

AutoSF: Searching Scoring Functions for Knowledge Graph Embedding

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Abstract—Scoring functions (SFs), which measure the plausibility of triplets in knowledge graph (KG), have become the crux of KG embedding. Lots of SFs, which target at capturing different kinds of relations in KGs, have been designed by humans in recent years. However, as relations can exhibit complex patterns that are hard to infer before training, none of them can consistently perform better than others on existing benchmark data sets. In this paper, inspired by the recent success of automated machine learning (AutoML), we propose to automatically design SFs (AutoSF) for distinct KGs by the AutoML techniques. However, it is non-trivial to explore domain-specific information here to make AutoSF efficient and effective. We firstly identify a unified representation over popularly used SFs, which helps to set up a search space for AutoSF. Then, we propose a greedy algorithm to search in such a space efficiently. The algorithm is further sped up by a filter and a predictor, which can avoid repeatedly training SFs with same expressive ability and help removing bad candidates during the search before model training. Finally, we perform extensive experiments on benchmark data sets. Results on link prediction and triplets classification show that the searched SFs by AutoSF, are KG dependent, new to the literature, and outperform the state-of-the-art SFs designed by humans.

I. INTRODUCTION

Knowledge Graph (KG) [28], [33], [40], as a special kind of graph structure with entities as nodes and relations as edges, is important to both data mining and machine learning, and has inspired various downstream applications, e.g., structured search [7], [33], [36], question answering [25] and recommendation [52]. In KGs, each edge is represented as a triplet with form (*head entity*, *relation*, *tail entity*), denoted as (h, r, t) . A fundamental issue is how to quantize the plausibility of triplets (h, r, t) s [14], [40]. KG embedding (KGE) has recently emerged and been developed as a promising method serving this purpose [6], [18], [19], [24], [29], [30], [39], [46], [54]. Basically, given a set of observed triplets, KGE attempts to learn low-dimensional vector representations of entities and relations so that the plausibility of triplets can be quantized. Scoring function (SF), which returns a score for (h, r, t) based on the embeddings, is used to measure the plausibility. Generally, SF is designed and chosen by humans and it has significant effects on embeddings' quality [20], [28], [40].

Ever since the invention of KGE, many SFs have been proposed in the literature. Let the embedded vectors of h , r and t be \mathbf{h} , \mathbf{r} and \mathbf{t} respectively. TransE [4], a representative embedding model, interprets the triplet (h, r, t) as a translation \mathbf{r} from head entity h to tail entity t , i.e. the embeddings satisfy

$\mathbf{h} + \mathbf{r} = \mathbf{t}$. Variants of TransE like TransH [43] and TransR [21], project the embedding vector into different spaces and enables the embedding to model relationships that are one-to-many, many-to-one or many-to-many. These models are categorized into translational distance models (TDMs). However, as proved in [41], [42], TDMs are not fully expressive and their empirical performance is inferior to other models. RESCAL [30], DistMult [46], ComplEx [39], Analogy [24] and more recently proposed Simple [18], [19], use a bilinear function $\mathbf{h}^T \mathbf{R} \mathbf{t}$ to model the plausibility of triplets, where \mathbf{R} is a square matrix related to relation embedding. These models belong to the bilinear model (BLMs). Different BLMs use different constraints to regularize the relation matrix \mathbf{R} in order to adapt to different datasets. Inspired by the success of deep networks [2], some neural network models (NNMs) have also been explored as SFs, e.g., MLP [7], NTM [34], Neural LP [47] and ConvE [6]. Even though neural networks are powerful and have strong expressive ability, the NNMs do not perform well in KGE domain because of not being well-regularized.

Among the existing SFs, BLM-based ones are the most powerful as indicated by both the state-of-the-art results [19] and theoretical guarantees on expressiveness [18], [41]. However, since different KGs have distinct patterns in relations [32], a SF which adapts well to one KG may not perform consistently well on other KGs. Besides, designing new SFs to outperform state-of-the-art SFs is challenging. Therefore, how to choose and design a good SF for a certain KG is a non-trivial and difficult problem.

Recently, automated machine learning (AutoML) [17], [50] has exhibited its power in many machine learning tasks and applications, e.g. model selection [13], image classification [23], [51] and recommendation [48]. In order to select proper models and hyper-parameters for different tasks, hyperparameter optimization (HPO) has been proposed [11], [13] to effectively and efficiently find better configurations, which previously requires great human efforts. Another hot trend in AutoML is to search better neural networks for deep learning models. Neural architecture search (NAS) [55] has identified networks with fewer parameters and better performance than networks designed by humans.

Inspired by the success of AutoML, we aim to design better and novel data-dependent SFs for KGE. Specifically, we propose automated scoring function (AutoSF) which can automatically search an SF for a given KG. It can not only

reduce human’s effort in designing new SFs, but also make adaptation to different KGs. However, it is not easy to achieve the above goal. When applying AutoML, two important perspectives, i.e. search space, which helps to figure out important properties of the underling problems, and search algorithm, which determines the efficiency of finding better points in the space, need serious consideration. In this work, we have made the following contributions to achieve the goals:

First, we make an important observation over existing SFs, which allows us to represent the BLM-based SFs in a unified form. Based on the unified representation, we formulate the design of SFs as an AutoML problem (i.e. AutoSF), and set up the corresponding search space. The space is not only specific enough to cover good SFs designed by humans, but also general enough to include novel SFs not visited in the literature.

Second, we observe it is common that different KGs have distinct properties on relations that are symmetric, asymmetric, inverse, etc. This inspires us to conduct domain-specific analysis on the KGE models, and design constraints to effectively guide subsequent searches in the space.

Third, we propose a progressive greedy algorithm to search through such a space. We further build a filter to avoid training redundant SFs and a predictor with specifically designed symmetry-related features (SRF) to select promising SFs. The search algorithm can significantly reduce the search space size by capturing the domain specific properties of candidate SFs.

Finally, experimental results on five popular benchmarks on link prediction and triplet classification tasks demonstrate that the SFs searched by AutoSF outperform the start-of-the-art SFs designed by humans. In addition, the searched SFs are KG dependent and new to the literature. We further conduct case study on the searched SFs to provide means for analyzing KGs, which can inspire better understanding of embedding techniques for future researches.

Notations. We denote vectors by lowercase boldface, and matrix by uppercase boldface. A KG contains a set of triplets $S = \{f(h, r, t)g\}$ with $h, t \in E$ and $r \in R$, where E and R are the set of entities and relations, respectively. For simplicity, the embeddings are represented by letters of indices in boldface, e.g. $\mathbf{h}, \mathbf{r}, \mathbf{t}$ are embeddings of h, r, t , respectively, and \mathbf{h}, \mathbf{t} share the same set of embedding parameters \mathbf{e} . $\mathbf{h}\mathbf{a}, \mathbf{b}, \mathbf{c}$ is the triple dot product and can be alternatively represented as $\mathbf{a}^T \text{diag}(\mathbf{b}) \mathbf{c}$, where $\text{diag}(\mathbf{b}) = \mathbf{D}^{\mathbf{b}} \in \mathbb{R}^{d \times d}$ is the diagonal matrix of \mathbf{b} . We denote the complex vector $\mathbf{v} = \mathbf{v}_{re} + i\mathbf{v}_{im} \in \mathbb{C}^d$ with $\mathbf{v}_{re}, \mathbf{v}_{im} \in \mathbb{R}^d$. The conjugate of a complex vector is $\text{conj}(\mathbf{v}) = \mathbf{v}_{re} - i\mathbf{v}_{im}$.

II. RELATED WORKS

A. Knowledge graph embedding (KGE)

Given a set of observed (positive) triplets, the goal of KGE is to learn low-dimensional vector representations of entities and relations so that the plausibility measured by $f(\mathbf{h}, \mathbf{r}, \mathbf{t})$ of

observed triplets (h, r, t) s are maximized while those of non-observed ones are minimized [40]. To build a KGE model, the most important thing is to design and choose a proper SF f , which measures the triplets’ plausibility based on embeddings. Since different SFs have different strengths and weaknesses, the choice of f is critical for the KGE’s performance [20], [40]. A large amount of KGE models with popular SFs follow the same framework (Alg.1) [40] using stochastic gradient descent. At step 5, negative triplets are sampled from $\bar{S}_{(h,r,t)}$, which contains all non-observed triplets for a current positive triplet (h, r, t) , by some fixed distributions [43] or dynamic sampling schemes [54]. Next, the gradients are computed based on the given SF and embeddings, and are used to update the model parameters (step 6). Hinge loss [4] and logistic loss [46] are popularly used as ℓ . In this paper, we use the multi-class loss [19] since it currently achieves the best performance and has little variance.

Algorithm 1 Stochastic training of KGE [40].

Input: training set $S = \{f(h, r, t)g\}$, scoring function f and loss function ℓ ;
1: initialize embeddings \mathbf{e}, \mathbf{r} for each $e \in E$ and $r \in R$.
2: **for** $i = 1, \dots, T$ **do**
3: sample a mini-batch $S_{\text{batch}} \subset S$ of size m ;
4: **for** each $(h, r, t) \in S_{\text{batch}}$ **do**
5: sample m negative triplets $\bar{S}_{(h,r,t)} = \{f(\bar{h}_j, r, t_j)g\}$ for the positive triplet (h, r, t) ;
6: update embedding parameters based on loss ℓ using selected positive and negative triplets;
7: **end for**
8: **end for**
9: **return** embeddings of entities in E and relations in R .

Existing human-designed SFs mainly fall into three types:
Translational distance models (TDMs): The translational approach exploits the distance-based SFs. Inspired by the word analogy results in word embeddings [2], the plausibility is measured based on the distance between two entities, after a translation carried out by the relation. In TransE [4], the SF is defined by the (negative) distance between $\mathbf{h} + \mathbf{r}$ and \mathbf{t} , i.e. $f(\mathbf{h}, \mathbf{r}, \mathbf{t}) = -\|\mathbf{h} + \mathbf{r} - \mathbf{t}\|_1$. Other TDMs-based SFs, e.g., TransH [43], TransR [12], enhance over TransE by introducing extra mapping matrices.

BiLinear models (BLMs): SFs in this group exploit the plausibility of a triplet by the product-based similarity. Generally, they share the form as $f(\mathbf{h}, \mathbf{r}, \mathbf{t}) = \mathbf{h}^T \mathbf{R} \mathbf{t}$ where $\mathbf{R} \in \mathbb{R}^{d \times d}$ is a matrix referring to the embedding of relation r [40], [41]. RESCAL [30] models the embedding of each relation by directly using \mathbf{R} . DistMult [46] overcomes the overfitting problem of RESCAL by constraining \mathbf{R} to be diagonal. ComplEx [39] allows \mathbf{R} and \mathbf{h}, \mathbf{t} to be complex values, which enables handling asymmetric relations. HolE [29] uses a circular correlation to replace the dot product operation, but is proven to be equivalent to ComplEx [16]. Other variants like Analogy [24], Simple [18] regularize the matrix \mathbf{R} in different ways.

Neural network models (NNMs): Neural models aim to

TABLE I

EXISTING SFs COVERED BY OUR SEARCH SPACE. FOR ANALOGY AND SIMPLE, THE EMBEDDING SPLITS INTO TWO PARTS, I.E. $\mathbf{h}^> = [\hat{\mathbf{h}}^>, \mathbf{h}^>]$ AND $d = \hat{d} + d$ (SAME FOR \mathbf{r} AND \mathbf{t}). THE RELATION TYPES ARE SUMMARIZED IN TAB. II.

scoring function	embeddings	definition	relation types that can model
DistMult [46]	symmetric $\mathbf{h}, \mathbf{r}, \mathbf{t} \in \mathbb{R}^d$	$\langle \mathbf{h}, \mathbf{r}, \mathbf{t} \rangle$	symmetric
ComplEx [39] / HolE [29]	$\mathbf{h}, \mathbf{r}, \mathbf{t} \in \mathbb{C}^d$	$\text{Re}(\langle \mathbf{h}, \mathbf{r}, \text{conj}(\mathbf{t}) \rangle)$	symmetric, anti-symmetric, asymmetric, inverse
Analogy [24]	$\hat{\mathbf{h}}, \hat{\mathbf{r}}, \hat{\mathbf{t}} \in \mathbb{R}^{\hat{d}}, \mathbf{h}, \mathbf{r}, \mathbf{t} \in \mathbb{R}^d$	$\langle \hat{\mathbf{h}}, \hat{\mathbf{r}}, \hat{\mathbf{t}} \rangle + \text{Re}(\langle \mathbf{h}, \mathbf{r}, \text{conj}(\mathbf{t}) \rangle)$	symmetric, anti-symmetric, asymmetric, inverse
SimplE [18] / CP [19]	$\hat{\mathbf{h}}, \hat{\mathbf{r}}, \hat{\mathbf{t}} \in \mathbb{R}^{\hat{d}}, \mathbf{h}, \mathbf{r}, \mathbf{t} \in \mathbb{R}^d$	$\langle \hat{\mathbf{h}}, \hat{\mathbf{r}}, \hat{\mathbf{t}} \rangle + \langle \mathbf{h}, \mathbf{r}, \mathbf{t} \rangle$	symmetric, anti-symmetric, asymmetric, inverse

output the probability of the triplets based on neural networks which take the entities' and relations' embeddings as inputs. MLP proposed in [7] and NTN proposed in [34] are representative neural models. Both of them use a large amount of parameters to combine entities' and relations' embeddings. ConvE [6] takes advantage of convolutional neural network to increase the interaction among different dimensions of the embeddings.

As proved in [41], TDMs have less expressive ability than BLMs, which further leads to their inferior empirical performance. Based on the power of deep networks, NNMs are also introduced for KGE. However, due to the huge model complexity and increasing difficulty of training, as well as the lack of domain-specific constraints, their performance is still worse than BLMs [6], [19]. Therefore, we focus on BLMs in the sequel. The most representative BLMs are listed in Tab. I.

B. Automated machine learning (AutoML)

Automated machine learning (AutoML) [17], [50] has recently exhibited its power in easing the usage of and designing better machine learning models. Basically, AutoML can be regarded as a bi-level optimization problem where we need to update model parameters by the training data sets and tune hyper-parameters by the validation data sets. Regarding the success of AutoML, there are two important perspectives:

Search space: This helps to figure out important properties of the underlying learning models and set up the search space for an AutoML problem. First, the space needs to be general enough to cover human wisdom as special cases. However, the space cannot be too general, otherwise searching in the space will be too expensive.

Search algorithm: Unlike convex optimization, there is no universal and efficient optimization tools. Once the search space is determined, efficient algorithms should be developed to search good points in the space.

We take NAS and HPO as examples. The search space in NAS is spanned by network operations, e.g., convolution with different sizes, skip-connections. Various tailor-made algorithms, such as reinforcement learning [55], evolution algorithms [44], and one-shot algorithms [23], [51], have been proposed for efficient optimization. For HPO, Bayesian optimization [11], [13] is usually customized to search the space made up by the hyper-parameters of the learning tools.

This paper is the first step towards automated embedding of knowledge graphs. However, such a step is not trivial since previous AutoML methods used in NAS and HPO cannot be

directly applied to KGE. The main problem is that we need to explore domain-specific properties in defining the search space and designing efficient search algorithm to achieve effectiveness with less cost.

III. THE SEARCH PROBLEM

As mentioned in Sec.II, new designs of SFs have continuously boosted the performance of KGEs in recent years. However, there is no absolute winner among the human-designed SFs. Besides, as different KGs usually exhibit distinct patterns in relations, how to choose a proper SF to achieve

TABLE II
COMMON RELATIONS IN KGs AND RESULTING REQUIREMENTS ON f , AND SEARCH CANDIDATES IN $g(\mathbf{r})$.

common relations	requirements on f	requirements on $g(\mathbf{r})$	examples from WN18/FB15K
symmetric [46]	$f(\mathbf{t}, \mathbf{r}, \mathbf{h}) = f(\mathbf{h}, \mathbf{r}, \mathbf{t})$	$g(\mathbf{r})^> = g(\mathbf{r})$	<i>IsSimilarTo</i> , <i>Spouse</i>
anti-symmetric [29], [39]	$f(\mathbf{t}, \mathbf{r}, \mathbf{h}) = f(\mathbf{h}, \mathbf{r}, \mathbf{t})$	$g(\mathbf{r})^> = g(\mathbf{r})$	<i>LargerThan</i> , <i>Hypernym</i>
general asymmetric [24]	$f(\mathbf{t}, \mathbf{r}, \mathbf{h}) \neq f(\mathbf{h}, \mathbf{r}, \mathbf{t})$	$g(\mathbf{r})^> \neq g(\mathbf{r})$	<i>LocatedIn</i> , <i>Profession</i>
inverse [18]	$f(\mathbf{t}, \mathbf{r}, \mathbf{h}) = f(\mathbf{h}, \mathbf{r}^\theta, \mathbf{t}), \mathbf{r} \neq \mathbf{r}^\theta$	$g(\mathbf{r})^> = g(\mathbf{r}^\theta)$	<i>Hypernym</i> , <i>Hyponym</i>

constraint in training procedure, which motivates us to use AutoML to search based on validation sets performance.

Remark III.2 (General Search Space). *Due to the recent success of deep networks [15] and the approximation ability of multilayer perceptron (MLP) [5], one may want to use a MLP as F for (2). However, the design of the MLP is also a searching problem, which is very time-consuming [55]. Besides, an arbitrarily large MLP will lead to an extremely large space. As verified in [49], the general approximator MLP is a bad choice for NAS and performs worse than those using reinforcement learning [55].*

IV. THE SEARCH STRATEGY

Here, we propose an efficient search strategy to address the AutoSF problem based on domain-specific properties in KGs.

A. Challenges in algorithm design

Same as the other AutoML problems, the search problem of AutoSF is black-box, the search space is huge, and each step in the search is very expensive since both model training and evaluation should be involved. These problems have previously been touched by algorithms such as reinforcement learning [55], Bayes optimization [13] and genetic programming [44]. However, they are not a good choice here since we have domain-specific problems, i.e. expressiveness and invariance, in KGE, which are more challenging.

1) *Expressiveness*: It is clear that not all SFs $g \in G$ (from Definition 2) are equally good. Expressiveness (Definition 3), which means f should be better able to handle common relations in KGs, is of big concern for SFs. Their consequent requirements on f and $g(\mathbf{r})$ are summarized in Tab. II.

Definition 3 (Expressiveness [18], [39], [41]). *If f can handle symmetric, anti-symmetric, general asymmetric, and inverse relations, then f is expressive.*

To ensure that f can handle those common relations, we propose Proposition 1.

Proposition 1. *If $g(\mathbf{r})$ can be symmetric for some $\mathbf{r} \in \mathbb{R}^d$, i.e. $g(\mathbf{r})^> = g(\mathbf{r})$, and skew-symmetric for some $\mathbf{r}^\theta \in \mathbb{R}^d$, i.e. $g(\mathbf{r}^\theta)^> = g(\mathbf{r}^\theta)$. Then the formulated f is expressive. (Proofs in Appendix A).*

With this Proposition and to avoid trivial solutions, we introduce the following constraints on g :

- (C1). $g(\mathbf{r})$ can be symmetric with proper \mathbf{r} and skew-symmetric with proper \mathbf{r}^θ .
- (C2). $g(\mathbf{r})$ has no zero rows/columns, covers all \mathbf{r}_1 to \mathbf{r}_4 , and has no repeated rows/columns.

For (C1), the symmetric property of $g(\mathbf{r})$ determines what kind of relation the given SF can model based on Proposition 1. For (C2), if there are zero rows/columns in g , the corresponding embedding dimensions will be useless. It means that these dimensions will never be optimized during training.

The above constraints are important for finding potentially good candidate $g \in G$, and they play a key role in filtering out the bad g 's for the design of an efficient search algorithm. As in Definition 3 and Proposition 1, we need to deal with Constraint (C1) for expressiveness. It is challenging since g only represents a structure, however the exact check of (C1) relies on the values in \mathbf{r} , which are unknown in advance. Fortunately, we can check it by value assignment. Take the SF in Fig. 2(a) for example. We can see that $g(\mathbf{r})$ can be symmetric by assigning $\mathbf{r}_3 = \mathbf{r}_1$ and $\mathbf{r}_4 = \mathbf{r}_2$ as in Fig. 2(b), and skew-symmetric by setting $\mathbf{r}_3 = -\mathbf{r}_2$ and $\mathbf{r}_4 = -\mathbf{r}_1$ like in Fig. 2(c). This is the key idea of addressing expressiveness.

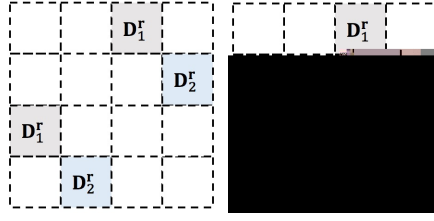
2) *Invariance*: As defined in Sec.III-B, the embeddings are split into 4 parts, i.e. $\mathbf{r} = [\mathbf{r}_1; \mathbf{r}_2; \mathbf{r}_3; \mathbf{r}_4]$. Prior to the embedding training, permuting the \mathbf{r}_i 's will lead to equivalent structure since "1,2,3,4" here are only identity of each component and these components are equivalent at this stage. For example, we can permute $\mathbf{r} = [\mathbf{r}_1; \mathbf{r}_2; \mathbf{r}_3; \mathbf{r}_4]$ into $\mathbf{r} = [\mathbf{r}_2; \mathbf{r}_1; \mathbf{r}_3; \mathbf{r}_4]$. Even though \mathbf{r}_1 and \mathbf{r}_2 change their position, the learned embedding could be the same by changing corresponding values after training. Therefore, the structure of SFs is invariance to permutation of \mathbf{r}_i 's. Similarly, since \mathbf{h} and \mathbf{t} share the same embedding parameters \mathbf{e} , the generated SFs are also equivalent by simultaneously permuting the \mathbf{h}_i 's and \mathbf{t}_i 's. Moreover, if we flip the signs of some \mathbf{r}_i , we can learn equal embeddings by flipping the true value of those \mathbf{r}_i after training. In summary, there exist three kinds of invariance: permuting entity embedding \mathbf{h}_i 's and \mathbf{t}_i 's, permuting relation embedding \mathbf{r}_i 's, and flipping signs. The example of three cases are given in Fig. 2(d-f). Let $\mathbf{h}, \mathbf{r}, \mathbf{t}$ be the embeddings of the SF g_1 and $\mathbf{h}, \mathbf{r}, \mathbf{t}$ be the embedding of another SF g_2 , and g_2 is formed through the invariance changes of g_1 . Then we will have $\mathbf{h}^>_{g_1}(\mathbf{r})\mathbf{t} = \mathbf{h}^>_{g_2}(\mathbf{r})\mathbf{t}$ after the model training. Therefore, it is tedious to train and evaluate the equivalents once we know the performance of one SF among them.

B. Progressive greedy search

As in Sec.III-B, adding one more block into g indicates adding one more nonzero multiplicative term into f , namely

$$f^{b+1} = f^b + s \mathbf{h} \mathbf{h}_i, \mathbf{r}_j, \mathbf{t}_k \mathbf{i}, \quad (7)$$

where $s \in [-1, 1]$ and $i, j, k \in \{1, 2, 3, 4\}$. In order to search efficiently, we propose a progressive greedy algorithm based on the inductive rule (7), which can significantly cut down the



(a) SimpleE.

(b) sym.

(c) skew-sym.

(d) permute \mathbf{h}_i and \mathbf{t}_i .(e) permute \mathbf{r}_i .

(f) flip sign.

Fig. 2. Illustration of *Invariance* and *Expressiveness*. (a) SimpleE model; (b) assign $\mathbf{r}_3 = \mathbf{r}_1, \mathbf{r}_4 = \mathbf{r}_2$; (c) assign $\mathbf{r}_3 = -\mathbf{r}_1, \mathbf{r}_4 = -\mathbf{r}_2$; (d) permute $[\mathbf{h}_1; \mathbf{h}_2; \mathbf{h}_3; \mathbf{h}_4]$ into $[\mathbf{h}_1; \mathbf{h}_3; \mathbf{h}_2; \mathbf{h}_4]$ and do the same for \mathbf{t} ; (e) permute $[\mathbf{r}_1; \mathbf{r}_2; \mathbf{r}_3; \mathbf{r}_4]$ into $[\mathbf{r}_1; \mathbf{r}_4; \mathbf{r}_2; \mathbf{r}_3]$; (f) flip the signs of \mathbf{r}_2 and \mathbf{r}_4 .

search space in a stage-wise manner. The intuition of using (7) to progressively generate SFs it to gradually adjust the relation matrix $g(\mathbf{r})$. However, greedy search usually leads to sub-optimal solutions [38], which can be more serious when faced with the expressive and invariance challenges in AutoSF. Therefore, we enhance the greedy search with a filter and a predictor to specifically deal with the expressiveness and invariance discussed in Sec.IV-A.

Algorithm 2 Progressive greedy search algorithm.

Input: B : number of nonzero blocks in g , a learnable predictor P ;
1: **for** b in 4, 6, 8, ..., B **do**
2: **repeat**
3: randomly select a top- K_1 model $f^{b-2} \in \mathcal{T}^{b-2}$;
4: generate 6 integers $i_1, j_1, k_1, i_2, j_2, k_2 \in \{1, 2, 3, 4\}$
 and s_1, s_2 from $\{-1, 1\}$, and form $f^b = f^{b-2} + s_1 \mathbf{h}_{i_1} \mathbf{r}_{j_1} \mathbf{t}_{k_1}^T + s_2 \mathbf{h}_{i_2} \mathbf{r}_{j_2} \mathbf{t}_{k_2}^T$;
5: **if** f^b satisfies filter Q **then** $H^b \leftarrow H^b \cup \{f^b\}$;
6: **until** $|H^b| = N$
7: select top- K_2 f^b s in H^b based on the predictor P ;
8: train embeddings with the selected f^b s;
9: evaluate the obtained embedding from the selected f^b s;
10: $\mathcal{T}^b \leftarrow \mathcal{T}^b \cup \{f^b\}$ and record f^b and their performance;
11: update the predictor P with records in \mathcal{T}^b .
12: **end for**
13: **return** desired SFs in \mathcal{T}^B .

1) *Complete procedures*: Alg.2 shows our progressive greedy algorithm. As in Definition 2, let the number of nonzero blocks in g be B and the SF in this group be f^B . The idea of progressive search is that given the desired B , we start from small blocks b and then gradually add in more blocks until $b = B$. Thus, we can greedily generate candidates based on the top SFs in \mathcal{T}^{b-2} at step 2-6 to reduce search space. Specifically, we greedily pick up the top- K_1 f^{b-2} in the previously evaluated models in \mathcal{T}^{b-2} . N candidates will then be generated by adding two more multiplicative term in step 4 to deal with Constraint (C1) since adding one block each step will result in simply lying on the diagonal. All the candidates are generated from $b = 4$ and are checked through the Filter Q (see Sec.IV-B2) to guarantee Constraint (C2) and avoid training equivalents. Next, we use the predictor P (see Sec.IV-B3) to further select K_2 promising candidates, which will then be trained and evaluated using Alg.1, in step 7 of Alg.2. The training data for P is gradually collected with the trained SFs in $\mathcal{T} = \mathcal{T}^4 \cup \mathcal{T}^6 \cup \dots$ at step 10.

2) *Invariance - Using a filter*: The filter Q we used in Alg.2 has two functions: 1) deal with Constraint (C2) and 2) remove equivalent structures due to invariance. Constraint (C2) is easy to check, given the structure of g , we can directly map it into a 4 × 4 substitute matrix and use $\mathbf{f}_0, \mathbf{r}_1, \mathbf{r}_2, \mathbf{r}_3, \mathbf{r}_4$ to represent $[g(\mathbf{r})]_{ij} \geq \mathbf{f}_0, \mathbf{r}_1, \mathbf{r}_2, \mathbf{r}_3, \mathbf{r}_4$. Then checking requirements in (C2) is a trivial task, i.e. checking if the 4 × 4 substitute matrix satisfies the Constraint (C2).

For the invariance, once a candidate f^b fulfilling Constraint (C2) is generated, we use the invariance property to generate a set of equivalents G_{f^b} . Specifically, we can permute the entity parts, relation parts, or flip signs to get $4! \times 4! \times 2^4 = 9,216$ equivalents of f^b . If $G_{f^b} \setminus H^b \setminus \mathcal{T}^b \neq \emptyset$, we throw f^b away since there are equivalent structures in the sampled set H^b and history record \mathcal{T}^b . This step can dramatically help us to reduce the cost in training equivalent structures. Take f^4 as an example, the whole space is reduced from $A_{16}^4 \times 2^4$ to 5 through the filter, namely there are only five good and unique candidates in f^4 . Besides, we add exception for the condition in step 5 of Alg.2 for f^4 since the number of candidates is smaller than N .

3) *Expressiveness - Constructing a predictor*: Even though the filter helps to throw away many unpromising candidates, it does not deal with Constraint (C1). Hence, after collecting N candidates, we use the predictor P to further select K_2 promising ones among them. Considering that the performance of SFs on a specific KG is closely related to how the SF is formed, we can use a learning model, i.e. the predictor P to predict the performance and select good candidates in advance. In general, we need to extract features for points which have been visited by the search algorithm, and then use a learning model to predict validation performance based on those features [13], [22]. The following are principles a good predictor needs to meet

- (P1). *Correlate well with true performance*: the predictor needs not to accurately predict the exact values of validation performance, instead it should rank good candidates over bad ones;
- (P2). *Learn from small samples*: as the real performance of each point in the search space is expensive to acquire, the complexity of the predictor should be low so that it can learn from few samples.

Based on Principle (P1), the extracted features from g should be closely related to the quality of defined SF. Meanwhile, the features should be cheap to construct, i.e. they

should not depend on values of r , which are unknown before training. For (P2), the number of features should be small to guarantee a simple predictor. Therefore, we are motivated to design the symmetry-related features (SRFs), which can effectively capture to what extent $g(r)$ can be symmetric or skew-symmetric (Proposition 2), and has low complexity.

Similar as the filter, we also use a 4×4 substitute matrix to represent g . As in Fig. 3, we use $v = [v_1; v_2; v_3; v_4]$ to represent $[r_1; r_2; r_3; r_4]$, then the symmetric and skew-symmetric property of g can be checked through $g(v) - g(v)^T$ and $g(v) + g(v)^T$. Since $g(v)$ is a simple 4×4 matrix, the checking procedure is very cheap. Then by assigning different values to v (details in Appendix C), a 22-dimensional SRF will be returned. Considering that the correlation of SRFs with SFs' performance is guaranteed under Proposition 2, we can use a simple two-layer MLP (22-2-1) as the predictor P . Other regression models with low complexity may also work here.

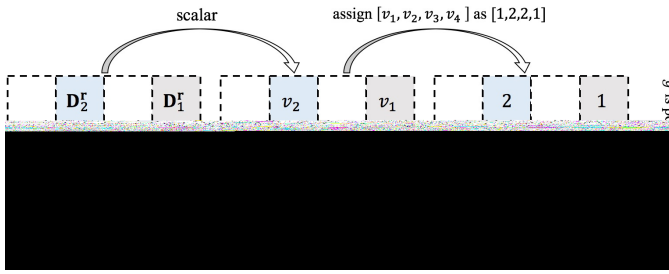


Fig. 3. Example of generating a feature of SRF.

Proposition 2. *The extracted SRFs (see Appendix C) are (i) invariant to both the permutations and flipping signs of blocks in R and (ii) give predictions related to symmetric or anti-symmetric properties.*

C. Search complexity analysis

There are 16 blocks and each can be filled with 9 different contents $f0, D_1^r, D_2^r, D_3^r, D_4^r, g$. Thus the whole space size is 9^{16} , which is extremely large. The greedy strategy, predictor and filter cut down the space in different perspectives. Specifically, in each greedy step:

Greedy: Considering that f^b is progressively generated on f^{b-2} for $b = 6, 8, \dots$, there can be $C_{16}^2 \cdot (b-2) \cdot 4^2 \cdot 2^2$ candidates ($C_{16}^2 \cdot (b-2)$ is to choose location, 4^2 is for the two r_i s and 2^2 is for signs). In comparison, there can be $C_{16}^b \cdot 4^b \cdot 2^b$ possible SFs in f^b . Take $b = 6$ for example, there are $2 \cdot 10^9$ possible candidates. Since f^6 is generated based on the 5 good candidates in f^4 , we reduce the space size from $2 \cdot 10^9$ to approximately $3 \cdot 10^4$ based on the greedy scheme.

Filter: The filter we designed is mainly used to deal with invariance properties. Permuting r_i 's leads to $4! = 24$ equivalent structures. Simultaneously permuting h_i 's and t_i 's also gives 24 equivalents. Besides, there are $2^4 = 16$ possible signs patterns. Therefore, given a $g(r)$, we can generate at most (there may exist same structures in this set) $24 \cdot 24 \cdot 16 = 9216$ equivalent SFs, which should perform the same. Besides, by constraining the SF under

Constraint (C2), many bad candidates can also be filtered out. Take f^4 as an example, only 5 candidates are selected to be trained among approximately 700k possible structures.

Predictor: Once N candidates are generated, the predictor will select K_2 ones based on their predicted performance. Thus, the reducing ratio of predictor is about N/K_2 .

While it is difficult to directly quantize to which extend the three steps together can help to reduce the search space, we can observe the significance of efficiency gained through each component. Besides, we perform an empirical study in Sec.V-E to show the performance gaining of these steps.

D. Comparison with existing AutoML approaches

The most related work in the AutoML literature is PNAS [22], which combines a greedy algorithm with a performance predictor to search a cell structure for the convolutional neural network (CNN). However, the filter is not used in PNAS as the search space for AutoSF is fundamentally different from that of CNN. Besides, PNAS adopts direct one-hot encoding for the predictor, which has a bad empirical performance here (see Sec.V-E1). As for the other AutoML approaches, even though the search problem of AutoSF is similarly defined as HPO [3], [9] and NAS [10], the search space and search algorithm of AutoSF are novel and specifically designed for KGE. There is no direct way for them to deal with the challenges in Sec.IV-A.

V. EMPIRICAL STUDY

All of the algorithms are written in python with PyTorch framework [31]. Experiments are run on 8 TITAN Xp GPUs.

A. Experiment setup

1) Datasets: Five data sets, i.e. WN18, FB15k, WN18RR, FB15k237 and YAGO3-10 are considered (statistics in Tab. III). WN18RR and FB15k237 are variants that remove near-duplicate or inverse-duplicate relations from WN18 and FB15k respectively, [37], [42]. YAGO3-10 is much larger than the others. These are benchmark datasets, and are popularly used to compare KGE models in the literature [4], [18], [19], [24], [39], [46]. The number of symmetric, anti-symmetric, inverse pairs and general asymmetric are computed in the following way: Given a relation r , let the number of positive triplets (h, r, t) be n_r . (i) If the number of (t, r, h) is larger than $0.9n_r$, then we regard it as symmetric; (ii) If the number of (t, r, h) is zero and the size of joint set of h and t is at least $0.1n_r$ (this is to ensure that they have same type), we regard as anti-symmetric; (iii) If there exist another relation r^0 that has at least $0.9n_r$ (t, r^0, h) , then r and r^0 are inverse pairs; (iv) others are regarded as general asymmetric. The threshold 0.9 and 0.1 are hand-made and just used to roughly (other values are fine) indicate the relation properties for each data set.

2) Hyper-parameters: Since the searched embedding models belong to BLMs, we fairly compare different SFs with a fixed set of hyper-parameters. In order to reduce training time, we set the dimension d as 64 during the search procedure. First, we use Simple [18] as the benchmark model and

TABLE III
STATISTICS OF THE DATA SETS USED IN EXPERIMENTS. “SYM” AND “ANTI-SYM” DENOTES THE SYMMETRIC AND ANTI-SYMMETRIC RELATIONS.

data set	#entity	#relation	#train	#valid	#test	#sym	#anti-sym	#inverse	#general
WN18 [4]	40,943	18	141,442	5,000	5,000	4	7	7	0
FB15k [4]	14,951	1,345	484,142	50,000	59,071	66	38	556	685
WN18RR [6]	40,943	11	86,835	3,034	3,134	4	3	1	3
FB15k237 [37]	14,541	237	272,115	17,535	20,466	33	5	20	179
YAGO3-10 [26]	123,188	37	1,079,040	5,000	5,000	8	0	1	28

TABLE IV
COMPARISON OF THE BEST SF IDENTIFIED BY AUTOSF AND THE STATE-OF-THE-ART SFs. THE BOLD NUMBER MEANS THE BEST PERFORMANCE, AND THE UNDERLINE MEANS THE SECOND BEST. DISTMULT, COMPLEX, ANALOGY AND SIMPLE ARE OBTAINED FROM OUR IMPLEMENTATION, OTHERS ARE COPIED FROM THE CORRESPONDING REFERENCE PAPER. STD IS LESS THAN 0.001, THUS NOT REPORTED.

type	model	WN18			FB15k			WN18RR			FB15k237			YAGO3-10		
		MRR	H@1	H@10	MRR	H@1	H@10	MRR	H@1	H@10	MRR	H@1	H@10	MRR	H@1	H@10
TDM	TransE [54]	0.500	—	94.1	0.495	—	77.4	0.178	—	45.1	0.256	—	41.9	—	—	—
	TransH [54]	0.521	—	94.5	0.452	—	76.6	0.186	—	45.1	0.233	—	40.1	—	—	—
	RotatE [35]	0.949	94.4	95.9	0.797	74.6	88.4	<u>0.476</u>	42.8	57.1	0.338	24.1	53.3	—	—	—
NNM	NTN [46]	0.53	—	66.1	0.25	—	41.4	—	—	—	—	—	—	—	—	—
	Neural LP [47]	0.94	—	94.5	0.76	—	83.7	—	—	—	0.24	—	36.2	—	—	—
	ConvE [6]	0.942	93.5	95.5	0.745	67.0	87.3	0.46	39.	48.	0.316	23.9	49.1	0.52	45.	66.
BLM	TuckER [1]	0.953	94.9	95.8	0.795	74.1	89.2	0.470	<u>44.3</u>	52.6	<u>0.358</u>	<u>26.6</u>	54.4	—	—	—
	HolEX [45]	0.938	93.0	94.9	0.800	75.0	88.6	—	—	—	—	—	—	—	—	—
	QuatE [53]	0.950	94.5	95.9	0.782	71.1	90.0	<u>0.488</u>	43.8	58.2	0.348	24.8	55.0	—	—	—
	DistMult	0.821	71.7	95.2	0.817	77.7	89.5	0.443	40.4	50.7	0.349	25.7	53.7	0.552	47.6	69.4
	ComplEx	0.951	94.5	95.7	<u>0.831</u>	79.6	<u>90.5</u>	0.471	43.0	55.1	0.347	25.4	54.1	<u>0.566</u>	<u>49.1</u>	70.9
	Analogy	0.950	94.6	95.7	0.829	79.3	<u>90.5</u>	0.472	43.3	55.8	0.348	25.6	<u>54.7</u>	<u>0.565</u>	<u>49.0</u>	<u>71.3</u>
	Simple/CP	0.950	94.5	<u>95.9</u>	0.830	<u>79.8</u>	<u>90.3</u>	0.468	42.9	55.2	0.350	26.0	54.4	0.565	<u>49.1</u>	71.0
	AnyBURL [27]	0.95	94.6	<u>95.9</u>	0.83	80.8	87.6	0.48	44.6	55.5	0.31	23.3	48.6	0.54	47.7	47.3
AutoSF		<u>0.952</u>	<u>94.7</u>	96.1	0.853	82.1	91.0	0.490	45.1	<u>56.7</u>	0.360	26.7	55.2	0.571	50.1	71.5

tune hyper-parameters with the help of HyperOpt, a hyper-parameter optimization framework based on TPE [3]. The searching ranges are given as follows: learning rate η in $[0, 1]$, L2 penalty λ in $[10^{-5}, 10^{-1}]$, decay rate in $[0.99, 1.0]$, batch size m in $\{256, 512, 1024\}$. All the models are trained until converge to avoid the influence of different convergence speed. Besides, we use Adagrad [8] as the optimizer since it tends to perform better as indicated in [19], [39]. Once a good hyper-parameter configuration is selected, we use it to train and evaluate different searched SFs. After the search procedure, we pick up the best SF evaluated by the MRR performance on the validation data set as the searched SF. When comparing the searched SFs with human-designed ones, we increase the dimension from 64 to $d \in \{256, 512, 1024, 2048\}$ as in [19]. As mentioned in [42], KGE models are sensitive to hyper-parameters. For a fair comparison, we use the same set of hyper-parameters to train and evaluated different models on each dataset.

3) *Meta Hyper-parameters*: The hyper-parameters K_1, K_2 and N have little influence to the search procedure. We use $K_1 = K_2 = 8$ and $N = 256$ for all data sets. Besides, steps 2-11 in Alg.2 run based on an inner loop. We train 8 models in parallel and iterate for 32 times (16 times for YAGO3-10), namely we evaluate 256 f^b s for each $b > 4$.

B. Comparison with existing SFs on link prediction

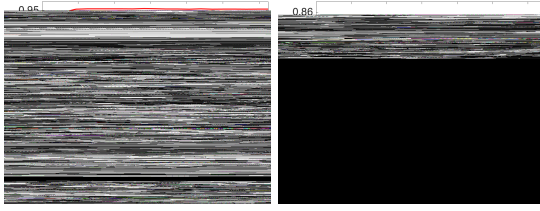
We compare our AutoSF with the state-of-the-art KGE models discussed in Sec.II-A, which are designed by humans, i.e. TransE [4], TransH [43], and RotatE [35] from TDMs; NTN [34], Neural LP [47], and ConvE [6] from NNMs; TuckER [1], HolE/HolEX [29], [45], Quat [53], DistMult [46],

ComplEx [39], Analogy [24] and Simple [18] from BLMs; and a rule-based method AnyBURL [27]. Hyper-parameters are selected by the MRR value on the validation set.

Following [6], [18], [24], [39], [46], we test KGE’s performance based on *link prediction*. For each triplet $(h, r, t) \in S$, where S is the validation or testing set, we compute the score of (h^0, r, t) for all $h^0 \in E$ and get the rank of h , the same for t based on scores of (h, r, t^0) over all $t^0 \in E$, r is not compared as in the literature [40]. Same as above mentioned papers, we adopt the following metrics: (i) Mean reciprocal ranking (MRR): $1/|S| \sum_{i=1}^{|S|} 1/\text{rank}_i$, where $\text{rank}_i, i \in 1, \dots, |S|$ is a set of ranking results and (ii) H@10: $1/|S| \sum_{i=1}^{|S|} \mathbb{I}(\text{rank}_i < 10)$, where $\mathbb{I}(\cdot)$ is the indicator function. We report the performance in a “filtered” setting as in [4], [43], where larger MRR and H@10 indicate higher embedding quality.

1) *Effectiveness*: A comparison of the testing performance of AutoSF and the current state-of-the-art SFs are shown in Tab. IV. Firstly, we can see that there is no absolute winner among the baseline SFs. For example, TuckER is the best on WN18, but is the worst among human-designed BLMs on FB15k. DistMult generally performs worse on the benchmarks except for FB15k237 since it does not follow Proposition 1. A single model is hard to adapt to different KGs. However, AutoSF performs consistently well among these five data sets. i.e. the best among FB15k, WN18RR, FB15k237 and YAGO3-10, and the runner-up on WN18.

Besides, we plot the learning curves of DistMult, Analogy, ComplEx, Simple and the best SF searched by AutoSF in Fig. 4. As shown, the searched SFs not only outperform baselines, but also converge faster, which may due to these SFs can better capture relations in these datasets.

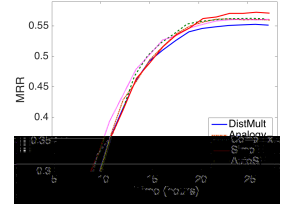


(a) WN18.

(b) FB15k.

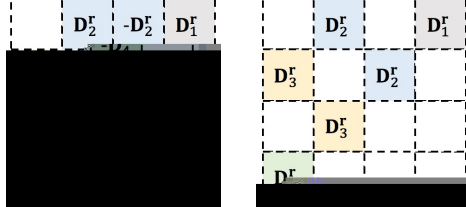
(c) WN18RR.

(d) FB15k237.



(e) YAGO3-10.

Fig. 4. Comparison on clock time (in hours) of model training v.s testing MRR between search SFs (by AutoSF) and human-designed ones.



(a) WN18.

(b) FB15k.

(c) WN18RR.

(d) FB15k237.

(e) YAGO3-10.

Fig. 5. A graphical illustration of SFs identified by our AutoSF on each data set.

2) *Case study: Distinctiveness*: To show the searched SFs are KG-dependent and novel to the literature, we plot them in Fig. 5. It is obvious that these SFs are different from each other, and they are not equivalent regarding invariance properties. As shown in Tab. III, WN18 and FB15k have many symmetric, anti-symmetric relations and inverse relation pairs, the best SF searched on them are very similar and have the same SRF. The other three data sets are more realistic and contains less symmetric, anti-symmetric and inverse relations, thus have different SRFs with fewer entry being non-zero.

The most special case is FB15k237, which can only be symmetric under (S11). Viewing the values in Tab. IV, we can see that the leading performance on FB15k237 is achieved by DistMult and AutoSF, both of which cannot be skew-symmetric. As given in the statistic information in Tab. III, FB15k237 has relatively fewer anti-symmetric relations. This may explain why skew-symmetric is not that important for $g(r)$. However, SRFs still work for these cases since it can be aware that skew-symmetric property is not that essential and focus more on searching different local structures.

TABLE V
MRRs OF APPLYING SF SEARCHED FROM ONE DATA SET (INDICATED BY EACH ROW) ON ANOTHER DATA SET (INDICATED BY EACH COLUMN).

	WN18	FB15k	WN18RR	FB15k237	YAGO3-10
WN18	0.952	0.841	0.473	0.349	0.561
FB15k	0.950	0.853	0.470	0.350	0.563
WN18RR	0.951	0.833	0.490	0.345	0.568
FB15k237	0.894	0.781	0.462	0.360	0.565
YAGO3-10	0.885	0.835	0.466	0.352	0.571

Besides, we pick up the best SF searched from one data set and test it on another data set in Tab. V. We can readily find that these SFs get the best performance on the data sets where they are searched. This again demonstrate that SFs found by AutoSF on different KGs are distinct from each other.

C. Comparison with existing SFs on triplet classification

To further demonstrate the effectiveness of the searched SFs, we do triplet classification as in [43]. This task is to

confirm whether a given (h, r, t) is correct or not and is more helpful in answering yes-or-no questions. The decision rule of classification is as follows: for each (h, r, t) , if its score is larger than the relation-specific threshold σ_r , which we predict to be positive, otherwise negative. The threshold σ_r is determined by maximizing the accuracy of the validation set. We test this task on FB15k, WN18RR and FB15k237, in which the positive and negative triplets are provided. As shown in Tab. VI, searched SFs consistently outperform human-designed BLMs.

TABLE VI
COMPARISON OF SEARCHED SFs WITH THE STATE-OF-THE-ART SFs ON ACCURACY (IN %) FOR TRIPLET CLASSIFICATION. STD < 0.2.

	FB15k	WN18RR	FB15k237
DistMult	80.8	84.6	79.8
Analogy	82.1	86.1	79.7
ComplEx	81.8	86.6	79.6
Simple	81.5	85.7	79.6
AutoSF	82.7	87.7	81.2

D. Comparison with other AutoML approaches

In this part, we compare AutoSF with the other search algorithms. WN18RR and FB15k237 are used here, and all algorithms share the same set of hyper-parameters. First, to show the effectiveness of the search space in BLM, we train a general approximator (*Gen-Approx*), i.e. MLP (in Appendix D), on the validation set. Then, AutoSF is compared with *Random* search and *Bayes* algorithm [3] on f^6 . As shown in Fig. 6, the general approximator performs much worse than BLM since it is too flexible to consider domain-specific constraints and easily overfits. For BLM settings, the Bayes algorithm can improve the efficiency upon random search. However, it will easily fall into local optimum and does not take the domain property into account. Among them, AutoSF is the most efficient and has the best any-time performance.

