 random training runs. For task **T2**, we sampled five sub-graphs from the citation network of Cora dataset and report their run time in Figure 4 and Table 6. For task **T3**, we sampled four sub-graphs from the entity resolution dataset and report their run time in Figure 5 and Table 7. For both tasks, the results indicate that the DeepPSL inference scales approximately linearly with number of ground rules.

The DeepPSL inference consists of two steps: The first step is a forward pass through neural network to evaluate neural predicates. The second step is standard PSL inference for determining inferred predicates, that is extremely efficient and scales linearly in number of ground rules. As discussed in Section 3.1 and [Bach *et al.*, 2017], PSL inference easily scales to systems with millions of ground rules by leveraging the continuous nature of HL-MRFs that cast MAP inference as a convex optimization problem.

For example, in task **T2**, the neural predicates NEURAL and SIMILAR are computed by a forward pass through the classifier and then LABEL is obtained by executing PSL inference. Since forward pass through the neural classifier takes negligible time as compared to PSL inference, the DeepPSL inference time will effectively be the same as PSL inference. Hence, DeepPSL inference scales linearly with number of ground rules.

The DeepPSL training algorithm, see Algorithm 1, executes DeepPSL inference step

$$\boldsymbol{y}_t = \operatorname*{arg\,min}_{\boldsymbol{u}} \ \tilde{f}(\boldsymbol{u}, \boldsymbol{w}_{t-1}, \boldsymbol{y})$$

and a gradient descent step

$$\begin{split} \boldsymbol{w}_t &= \eta \nabla_{\boldsymbol{w}_t} \big[ \tilde{f}(\boldsymbol{u}, \boldsymbol{w}_t, \boldsymbol{y}_t - \alpha \nabla_{\boldsymbol{y}} L(\hat{\boldsymbol{y}}, \boldsymbol{y}_t)) \\ &- \tilde{f}(\boldsymbol{u}, \boldsymbol{w}_t, \boldsymbol{y}_t) + \mu \Omega(\boldsymbol{u}, \boldsymbol{w}_t, \boldsymbol{y}_t) \big] \end{split}$$

for neural network training and rule weights learning.

From Figure 4 and Figure 5, we note that DeepPSL time for every epoch also scales linearly with number of ground rules. It is evident from the experiments, see Table 4, that the run time of the gradient descent step is negligible as compared to DeepPSL inference step. Hence, from a computation perspective, the training of DeepPSL is equivalent to performing repeated DeepPSL inferences. The inference time for DeepPSL is approximately the same as PSL, and therefore, the training time for DeepPSL will be approximately equal to the time for repeated PSL inferences. Since PSL inference scales linearly with number of ground rules, DeepPSL training also approximately scales linearly with number of ground rules as observed in experiments, see Figure 4 and Figure 5.

#### **B** DeepPSL Hyper-parameters Study

We conduct experiments for studying effect of various hyperparameters on the convergence of DeepPSL. The following



Figure 4: Computation time of task T2 on Cora dataset

Table 6: Run time statistics of task T2 on Cora dataset

Nodes	Edges	Ground rules	Inf*	$\text{GDs}^{\dagger}$	Total <sup>‡</sup>
500	418	30352	1.98	0.10	2.08
1000	1476	69664	3.26	0.17	3.43
1500	3334	120176	6.06	0.36	6.42
2000	6572	190008	9.15	0.7	9.85
2708	10556	280476	10.73	1.02	11.75

\* DeepPSL Inference time per epochs (in sec).

<sup>†</sup> Gradient descent step time per epochs (in sec).

<sup>‡</sup> Total training time per epochs (in sec).

hyperparameters are considered for the study while training DeepPSL on task **T2** with Cora dataset:

**1.** random initialization of neural network weights and rule weights,

**2.** regularization parameter  $\mu$ ,

**3.** neural network weights's learning rate  $\eta_{\omega}$ ,

4. neural network's hidden nodes and dropout rate.

# B.1 Initialization of neural weights and rule weights

The DeepPSL rule weights are randomly initialized by drawing a sample from normal distribution  $\mathcal{N}(1, 0.1)$  and the neural network weights are initialized using standard techniques. We train the DeepPSL system for task **T2** on Split 2 of the

Table 7: Run time statistics of task T3

Ground rules	Inf Time*	GDs Time^{\dagger}	Total Time $^{\ddagger}$
551220	12.1	1.5	13.6
1011922	25	2.5	27.5
2083797	60.7	4.8	65.5
3143623	81.8	5.5	87.3

\* DeepPSL Inference time per epochs (in sec).

<sup>†</sup> Gradient descent step time per epochs (in sec).

<sup>‡</sup> Total training time per epochs (in sec).



Figure 5: Computation time of task T3

Cora dataset. We take average of cross-entropy loss and validation accuracy over 10 random runs with random weights initialization and random data splits, see Figure 6. These experiments show that DeepPSL is quite robust to random initialization of neural and rule weights.



Figure 6: Task T2 on Split 2 of Cora dataset

#### **B.2** Regularization parameter

In this experiment, we varied the regularization hyperparameter  $\mu$  among the set {0.04, 0.12, 0.2} and found that DeepPSL trains effectively and reaches the desired performance across a broad range of the regularization parameter. As shown in Figure 7, the validation accuracy and cross-entropy loss indicate that DeepPSL does not become trapped in local minima.

#### **B.3** Learning rate of neural network weights

In this experiment, we varied the learning rate of neural network weights  $\eta_{\omega} \in \{0.004, 0.012, 0.1\}$  to examine the robustness of DeepPSL training across different learning rates. We found that while small learning rates slow down training, large learning rates can lead to oscillations, as seen in Figure 8. However, the validation accuracy and cross-entropy loss, see Figure 8, indicate that DeepPSL training is robust for a wide range of learning rate parameters.



Figure 7: Task T2 on Split 1 of Cora dataset



Figure 8: Task T2 on Split 1 of Cora dataset

#### **B.4** Neural network hyperparameters

We varied the number of hidden nodes and dropout rate of the neural classifier and found that DeepPSL training is relatively robust to these hyperparameters, as shown in Table 8. However, the performance of DeepPSL decreases with fewer hidden nodes and higher dropout rates, indicating that this combination of hyperparameters is not sufficient to capture the underlying structure of the data.

Table 8: Classification accuracy of DeepPSL on task **T2** (Split 1 of Cora dataset)

Nds <sup>†</sup> Drp <sup>*</sup>	8	16	24
0	$80.12\pm0.70$	$80.42\pm0.49$	$80.64 \pm 0.35$
0.2	$79.97 \pm 0.60$	$81.85\pm0.43$	$81.76 \pm 0.45$
0.4	$74.23 \pm 0.78$	$79.16 \pm 0.57$	$80.47\pm0.70$

\* Dropout rate in neural classifier.

<sup>†</sup> Hidden node of neural classifier.

#### **C** Toy example for DeepPSL

To explain DeepPSL in a simple and intuitive fashion, we discuss a toy version of the entity resolution problem (explained in Section 5.3) to first elucidate the core concepts of PSL and then introduce the DeepPSL framework. In the toy problem, the goal is to identify the duplicate author references based on similarity of author names. Using the following PSL rules, we express the knowledge that two authors are same if they have similar names and that there is a transitive relation.



Figure 9: DeepPSL rule set for entity resolution

Here, SAMEAUTHOR is an inferred predicate and, AUTHORNAME, SIMNAME are observed predicates. Consider the scenario where there are 3 authors mention with names,

$$\mathcal{L} = \{$$
"Sheldon Cooper", "Cooper Sheldon", "Sheldon C" $\}$ 

with their corresponding IDs in

$$\mathcal{D} = \{ \text{AID1}, \text{AID2}, \text{AID3} \}$$

respectively. The base of ground atoms is given by

$$\mathcal{A} = \begin{cases} x_{ij} = \text{AUTHORNAME}(i, j) \text{ for } i \in \mathcal{D} \text{ and } j \in \mathcal{L}, \\ \bar{x}_{ij} = \text{SIMNAME}(i, j) \quad \text{for } i, j \in \mathcal{L}, \\ y_{ij} = \text{SAMEAUTHOR}(i, j) \text{ for } i, j \in \mathcal{D} \text{ and } i \neq j \end{cases}$$
(15)

The above rules will induce to the following ground rules:

$$\begin{aligned} \theta_1: & x_{ij} \& x_{kl} \& \bar{x}_{jl} \to y_{ik} \\ & \text{for } i, k \in \mathcal{D}, \ i \neq k, \text{ and } j, l \in \mathcal{L} \end{aligned}$$
 (16)

$$\theta_2: \quad y_{ij} \& y_{jk} \to y_{ik}$$
  
for  $i, j, k \in \mathcal{D}$  and  $i \neq j \neq k$  (17)

which is equivalent to the following:

$$\theta_1 : !x_{ij} \parallel !x_{kl} \parallel !\bar{x}_{jl} \parallel y_{ik}, \qquad \theta_2 : !y_{ij} \parallel !y_{jk} \parallel y_{ik}$$
(18)

The disjunction expression (18) is converted to a soft logic using Lukasiewicz t-norm and its corresponding co-norm as the relaxation of the logical AND and OR respectively. Additionally, negation of an atom, "a" is considered as "1 - a". Therefore, the satisfaction of the rules (18) are determined by the following expressions:

$$\min\left\{ (1 - x_{ij}) + (1 - x_{kl}) + (1 - \bar{x}_{jl}) + y_{ik}, 1 \right\},$$

$$\min\left\{ (1 - y_{ij}) + (1 - y_{jk}) + y_{ik}, 1 \right\}.$$
(19)

The rule satisfaction (19) are rewritten as the distance to satisfaction leading to the following hinge-loss potentials:

$$\begin{cases} \phi_{ijkl} = (\max\{x_{ij} + x_{kl} + \bar{x}_{jl} - y_{ik} - 2, 0\})^p \\ \bar{\phi}_{ijk} = (\max\{y_{ij} + y_{jk} - y_{ik} - 1, 0\})^p \end{cases}$$

for  $p \in \{1, 2\}$ . With  $\boldsymbol{x} := (x_{ij}, \bar{x}_{ij}), \boldsymbol{y} := (y_{ij}), \boldsymbol{\theta} := (\theta_1, \theta_2)$ , the energy function is defined as

$$f_{\boldsymbol{\theta}}(\boldsymbol{x}, \boldsymbol{y}) = \theta_1 \sum_{\substack{i,k \in \mathcal{D} \\ i \neq k \\ j,l \in \mathcal{L}}} \phi_{ijkl} + \theta_2 \sum_{\substack{i,j,k \in \mathcal{D} \\ i \neq j \neq k}} \bar{\phi}_{ijk}$$
(20)

The inferred predicates y are obtained by solving a convex PSL inference optimization (4).

For DeepPSL, the string similarity values need not be supplied beforehand. A siamese network can be plugged into the SimName predicate which computes the similarity of strings. We pass the string pairs to the neural network, get the similarity values and feed them as groundings for SimName predicate. Once the groundings are obtained, similar to PSL we minimize the total hinge loss to obtain the inferred values.