AmpliGraph

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Luca Costabello - Accenture Labs Dublin

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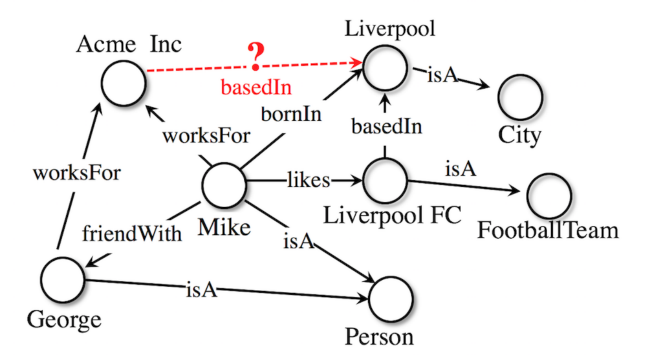
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Open source Python library that predicts links between concepts in a knowledge graph.



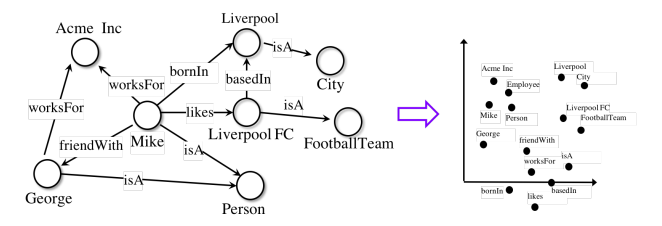
AmpliGraph is a suite of neural machine learning models for relational Learning, a branch of machine learning that deals with supervised learning on knowledge graphs.



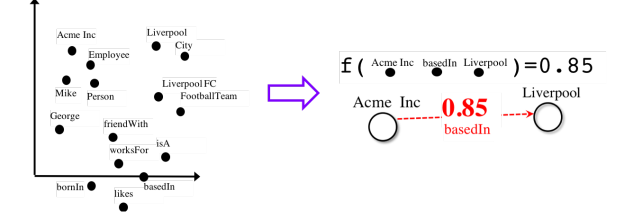
Use AmpliGraph if you need to:

- Discover new knowledge from an existing knowledge graph.
- Complete large knowledge graphs with missing statements.
- Generate stand-alone knowledge graph embeddings.
- Develop and evaluate a new relational model.

AmpliGraph's machine learning models generate **knowledge graph embeddings**, vector representations of concepts in a metric space:



It then combines embeddings with model-specific scoring functions to predict unseen and novel links:



CHAPTER 1

Key Features

- **Intuitive APIs**: AmpliGraph APIs are designed to reduce the code amount required to learn models that predict links in knowledge graphs.
- **GPU-Ready**: AmpliGraph is based on TensorFlow, and it is designed to run seamlessly on CPU and GPU devices to speed-up training.
- Extensible: Roll your own knowledge graph embeddings model by extending AmpliGraph base estimators.

CHAPTER 2

Modules

AmpliGraph includes the following submodules:

- Datasets: helper functions to load datasets (knowledge graphs).
- Models: knowledge graph embedding models. AmpliGraph contains TransE, DistMult, ComplEx, HolE, ConvE, ConvKB (More to come!)
- Evaluation: metrics and evaluation protocols to assess the predictive power of the models.
- **Discovery**: High-level convenience APIs for knowledge discovery (discover new facts, cluster entities, predict near duplicates).

CHAPTER 3

How to Cite

If you like AmpliGraph and you use it in your project, why not starring the project on GitHub!

If you instead use AmpliGraph in an academic publication, cite as:

```
@misc{ampligraph,
  author= {Luca Costabello and
      Sumit Pai and
      Chan Le Van and
      Rory McGrath and
      Nick McCarthy and
      Pedro Tabacof},
  title = {{AmpliGraph: a Library for Representation Learning on Knowledge Graphs}},
  month = mar,
  year = 2019,
  doi = {10.5281/zenodo.2595043},
  url = {https://doi.org/10.5281/zenodo.2595043}
```

3.1 Installation

3.1.1 Prerequisites

- Linux, macOS, Windows
- Python 3.7

Provision a Virtual Environment

Create and activate a virtual environment (conda)

```
conda create --name ampligraph python=3.7
source activate ampligraph
```

Install TensorFlow

AmpliGraph is built on TensorFlow 1.x. Install from pip or conda:

CPU-only

```
pip install "tensorflow>=1.15.2,<2.0"</pre>
```

or

```
conda install tensorflow'>=1.15.2,<2.0.0'</pre>
```

GPU support

```
pip install "tensorflow-gpu>=1.15.2,<2.0"</pre>
```

or

```
conda install tensorflow-gpu'>=1.15.2,<2.0.0'</pre>
```

3.1.2 Install AmpliGraph

Install the latest stable release from pip:

```
pip install ampligraph
```

If instead you want the most recent development version, you can clone the repository and install from source as below (also see the *How to Contribute guide* for details):

```
git clone https://github.com/Accenture/AmpliGraph.git
cd AmpliGraph
git checkout develop
pip install -e .
```

3.1.3 Sanity Check

```
>> import ampligraph
>> ampligraph.__version__
'1.4.0'
```

3.2 Background

For a comprehensive theoretical and hands-on overview of KGE models and hands-on AmpliGraph, check out our ECAI-20 Tutorial (Slides + Recording + Colab Notebook).

Knowledge graphs are graph-based knowledge bases whose facts are modeled as relationships between entities. Knowledge graph research led to broad-scope graphs such as DBpedia [ABK+07], WordNet [Pri10], and YAGO [SKW07]. Countless domain-specific knowledge graphs have also been published on the web, giving birth to the so-called Web of Data [BHBL11].

Formally, a knowledge graph $\mathcal{G} = \{(sub, pred, obj)\} \subseteq \mathcal{E} \times \mathcal{R} \times \mathcal{E} \text{ is a set of } (sub, pred, obj) \text{ triples, each including a subject } sub \in \mathcal{E}, \text{ a predicate } pred \in \mathcal{R}, \text{ and an object } obj \in \mathcal{E}. \mathcal{E} \text{ and } \mathcal{R} \text{ are the sets of all entities and relation types of } \mathcal{G}.$

Knowledge graph embedding models are neural architectures that encode concepts from a knowledge graph (i.e. entities \mathcal{E} and relation types \mathcal{R}) into low-dimensional, continuous vectors $\in \mathcal{R}^k$. Such textit{knowledge graph embeddings} have applications in knowledge graph completion, entity resolution, and link-based clustering, just to cite a few [NMTG16]. Knowledge graph embeddings are learned by training a neural architecture over a graph. Although such architectures vary, the training phase always consists in minimizing a loss function \mathcal{L} that includes a *scoring function* $f_m(t)$, i.e. a model-specific function that assigns a score to a triple t = (sub, pred, obj).

The goal of the optimization procedure is learning optimal embeddings, such that the scoring function is able to assign high scores to positive statements and low scores to statements unlikely to be true. Existing models propose scoring functions that combine the embeddings \mathbf{e}_{sub} , \mathbf{e}_{pred} , $\mathbf{e}_{obj} \in \mathcal{R}^k$ of the subject, predicate, and object of triple t = (sub, pred, obj) using different intuitions: TransE [BUGD+13] relies on distances, DistMult [YYH+14] and ComplEx [TWR+16] are bilinear-diagonal models, HolE [NRP+16] uses circular correlation. While the above models can be interpreted as multilayer perceptrons, others such as ConvE include convolutional layers [DMSR18].

As example, the scoring function of TransE computes a similarity between the embedding of the subject \mathbf{e}_{sub} translated by the embedding of the predicate \mathbf{e}_{pred} and the embedding of the object \mathbf{e}_{obj} , using the L_1 or L_2 norm $|| \cdot ||$:

$$f_{TransE} = -||\mathbf{e}_{sub} + \mathbf{e}_{pred} - \mathbf{e}_{obj}||_n$$

Such scoring function is then used on positive and negative triples t^+ , t^- in the loss function. This can be for example a pairwise margin-based loss, as shown in the equation below:

$$\mathcal{L}(\Theta) = \sum_{t^+ \in \mathcal{G}} \sum_{t^- \in \mathcal{N}} max(0, [\gamma + f_m(t^-; \Theta) - f_m(t^+; \Theta)])$$

where Θ are the embeddings learned by the model, f_m is the model-specific scoring function, $\gamma \in \mathcal{R}$ is the margin and \mathcal{N} is a set of negative triples generated with a corruption heuristic [BUGD+13].

3.3 API

AmpliGraph includes the following submodules:

3.3.1 Datasets

Helper functions to load knowledge graphs.

Note: It is recommended to set the AMPLIGRAPH_DATA_HOME environment variable:

```
export AMPLIGRAPH_DATA_HOME=/YOUR/PATH/TO/datasets
```

When attempting to load a dataset, the module will first check if AMPLIGRAPH_DATA_HOME is set. If it is, it will search this location for the required dataset. If the dataset is not found it will be downloaded and placed in this directory.

If AMPLIGRAPH_DATA_HOME has not been set the databases will be saved in the following directory:

~/ampligraph_datasets

Benchmark Datasets Loaders

Use these helper functions to load datasets used in graph representation learning literature. The functions will **automatically download** the datasets if they are not present in ~/ampligraph_datasets or at the location set in AMPLIGRAPH_DATA_HOME.

load_fb15k_237([check_md5hash,])	Load the FB15k-237 dataset
<pre>load_wn18rr([check_md5hash, clean_unseen,])</pre>	Load the WN18RR dataset
load_yago3_10([check_md5hash, clean_unseen,	Load the YAGO3-10 dataset
])	
<pre>load_fb15k([check_md5hash, add_reciprocal_rels])</pre>	Load the FB15k dataset
<pre>load_wn18([check_md5hash, add_reciprocal_rels])</pre>	Load the WN18 dataset
<pre>load_wn11([check_md5hash, clean_unseen,])</pre>	Load the WordNet11 (WN11) dataset
<pre>load_fb13([check_md5hash, clean_unseen,])</pre>	Load the Freebase13 (FB13) dataset

load_fb15k_237

ampligraph.datasets.load_fb15k_237(check_md5hash=False,

clean_unseen=True,

Load the FB15k-237 dataset

FB15k-237 is a reduced version of FB15K. It was first proposed by [TCP+15].

The FB15k-237 dataset is loaded from file if it exists at the AMPLIGRAPH_DATA_HOME location. If AMPLIGRAPH_DATA_HOME is not set the the default ~/ampligraph_datasets is checked.

add_reciprocal_rels=False)

If the dataset is not found at either location it is downloaded and placed in <code>AMPLIGRAPH_DATA_HOME</code> or $\sim/ampligraph_datasets$.

The dataset is divided in three splits:

- train
- valid
- test

Dataset	Train	Valid	Test	Entities	Relations
FB15K-237	272,115	17,535	20,466	14,541	237

Warning: FB15K-237's validation set contains 8 unseen entities over 9 triples. The test set has 29 unseen entities, distributed over 28 triples.

Parameters

- **check_md5hash** (*boolean*) If True check the md5hash of the files. Defaults to False.
- **clean_unseen** (*bool*) If True, filters triples in validation and test sets that include entities not present in the training set.

• **add_reciprocal_rels** (*bool*) – Flag which specifies whether to add reciprocal relations. For every <s, p, o> in the dataset this creates a corresponding triple with reciprocal relation <0, p_reciprocal, s>. (default: False).

Returns splits – The dataset splits: {'train': train, 'valid': valid, 'test': test}. Each split is an ndarray of shape [n, 3].

Return type dict

Examples

load_wn18rr

```
ampligraph.datasets.load_wn18rr(check_md5hash=False,
```

add_reciprocal_rels=False)

clean_unseen=True,

Load the WN18RR dataset

The dataset is described in [DMSR18].

The WN18RR dataset is loaded from file if it exists at the AMPLIGRAPH_DATA_HOME location. If AMPLIGRAPH_DATA_HOME is not set the default ~/ampligraph_datasets is checked.

If the dataset is not found at either location it is downloaded and placed in <code>AMPLIGRAPH_DATA_HOME</code> or $\sim/ampligraph_datasets$.

It is divided in three splits:

- train
- valid
- test

Dataset	Train	Valid	Test	Entities	Relations
WN18RR	86,835	3,034	3,134	40,943	11

Warning: WN18RR's validation set contains 198 unseen entities over 210 triples. The test set has 209 unseen entities, distributed over 210 triples.

Parameters

- **clean_unseen** (*bool*) If True, filters triples in validation and test sets that include entities not present in the training set.
- **check_md5hash** (*bool*) If True check the md5hash of the datset files. Defaults to False.
- add_reciprocal_rels (bool) Flag which specifies whether to add reciprocal relations. For every <s, p, o> in the dataset this creates a corresponding triple with reciprocal relation <0, p_reciprocal, s>. (default: False).

Returns splits – The dataset splits: {'train': train, 'valid': valid, 'test': test}. Each split is an ndarray of shape [n, 3].

Return type dict

Examples

```
>>> from ampligraph.datasets import load_wn18rr
>>> X = load_wn18rr()
>>> X["valid"][0]
array(['02174461', '_hypernym', '02176268'], dtype=object)
```

load_yago3_10

Load the YAGO3-10 dataset

The dataset is a split of YAGO3 [MBS13], and has been first presented in [DMSR18].

The YAGO3-10 dataset is loaded from file if it exists at the AMPLIGRAPH_DATA_HOME location. If AMPLIGRAPH_DATA_HOME is not set the the default ~/ampligraph_datasets is checked.

If the dataset is not found at either location it is downloaded and placed in <code>AMPLIGRAPH_DATA_HOME</code> or $\sim/ampligraph_datasets$.

It is divided in three splits:

- train
- valid
- test

Dataset	Train	Valid	Test	Entities	Relations
YAGO3-10	1,079,040	5,000	5,000	123,182	37

Parameters

- **check_md5hash** (*boolean*) If True check the md5hash of the files. Defaults to False.
- **clean_unseen** (*bool*) If True, filters triples in validation and test sets that include entities not present in the training set.
- **add_reciprocal_rels** (*bool*) Flag which specifies whether to add reciprocal relations. For every <s, p, o> in the dataset this creates a corresponding triple with reciprocal relation <0, p_reciprocal, s>. (default: False).
- **Returns splits** The dataset splits: {'train': train, 'valid': valid, 'test': test}. Each split is an ndarray of shape [n, 3].

Return type dict

Examples

```
>>> from ampligraph.datasets import load_yago3_10
>>> X = load_yago3_10()
>>> X["valid"][0]
array(['Mikheil_Khutsishvili', 'playsFor', 'FC_Merani_Tbilisi'], dtype=object)
```

load_fb15k

ampligraph.datasets.load_fb15k (check_md5hash=False, add_reciprocal_rels=False)
Load the FB15k dataset

Warning: The dataset includes a large number of inverse relations that spilled to the test set, and its use in experiments has been deprecated. Use FB15k-237 instead.

FB15k is a split of Freebase, first proposed by [BUGD+13].

The FB15k dataset is loaded from file if it exists at the AMPLIGRAPH_DATA_HOME location. If AMPLIGRAPH_DATA_HOME is not set the default ~/ampligraph_datasets is checked.

If the dataset is not found at either location it is downloaded and placed in <code>AMPLIGRAPH_DATA_HOME</code> or $\sim/ampligraph_datasets$.

The dataset is divided in three splits:

- train
- valid
- test

Dataset	Train	Valid	Test	Entities	Relations
FB15K	483,142	50,000	59,071	14,951	1,345

Parameters

- **check_md5hash** (*boolean*) If True check the md5hash of the files. Defaults to False.
- **add_reciprocal_rels** (*bool*) Flag which specifies whether to add reciprocal relations. For every <s, p, o> in the dataset this creates a corresponding triple with reciprocal relation <0, p_reciprocal, s>. (default: False).
- **Returns splits** The dataset splits: {'train': train, 'valid': valid, 'test': test}. Each split is an ndarray of shape [n, 3].

Return type dict

Examples

```
>>> from ampligraph.datasets import load_fb15k
>>> X = load_fb15k()
>>> X['test'][:3]
array([['/m/01qscs',
```

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load_wn18

```
ampligraph.datasets.load_wn18(check_md5hash=False, add_reciprocal_rels=False)
Load the WN18 dataset
```

Warning: The dataset includes a large number of inverse relations that spilled to the test set, and its use in experiments has been deprecated. Use WN18RR instead.

WN18 is a subset of Wordnet. It was first presented by [BUGD+13].

The WN18 dataset is loaded from file if it exists at the AMPLIGRAPH_DATA_HOME location. If AMPLIGRAPH_DATA_HOME is not set the the default ~/ampligraph_datasets is checked.

If the dataset is not found at either location it is downloaded and placed in <code>AMPLIGRAPH_DATA_HOME</code> or \sim /ampligraph_datasets.

The dataset is divided in three splits:

- train: 141,442 triples
- valid 5,000 triples
- test 5,000 triples

Dataset	Train	Valid	Test	Entities	Relations
WN18	141,442	5,000	5,000	40,943	18

Parameters

- check_md5hash (bool) If True check the md5hash of the files. Defaults to False.
- add_reciprocal_rels (bool) Flag which specifies whether to add reciprocal relations. For every <s, p, o> in the dataset this creates a corresponding triple with reciprocal relation <0, p_reciprocal, s>. (default: False).
- **Returns splits** The dataset splits { 'train': train, 'valid': valid, 'test': test}. Each split is an ndarray of shape [n, 3].

Return type dict

Examples

```
>>> from ampligraph.datasets import load_wn18
>>> X = load_wn18()
>>> X['test'][:3]
```

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clean_unseen=True,

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```
array([['06845599', '_member_of_domain_usage', '03754979'],
['00789448', '_verb_group', '01062739'],
['10217831', '_hyponym', '10682169']], dtype=object)
```

load_wn11

Load the WordNet11 (WN11) dataset

WordNet was originally proposed in WordNet: a lexical database for English [Mil95].

WN11 dataset is loaded from file if it exists at the AMPLIGRAPH_DATA_HOME location. If AMPLIGRAPH_DATA_HOME is not set the default ~/ampligraph_datasets is checked.

If the dataset is not found at either location, it is downloaded and placed in AMPLIGRAPH_DATA_HOME or ~/ampligraph_datasets.

It is divided in three splits:

- train
- valid
- test

Both the validation and test splits are associated with labels (binary ndarrays), with *True* for positive statements and *False* for negatives:

- valid_labels
- test_labels

Dataset	Train	Valid Pos	Valid Neg	Test Pos	Test Neg	Entities	Relations
WN11	110361	2606	2609	10493	10542	38588	11

Parameters

- **check_md5hash** (*boolean*) If True check the md5hash of the files. Defaults to False.
- **clean_unseen** (*bool*) If True, filters triples in validation and test sets that include entities not present in the training set.
- add_reciprocal_rels (bool) Flag which specifies whether to add reciprocal relations. For every <s, p, o> in the dataset this creates a corresponding triple with reciprocal relation <0, p_reciprocal, s>. (default: False).
- **Returns splits** The dataset splits: { 'train': train, 'valid': valid, 'valid_labels': valid_labels, 'test': test, 'test_labels': test_labels}. Each split containing a dataset is an ndarray of shape [n, 3]. The labels are ndarray of shape [n].

Return type dict

Examples

```
>>> from ampligraph.datasets import load_wn11
>>> X = load_wn11()
>>> X["valid"][0]
array(['__genus_xylomelum_1', '_type_of', '__dicot_genus_1'], dtype=object)
>>> X["valid_labels"][0:3]
array([ True, False, True])
```

load_fb13

Load the Freebase13 (FB13) dataset

FB13 is a subset of Freebase [BEP+08] and was initially presented in *Reasoning With Neural Tensor Networks for Knowledge Base Completion* [SCMN13].

FB13 dataset is loaded from file if it exists at the AMPLIGRAPH_DATA_HOME location. If AMPLIGRAPH_DATA_HOME is not set the the default ~/ampligraph_datasets is checked.

If the dataset is not found at either location, it is downloaded and placed in <code>AMPLIGRAPH_DATA_HOME</code> or \sim /ampligraph_datasets.

It is divided in three splits:

- train
- valid
- test

Both the validation and test splits are associated with labels (binary ndarrays), with *True* for positive statements and *False* for negatives:

- valid_labels
- test_labels

Dataset	Train	Valid Pos	Valid Neg	Test Pos	Test Neg	Entities	Relations
FB13	316232	5908	5908	23733	23731	75043	13

Parameters

- check_md5hash (boolean) If True check the md5hash of the files. Defaults to False.
- **clean_unseen** (*bool*) If True, filters triples in validation and test sets that include entities not present in the training set.
- **add_reciprocal_rels** (bool) Flag which specifies whether to add reciprocal relations. For every <s, p, o> in the dataset this creates a corresponding triple with reciprocal relation <0, p_reciprocal, s>. (default: False).
- **Returns splits** The dataset splits: { 'train': train, 'valid': valid, 'valid_labels': valid_labels, 'test': test, 'test_labels': test_labels}. Each split containing a dataset is an ndarray of shape [n, 3]. The labels are ndarray of shape [n].

Return type dict

Examples

```
>>> from ampligraph.datasets import load_fb13
>>> X = load_fb13()
>>> X["valid"][0]
array(['cornelie_van_zanten', 'gender', 'female'], dtype=object)
>>> X["valid_labels"][0:3]
array([True, False, True], dtype=object)
```

Datasets Summary

Dataset	Train	Valid	Test	Entities	Relations
FB15K-237	272,115	17,535	20,466	14,541	237
WN18RR	86,835	3,034	3,134	40,943	11
FB15K	483,142	50,000	59,071	14,951	1,345
WN18	141,442	5,000	5,000	40,943	18
YAGO3-10	1,079,040	5,000	5,000	123,182	37
WN11	110,361	5,215	21,035	38,194	11
FB13	316,232	11,816	47,464	75,043	13

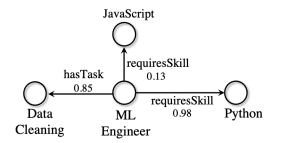
Warning: WN18 and FB15k include a large number of inverse relations, and its use in experiments has been deprecated. Use WN18RR and FB15K-237 instead.

Warning: FB15K-237's validation set contains 8 unseen entities over 9 triples. The test set has 29 unseen entities, distributed over 28 triples. WN18RR's validation set contains 198 unseen entities over 210 triples. The test set has 209 unseen entities, distributed over 210 triples.

Note: WN11 and FB13 also provide true and negative labels for the triples in the validation and tests sets. In both cases the positive base rate is close to 50%.

Benchmark Datasets Loaders (Knowledge Graphs With Numeric Values on Edges)

These helper functions load benchmark datasets **with numeric values on edges**, as described in [PC21] (the figure below shows a toy example).



Hint: To process a knowledge graphs with numeric values associated to edges, enable the FocusE layer when training

a knowledge graph embedding model [PC21].

The functions will **automatically download** the datasets if they are not present in ~/ampligraph_datasets or at the location set in AMPLIGRAPH_DATA_HOME.

<pre>load_onet20k([check_md5hash, clea</pre>	n_unseen,	Load the O*NET20K dataset
])		
<pre>load_ppi5k([check_md5hash, clean_unset</pre>	en,])	Load the PPI5K dataset
load_n127k([check_md5hash, clean_unser	en,])	Load the NL27K dataset
load_cn15k([check_md5hash, clean_unser	en,])	Load the CN15K dataset

load_onet20k

Load the O*NET20K dataset

O*NET20K was originally proposed in [PC21]. It a subset of O*NET, a dataset that includes job descriptions, skills and labeled, binary relations between such concepts. Each triple is labeled with a numeric value that indicates the importance of that link.

ONET*20K dataset is loaded from file if it exists at the AMPLIGRAPH_DATA_HOME location. If AMPLIGRAPH_DATA_HOME is not set the the default ~/ampligraph_datasets is checked.

If the dataset is not found at either location, it is downloaded and placed in AMPLIGRAPH_DATA_HOME or ~/ampligraph_datasets.

It is divided in three splits:

- train
- valid
- test

Each triple in these splits is associated to a numeric value which represents the importance/relevance of the link.

Dataset	Train	Valid	Test	Entities	Relations
ONET*20K	461,932	850	2,000	20,643	19

Parameters

- **check_md5hash** (*boolean*) If True check the md5hash of the files. Defaults to False.
- **clean_unseen** (*bool*) If True, filters triples in validation and test sets that include entities not present in the training set.
- **split_test_into_top_bottom** (*bool*) Splits the test set by numeric values and returns test_top_split and test_bottom_split by splitting based on sorted numeric values and returning top and bottom k% triples, where k is specified by *split_threshold* argument
- **split_threshold**(*float*) specifies the top and bottom percentage of triples to return

Returns

splits – The dataset splits: {'train': train, 'valid': valid, 'test': test, 'test_topk': test_topk, 'test_bottomk': test_bottomk, 'train_numeric_values': train_numeric_values,

'valid_numeric_values':valid_numeric_values, 'test_numeric_values': test_numeric_values': test_topk_numeric_values': test_topk_numeric_values, 'test_bottomk_numeric_values': test_bottomk_numeric_values}.

Each *_numeric_values split contains numeric values associated to the corresponding dataset split and is a ndarray of shape [n].

Each dataset split is a ndarray of shape [n,3].

The *_topk and *_bottomk splits are only returned when split_test_into_top_bottom=True.

Return type dict

Examples

```
>>> from ampligraph.datasets import load_onet20k
>>> X = load_onet20k()
```

load_ppi5k

Load the PPI5K dataset

Originally proposed in [CCS+19], PPI5K is a subset of the protein-protein interactions (PPI) knowledge graph [SM+16]. Numeric values represent the confidence of the link based on existing scientific literature evidence.

PPI5K is loaded from file if it exists at the AMPLIGRAPH_DATA_HOME location. If AMPLIGRAPH_DATA_HOME is not set the the default ~/ampligraph_datasets is checked.

If the dataset is not found at either location, it is downloaded and placed in AMPLIGRAPH_DATA_HOME or ~/ampligraph_datasets.

It is divided into three splits:

- train
- valid
- test

Each triple in these splits is associated to a numeric value which models additional information on the fact (importance, relevance of the link).

Dataset	Train	Valid	Test	Entities	Relations
PPI5K	230929	19017	21720	4999	7

Parameters

- **check_md5hash** (*boolean*) If True check the md5hash of the files. Defaults to False.
- **clean_unseen** (*bool*) If True, filters triples in validation and test sets that include entities not present in the training set.
- **split_test_into_top_bottom** (*bool*) When set to True, the function also returns subsets of the test set that includes only the top-k or bottom-k numeric-enriched triples. splits test_topk, test_bottomk and their numeric values. Such splits are

generated by sorting Splits the test set by numeric values and returns test_top_split and test_bottom_split by splitting based on sorted numeric values and returning top and bottom k% triples, where 'k' is specified by the split_threshold argument.

• **split_threshold**(*float*) – specifies the top and bottom percentage of triples to return

Returns

splits - The dataset splits: {'train': train, 'valid': valid, 'test': test, 'test_topk': test_topk, 'test_bottomk': test_bottomk, 'train_numeric_values': train_numeric_values, 'valid_numeric_values':valid_numeric_values, 'test_numeric_values': test_numeric_values, 'test_topk_numeric_values': test_topk_numeric_values, 'test_bottomk_numeric_values': test_bottomk_numeric_values}.

Each *_numeric_values split contains numeric values associated to the corresponding dataset split and is a ndarray of shape [n].

Each dataset split is a ndarray of shape [n,3].

The *_topk and *_bottomk splits are only returned when split_test_into_top_bottom=True.

Return type dict

Examples

```
>>> from ampligraph.datasets import load_ppi5k
>>> X = load_ppi5k()
```

load_nl27k

Load the NL27K dataset

NL27K was originally proposed in [CCS+19]. It is a subset of the Never Ending Language Learning (NELL) dataset [MCH+18], which collects data from web pages. Numeric values on triples represent link uncertainty.

NL27K is loaded from file if it exists at the AMPLIGRAPH_DATA_HOME location. If AMPLIGRAPH_DATA_HOME is not set the the default ~/ampligraph_datasets is checked.

If the dataset is not found at either location, it is downloaded and placed in AMPLIGRAPH_DATA_HOME or ~/ampligraph_datasets.

It is divided into three splits:

- train
- valid
- test

Each triple in these splits is associated to a numeric value which represents the importance/relevance of the link.

Dataset	Train	Valid	Test	Entities	Relations
NL27K	149100	12274	14026	27221	405

Parameters

- **check_md5hash** (*boolean*) If True check the md5hash of the files. Defaults to False.
- **clean_unseen** (*bool*) If True, filters triples in validation and test sets that include entities not present in the training set.
- **split_test_into_top_bottom** (bool) Splits the test set by numeric values and returns test_top_split and test_bottom_split by splitting based on sorted numeric values and returning top and bottom k% triples, where k is specified by *split_threshold* argument
- **split_threshold** (*float*) specifies the top and bottom percentage of triples to return

Returns

splits - The dataset splits: {'train': train, 'valid': valid, 'test': test, 'test_topk': test_topk, 'test_bottomk': test_bottomk, 'train_numeric_values': train_numeric_values, 'valid_numeric_values':valid_numeric_values, 'test_numeric_values': test_numeric_values, 'test_topk_numeric_values': test_topk_numeric_values, 'test_bottomk_numeric_values': test_bottomk_numeric_values}.

Each *_numeric_values split contains numeric values associated to the corresponding dataset split and is a ndarray of shape [n].

Each dataset split is a ndarray of shape [n,3].

The *_topk and *_bottomk splits are only returned when split_test_into_top_bottom=True.

Return type dict

Examples

```
>>> from ampligraph.datasets import load_nl27k
>>> X = load_nl27k()
```

load_cn15k

Load the CN15K dataset

CN15K was originally proposed in [CCS+19], it is a subset of ConceptNet [SCH16], a common-sense knowledge graph built to represent general human knowledge. Numeric values on triples represent uncertainty.

CN15k dataset is loaded from file if it exists at the AMPLIGRAPH_DATA_HOME location. If AMPLIGRAPH_DATA_HOME is not set the default ~/ampligraph_datasets is checked.

If the dataset is not found at either location, it is downloaded and placed in AMPLIGRAPH_DATA_HOME or ~/ampligraph_datasets.

It is divided into three splits:

- train
- valid
- test

Each triple in these splits is associated to a numeric value which represents the importance/relevance of the link.

Dataset	Train	Valid	Test	Entities	Relations
CN15K	199417	16829	19224	15000	36

Parameters

- **check_md5hash** (*boolean*) If True check the md5hash of the files. Defaults to False.
- **clean_unseen** (*bool*) If True, filters triples in validation and test sets that include entities not present in the training set.
- **split_test_into_top_bottom** (bool) Splits the test set by numeric values and returns test_top_split and test_bottom_split by splitting based on sorted numeric values and returning top and bottom k% triples, where k is specified by *split_threshold* argument
- **split_threshold**(*float*) specifies the top and bottom percentage of triples to return

Returns

splits - The dataset splits: {'train': train, 'valid': valid, 'test': test, 'test_topk': test_topk, 'test_bottomk': test_bottomk, 'train_numeric_values': train_numeric_values, 'valid_numeric_values':valid_numeric_values, 'test_numeric_values': test_numeric_values, 'test_topk_numeric_values': test_topk_numeric_values, 'test_bottomk_numeric_values': test_bottomk_numeric_values}.

Each *_numeric_values split contains numeric values associated to the corresponding dataset split and is a ndarray of shape [n].

Each dataset split is a ndarray of shape [n,3].

The *_topk and *_bottomk splits are only returned when split_test_into_top_bottom=True.

Return type dict

Examples

```
>>> from ampligraph.datasets import load_cn15k
>>> X = load_cn15k()
```

Datasets Summary (KGs with numeric values on edges)

Dataset	Train	Valid	Test	Entities	Relations
O*NET20K	461,932	138	2,000	20,643	19
PPI5K	230,929	19,017	21,720	4,999	7
NL27K	149,100	12,274	14,026	27,221	405
CN15K	199,417	16,829	19,224	15,000	36

Loaders for Custom Knowledge Graphs

Functions to load custom knowledge graphs from disk.

<pre>load_from_csv(directory_path, file_name[,])</pre>	Load a knowledge graph from a csv file	
<pre>load_from_ntriples(folder_name, file_name[,</pre>	Load RDF ntriples	
])		
<pre>load_from_rdf(folder_name, file_name[,])</pre>	Load an RDF file	

load_from_csv

Load a knowledge graph from a csv file

Loads a knowledge graph serialized in a csv file as:

```
subj1 relationX obj1
subj1 relationY obj2
subj3 relationZ obj2
subj4 relationY obj2
...
```

Note: The function filters duplicated statements.

Note: It is recommended to use *ampligraph.evaluation.train_test_split_no_unseen()* to split custom knowledge graphs into train, validation, and test sets. Using this function will lead to validation, test sets that do not include triples with entities that do not occur in the training set.

Parameters

- **directory_path** (*str*) Folder where the input file is stored.
- **file_name** (*str*) File name.
- **sep** (*str*) The subject-predicate-object separator (default).
- header (*int*, *None*) The row of the header of the csv file. Same as pandas.read_csv header param.
- add_reciprocal_rels (bool) Flag which specifies whether to add reciprocal relations. For every <s, p, o> in the dataset this creates a corresponding triple with reciprocal relation <0, p_reciprocal, s>. (default: False)

Returns triples – The actual triples of the file.

Return type ndarray, shape [n, 3]

Examples

```
>>> from ampligraph.datasets import load_from_csv
>>> X = load_from_csv('folder', 'dataset.csv', sep=',')
>>> X[:3]
array([['a', 'y', 'b'],
        ['b', 'y', 'a'],
```

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```
['a', 'y', 'c']],
dtype='<U1')
```

load_from_ntriples

Load RDF ntriples

Loads an RDF knowledge graph serialized as ntriples, without building an RDF graph in memory. This function should be preferred over load_from_rdf(), since it does not load the graph into an rdflib model (and it is therefore faster by order of magnitudes). Nevertheless, it requires a ntriples serialization as in the example below:

```
_:alice <http://xmlns.com/foaf/0.1/knows> _:bob .
_:bob <http://xmlns.com/foaf/0.1/knows> _:alice .
```

Note: It is recommended to use *ampligraph.evaluation.train_test_split_no_unseen()* to split custom knowledge graphs into train, validation, and test sets. Using this function will lead to validation, test sets that do not include triples with entities that do not occur in the training set.

Parameters

- **folder_name** (*str*) base folder where the file is stored.
- **file_name** (*str*) file name
- **data_home** (*str*) The path to the folder that contains the datasets.
- **add_reciprocal_rels** (*bool*) Flag which specifies whether to add reciprocal relations. For every <s, p, o> in the dataset this creates a corresponding triple with reciprocal relation <0, p_reciprocal, s>. (default: False).

Returns triples – the actual triples of the file.

Return type ndarray, shape [n, 3]

load_from_rdf

Load an RDF file

Loads an RDF knowledge graph using rdflib APIs. Multiple RDF serialization formats are supported (nt, ttl, rdf/xml, etc). The entire graph will be loaded in memory, and converted into an rdflib *Graph* object.

Warning: Large RDF graphs should be serialized to ntriples beforehand and loaded with load_from_ntriples() instead.

Note: It is recommended to use *ampligraph.evaluation.train_test_split_no_unseen()* to split custom knowledge graphs into train, validation, and test sets. Using this function will lead to validation, test sets that do not include triples with entities that do not occur in the training set.

Parameters

- **folder_name** (*str*) Base folder where the file is stored.
- **file_name** (*str*) File name.
- **rdf_format** (*str*) The RDF serialization format (nt, ttl, rdf/xml see rdflib documentation).
- **data_home** (*str*) The path to the folder that contains the datasets.
- add_reciprocal_rels (bool) Flag which specifies whether to add reciprocal relations. For every <s, p, o> in the dataset this creates a corresponding triple with reciprocal relation <0, p_reciprocal, s>. (default: False).

Returns triples – the actual triples of the file.

Return type ndarray, shape [n, 3]

Hint: AmpliGraph includes a helper function to split a generic knowledge graphs into **training**, **validation**, and **test** sets. See *ampligraph.evaluation.train_test_split_no_unseen()*.

3.3.2 Models

Knowledge Graph Embedding Models

RandomBaseline([seed, verbose])	Random baseline
<pre>TransE([k, eta, epochs, batches_count,])</pre>	Translating Embeddings (TransE)
<pre>DistMult([k, eta, epochs, batches_count,])</pre>	The DistMult model
<pre>ComplEx([k, eta, epochs, batches_count,])</pre>	Complex embeddings (ComplEx)
<i>HolE</i> ([k, eta, epochs, batches_count, seed,])	Holographic Embeddings
<pre>ConvE([k, eta, epochs, batches_count, seed,])</pre>	Convolutional 2D KG Embeddings
<i>ConvKB</i> ([k, eta, epochs, batches_count,])	Convolution-based model

RandomBaseline

class ampligraph.latent_features.RandomBaseline(seed=0, verbose=False)
 Random baseline

A dummy model that assigns a pseudo-random score included between 0 and 1, drawn from a uniform distribution.

The model is useful whenever you need to compare the performance of another model on a custom knowledge graph, and no other baseline is available.

Note: Although the model still requires invoking the fit () method, no actual training will be carried out.

Examples

```
>>> import numpy as np
>>> from ampligraph.latent_features import RandomBaseline
>>> model = RandomBaseline()
>>> X = np.array([['a', 'y', 'b'],
                  ['b', 'y', 'a'],
>>>
                  ['a', 'y', 'c'],
>>>
                  ['c', 'y', 'a'],
>>>
                  ['a', 'y', 'd'],
>>>
                  ['c', 'y', 'd'],
>>>
                  ['b', 'y', 'c'],
>>>
                  ['f', 'y', 'e']])
>>>
>>> model.fit(X)
>>> model.predict(np.array([['f', 'y', 'e'], ['b', 'y', 'd']]))
[0.5488135039273248, 0.7151893663724195]
```

Methods

init([seed, verbose])	Initialize the model
<pre>fit(X[, early_stopping,])</pre>	Train the random model.
<pre>predict(X[, from_idx])</pre>	Predict the scores of triples using a trained embed-
	ding model.
<pre>get_hyperparameter_dict()</pre>	Returns hyperparameters of the model.

____init___ (seed=0, verbose=False) Initialize the model

Parameters

- **seed** (*int*) The seed used by the internal random numbers generator.
- **verbose** (bool) Verbose mode.
- fit (X, early_stopping=False, early_stopping_params={}, focusE_numeric_edge_values=None, tensorboard_logs_path=None) Train the random model.

There is no actual training involved in practice and the early stopping parameters won't have any effect.

Parameters

- X (ndarray, shape [n, 3]) The training triples
- **early_stopping** (bool) Flag to enable early stopping (default:False).

If set to True, the training loop adopts the following early stopping heuristic:

- The model will be trained regardless of early stopping for burn_in epochs.
- Every check_interval epochs the method will compute the metric specified in criteria.

If such metric decreases for stop_interval checks, we stop training early.

Note the metric is computed on x_valid. This is usually a validation set that you held out.

Also, because criteria is a ranking metric, it requires generating negatives. Entities used to generate corruptions can be specified, as long as the side(s) of a triple to corrupt. The method supports filtered metrics, by passing an array of positives to x_filter . This will be used to filter the negatives generated on the fly (i.e. the corruptions).

Note: Keep in mind the early stopping criteria may introduce a certain overhead (caused by the metric computation). The goal is to strike a good trade-off between such overhead and saving training epochs.

A common approach is to use MRR unfiltered:

early_stopping_params={x_valid=X['valid'], 'criteria': 'mrr'}

Note the size of validation set also contributes to such overhead. In most cases a smaller validation set would be enough.

• **early_stopping_params** (*dictionary*) – Dictionary of hyperparameters for the early stopping heuristics.

The following string keys are supported:

- 'x_valid': ndarray, shape [n, 3] : Validation set to be used for early stopping.
- 'criteria': string : criteria for early stopping 'hits10', 'hits3', 'hits1' or 'mrr'(default).
- 'x_filter': ndarray, shape [n, 3] : Positive triples to use as filter if a 'filtered' early stopping criteria is desired (i.e. filtered-MRR if 'criteria':'mrr'). Note this will affect training time (no filter by default).
- 'burn_in': int : Number of epochs to pass before kicking in early stopping (de-fault: 100).
- check_interval': int : Early stopping interval after burn-in (default:10).
- 'stop_interval': int : Stop if criteria is performing worse over n consecutive checks (default: 3)
- 'corruption_entities': List of entities to be used for corruptions. If 'all', it uses all entities (default: 'all')
- 'corrupt_side': Specifies which side to corrupt. 's', 'o', 's+o' (default)

Example: early_stopping_params={x_valid=X['valid'],
'criteria': 'mrr'}

- **focusE_numeric_edge_values** (*nd array (n, 1)*) Numeric values associated with links. Semantically, the numeric value can signify importance, uncertainity, significance, confidence, etc. If the numeric value is unknown pass a NaN weight. The model will uniformly randomly assign a numeric value. One can also think about assigning numeric values by looking at the distribution of it per predicate.
- tensorboard_logs_path (*str or None*) Path to store tensorboard logs, e.g. average training loss tracking per epoch (default: None indicating no logs will be collected). When provided it will create a folder under provided path and save tensorboard files there. To then view the loss in the terminal run: tensorboard --logdir <tensorboard_logs_path>.

predict (X, from_idx=False)

Predict the scores of triples using a trained embedding model. The function returns raw scores generated by the model.

Note: To obtain probability estimates, calibrate the model with *calibrate()*, then call *predict_proba()*.

Parameters

- X (ndarray, shape [n, 3]) The triples to score.
- from_idx (bool) If True, will skip conversion to internal IDs. (default: False).

Returns scores_predict – The predicted scores for input triples X.

Return type ndarray, shape [n]

get_hyperparameter_dict()

Returns hyperparameters of the model.

Returns hyperparam_dict – Dictionary of hyperparameters that were used for training.

Return type dict

TransE

class	ampligraph.laten	t_features. Tran	sE (<i>k=100</i> ,	<i>eta=2</i> ,	epochs=100,
			batches_count=	=100, seed=	0, embed-
			ding_model_pa	rams={'corrupt_s	ides': ['s,o'],
			'negative_corrı	ption_entities': '	all', 'norm': 1,
			'normalize_ent	_emb': False}, o _f	otimizer='adam',
			optimizer_para	ms={'lr': 0.00	05}, loss='nll',
			loss_params={	}, regularizer=1	Vone, regular-
			izer_params={	}, initializer='x	avier', initial-
-			izer_params={	'uniform': False},	verbose=False)

Translating Embeddings (TransE)

The model as described in [BUGD+13].

The scoring function of TransE computes a similarity between the embedding of the subject \mathbf{e}_{sub} translated by the embedding of the predicate \mathbf{e}_{pred} and the embedding of the object \mathbf{e}_{obj} , using the L_1 or L_2 norm $|| \cdot ||$:

$$f_{TransE} = -||\mathbf{e}_{sub} + \mathbf{e}_{pred} - \mathbf{e}_{obj}||_m$$

Such scoring function is then used on positive and negative triples t^+, t^- in the loss function.

Examples

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```
>>>
                  ['c', 'y', 'a'],
>>>
                  ['a', 'y', 'd'],
                  ['c', 'y', 'd'],
>>>
                  ['b', 'y', 'c'],
>>>
                  ['f', 'y', 'e']])
>>>
>>> model.fit(X)
>>> model.predict(np.array([['f', 'y', 'e'], ['b', 'y', 'd']]))
[-4.6903257, -3.9047198]
>>> model.get_embeddings(['f','e'], embedding_type='entity')
array([[ 0.10673896, -0.28916815, 0.6278883 , -0.1194713 , -0.10372276,
-0.37258488, 0.06460134, -0.27879423, 0.25456288, 0.18665907],
[-0.64494324, -0.12939683, 0.3181001, 0.16745451, -0.03766293,
0.24314676, -0.23038973, -0.658638 , 0.5680542 , -0.05401703]],
dtype=float32)
```

Methods

init([k, eta, epochs, batches_count,])	Initialize an EmbeddingModel.
<pre>fit(X[, early_stopping,])</pre>	Train an Translating Embeddings model.
<pre>get_embeddings(entities[, embedding_type])</pre>	Get the embeddings of entities or relations.
get_hyperparameter_dict()	Returns hyperparameters of the model.
<pre>predict(X[, from_idx])</pre>	Predict the scores of triples using a trained embed-
	ding model.
<pre>calibrate(X_pos[, X_neg,])</pre>	Calibrate predictions
predict_proba(X)	Predicts probabilities using the Platt scaling model
	(after calibration).

Initialize an EmbeddingModel.

Also creates a new Tensorflow session for training.

Parameters

- **k** (*int*) Embedding space dimensionality.
- **eta** (*int*) The number of negatives that must be generated at runtime during training for each positive.
- epochs (*int*) The iterations of the training loop.
- **batches_count** (*int*) The number of batches in which the training set must be split during the training loop.
- **seed** (*int*) The seed used by the internal random numbers generator.
- **embedding_model_params** (*dict*) TransE-specific hyperparams, passed to the model as a dictionary.

Supported keys:

- 'norm' (int): the norm to be used in the scoring function (1 or 2-norm - default: 1).

- 'normalize_ent_emb' (bool): flag to indicate whether to normalize entity embeddings after each batch update (default: False).
- negative_corruption_entities : entities to be used for generation of corruptions while training. It can take the following values : all (default: all entities), batch (entities present in each batch), list of entities or an int (which indicates how many entities that should be used for corruption generation).
- corrupt_sides : Specifies how to generate corruptions for training. Takes values s, o, s+o or any combination passed as a list.
- 'non_linearity': can be one of the following values linear, softplus, sigmoid, tanh
- 'stop_epoch': specifies how long to decay (linearly) the numeric values from 1 to original value

until it reachs original value. - **'structural_wt'**: structural influence hyperparameter [0, 1] that modulates the influence of graph topology. - **'normalize_numeric_values'**: normalize the numeric values, such that they are scaled between [0, 1]

The last 4 parameters are related to FocusE layers.

```
Example: embedding_model_params={'norm': 1,
'normalize_ent_emb': False}
```

- **optimizer** (*string*) The optimizer used to minimize the loss function. Choose between 'sgd', 'adagrad', 'adam', 'momentum'.
- **optimizer_params** (*dict*) Arguments specific to the optimizer, passed as a dictionary.

Supported keys:

- 'lr' (float): learning rate (used by all the optimizers). Default: 0.1.
- 'momentum' (float): learning momentum (only used when optimizer=momentum). Default: 0.9.

Example: optimizer_params={ 'lr': 0.01}

- **loss** (*string*) The type of loss function to use during training.
 - pairwise the model will use pairwise margin-based loss function.
 - nll the model will use negative loss likelihood.
 - absolute_margin the model will use absolute margin likelihood.
 - self_adversarial the model will use adversarial sampling loss function.
 - multiclass_nll the model will use multiclass nll loss. Switch to multiclass loss defined in [aC15] by passing 'corrupt_sides' as ['s','o'] to embedding_model_params. To use loss defined in [KBK17] pass 'corrupt_sides' as 'o' to embedding_model_params.
- **loss_params** (*dict*) Dictionary of loss-specific hyperparameters. See *loss functions* documentation for additional details.

Example: optimizer_params={'lr': 0.01} if loss='pairwise'.

• **regularizer** (*string*) – The regularization strategy to use with the loss function.

- None: the model will not use any regularizer (default)
- 'LP': the model will use L1, L2 or L3 based on the value of regularizer params['p'] (see below).
- regularizer_params (dict) Dictionary of regularizer-specific hyperparameters. See the *regularizers* documentation for additional details.

Example: regularizer_params={'lambda': 1e-5, 'p': 2} if regularizer='LP'.

- **initializer** (*string*) The type of initializer to use.
 - normal: The embeddings will be initialized from a normal distribution
 - uniform: The embeddings will be initialized from a uniform distribution
 - xavier: The embeddings will be initialized using xavier strategy (default)
- initializer_params (dict) Dictionary of initializer-specific hyperparameters. See the *initializer* documentation for additional details.

```
Example: initializer params={'mean': 0, 'std': 0.001} if
initializer='normal'.
```

verbose [bool] Verbose mode

fit (X, early stopping=False, early stopping params={}, focusE numeric edge values=None, tensorboard logs path=None) Train an Translating Embeddings model.

The model is trained on a training set X using the training protocol described in [TWR+16].

Parameters

- X (ndarray, shape [n, 3]) The training triples
- **early_stopping** (*bool*) Flag to enable early stopping (default:False).

If set to True, the training loop adopts the following early stopping heuristic:

- The model will be trained regardless of early stopping for burn_in epochs.
- Every check_interval epochs the method will compute the metric specified in criteria.

If such metric decreases for stop interval checks, we stop training early.

Note the metric is computed on x_valid. This is usually a validation set that you held out.

Also, because criteria is a ranking metric, it requires generating negatives. Entities used to generate corruptions can be specified, as long as the side(s) of a triple to corrupt. The method supports filtered metrics, by passing an array of positives to x_filter. This will be used to filter the negatives generated on the fly (i.e. the corruptions).

Note: Keep in mind the early stopping criteria may introduce a certain overhead (caused by the metric computation). The goal is to strike a good trade-off between such overhead and saving training epochs.

A common approach is to use MRR unfiltered:

early_stopping_params={x_valid=X['valid'], 'criteria': 'mrr'}

Note the size of validation set also contributes to such overhead. In most cases a smaller validation set would be enough.

• **early_stopping_params** (*dictionary*) – Dictionary of hyperparameters for the early stopping heuristics.

The following string keys are supported:

- 'x_valid': ndarray, shape [n, 3] : Validation set to be used for early stopping.
- 'criteria': string : criteria for early stopping 'hits10', 'hits3', 'hits1' or 'mrr'(default).
- 'x_filter': ndarray, shape [n, 3] : Positive triples to use as filter if a 'filtered' early stopping criteria is desired (i.e. filtered-MRR if 'criteria': 'mrr'). Note this will affect training time (no filter by default).
- 'burn_in': int : Number of epochs to pass before kicking in early stopping (default: 100).
- check_interval': int : Early stopping interval after burn-in (default:10).
- 'stop_interval': int : Stop if criteria is performing worse over n consecutive checks (default: 3)
- 'corruption_entities': List of entities to be used for corruptions. If 'all', it uses all entities (default: 'all')
- 'corrupt_side': Specifies which side to corrupt. 's', 'o', 's+o' (default)

Example: early_stopping_params={x_valid=X['valid'],
'criteria': 'mrr'}

• **focusE_numeric_edge_values** (*ndarray*, *shape* [*n*]) – If processing a knowledge graph with numeric values associated with links, this is the vector of such numbers. Passing this argument will activate the *FocusE layer* [PC21]. Semantically, numeric values can signify importance, uncertainity, significance, confidence, etc. Values can be any number, and will be automatically normalised to the [0, 1] range, on a predicate-specific basis. If the numeric value is unknown pass a np.NaN value. The model will uniformly randomly assign a numeric value.

Note: The following toy example shows how to enable the FocusE layer to process edges with numeric literals:

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• tensorboard_logs_path (*str or None*) – Path to store tensorboard logs, e.g. average training loss tracking per epoch (default: None indicating no logs will be collected). When provided it will create a folder under provided path and save tensorboard files there. To then view the loss in the terminal run: tensorboard --logdir <tensorboard_logs_path>.

get_embeddings (entities, embedding_type='entity')

Get the embeddings of entities or relations.

Note: Use *ampligraph.utils.create_tensorboard_visualizations()* to visualize the embeddings with TensorBoard.

Parameters

- entities (array-like, dtype=int, shape=[n]) The entities (or relations) of interest. Element of the vector must be the original string literals, and not internal IDs.
- **embedding_type** (*string*) If 'entity', entities argument will be considered as a list of knowledge graph entities (i.e. nodes). If set to 'relation', they will be treated as relation types instead (i.e. predicates).

Returns embeddings - An array of k-dimensional embeddings.

Return type ndarray, shape [n, k]

get_hyperparameter_dict()

Returns hyperparameters of the model.

Returns hyperparam_dict – Dictionary of hyperparameters that were used for training.

Return type dict

predict (X, from_idx=False)

Predict the scores of triples using a trained embedding model. The function returns raw scores generated by the model.

Note: To obtain probability estimates, calibrate the model with *calibrate()*, then call *predict_proba()*.

Parameters

• X (ndarray, shape [n, 3]) – The triples to score.

• from_idx (bool) – If True, will skip conversion to internal IDs. (default: False).

Returns scores_predict – The predicted scores for input triples X.

Return type ndarray, shape [n]

```
calibrate (X_pos, X_neg=None, positive_base_rate=None, batches_count=100, epochs=50)
Calibrate predictions
```

The method implements the heuristics described in [TC20], using Platt scaling [P+99].

The calibrated predictions can be obtained with *predict_proba()* after calibration is done.

Ideally, calibration should be performed on a validation set that was not used to train the embeddings.

There are two modes of operation, depending on the availability of negative triples:

- 1. Both positive and negative triples are provided via X_pos and X_neg respectively. The optimization is done using a second-order method (limited-memory BFGS), therefore no hyperparameter needs to be specified.
- 2. Only positive triples are provided, and the negative triples are generated by corruptions just like it is done in training or evaluation. The optimization is done using a first-order method (ADAM), therefore batches_count and epochs must be specified.

Calibration is highly dependent on the base rate of positive triples. Therefore, for mode (2) of operation, the user is required to provide the positive_base_rate argument. For mode (1), that can be inferred automatically by the relative sizes of the positive and negative sets, but the user can override that by providing a value to positive_base_rate.

Defining the positive base rate is the biggest challenge when calibrating without negatives. That depends on the user choice of which triples will be evaluated during test time. Let's take WN11 as an example: it has around 50% positives triples on both the validation set and test set, so naturally the positive base rate is 50%. However, should the user resample it to have 75% positives and 25% negatives, its previous calibration will be degraded. The user must recalibrate the model now with a 75% positive base rate. Therefore, this parameter depends on how the user handles the dataset and cannot be determined automatically or a priori.

Note: Incompatible with large graph mode (i.e. if self.dealing_with_large_graphs=True).

Note: Experiments for the ICLR-21 calibration paper are available here [TC20].

Parameters

- X_pos (ndarray (shape [n, 3])) Numpy array of positive triples.
- X_neg (ndarray (shape [n, 3])) Numpy array of negative triples.

If *None*, the negative triples are generated via corruptions and the user must provide a positive base rate instead.

• **positive_base_rate** (*float*) – Base rate of positive statements.

For example, if we assume there is a fifty-fifty chance of any query to be true, the base rate would be 50%.

If X_neg is provided and this is *None*, the relative sizes of X_pos and X_neg will be used to determine the base rate. For example, if we have 50 positive triples and

200 negative triples, the positive base rate will be assumed to be 50/(50+200) = 1/5 = 0.2.

This must be a value between 0 and 1.

- **batches_count** (*int*) Number of batches to complete one epoch of the Platt scaling training. Only applies when X_neg is *None*.
- **epochs** (*int*) Number of epochs used to train the Platt scaling model. Only applies when X_neg is *None*.

Examples

```
>>> import numpy as np
>>> from sklearn.metrics import brier_score_loss, log_loss
>>> from scipy.special import expit
>>>
>>> from ampligraph.datasets import load_wn11
>>> from ampligraph.latent_features.models import TransE
>>>
>>> X = load_wn11()
>>> X_valid_pos = X['valid'][X['valid_labels']]
>>> X_valid_neg = X['valid'][~X['valid_labels']]
>>>
>>> model = TransE(batches_count=64, seed=0, epochs=500, k=100, eta=20,
>>>
                   optimizer='adam', optimizer_params={'lr':0.0001},
                   loss='pairwise', verbose=True)
>>>
>>>
>>> model.fit(X['train'])
>>>
>>> # Raw scores
>>> scores = model.predict(X['test'])
>>>
>>> # Calibrate with positives and negatives
>>> model.calibrate(X_valid_pos, X_valid_neg, positive_base_rate=None)
>>> probas_pos_neg = model.predict_proba(X['test'])
>>>
>>> # Calibrate with just positives and base rate of 50%
>>> model.calibrate(X_valid_pos, positive_base_rate=0.5)
>>> probas_pos = model.predict_proba(X['test'])
>>>
>>> # Calibration evaluation with the Brier score loss (the smaller, the
\rightarrow better)
>>> print("Brier scores")
>>> print("Raw scores:", brier_score_loss(X['test_labels'], expit(scores)))
>>> print("Positive and negative calibration:", brier_score_loss(X['test_
→labels'], probas_pos_neg))
>>> print("Positive only calibration:", brier_score_loss(X['test_labels'],_

→probas_pos))
Brier scores
Raw scores: 0.4925058891371126
Positive and negative calibration: 0.20434617882733366
Positive only calibration: 0.22597599585144656
```

$predict_proba(X)$

Predicts probabilities using the Platt scaling model (after calibration).

Model must be calibrated beforehand with the calibrate method.

Parameters X (*ndarray* (*shape* [n, 3])) – Numpy array of triples to be evaluated.
Returns probas – Probability of each triple to be true according to the Platt scaling calibration.

Return type ndarray (shape [n])

DistMult

```
class ampligraph.latent_features.DistMult(k=100,
                                                                                        epochs=100,
                                                                         eta=2.
                                                         batches count=100,
                                                                                seed=0,
                                                                                             embed-
                                                         ding_model_params={'corrupt_sides': ['s,o'],
                                                         'negative corruption entities':
                                                                                         'all',
                                                                                               'nor-
                                                         malize_ent_emb': False}, optimizer='adam',
                                                         optimizer_params={'lr': 0.0005}, loss='nll',
                                                         loss_params={},
                                                                           regularizer=None,
                                                                                                 reg-
                                                         ularizer_params={},
                                                                                  initializer='xavier',
                                                                                        False}, ver-
                                                         initializer_params={'uniform':
                                                         bose=False)
```

The DistMult model

The model as described in [YYH+14].

The bilinear diagonal DistMult model uses the trilinear dot product as scoring function:

$$f_{DistMult} = \langle \mathbf{r}_p, \mathbf{e}_s, \mathbf{e}_o \rangle$$

where \mathbf{e}_s is the embedding of the subject, \mathbf{r}_p the embedding of the predicate and \mathbf{e}_o the embedding of the object.

Examples

```
>>> import numpy as np
>>> from ampligraph.latent_features import DistMult
>>> model = DistMult(batches_count=1, seed=555, epochs=20, k=10, loss='pairwise',
            loss_params={'margin':5})
>>>
>>> X = np.array([['a', 'y', 'b'],
                        'y', 'a'],
>>>
                  ['b',
>>>
                  ['a',
                        'y', 'c'],
                  ['c',
>>>
                        'y', 'a'],
                  ['a',
                            'd'],
                        'y',
>>>
                  ['c',
                        'y', 'd'],
>>>
                  ['b', 'y', 'c'],
>>>
>>>
                  ['f', 'y', 'e']])
>>> model.fit(X)
>>> model.predict(np.array([['f', 'y', 'e'], ['b', 'y', 'd']]))
[-0.13863425, -0.09917116]
>>> model.get_embeddings(['f','e'], embedding_type='entity')
array([[ 0.10137264, -0.28248304, 0.6153027, -0.13133956, -0.11675504,
-0.37876177, 0.06027773, -0.26390398, 0.254603 , 0.1888549 ],
[-0.6467299, -0.13729756, 0.3074872, 0.16966867, -0.04098966,
0.25289047, -0.2212451, -0.6527815, 0.5657673, -0.03876532]],
dtype=float32)
```

Methods

	Initialize an EmbeddingModel
<pre>fit(X[, early_stopping,])</pre>	Train an DistMult.
<pre>get_embeddings(entities[, embedding_type])</pre>	Get the embeddings of entities or relations.
get_hyperparameter_dict()	Returns hyperparameters of the model.
<pre>predict(X[, from_idx])</pre>	Predict the scores of triples using a trained embed-
	ding model.
<pre>calibrate(X_pos[, X_neg,])</pre>	Calibrate predictions
predict_proba(X)	Predicts probabilities using the Platt scaling model
	(after calibration).

__init__ (k=100, eta=2, epochs=100, batches_count=100, seed=0, embedding_model_params={'corrupt_sides': ['s,o'], 'negative_corruption_entities': 'all', 'normalize_ent_emb': False}, optimizer='adam', optimizer_params={'lr': 0.0005}, loss='nll', loss_params={}, regularizer=None, regularizer_params={}, initializer='xavier', initializer_params={'uniform': False}, verbose=False) Initialize an EmbeddingModel

Also creates a new Tensorflow session for training.

Parameters

- **k** (*int*) Embedding space dimensionality
- **eta** (*int*) The number of negatives that must be generated at runtime during training for each positive.
- epochs (*int*) The iterations of the training loop.
- **batches_count** (*int*) The number of batches in which the training set must be split during the training loop.
- **seed** (*int*) The seed used by the internal random numbers generator.
- **embedding_model_params** (*dict*) DistMult-specific hyperparams, passed to the model as a dictionary.

Supported keys:

- 'normalize_ent_emb' (bool): flag to indicate whether to normalize entity embeddings after each batch update (default: False).
- 'negative_corruption_entities' Entities to be used for generation of corruptions while training. It can take the following values : all (default: all entities), batch (entities present in each batch), list of entities or an int (which indicates how many entities that should be used for corruption generation).
- corrupt_sides : Specifies how to generate corruptions for training. Takes values s,
 o, s+o or any combination passed as a list
- 'non_linearity': can be one of the following values linear, softplus, sigmoid, tanh
- 'stop_epoch': specifies how long to decay (linearly) the numeric values from 1 to original value

until it reachs original value. - **'structural_wt'**: structural influence hyperparameter [0, 1] that modulates the influence of graph topology. - **'normalize_numeric_values'**: normalize the numeric values, such that they are scaled between [0, 1]

The last 4 parameters are related to FocusE layers.

Example: embedding_model_params={'normalize_ent_emb': False}

- **optimizer** (*string*) The optimizer used to minimize the loss function. Choose between 'sgd', 'adagrad', 'adam', 'momentum'.
- **optimizer_params** (*dict*) Arguments specific to the optimizer, passed as a dictionary.

Supported keys:

- 'lr' (float): learning rate (used by all the optimizers). Default: 0.1.
- 'momentum' (float): learning momentum (only used when optimizer=momentum). Default: 0.9.

Example: optimizer_params={ 'lr': 0.01}

- **loss** (*string*) The type of loss function to use during training.
 - pairwise the model will use pairwise margin-based loss function.
 - nll the model will use negative loss likelihood.
 - absolute_margin the model will use absolute margin likelihood.
 - self_adversarial the model will use adversarial sampling loss function.
 - multiclass_nll the model will use multiclass nll loss. Switch to multiclass loss defined in [aC15] by passing 'corrupt_sides' as ['s','o'] to embedding_model_params. To use loss defined in [KBK17] pass 'corrupt_sides' as 'o' to embedding_model_params.
- **loss_params** (*dict*) Dictionary of loss-specific hyperparameters. See *loss functions* documentation for additional details.

Example: optimizer_params={'lr': 0.01} if loss='pairwise'.

- **regularizer** (*string*) The regularization strategy to use with the loss function.
 - None: the model will not use any regularizer (default)
 - 'LP': the model will use L1, L2 or L3 based on the value of regularizer_params['p'] (see below).
- **regularizer_params** (*dict*) Dictionary of regularizer-specific hyperparameters. See the *regularizers* documentation for additional details.

```
Example: regularizer_params={'lambda': 1e-5, 'p': 2} if
regularizer='LP'.
```

- **initializer** (*string*) The type of initializer to use.
 - normal: The embeddings will be initialized from a normal distribution
 - uniform: The embeddings will be initialized from a uniform distribution
 - xavier: The embeddings will be initialized using xavier strategy (default)
- **initializer_params** (*dict*) Dictionary of initializer-specific hyperparameters. See the *initializer* documentation for additional details.

```
Example: initializer_params={'mean': 0, 'std': 0.001} if
initializer='normal'.
```

• **verbose** (*bool*) – Verbose mode.

fit (X, early_stopping=False, early_stopping_params={}, focusE_numeric_edge_values=None, tensorboard_logs_path=None) Train an DistMult.

The model is trained on a training set X using the training protocol described in [TWR+16].

Parameters

- X (ndarray, shape [n, 3]) The training triples
- **early_stopping** (bool) Flag to enable early stopping (default:False).

If set to True, the training loop adopts the following early stopping heuristic:

- The model will be trained regardless of early stopping for burn_in epochs.
- Every check_interval epochs the method will compute the metric specified in criteria.

If such metric decreases for stop_interval checks, we stop training early.

Note the metric is computed on x_valid . This is usually a validation set that you held out.

Also, because criteria is a ranking metric, it requires generating negatives. Entities used to generate corruptions can be specified, as long as the side(s) of a triple to corrupt. The method supports filtered metrics, by passing an array of positives to x_filter . This will be used to filter the negatives generated on the fly (i.e. the corruptions).

Note: Keep in mind the early stopping criteria may introduce a certain overhead (caused by the metric computation). The goal is to strike a good trade-off between such overhead and saving training epochs.

A common approach is to use MRR unfiltered:

early_stopping_params={x_valid=X['valid'], 'criteria': 'mrr'}

Note the size of validation set also contributes to such overhead. In most cases a smaller validation set would be enough.

• **early_stopping_params** (*dictionary*) – Dictionary of hyperparameters for the early stopping heuristics.

The following string keys are supported:

- 'x_valid': ndarray, shape [n, 3] : Validation set to be used for early stopping.
- 'criteria': string : criteria for early stopping 'hits10', 'hits3', 'hits1' or 'mrr'(default).
- 'x_filter': ndarray, shape [n, 3] : Positive triples to use as filter if a 'filtered' early stopping criteria is desired (i.e. filtered-MRR if 'criteria': 'mrr'). Note this will affect training time (no filter by default).
- 'burn_in': int : Number of epochs to pass before kicking in early stopping (default: 100).
- check_interval': int : Early stopping interval after burn-in (default:10).
- 'stop_interval': int : Stop if criteria is performing worse over n consecutive checks (default: 3)

- 'corruption_entities': List of entities to be used for corruptions. If 'all', it uses all entities (default: 'all')
- 'corrupt_side': Specifies which side to corrupt. 's', 'o', 's+o' (default)

```
Example: early_stopping_params={x_valid=X['valid'],
'criteria': 'mrr'}
```

• **focusE_numeric_edge_values** (*ndarray*, *shape* [*n*]) – If processing a knowledge graph with numeric values associated with links, this is the vector of such numbers. Passing this argument will activate the *FocusE layer* [PC21]. Semantically, numeric values can signify importance, uncertainity, significance, confidence, etc. Values can be any number, and will be automatically normalised to the [0, 1] range, on a predicate-specific basis. If the numeric value is unknown pass a np.NaN value. The model will uniformly randomly assign a numeric value.

Note: The following toy example shows how to enable the FocusE layer to process edges with numeric literals:

```
import numpy as np
from ampligraph.latent_features import DistMult
model = DistMult(batches_count=1, seed=555, epochs=20,
                 k=10, loss='pairwise',
                 loss_params={'margin':5})
X = np.array([['a', 'y', 'b']],
              ['b', 'y', 'a'],
              ['a', 'y', 'c'],
              ['c', 'y', 'a'],
              ['a', 'y', 'd'],
              ['c', 'y', 'd'],
              ['b', 'y', 'c'],
              ['f', 'y', 'e']])
# Numeric values below are associate to each triple in X.
# They can be any number and will be automatically
# normalised to the [0, 1] range, on a
# predicate-specific basis.
X_edge_values = np.array([5.34, -1.75, 0.33, 5.12,
                          np.nan, 3.17, 2.76, 0.41])
model.fit(X, focusE_numeric_edge_values=X_edge_values)
```

tensorboard_logs_path: str or None Path to store tensorboard logs, e.g. average training loss tracking per epoch (default: None indicating no logs will be collected). When provided it will create a folder under provided path and save tensorboard files there. To then view the loss in the terminal run: tensorboard --logdir <tensorboard_logs_path>.

get_embeddings (*entities*, *embedding_type='entity'*) Get the embeddings of entities or relations.

Note: Use *ampligraph.utils.create_tensorboard_visualizations()* to visualize the embeddings with TensorBoard.

Parameters

- entities (array-like, dtype=int, shape=[n]) The entities (or relations) of interest. Element of the vector must be the original string literals, and not internal IDs.
- **embedding_type** (*string*) If 'entity', entities argument will be considered as a list of knowledge graph entities (i.e. nodes). If set to 'relation', they will be treated as relation types instead (i.e. predicates).

Returns embeddings – An array of k-dimensional embeddings.

Return type ndarray, shape [n, k]

get_hyperparameter_dict()

Returns hyperparameters of the model.

Returns hyperparam_dict – Dictionary of hyperparameters that were used for training.

Return type dict

predict (X, from_idx=False)

Predict the scores of triples using a trained embedding model. The function returns raw scores generated by the model.

Note: To obtain probability estimates, calibrate the model with *calibrate()*, then call *predict_proba()*.

Parameters

- X (ndarray, shape [n, 3]) The triples to score.
- from_idx (bool) If True, will skip conversion to internal IDs. (default: False).

Returns scores_predict – The predicted scores for input triples X.

Return type ndarray, shape [n]

calibrate (*X_pos*, *X_neg=None*, *positive_base_rate=None*, *batches_count=100*, *epochs=50*) Calibrate predictions

The method implements the heuristics described in [TC20], using Platt scaling [P+99].

The calibrated predictions can be obtained with *predict_proba()* after calibration is done.

Ideally, calibration should be performed on a validation set that was not used to train the embeddings.

There are two modes of operation, depending on the availability of negative triples:

- 1. Both positive and negative triples are provided via X_pos and X_neg respectively. The optimization is done using a second-order method (limited-memory BFGS), therefore no hyperparameter needs to be specified.
- 2. Only positive triples are provided, and the negative triples are generated by corruptions just like it is done in training or evaluation. The optimization is done using a first-order method (ADAM), therefore batches_count and epochs must be specified.

Calibration is highly dependent on the base rate of positive triples. Therefore, for mode (2) of operation, the user is required to provide the positive_base_rate argument. For mode (1), that can be inferred automatically by the relative sizes of the positive and negative sets, but the user can override that by providing a value to positive_base_rate.

Defining the positive base rate is the biggest challenge when calibrating without negatives. That depends on the user choice of which triples will be evaluated during test time. Let's take WN11 as an example: it has around 50% positives triples on both the validation set and test set, so naturally the positive base rate is 50%. However, should the user resample it to have 75% positives and 25% negatives, its previous calibration will be degraded. The user must recalibrate the model now with a 75% positive base rate. Therefore, this parameter depends on how the user handles the dataset and cannot be determined automatically or a priori.

Note: Incompatible with large graph mode (i.e. if self.dealing_with_large_graphs=True).

Note: Experiments for the ICLR-21 calibration paper are available here [TC20].

Parameters

- X_pos (ndarray (shape [n, 3])) Numpy array of positive triples.
- X_neg (ndarray (shape [n, 3])) Numpy array of negative triples.

If *None*, the negative triples are generated via corruptions and the user must provide a positive base rate instead.

• **positive_base_rate** (*float*) – Base rate of positive statements.

For example, if we assume there is a fifty-fifty chance of any query to be true, the base rate would be 50%.

If X_neg is provided and this is *None*, the relative sizes of X_pos and X_neg will be used to determine the base rate. For example, if we have 50 positive triples and 200 negative triples, the positive base rate will be assumed to be 50/(50+200) = 1/5 = 0.2.

This must be a value between 0 and 1.

- **batches_count** (*int*) Number of batches to complete one epoch of the Platt scaling training. Only applies when X_neg is *None*.
- **epochs** (*int*) Number of epochs used to train the Platt scaling model. Only applies when X_neg is *None*.

Examples

```
>>> import numpy as np
>>> from sklearn.metrics import brier_score_loss, log_loss
>>> from scipy.special import expit
>>>
>>> from ampligraph.datasets import load_wn11
>>> from ampligraph.latent_features.models import TransE
>>>
>>> X = load_wn11()
>>> X_valid_pos = X['valid'][X['valid_labels']]
>>> X_valid_neg = X['valid'][~X['valid_labels']]
>>>
>>> model = TransE(batches_count=64, seed=0, epochs=500, k=100, eta=20,
                   optimizer='adam', optimizer_params={'lr':0.0001},
>>>
                   loss='pairwise', verbose=True)
>>>
```

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```
>>>
>>> model.fit(X['train'])
>>>
>>> # Raw scores
>>> scores = model.predict(X['test'])
>>>
>>> # Calibrate with positives and negatives
>>> model.calibrate(X_valid_pos, X_valid_neg, positive_base_rate=None)
>>> probas_pos_neg = model.predict_proba(X['test'])
>>>
>>> # Calibrate with just positives and base rate of 50%
>>> model.calibrate(X_valid_pos, positive_base_rate=0.5)
>>> probas_pos = model.predict_proba(X['test'])
>>>
>>> # Calibration evaluation with the Brier score loss (the smaller, the
\rightarrow better)
>>> print("Brier scores")
>>> print("Raw scores:", brier_score_loss(X['test_labels'], expit(scores)))
>>> print("Positive and negative calibration:", brier_score_loss(X['test_
→labels'], probas_pos_neg))
>>> print("Positive only calibration:", brier_score_loss(X['test_labels'],_
\rightarrow probas_pos))
Brier scores
Raw scores: 0.4925058891371126
Positive and negative calibration: 0.20434617882733366
Positive only calibration: 0.22597599585144656
```

$predict_proba(X)$

Predicts probabilities using the Platt scaling model (after calibration).

Model must be calibrated beforehand with the calibrate method.

Parameters X (*ndarray* (*shape* [*n*, 3])) – Numpy array of triples to be evaluated.

Returns probas – Probability of each triple to be true according to the Platt scaling calibration.

Return type ndarray (shape [n])

ComplEx

```
class ampligraph.latent features.ComplEx (k=100,
                                                                        eta=2,
                                                                                        epochs=100,
                                                       batches count=100,
                                                                               seed=0.
                                                                                             embed-
                                                       ding model params={'corrupt sides':
                                                                                             ['s, o'],
                                                       'negative_corruption_entities':
                                                                                        `all`},
                                                                                                op-
                                                       timizer='adam',
                                                                             optimizer params={'lr':
                                                       0.0005}, loss='nll', loss_params={}, regular-
                                                       izer=None.
                                                                    regularizer params={},
                                                                                             initial-
                                                       izer='xavier',
                                                                       initializer_params={'uniform':
                                                       False, verbose=False)
```

Complex embeddings (ComplEx)

The ComplEx model [TWR+16] is an extension of the *ampligraph.latent_features.DistMult* bilinear diagonal model. ComplEx scoring function is based on the trilinear Hermitian dot product in C:

$$f_{ComplEx} = Re(\langle \mathbf{r}_p, \mathbf{e}_s, \overline{\mathbf{e}_o} \rangle)$$

ComplEx can be improved if used alongside the nuclear 3-norm (the **ComplEx-N3** model [LUO18]), which can be easily added to the loss function via the regularizer hyperparameter with p=3 and a chosen regularisation weight (represented by lambda), as shown in the example below. See also *ampligraph*. *latent_features.LPRegularizer()*.

Note: Since ComplEx embeddings belong to C, this model uses twice as many parameters as *ampligraph*. *latent_features.DistMult*.

Examples

```
>>> import numpy as np
>>> from ampligraph.latent features import ComplEx
>>>
>>> model = ComplEx (batches_count=2, seed=555, epochs=100, k=20, eta=5,
>>>
               loss='pairwise', loss_params={'margin':1},
               regularizer='LP', regularizer_params={'p': 2, 'lambda':0.1})
>>>
>>> X = np.array([['a', 'y', 'b'],
                  ['b', 'y', 'a'],
>>>
                  ['a', 'y', 'c'],
>>>
                  ['c', 'y', 'a'],
>>>
                  ['a', 'y', 'd'],
>>>
                  ['c', 'y', 'd'],
>>>
                  ['b', 'y', 'c'],
>>>
                  ['f', 'y', 'e']])
>>>
>>> model.fit(X)
>>> model.predict(np.array([['f', 'y', 'e'], ['b', 'y', 'd']]))
[[0.019520484], [-0.14998421]]
>>> model.get_embeddings(['f','e'], embedding_type='entity')
array([[-0.33021057, 0.26524785, 0.0446662, -0.07932718, -0.15453218,
    -0.22342539, -0.03382565, 0.17444217, 0.03009969, -0.33569157,
    0.3200497, 0.03803705, 0.05536304, -0.00929996, 0.24446663,
    0.34408194, 0.16192885, -0.15033236, -0.19703785, -0.00783876,
    0.1495124 , -0.3578853 , -0.04975723 , -0.03930473 , 0.1663541 ,
    -0.24731971, -0.141296 , 0.03150219, 0.15328223, -0.18549544,
    -0.39240393, -0.10824018, 0.03394471, -0.11075485, 0.1367736,
    0.10059565, -0.32808647, -0.00472086, 0.14231135, -0.13876757],
   [-0.09483694, 0.3531292, 0.04992269, -0.07774793, 0.1635035,
    0.30610007, 0.3666711 , -0.13785957, -0.3143734 , -0.36909637,
    -0.13792469, -0.07069954, -0.0368113 , -0.16743314, 0.4090072 ,
   -0.03407392, 0.3113114, -0.08418448, 0.21435146, 0.12006859,
    0.08447982, -0.02025972, 0.38752195, 0.11451488, -0.0258422,
    -0.10990044, -0.22661531, -0.00478273, -0.0238297 , -0.14207476,
    0.11064807, 0.20135397, 0.22501846, -0.1731076, -0.2770435,
    0.30784574, -0.15043163, -0.11599299, 0.05718031, -0.1300622 ]],
  dtype=float32)
```

Methods

init([k, eta, epochs, batches_count,])	Initialize an EmbeddingModel
<pre>fit(X[, early_stopping,])</pre>	Train a ComplEx model.
<pre>get_embeddings(entities[, embedding_type])</pre>	Get the embeddings of entities or relations.
	Continued on next page

Table 6 – continued from previous page	
<pre>get_hyperparameter_dict()</pre>	Returns hyperparameters of the model.
<pre>predict(X[, from_idx])</pre>	Predict the scores of triples using a trained embed-
	ding model.
<pre>calibrate(X_pos[, X_neg,])</pre>	Calibrate predictions
predict_proba(X)	Predicts probabilities using the Platt scaling model
	(after calibration).

Table 8 - continued from previous page

__init___(k=100, eta=2, epochs=100, batches_count=100, seed=0, embedding_model_params={'corrupt_sides': ['s,o'], 'negative_corruption_entities': 'all'}, optimizer='adam', optimizer_params={'lr': 0.0005}, loss='nll', loss_params={}, regularizer=None, regularizer_params={}, initializer='xavier', initializer_params={'uniform': False}, verbose=False)

Initialize an EmbeddingModel

Also creates a new Tensorflow session for training.

Parameters

- **k** (*int*) Embedding space dimensionality
- **eta** (*int*) The number of negatives that must be generated at runtime during training for each positive.
- epochs (*int*) The iterations of the training loop.
- **batches_count** (*int*) The number of batches in which the training set must be split during the training loop.
- **seed** (*int*) The seed used by the internal random numbers generator.
- **embedding_model_params** (*dict*) ComplEx-specific hyperparams:
 - 'negative_corruption_entities' Entities to be used for generation of corruptions while training. It can take the following values : all (default: all entities), batch (entities present in each batch), list of entities or an int (which indicates how many entities that should be used for corruption generation).
 - corrupt_sides : Specifies how to generate corruptions for training. Takes values s,
 o, s+o or any combination passed as a list
 - 'non_linearity': can be one of the following values linear, softplus, sigmoid, tanh
 - 'stop_epoch': specifies how long to decay (linearly) the numeric values from 1 to original value

until it reachs original value. - **'structural_wt'**: structural influence hyperparameter [0, 1] that modulates the influence of graph topology. - **'normalize_numeric_values'**: normalize the numeric values, such that they are scaled between [0, 1]

The last 4 parameters are related to FocusE layers.

- **optimizer** (*string*) The optimizer used to minimize the loss function. Choose between 'sgd', 'adagrad', 'adam', 'momentum'.
- **optimizer_params** (*dict*) Arguments specific to the optimizer, passed as a dictionary.

Supported keys:

- 'lr' (float): learning rate (used by all the optimizers). Default: 0.1.

- 'momentum' (float): learning momentum (only used when optimizer=momentum). Default: 0.9.

Example: optimizer_params={ 'lr': 0.01}

- **loss** (*string*) The type of loss function to use during training.
 - pairwise the model will use pairwise margin-based loss function.
 - nll the model will use negative loss likelihood.
 - absolute_margin the model will use absolute margin likelihood.
 - self_adversarial the model will use adversarial sampling loss function.
 - multiclass_nll the model will use multiclass nll loss. Switch to multiclass loss defined in [aC15] by passing 'corrupt_sides' as ['s','o'] to embedding_model_params. To use loss defined in [KBK17] pass 'corrupt_sides' as 'o' to embedding_model_params.
- **loss_params** (*dict*) Dictionary of loss-specific hyperparameters. See *loss functions* documentation for additional details.

Example: optimizer_params={'lr': 0.01} if loss='pairwise'.

- **regularizer** (*string*) The regularization strategy to use with the loss function.
 - None: the model will not use any regularizer (default)
 - 'LP': the model will use L1, L2 or L3 based on the value of regularizer_params['p'] (see below).
- **regularizer_params** (*dict*) Dictionary of regularizer-specific hyperparameters. See the *regularizers* documentation for additional details.

Example: regularizer_params={'lambda': 1e-5, 'p': 2} if
regularizer='LP'.

- initializer (*string*) The type of initializer to use.
 - normal: The embeddings will be initialized from a normal distribution
 - uniform: The embeddings will be initialized from a uniform distribution
 - xavier: The embeddings will be initialized using xavier strategy (default)
- **initializer_params** (*dict*) Dictionary of initializer-specific hyperparameters. See the *initializer* documentation for additional details.

Example: initializer_params={'mean': 0, 'std': 0.001} if initializer='normal'.

- **verbose** (*bool*) Verbose mode.
- fit (X, early_stopping=False, early_stopping_params={}, focusE_numeric_edge_values=None, tensorboard_logs_path=None) Train a ComplEx model.

The model is trained on a training set X using the training protocol described in [TWR+16].

Parameters

- X (ndarray, shape [n, 3]) The training triples
- **early_stopping** (bool) Flag to enable early stopping (default:False).

If set to True, the training loop adopts the following early stopping heuristic:

- The model will be trained regardless of early stopping for burn_in epochs.
- Every check_interval epochs the method will compute the metric specified in criteria.

If such metric decreases for stop_interval checks, we stop training early.

Note the metric is computed on x_valid. This is usually a validation set that you held out.

Also, because criteria is a ranking metric, it requires generating negatives. Entities used to generate corruptions can be specified, as long as the side(s) of a triple to corrupt. The method supports filtered metrics, by passing an array of positives to x_filter . This will be used to filter the negatives generated on the fly (i.e. the corruptions).

Note: Keep in mind the early stopping criteria may introduce a certain overhead (caused by the metric computation). The goal is to strike a good trade-off between such overhead and saving training epochs.

A common approach is to use MRR unfiltered:

early_stopping_params={x_valid=X['valid'], 'criteria': 'mrr'}

Note the size of validation set also contributes to such overhead. In most cases a smaller validation set would be enough.

• **early_stopping_params** (*dictionary*) – Dictionary of hyperparameters for the early stopping heuristics.

The following string keys are supported:

- 'x_valid': ndarray, shape [n, 3] : Validation set to be used for early stopping.
- 'criteria': string : criteria for early stopping 'hits10', 'hits3', 'hits1' or 'mrr'(default).
- 'x_filter': ndarray, shape [n, 3]: Positive triples to use as filter if a 'filtered' early stopping criteria is desired (i.e. filtered-MRR if 'criteria':'mrr'). Note this will affect training time (no filter by default).
- 'burn_in': int : Number of epochs to pass before kicking in early stopping (default: 100).
- check_interval': int : Early stopping interval after burn-in (default:10).
- 'stop_interval': int : Stop if criteria is performing worse over n consecutive checks (default: 3)
- 'corruption_entities': List of entities to be used for corruptions. If 'all', it uses all entities (default: 'all')
- 'corrupt_side': Specifies which side to corrupt. 's', 'o', 's+o' (default)

Example: early_stopping_params={x_valid=X['valid'],
'criteria': 'mrr'}

• **focusE_numeric_edge_values** (*ndarray*, *shape* [*n*]) – If processing a knowledge graph with numeric values associated with links, this is the vector of such numbers. Passing this argument will activate the *FocusE layer* [PC21]. Semantically, numeric values can signify importance, uncertainity, significance, confidence, etc.

Values can be any number, and will be automatically normalised to the [0, 1] range, on a predicate-specific basis. If the numeric value is unknown pass a np.NaN value. The model will uniformly randomly assign a numeric value.

Note: The following toy example shows how to enable the FocusE layer to process edges with numeric literals:

```
import numpy as np
from ampligraph.latent_features import ComplEx
model = ComplEx(batches_count=1, seed=555, epochs=20,
               k=10, loss='pairwise',
               loss_params={'margin':5})
X = np.array([['a', 'y', 'b']],
              ['b', 'y', 'a'],
              ['a', 'y', 'c'],
              ['c', 'y', 'a'],
              ['a', 'y', 'd'],
              ['c', 'y', 'd'],
              ['b', 'y', 'c'],
              ['f', 'y', 'e']])
# Numeric values below are associate to each triple in X.
# They can be any number and will be automatically
# normalised to the [0, 1] range, on a
# predicate-specific basis.
X_edge_values = np.array([5.34, -1.75, 0.33, 5.12,
                          np.nan, 3.17, 2.76, 0.41])
model.fit(X, focusE_numeric_edge_values=X_edge_values)
```

- **tensorboard_logs_path: str or None** Path to store tensorboard logs, e.g. average training loss tracking per epoch (default: None indicating no logs will be collected). When provided it will create a folder under provided path and save tensorboard files there. To then view the loss in the terminal run: tensorboard --logdir <tensorboard_logs_path>.
- **get_embeddings** (*entities*, *embedding_type='entity'*) Get the embeddings of entities or relations.

Note: Use ${\tt ampligraph.utils.create_tensorboard_visualizations()}$ to visualize the embeddings with TensorBoard.

Parameters

- entities (array-like, dtype=int, shape=[n]) The entities (or relations) of interest. Element of the vector must be the original string literals, and not internal IDs.
- **embedding_type** (*string*) If 'entity', entities argument will be considered as a list of knowledge graph entities (i.e. nodes). If set to 'relation', they will be treated as relation types instead (i.e. predicates).

Returns embeddings – An array of k-dimensional embeddings.

Return type ndarray, shape [n, k]

get_hyperparameter_dict()

Returns hyperparameters of the model.

Returns hyperparam_dict – Dictionary of hyperparameters that were used for training.

Return type dict

predict (X, from_idx=False)

Predict the scores of triples using a trained embedding model. The function returns raw scores generated by the model.

Note: To obtain probability estimates, calibrate the model with *calibrate()*, then call *predict_proba()*.

Parameters

- X (ndarray, shape [n, 3]) The triples to score.
- from_idx (bool) If True, will skip conversion to internal IDs. (default: False).

Returns scores_predict – The predicted scores for input triples X.

Return type ndarray, shape [n]

calibrate (*X_pos*, *X_neg=None*, *positive_base_rate=None*, *batches_count=100*, *epochs=50*) Calibrate predictions

The method implements the heuristics described in [TC20], using Platt scaling [P+99].

The calibrated predictions can be obtained with *predict_proba()* after calibration is done.

Ideally, calibration should be performed on a validation set that was not used to train the embeddings.

There are two modes of operation, depending on the availability of negative triples:

- 1. Both positive and negative triples are provided via X_pos and X_neg respectively. The optimization is done using a second-order method (limited-memory BFGS), therefore no hyperparameter needs to be specified.
- 2. Only positive triples are provided, and the negative triples are generated by corruptions just like it is done in training or evaluation. The optimization is done using a first-order method (ADAM), therefore batches_count and epochs must be specified.

Calibration is highly dependent on the base rate of positive triples. Therefore, for mode (2) of operation, the user is required to provide the positive_base_rate argument. For mode (1), that can be inferred automatically by the relative sizes of the positive and negative sets, but the user can override that by providing a value to positive_base_rate.

Defining the positive base rate is the biggest challenge when calibrating without negatives. That depends on the user choice of which triples will be evaluated during test time. Let's take WN11 as an example: it has around 50% positives triples on both the validation set and test set, so naturally the positive base rate is 50%. However, should the user resample it to have 75% positives and 25% negatives, its previous calibration will be degraded. The user must recalibrate the model now with a 75% positive base rate. Therefore, this parameter depends on how the user handles the dataset and cannot be determined automatically or a priori.

Note: Incompatible with large graph mode (i.e. if self.dealing_with_large_graphs=True).

Note: Experiments for the ICLR-21 calibration paper are available here [TC20].

Parameters

- **X_pos** (*ndarray* (*shape* [*n*, 3])) Numpy array of positive triples.
- X_neg (ndarray (shape [n, 3])) Numpy array of negative triples.

If *None*, the negative triples are generated via corruptions and the user must provide a positive base rate instead.

• **positive_base_rate** (*float*) – Base rate of positive statements.

For example, if we assume there is a fifty-fifty chance of any query to be true, the base rate would be 50%.

If X_neg is provided and this is *None*, the relative sizes of X_pos and X_neg will be used to determine the base rate. For example, if we have 50 positive triples and 200 negative triples, the positive base rate will be assumed to be 50/(50+200) = 1/5 = 0.2.

This must be a value between 0 and 1.

- **batches_count** (*int*) Number of batches to complete one epoch of the Platt scaling training. Only applies when X_neg is *None*.
- **epochs** (*int*) Number of epochs used to train the Platt scaling model. Only applies when X_neg is *None*.

Examples

```
>>> import numpy as np
>>> from sklearn.metrics import brier_score_loss, log_loss
>>> from scipy.special import expit
>>>
>>> from ampligraph.datasets import load_wn11
>>> from ampligraph.latent_features.models import TransE
>>>
>>> X = load_wn11()
>>> X_valid_pos = X['valid'][X['valid_labels']]
>>> X_valid_neg = X['valid'][~X['valid_labels']]
>>>
>>> model = TransE(batches_count=64, seed=0, epochs=500, k=100, eta=20,
                   optimizer='adam', optimizer_params={'lr':0.0001},
>>>
>>>
                   loss='pairwise', verbose=True)
>>>
>>> model.fit(X['train'])
>>>
>>> # Raw scores
>>> scores = model.predict(X['test'])
>>>
>>> # Calibrate with positives and negatives
>>> model.calibrate(X_valid_pos, X_valid_neg, positive_base_rate=None)
>>> probas_pos_neg = model.predict_proba(X['test'])
>>>
>>> # Calibrate with just positives and base rate of 50%
>>> model.calibrate(X_valid_pos, positive_base_rate=0.5)
```

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$predict_proba(X)$

Predicts probabilities using the Platt scaling model (after calibration).

Model must be calibrated beforehand with the calibrate method.

Parameters X (*ndarray* (*shape* [*n*, 3])) – Numpy array of triples to be evaluated.

Returns probas – Probability of each triple to be true according to the Platt scaling calibration.

Return type ndarray (shape [n])

HolE

class ampligraph.latent features.**HolE** (*k=100*, epochs=100. eta=2. seed=0. batches_count=100, embedding_model_params={'corrupt_sides': ['s, o'],'negative_corruption_entities': 'all'}, optimizer='adam', optimizer_params={'lr': $0.0005\},$ loss='nll', loss_params={}, regularizer=None, regularizer_params={}, initializer='xavier', initial*izer_params={'uniform': False}, verbose=False)*

Holographic Embeddings

The HolE model [NRP+16] as re-defined by Hayashi et al. [HS17]:

$$f_{HolE} = \frac{2}{n} f_{ComplEx}$$

Examples

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```
>>> ['c', 'y', 'a'],
>>> ['a', 'y', 'd'],
>>> ['c', 'y', 'd'],
>>> ['b', 'y', 'c'],
>>> ['f', 'y', 'e']])
>>> model.fit(X)
>>> model.predict(np.array([['f', 'y', 'e'], ['b', 'y', 'd']]))
[[0.009254738], [0.00023370088]]
```

Methods

	Initialize an EmbeddingModel
<pre>fit(X[, early_stopping,])</pre>	Train a HolE model.
<pre>get_embeddings(entities[, embedding_type])</pre>	Get the embeddings of entities or relations.
<pre>get_hyperparameter_dict()</pre>	Returns hyperparameters of the model.
<pre>predict(X[, from_idx])</pre>	Predict the scores of triples using a trained embed-
	ding model.
<pre>calibrate(X_pos[, X_neg,])</pre>	Calibrate predictions
predict_proba(X)	Predicts probabilities using the Platt scaling model
	(after calibration).

__init___(k=100, eta=2, epochs=100, batches_count=100, seed=0, embedding_model_params={'corrupt_sides': ['s,o'], 'negative_corruption_entities': 'all'}, optimizer='adam', optimizer_params={'lr': 0.0005}, loss='nll', loss_params={}, regularizer=None, regularizer_params={}, initializer='xavier', initializer_params={'uniform': False}, verbose=False)

Initialize an EmbeddingModel

Also creates a new Tensorflow session for training.

Parameters

- **k** (*int*) Embedding space dimensionality
- **eta** (*int*) The number of negatives that must be generated at runtime during training for each positive.
- **epochs** (*int*) The iterations of the training loop.
- **batches_count** (*int*) The number of batches in which the training set must be split during the training loop.
- **seed** (*int*) The seed used by the internal random numbers generator.
- **embedding_model_params** (*dict*) HolE-specific hyperparams:
 - negative_corruption_entities Entities to be used for generation of corruptions while training. It can take the following values : all (default: all entities), batch (entities present in each batch), list of entities or an int (which indicates how many entities that should be used for corruption generation).
 - corrupt_sides : Specifies how to generate corruptions for training. Takes values s, o, s+o or any combination passed as a list.
 - 'non_linearity': can be one of the following values linear, softplus, sigmoid, tanh

 - 'stop_epoch': specifies how long to decay (linearly) the numeric values from 1 to original value

until it reachs original value. - **'structural_wt'**: structural influence hyperparameter [0, 1] that modulates the influence of graph topology. - **'normalize_numeric_values'**: normalize the numeric values, such that they are scaled between [0, 1]

The last 4 parameters are related to FocusE layers.

- **optimizer** (*string*) The optimizer used to minimize the loss function. Choose between 'sgd', 'adagrad', 'adam', 'momentum'.
- **optimizer_params** (*dict*) Arguments specific to the optimizer, passed as a dictionary.

Supported keys:

- 'lr' (float): learning rate (used by all the optimizers). Default: 0.1.
- 'momentum' (float): learning momentum (only used when optimizer=momentum). Default: 0.9.

Example: optimizer_params={ 'lr': 0.01}

- **loss** (*string*) The type of loss function to use during training.
 - pairwise the model will use pairwise margin-based loss function.
 - nll the model will use negative loss likelihood.
 - absolute_margin the model will use absolute margin likelihood.
 - self_adversarial the model will use adversarial sampling loss function.
 - multiclass_nll the model will use multiclass nll loss. Switch to multiclass loss defined in [aC15] by passing 'corrupt_sides' as ['s','o'] to embedding_model_params. To use loss defined in [KBK17] pass 'corrupt_sides' as 'o' to embedding_model_params.
- **loss_params** (*dict*) Dictionary of loss-specific hyperparameters. See *loss functions* documentation for additional details.

Example: optimizer_params={'lr': 0.01} if loss='pairwise'.

- **regularizer** (*string*) The regularization strategy to use with the loss function.
 - None: the model will not use any regularizer (default)
 - 'LP': the model will use L1, L2 or L3 based on the value of regularizer_params['p'] (see below).
- **regularizer_params** (*dict*) Dictionary of regularizer-specific hyperparameters. See the *regularizers* documentation for additional details.

```
Example: regularizer_params={'lambda': 1e-5, 'p': 2} if
regularizer='LP'.
```

- **initializer** (*string*) The type of initializer to use.
 - normal: The embeddings will be initialized from a normal distribution
 - uniform: The embeddings will be initialized from a uniform distribution
 - xavier: The embeddings will be initialized using xavier strategy (default)

• **initializer_params** (*dict*) – Dictionary of initializer-specific hyperparameters. See the *initializer* documentation for additional details.

```
Example: initializer_params={'mean': 0, 'std': 0.001} if
initializer='normal'.
```

- **verbose** (bool) Verbose mode.
- fit (X, early_stopping=False, early_stopping_params={}, focusE_numeric_edge_values=None, tensorboard_logs_path=None) Train a HolE model.

The model is trained on a training set X using the training protocol described in [NRP+16].

Parameters

- **X** (*ndarray*, *shape* [*n*, 3]) The training triples
- **early_stopping** (bool) Flag to enable early stopping (default:False).

If set to True, the training loop adopts the following early stopping heuristic:

- The model will be trained regardless of early stopping for burn_in epochs.
- Every check_interval epochs the method will compute the metric specified in criteria.

If such metric decreases for stop_interval checks, we stop training early.

Note the metric is computed on x_valid. This is usually a validation set that you held out.

Also, because criteria is a ranking metric, it requires generating negatives. Entities used to generate corruptions can be specified, as long as the side(s) of a triple to corrupt. The method supports filtered metrics, by passing an array of positives to x_filter . This will be used to filter the negatives generated on the fly (i.e. the corruptions).

Note: Keep in mind the early stopping criteria may introduce a certain overhead (caused by the metric computation). The goal is to strike a good trade-off between such overhead and saving training epochs.

A common approach is to use MRR unfiltered:

early_stopping_params={x_valid=X['valid'], 'criteria': 'mrr'}

Note the size of validation set also contributes to such overhead. In most cases a smaller validation set would be enough.

• **early_stopping_params** (*dictionary*) – Dictionary of hyperparameters for the early stopping heuristics.

The following string keys are supported:

- 'x_valid': ndarray, shape [n, 3] : Validation set to be used for early stopping.
- 'criteria': string : criteria for early stopping 'hits10', 'hits3', 'hits1' or 'mrr'(default).
- 'x_filter': ndarray, shape [n, 3] : Positive triples to use as filter if a 'filtered' early stopping criteria is desired (i.e. filtered-MRR if 'criteria':'mrr'). Note this will affect training time (no filter by default).

- 'burn_in': int : Number of epochs to pass before kicking in early stopping (default: 100).
- check_interval': int : Early stopping interval after burn-in (default:10).
- 'stop_interval': int : Stop if criteria is performing worse over n consecutive checks (default: 3)
- 'corruption_entities': List of entities to be used for corruptions. If 'all', it uses all entities (default: 'all')
- 'corrupt_side': Specifies which side to corrupt. 's', 'o', 's+o' (default)

Example: early_stopping_params={x_valid=X['valid'],
'criteria': 'mrr'}

- **focusE_numeric_edge_values** (*nd array (n, 1)*)-Numeric values associated with links. Semantically, the numeric value can signify importance, uncertainity, significance, confidence, etc. If the numeric value is unknown pass a NaN weight. The model will uniformly randomly assign a numeric value. One can also think about assigning numeric values by looking at the distribution of it per predicate.
- tensorboard_logs_path (*str or None*) Path to store tensorboard logs, e.g. average training loss tracking per epoch (default: None indicating no logs will be collected). When provided it will create a folder under provided path and save tensorboard files there. To then view the loss in the terminal run: tensorboard --logdir <tensorboard_logs_path>.
- **get_embeddings** (*entities*, *embedding_type='entity'*) Get the embeddings of entities or relations.

Note: Use *ampligraph.utils.create_tensorboard_visualizations()* to visualize the embeddings with TensorBoard.

Parameters

- **entities** (*array-like*, *dtype=int*, *shape=[n]*) The entities (or relations) of interest. Element of the vector must be the original string literals, and not internal IDs.
- **embedding_type** (*string*) If 'entity', entities argument will be considered as a list of knowledge graph entities (i.e. nodes). If set to 'relation', they will be treated as relation types instead (i.e. predicates).

Returns embeddings - An array of k-dimensional embeddings.

Return type ndarray, shape [n, k]

```
get_hyperparameter_dict()
```

Returns hyperparameters of the model.

Returns hyperparam_dict – Dictionary of hyperparameters that were used for training.

Return type dict

predict (X, from_idx=False)

Predict the scores of triples using a trained embedding model. The function returns raw scores generated by the model.

Note: To obtain probability estimates, calibrate the model with *calibrate()*, then call *predict_proba()*.

Parameters

- X (ndarray, shape [n, 3]) The triples to score.
- from_idx (bool) If True, will skip conversion to internal IDs. (default: False).

Returns scores_predict – The predicted scores for input triples X.

Return type ndarray, shape [n]

calibrate (*X_pos*, *X_neg=None*, *positive_base_rate=None*, *batches_count=100*, *epochs=50*) Calibrate predictions

The method implements the heuristics described in [TC20], using Platt scaling [P+99].

The calibrated predictions can be obtained with *predict_proba()* after calibration is done.

Ideally, calibration should be performed on a validation set that was not used to train the embeddings.

There are two modes of operation, depending on the availability of negative triples:

- 1. Both positive and negative triples are provided via X_pos and X_neg respectively. The optimization is done using a second-order method (limited-memory BFGS), therefore no hyperparameter needs to be specified.
- 2. Only positive triples are provided, and the negative triples are generated by corruptions just like it is done in training or evaluation. The optimization is done using a first-order method (ADAM), therefore batches_count and epochs must be specified.

Calibration is highly dependent on the base rate of positive triples. Therefore, for mode (2) of operation, the user is required to provide the positive_base_rate argument. For mode (1), that can be inferred automatically by the relative sizes of the positive and negative sets, but the user can override that by providing a value to positive_base_rate.

Defining the positive base rate is the biggest challenge when calibrating without negatives. That depends on the user choice of which triples will be evaluated during test time. Let's take WN11 as an example: it has around 50% positives triples on both the validation set and test set, so naturally the positive base rate is 50%. However, should the user resample it to have 75% positives and 25% negatives, its previous calibration will be degraded. The user must recalibrate the model now with a 75% positive base rate. Therefore, this parameter depends on how the user handles the dataset and cannot be determined automatically or a priori.

Note: Incompatible with large graph mode (i.e. if self.dealing_with_large_graphs=True).

Note: Experiments for the ICLR-21 calibration paper are available here [TC20].

Parameters

- **X_pos** (*ndarray* (*shape* [*n*, 3])) Numpy array of positive triples.
- **X_neg** (*ndarray* (*shape* [*n*, 3])) Numpy array of negative triples.

If *None*, the negative triples are generated via corruptions and the user must provide a positive base rate instead.

• **positive_base_rate** (*float*) – Base rate of positive statements.

For example, if we assume there is a fifty-fifty chance of any query to be true, the base rate would be 50%.

If X_neg is provided and this is *None*, the relative sizes of X_pos and X_neg will be used to determine the base rate. For example, if we have 50 positive triples and 200 negative triples, the positive base rate will be assumed to be 50/(50+200) = 1/5 = 0.2.

This must be a value between 0 and 1.

- **batches_count** (*int*) Number of batches to complete one epoch of the Platt scaling training. Only applies when X_neg is *None*.
- **epochs** (*int*) Number of epochs used to train the Platt scaling model. Only applies when X_neg is *None*.

Examples

```
>>> import numpy as np
>>> from sklearn.metrics import brier score loss, log loss
>>> from scipy.special import expit
>>>
>>> from ampligraph.datasets import load_wn11
>>> from ampligraph.latent_features.models import TransE
>>>
>>> X = load wn11()
>>> X_valid_pos = X['valid'][X['valid_labels']]
>>> X_valid_neg = X['valid'][~X['valid_labels']]
>>>
>>> model = TransE(batches_count=64, seed=0, epochs=500, k=100, eta=20,
                   optimizer='adam', optimizer_params={'lr':0.0001},
>>>
                   loss='pairwise', verbose=True)
>>>
>>>
>>> model.fit(X['train'])
>>>
>>> # Raw scores
>>> scores = model.predict(X['test'])
>>>
>>> # Calibrate with positives and negatives
>>> model.calibrate(X_valid_pos, X_valid_neq, positive_base_rate=None)
>>> probas_pos_neg = model.predict_proba(X['test'])
>>>
>>> # Calibrate with just positives and base rate of 50%
>>> model.calibrate(X_valid_pos, positive_base_rate=0.5)
>>> probas_pos = model.predict_proba(X['test'])
>>>
>>> # Calibration evaluation with the Brier score loss (the smaller, the
\rightarrow better)
>>> print("Brier scores")
>>> print("Raw scores:", brier_score_loss(X['test_labels'], expit(scores)))
>>> print("Positive and negative calibration:", brier_score_loss(X['test_
→labels'], probas pos neg))
>>> print("Positive only calibration:", brier_score_loss(X['test_labels'],...

→probas_pos))
```

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```
Brier scores
Raw scores: 0.4925058891371126
Positive and negative calibration: 0.20434617882733366
Positive only calibration: 0.22597599585144656
```

$predict_proba(X)$

Predicts probabilities using the Platt scaling model (after calibration).

Model must be calibrated beforehand with the calibrate method.

Parameters X (*ndarray* (*shape* [*n*, 3])) – Numpy array of triples to be evaluated.

Returns probas – Probability of each triple to be true according to the Platt scaling calibration.

Return type ndarray (shape [n])

ConvE

```
class ampligraph.latent_features.ConvE(k=100, eta=2, epochs=100, batches_count=100,
                                                   seed=0, embedding_model_params={'conv_filters':
                                                    32, 'conv_kernel_size': 3, 'dropout_conv': 0.3,
                                                    'dropout_dense':
                                                                     0.2, 'dropout_embed':
                                                                                               0.2,
                                                    'use_batchnorm': True, 'use_bias': True}, opti-
                                                   mizer='adam', optimizer_params={'lr': 0.0005},
                                                   loss='bce',
                                                                    loss_params={'label_smoothing':
                                                   0.1.
                                                           'label_weighting':
                                                                                 False},
                                                                                           regular-
                                                   izer=None,
                                                                  regularizer_params={},
                                                                                            initial-
                                                   izer='xavier',
                                                                      initializer_params={'uniform':
                                                    False}, low_memory=False, verbose=False)
     Convolutional 2D KG Embeddings
```

The ConvE model [DMSR18].

ConvE uses convolutional layers. g is a non-linear activation function, * is the linear convolution operator, vec indicates 2D reshaping.

 $f_{ConvE} = \langle \sigma \left(vec \left(g \left(\left[\overline{\mathbf{e}_s}; \overline{\mathbf{r}_p} \right] * \Omega \right) \right) \mathbf{W} \right) \right) \mathbf{e}_o \rangle$

Note: ConvE does not handle 's+o' corruptions currently, nor large_graph mode.

Examples

```
>>> import numpy as np
>>> from ampligraph.latent_features import ConvE
>>> model = ConvE(batches_count=1, seed=22, epochs=5, k=100)
>>>
X = np.array([['a', 'y', 'b'],
>>> ['b', 'y', 'a'],
>>> ['a', 'y', 'c'],
>>> ['c', 'y', 'a'],
>>> ['a', 'y', 'd'],
```

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```
>>> ['c', 'y', 'd'],
>>> ['b', 'y', 'c'],
>>> ['f', 'y', 'e']])
>>> model.fit(X)
>>> model.predict(np.array([['f', 'y', 'e'], ['b', 'y', 'd']]))
[0.42921206 0.38998795]
```

Attributes

	Initialize a ConvE model
<pre>fit(X[, early_stopping, early_stopping_params])</pre>	Train a ConvE (with optional early stopping).
<pre>get_embeddings(entities[, embedding_type])</pre>	Get the embeddings of entities or relations.
get_hyperparameter_dict()	Returns hyperparameters of the model.
<pre>predict(X[, from_idx])</pre>	Predict the scores of triples using a trained embed-
	ding model.
<pre>calibrate(X_pos[, X_neg,])</pre>	Calibrate predictions
predict_proba(X)	Predicts probabilities using the Platt scaling model
	(after calibration).

___init___(*k=100*, eta=2, epochs=100, *batches_count=100*, seed=0, embed*ding_model_params={'conv_filters':* 32, 'conv_kernel_size': 3. 'dropout conv': 0.3, 'dropout_dense': 0.2, 'dropout_embed': 0.2, 'use_batchnorm': True, 'use_bias': True}, optimizer='adam', optimizer_params={'lr': 0.0005}, loss='bce', loss_params={'label_smoothing': 0.1, 'label_weighting': False}, regularizer=None, regularizer params={}, initializer='xavier', initializer params={'uniform': False}. *low memory=False*, *verbose=False*)

Initialize a ConvE model

Also creates a new Tensorflow session for training.

Parameters

- **k** (*int*) Embedding space dimensionality.
- **eta** (*int*) The number of negatives that must be generated at runtime during training for each positive. Note: This parameter is not used in ConvE.
- **epochs** (*int*) The iterations of the training loop.
- **batches_count** (*int*) The number of batches in which the training set must be split during the training loop.
- **seed** (*int*) The seed used by the internal random numbers generator.
- **embedding_model_params** (*dict*) ConvE-specific hyperparams:
 - conv_filters (int): Number of convolution feature maps. Default: 32
 - conv_kernel_size (int): Convolution kernel size. Default: 3
 - dropout_embed (floatlNone): Dropout on the embedding layer. Default: 0.2
 - dropout_conv (float|None): Dropout on the convolution maps. Default: 0.3
 - dropout_dense (floatlNone): Dropout on the dense layer. Default: 0.2
 - use_bias (bool): Use bias layer. Default: True

- use_batchnorm (bool): Use batch normalization after input, convolution, dense layers. Default: True
- **optimizer** (*string*) The optimizer used to minimize the loss function. Choose between 'sgd', 'adagrad', 'adam', 'momentum'.
- **optimizer_params** (*dict*) Arguments specific to the optimizer, passed as a dictionary.

Supported keys:

- 'lr' (float): learning rate (used by all the optimizers). Default: 0.1.
- 'momentum' (float): learning momentum (only used when optimizer=momentum). Default: 0.9.

Example: optimizer_params={'lr': 0.01}

- **loss** (*string*) The type of loss function to use during training.
 - bce the model will use binary cross entropy loss function.
- **loss_params** (*dict*) Dictionary of loss-specific hyperparameters. See *loss functions* documentation for additional details.

Supported keys:

- 'lr' (float): learning rate (used by all the optimizers). Default: 0.1.
- 'momentum' (float): learning momentum (only used when optimizer=momentum). Default: 0.9.
- 'label_smoothing' (float): applies label smoothing to one-hot outputs. Default: 0.1.
- 'label_weighting' (bool): applies label weighting to one-hot outputs. Default: True

Example: optimizer_params={'lr': 0.01, 'label_smoothing':
0.1}

- **regularizer** (*string*) The regularization strategy to use with the loss function.
 - None: the model will not use any regularizer (default)
 - LP: the model will use L1, L2 or L3 based on the value of regularizer_params['p'] (see below).
- **regularizer_params** (*dict*) Dictionary of regularizer-specific hyperparameters. See the *regularizers* documentation for additional details.

```
Example: regularizer_params={'lambda': 1e-5, 'p': 2} if
regularizer='LP'.
```

- **initializer** (*string*) The type of initializer to use.
 - normal: The embeddings will be initialized from a normal distribution
 - uniform: The embeddings will be initialized from a uniform distribution
 - xavier: The embeddings will be initialized using xavier strategy (default)
- **initializer_params** (*dict*) Dictionary of initializer-specific hyperparameters. See the *initializer* documentation for additional details.

Example: initializer_params={'mean': 0, 'std': 0.001} if initializer='normal'.

- **verbose** (bool) Verbose mode.
- **low_memory** (*bool*) Train ConvE with a (slower) low_memory option. If MemoryError is still encountered, try raising the batches_count value. Default: False.

```
fit (X, early_stopping=False, early_stopping_params={})
```

Train a ConvE (with optional early stopping).

The model is trained on a training set X using the training protocol described in [DMSR18].

Parameters

- X (ndarray (shape [n, 3]) or object of ConvEDatasetAdapter) Numpy array of training triples OR handle of Dataset adapter which would help retrieve data.
- **early_stopping** (bool) Flag to enable early stopping (default:False)
- **early_stopping_params** (*dictionary*) Dictionary of hyperparameters for the early stopping heuristics.

The following string keys are supported:

- 'x_valid': ndarray (shape [n, 3]) or object of AmpligraphDatasetAdapter : Numpy array of validation triples OR handle of Dataset adapter which would help retrieve data.
- 'criteria': string : criteria for early stopping 'hits10', 'hits3', 'hits1' or 'mrr'(default).
- 'x_filter': ndarray, shape [n, 3] [Positive triples to use as filter if a 'filtered' early] stopping criteria is desired (i.e. filtered-MRR if 'criteria':'mrr'). Note this will affect training time (no filter by default). If the filter has already been set in the adapter, pass True
- 'burn_in': int : Number of epochs to pass before kicking in early stopping (default: 100).
- check_interval': int : Early stopping interval after burn-in (default:10).
- 'stop_interval': int : Stop if criteria is performing worse over n consecutive checks (default: 3)
- 'corruption_entities': List of entities to be used for corruptions. If 'all', it uses all entities (default: 'all')
- 'corrupt_side': Specifies which side to corrupt. 'o' (default). Note: ConvE does not currently support subject corruptions in early stopping.

```
Example: early_stopping_params={x_valid=X['valid'],
'criteria': 'mrr'}
```

get_embeddings (*entities*, *embedding_type='entity'*) Get the embeddings of entities or relations.

Note: Use *ampligraph.utils.create_tensorboard_visualizations()* to visualize the embeddings with TensorBoard.

Parameters

- **entities** (*array-like*, *dtype=int*, *shape=[n]*) The entities (or relations) of interest. Element of the vector must be the original string literals, and not internal IDs.
- **embedding_type** (*string*) If 'entity', entities argument will be considered as a list of knowledge graph entities (i.e. nodes). If set to 'relation', they will be treated as relation types instead (i.e. predicates).

Returns embeddings – An array of k-dimensional embeddings.

Return type ndarray, shape [n, k]

get_hyperparameter_dict()

Returns hyperparameters of the model.

Returns hyperparam_dict – Dictionary of hyperparameters that were used for training.

Return type dict

predict (X, from_idx=False)

Predict the scores of triples using a trained embedding model. The function returns raw scores generated by the model.

Note: To obtain probability estimates, calibrate the model with *calibrate()*, then call *predict_proba()*.

Parameters

- X (ndarray, shape [n, 3]) The triples to score.
- **from_idx** (bool) If True, will skip conversion to internal IDs. (default: False).

Returns scores_predict – The predicted scores for input triples X.

Return type ndarray, shape [n]

calibrate (*X_pos*, *X_neg=None*, *positive_base_rate=None*, *batches_count=100*, *epochs=50*) Calibrate predictions

The method implements the heuristics described in [TC20], using Platt scaling [P+99].

The calibrated predictions can be obtained with *predict_proba()* after calibration is done.

Ideally, calibration should be performed on a validation set that was not used to train the embeddings.

There are two modes of operation, depending on the availability of negative triples:

- 1. Both positive and negative triples are provided via X_pos and X_neg respectively. The optimization is done using a second-order method (limited-memory BFGS), therefore no hyperparameter needs to be specified.
- 2. Only positive triples are provided, and the negative triples are generated by corruptions just like it is done in training or evaluation. The optimization is done using a first-order method (ADAM), therefore batches_count and epochs must be specified.

Calibration is highly dependent on the base rate of positive triples. Therefore, for mode (2) of operation, the user is required to provide the positive_base_rate argument. For mode (1), that can be inferred automatically by the relative sizes of the positive and negative sets, but the user can override that by providing a value to positive_base_rate.

Defining the positive base rate is the biggest challenge when calibrating without negatives. That depends on the user choice of which triples will be evaluated during test time. Let's take WN11 as an example: it has around 50% positives triples on both the validation set and test set, so naturally the positive base rate is 50%. However, should the user resample it to have 75% positives and 25% negatives, its previous calibration will be degraded. The user must recalibrate the model now with a 75% positive base rate. Therefore, this parameter depends on how the user handles the dataset and cannot be determined automatically or a priori.

Note: Incompatible with large graph mode (i.e. if self.dealing_with_large_graphs=True).

Note: Experiments for the ICLR-21 calibration paper are available here [TC20].

Parameters

- X_pos (ndarray (shape [n, 3])) Numpy array of positive triples.
- X_neg (ndarray (shape [n, 3])) Numpy array of negative triples.

If *None*, the negative triples are generated via corruptions and the user must provide a positive base rate instead.

• **positive_base_rate** (*float*) – Base rate of positive statements.

For example, if we assume there is a fifty-fifty chance of any query to be true, the base rate would be 50%.

If X_neg is provided and this is *None*, the relative sizes of X_pos and X_neg will be used to determine the base rate. For example, if we have 50 positive triples and 200 negative triples, the positive base rate will be assumed to be 50/(50+200) = 1/5 = 0.2.

This must be a value between 0 and 1.

- **batches_count** (*int*) Number of batches to complete one epoch of the Platt scaling training. Only applies when X_neg is *None*.
- **epochs** (*int*) Number of epochs used to train the Platt scaling model. Only applies when X_neg is *None*.

Examples

```
>>> import numpy as np
>>> from sklearn.metrics import brier_score_loss, log_loss
>>> from scipy.special import expit
>>>
>>> from ampligraph.datasets import load_wn11
>>> from ampligraph.latent_features.models import TransE
>>>
>>> X = load_wn11()
>>> X_valid_pos = X['valid'][X['valid_labels']]
>>> X_valid_neg = X['valid'][~X['valid_labels']]
>>>
>>> model = TransE(batches_count=64, seed=0, epochs=500, k=100, eta=20,
                   optimizer='adam', optimizer_params={'lr':0.0001},
>>>
                   loss='pairwise', verbose=True)
>>>
```

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```
>>>
>>> model.fit(X['train'])
>>>
>>> # Raw scores
>>> scores = model.predict(X['test'])
>>>
>>> # Calibrate with positives and negatives
>>> model.calibrate(X_valid_pos, X_valid_neg, positive_base_rate=None)
>>> probas_pos_neg = model.predict_proba(X['test'])
>>>
>>> # Calibrate with just positives and base rate of 50%
>>> model.calibrate(X_valid_pos, positive_base_rate=0.5)
>>> probas_pos = model.predict_proba(X['test'])
>>>
>>> # Calibration evaluation with the Brier score loss (the smaller, the
\rightarrow better)
>>> print("Brier scores")
>>> print("Raw scores:", brier_score_loss(X['test_labels'], expit(scores)))
>>> print("Positive and negative calibration:", brier_score_loss(X['test_
→labels'], probas_pos_neg))
>>> print("Positive only calibration:", brier_score_loss(X['test_labels'],_
\rightarrow probas_pos))
Brier scores
Raw scores: 0.4925058891371126
Positive and negative calibration: 0.20434617882733366
Positive only calibration: 0.22597599585144656
```

$predict_proba(X)$

Predicts probabilities using the Platt scaling model (after calibration).

Model must be calibrated beforehand with the calibrate method.

Parameters X (*ndarray* (*shape* [*n*, 3])) – Numpy array of triples to be evaluated.

Returns probas – Probability of each triple to be true according to the Platt scaling calibration.

Return type ndarray (shape [n])

ConvKB

Convolution-based model

The ConvKB model [NNNP18]:

```
f_{ConvKB} = concat \left( g \left( \left[ \mathbf{e}_s, \mathbf{r}_p, \mathbf{e}_o \right] \right) * \Omega \right) \right) \cdot W
```

where g is a non-linear function, * is the convolution operator, \cdot is the dot product, *concat* is the concatenation operator and Ω is a set of filters.

Note: The evaluation protocol implemented in *ampligraph.evaluation. evaluate_performance()* assigns the worst rank to a positive test triple in case of a tie with negatives. This is the agreed upon behaviour in literature. The original ConvKB implementation [NNNP18] assigns instead the top rank, hence leading to results which are not directly comparable with literature . We report results obtained with the agreed-upon protocol (tie=worst rank). Note that under these conditions the model *does not reach the state-of-the-art results claimed in the original paper.*

Examples

```
>>> from ampligraph.latent_features import ConvKB
>>> from ampligraph.datasets import load_wn18
>>> model = ConvKB(batches_count=2, seed=22, epochs=1, k=10, eta=1,
                  embedding_model_params={'num_filters': 32, 'filter_sizes': [1],
>>>
>>>
                                           'dropout': 0.1},
>>>
                  optimizer='adam', optimizer_params={'lr': 0.001},
>>>
                  loss='pairwise', loss_params={}, verbose=True)
>>>
>>> X = load_wn18()
>>>
>>> model.fit(X['train'])
>>>
>>> print(model.predict(X['test'][:5]))
[[0.2803744], [0.0866661], [0.012815937], [-0.004235901], [-0.010947697]]
```

Methods

init([k, eta, epochs, batches_count,])	Initialize an EmbeddingModel
<pre>fit(X[, early_stopping,])</pre>	Train a ConvKB model (with optional early stop-
	ping).
<pre>get_embeddings(entities[, embedding_type])</pre>	Get the embeddings of entities or relations.
<pre>get_hyperparameter_dict()</pre>	Returns hyperparameters of the model.
<pre>predict(X[, from_idx])</pre>	Predict the scores of triples using a trained embed-
	ding model.
<pre>calibrate(X_pos[, X_neg,])</pre>	Calibrate predictions
predict_proba(X)	Predicts probabilities using the Platt scaling model
	(after calibration).

__init___(k=100, eta=2, epochs=100, batches_count=100, seed=0, embedding_model_params={'dropout': 0.1, 'filter_sizes': [1], 'num_filters': 32}, optimizer='adam', optimizer_params={'lr': 0.0005}, loss='nll', loss_params={}, regularizer=None, regularizer_params={}, initializer='xavier', initializer_params={'uniform': False}, large_graphs=False, verbose=False) Initialize an EmbeddingModel

Parameters

- **k** (*int*) Embedding space dimensionality.
- **eta** (*int*) The number of negatives that must be generated at runtime during training for each positive.
- **epochs** (*int*) The iterations of the training loop.

- **batches_count** (*int*) The number of batches in which the training set must be split during the training loop.
- **seed** (*int*) The seed used by the internal random numbers generator.
- embedding_model_params (dict) ConvKB-specific hyperparams: num_filters - Number of feature maps per convolution kernel. Default: 32 - filter_sizes - List of convolution kernel sizes. Default: [1] - dropout - Dropout on the embedding layer. Default: 0.0 - 'non_linearity': can be one of the following values linear, softplus, sigmoid, tanh - 'stop_epoch': specifies how long to decay (linearly) the numeric values from 1 to original value until it reachs original value. - 'structural_wt': structural influence hyperparameter [0, 1] that modulates the influence of graph topology. - 'normalize_numeric_values': normalize the numeric values, such that they are scaled between [0, 1]

The last 4 parameters are related to FocusE layers.

- **optimizer** (*string*) The optimizer used to minimize the loss function. Choose between 'sgd', 'adagrad', 'adam', 'momentum'.
- **optimizer_params** (*dict*) Arguments specific to the optimizer, passed as a dictionary.

Supported keys:

- 'lr' (float): learning rate (used by all the optimizers). Default: 0.1.
- 'momentum' (float): learning momentum (only used when optimizer=momentum). Default: 0.9.

Example: optimizer_params={'lr': 0.01}

- **loss** (*string*) The type of loss function to use during training.
- **loss_params** (*dict*) Dictionary of loss-specific hyperparameters. See *loss functions* documentation for additional details.

Supported keys:

- 'lr' (float): learning rate (used by all the optimizers). Default: 0.1.
- 'momentum' (float): learning momentum (only used when optimizer=momentum). Default: 0.9.

Example: optimizer_params={'lr': 0.01,
'label_smoothing': 0.1}

- **regularizer** (*string*) The regularization strategy to use with the loss function.
 - None: the model will not use any regularizer (default)
 - LP: the model will use L1, L2 or L3 based on the value of regularizer_params['p'] (see below).
- **regularizer_params** (*dict*) Dictionary of regularizer-specific hyperparameters. See the *regularizers* documentation for additional details.

Example: regularizer_params={'lambda': 1e-5, 'p': 2} if
regularizer='LP'.

- **initializer** (*string*) The type of initializer to use.
 - normal: The embeddings will be initialized from a normal distribution

- uniform: The embeddings will be initialized from a uniform distribution
- xavier: The embeddings will be initialized using xavier strategy (default)
- **initializer_params** (*dict*) Dictionary of initializer-specific hyperparameters. See the *initializer* documentation for additional details.

Example: initializer_params={'mean': 0, 'std': 0.001} if initializer='normal'.

- **large_graphs** (*bool*) Avoid loading entire dataset onto GPU when dealing with large graphs.
- **verbose** (bool) Verbose mode.
- fit (X, early_stopping=False, early_stopping_params={}, focusE_numeric_edge_values=None, tensorboard_logs_path=None) Train a ConvKB model (with optional early stopping).

The model is trained on a training set X using the training protocol described in [TWR+16].

Parameters

- **X**(*ndarray*, *shape* [*n*, 3]) The training triples
- **early_stopping** (*bool*) Flag to enable early stopping (default:False).

If set to True, the training loop adopts the following early stopping heuristic:

- The model will be trained regardless of early stopping for burn_in epochs.
- Every check_interval epochs the method will compute the metric specified in criteria.

If such metric decreases for stop_interval checks, we stop training early.

Note the metric is computed on x_valid. This is usually a validation set that you held out.

Also, because criteria is a ranking metric, it requires generating negatives. Entities used to generate corruptions can be specified, as long as the side(s) of a triple to corrupt. The method supports filtered metrics, by passing an array of positives to x_filter. This will be used to filter the negatives generated on the fly (i.e. the corruptions).

Note: Keep in mind the early stopping criteria may introduce a certain overhead (caused by the metric computation). The goal is to strike a good trade-off between such overhead and saving training epochs.

A common approach is to use MRR unfiltered:

```
early_stopping_params={x_valid=X['valid'], 'criteria':
'mrr'}
```

Note the size of validation set also contributes to such overhead. In most cases a smaller validation set would be enough.

• **early_stopping_params** (*dictionary*) – Dictionary of hyperparameters for the early stopping heuristics.

The following string keys are supported:

- 'x_valid': ndarray, shape [n, 3] : Validation set to be used for early stopping.
- 'criteria': string : criteria for early stopping 'hits10', 'hits3', 'hits1' or 'mrr'(default).
- 'x_filter': ndarray, shape [n, 3] : Positive triples to use as filter if a 'filtered' early stopping criteria is desired (i.e. filtered-MRR if 'criteria':'mrr'). Note this will affect training time (no filter by default).
- 'burn_in': int : Number of epochs to pass before kicking in early stopping (default: 100).
- check_interval': int : Early stopping interval after burn-in (default:10).
- 'stop_interval': int : Stop if criteria is performing worse over n consecutive checks (default: 3)
- 'corruption_entities': List of entities to be used for corruptions. If 'all', it uses all entities (default: 'all')
- 'corrupt_side': Specifies which side to corrupt. 's', 'o', 's+o' (default)

Example: early_stopping_params={x_valid=X['valid'],
'criteria': 'mrr'}

- **focusE_numeric_edge_values** (*nd array (n, 1)*) Numeric values associated with links. Semantically, the numeric value can signify importance, uncertainity, significance, confidence, etc. If the numeric value is unknown pass a NaN weight. The model will uniformly randomly assign a numeric value. One can also think about assigning numeric values by looking at the distribution of it per predicate.
- tensorboard_logs_path(*str or None*)-Path to store tensorboard logs, e.g. average training loss tracking per epoch (default: None indicating no logs will be collected). When provided it will create a folder under provided path and save tensorboard files there. To then view the loss in the terminal run: tensorboard --logdir <tensorboard_logs_path>.
- **get_embeddings** (*entities*, *embedding_type='entity'*) Get the embeddings of entities or relations.

Note: Use *ampligraph.utils.create_tensorboard_visualizations()* to visualize the embeddings with TensorBoard.

Parameters

- **entities** (*array-like*, *dtype=int*, *shape=[n]*) The entities (or relations) of interest. Element of the vector must be the original string literals, and not internal IDs.
- **embedding_type** (*string*) If 'entity', entities argument will be considered as a list of knowledge graph entities (i.e. nodes). If set to 'relation', they will be treated as relation types instead (i.e. predicates).

Returns embeddings – An array of k-dimensional embeddings.

Return type ndarray, shape [n, k]

get_hyperparameter_dict()

Returns hyperparameters of the model.

Returns hyperparam_dict – Dictionary of hyperparameters that were used for training.

Return type dict

predict (X, from_idx=False)

Predict the scores of triples using a trained embedding model. The function returns raw scores generated by the model.

Note: To obtain probability estimates, calibrate the model with *calibrate()*, then call *predict_proba()*.

Parameters

- X (ndarray, shape [n, 3]) The triples to score.
- from_idx (bool) If True, will skip conversion to internal IDs. (default: False).

Returns scores_predict – The predicted scores for input triples X.

Return type ndarray, shape [n]

calibrate (*X_pos*, *X_neg=None*, *positive_base_rate=None*, *batches_count=100*, *epochs=50*) Calibrate predictions

The method implements the heuristics described in [TC20], using Platt scaling [P+99].

The calibrated predictions can be obtained with *predict_proba()* after calibration is done.

Ideally, calibration should be performed on a validation set that was not used to train the embeddings.

There are two modes of operation, depending on the availability of negative triples:

- 1. Both positive and negative triples are provided via X_pos and X_neg respectively. The optimization is done using a second-order method (limited-memory BFGS), therefore no hyperparameter needs to be specified.
- 2. Only positive triples are provided, and the negative triples are generated by corruptions just like it is done in training or evaluation. The optimization is done using a first-order method (ADAM), therefore batches_count and epochs must be specified.

Calibration is highly dependent on the base rate of positive triples. Therefore, for mode (2) of operation, the user is required to provide the positive_base_rate argument. For mode (1), that can be inferred automatically by the relative sizes of the positive and negative sets, but the user can override that by providing a value to positive_base_rate.

Defining the positive base rate is the biggest challenge when calibrating without negatives. That depends on the user choice of which triples will be evaluated during test time. Let's take WN11 as an example: it has around 50% positives triples on both the validation set and test set, so naturally the positive base rate is 50%. However, should the user resample it to have 75% positives and 25% negatives, its previous calibration will be degraded. The user must recalibrate the model now with a 75% positive base rate. Therefore, this parameter depends on how the user handles the dataset and cannot be determined automatically or a priori.

Note: Incompatible with large graph mode (i.e. if self.dealing_with_large_graphs=True).

Note: Experiments for the ICLR-21 calibration paper are available here [TC20].

Parameters

- **X_pos** (*ndarray* (*shape* [*n*, 3])) Numpy array of positive triples.
- **X_neg** (*ndarray* (*shape* [*n*, 3])) Numpy array of negative triples.

If *None*, the negative triples are generated via corruptions and the user must provide a positive base rate instead.

• **positive_base_rate** (*float*) – Base rate of positive statements.

For example, if we assume there is a fifty-fifty chance of any query to be true, the base rate would be 50%.

If X_neg is provided and this is *None*, the relative sizes of X_pos and X_neg will be used to determine the base rate. For example, if we have 50 positive triples and 200 negative triples, the positive base rate will be assumed to be 50/(50+200) = 1/5 = 0.2.

This must be a value between 0 and 1.

- **batches_count** (*int*) Number of batches to complete one epoch of the Platt scaling training. Only applies when X_neg is *None*.
- **epochs** (*int*) Number of epochs used to train the Platt scaling model. Only applies when X_neg is *None*.

Examples

```
>>> import numpy as np
>>> from sklearn.metrics import brier_score_loss, log_loss
>>> from scipy.special import expit
>>>
>>> from ampligraph.datasets import load_wn11
>>> from ampligraph.latent_features.models import TransE
>>>
>>> X = load_wn11()
>>> X_valid_pos = X['valid'][X['valid_labels']]
>>> X_valid_neg = X['valid'][~X['valid_labels']]
>>>
>>> model = TransE(batches_count=64, seed=0, epochs=500, k=100, eta=20,
                   optimizer='adam', optimizer_params={'lr':0.0001},
>>>
>>>
                   loss='pairwise', verbose=True)
>>>
>>> model.fit(X['train'])
>>>
>>> # Raw scores
>>> scores = model.predict(X['test'])
>>>
>>> # Calibrate with positives and negatives
>>> model.calibrate(X_valid_pos, X_valid_neg, positive_base_rate=None)
>>> probas_pos_neg = model.predict_proba(X['test'])
>>>
>>> # Calibrate with just positives and base rate of 50%
>>> model.calibrate(X_valid_pos, positive_base_rate=0.5)
```

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$predict_proba(X)$

Predicts probabilities using the Platt scaling model (after calibration).

Model must be calibrated beforehand with the calibrate method.

Parameters X (*ndarray* (*shape* [*n*, 3])) – Numpy array of triples to be evaluated.

Returns probas – Probability of each triple to be true according to the Platt scaling calibration.

Return type ndarray (shape [n])

Anatomy of a Model

Knowledge graph embeddings are learned by training a neural architecture over a graph. Although such architectures vary, the training phase always consists in minimizing a *loss function* \mathcal{L} that includes a *scoring function* $f_m(t)$, i.e. a model-specific function that assigns a score to a triple t = (sub, pred, obj).

AmpliGraph models include the following components:

- Scoring function f(t)
- Loss function L
- Optimization algorithm
- Regularizer
- Initializer
- Negatives generation strategy
- Support for Numeric Literals on Edges

AmpliGraph comes with a number of such components. They can be used in any combination to come up with a model that performs sufficiently well for the dataset of choice.

AmpliGraph features a number of abstract classes that can be extended to design new models:

EmbeddingModel([k, eta, epochs,])	Abstract class for embedding models
Loss(eta, hyperparam_dict[, verbose])	Abstract class for loss function.
Regularizer(hyperparam_dict[, verbose])	Abstract class for Regularizer.
<pre>Initializer([initializer_params, verbose, seed])</pre>	Abstract class for initializer .

EmbeddingModel

class ampligraph.latent_features.EmbeddingMode	1 (<i>k</i> =100,	<i>eta</i> =2,	epochs=100,
	batches_cour	nt=100,	seed=0,
	embedding_n	nodel_params	s={},
	optimizer='a	dam',	opti-
	mizer_param	s={'lr':	0.0005},
	loss='nll',	loss_params	={}, regu-
	larizer=None	e, regularizer <u></u>	_params={},
	initializer='x	cavier',	initial-
	izer_params:	={'uniform':	False},
	large_graphs	s=False, verbo	ose=False)

Abstract class for embedding models

AmpliGraph neural knowledge graph embeddings models extend this class and its core methods.

Methods

fit(X[, early_stopping,])Train an EmbeddingModel (with optional early sping).get_embeddings(entities[, embedding_type])Get the embeddings of entities or relations.get_hyperparameter_dict()Returns hyperparameters of the model.predict(X[, from_idx])Predict the scores of triples using a trained emding model.calibrate(X_pos[, X_neg,])Calibrate predictionspredict_proba(X)Predicts probabilities using the Platt scaling model	bed-
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get_embedding_model_params(output_dict) Save the model parameters in the dictionary.	iza-
restore_model_params(in_dict) Load the model parameters from the input diction	
save_trained_params() After model fitting, save all the trained parameter	rs in
trained_model_params in some order.	
_load_model_from_trained_params() Load the model from trained params.	
initialize_early_stopping() Initializes and creates evaluation graph for e	arly
stopping.	
_perform_early_stopping_test(epoch) Performs regular validation checks and stop ear	ly if
the criteria is achieved.	
<pre>configure_evaluation_protocol([config]) Set the configuration for evaluation</pre>	
set_filter_for_eval() Configures to use filter	
_initialize_eval_graph([mode]) Initialize the evaluation graph.	
end_evaluation() End the evaluation and close the Tensorflow ses	

Also creates a new Tensorflow session for training.

Parameters

- **k** (*int*) Embedding space dimensionality.
- **eta** (*int*) The number of negatives that must be generated at runtime during training for each positive.
- epochs (*int*) The iterations of the training loop.
- **batches_count** (*int*) The number of batches in which the training set must be split during the training loop.
- **seed** (*int*) The seed used by the internal random numbers generator.
- **embedding_model_params** (*dict*) Model-specific hyperparams, passed to the model as a dictionary. Refer to model-specific documentation for details.

For FocusE Layer, following hyper-params can be passed:

- 'non_linearity': can be one of the following values linear, softplus, sigmoid, tanh
- 'stop_epoch': specifies how long to decay (linearly) the numeric values from 1 to original value

until it reachs original value. - **'structural_wt'**: structural influence hyperparameter [0, 1] that modulates the influence of graph topology. - **'normalize_numeric_values'**: normalize the numeric values, such that they are scaled between [0, 1]

- **optimizer** (*string*) The optimizer used to minimize the loss function. Choose between 'sgd', 'adagrad', 'adam', 'momentum'.
- **optimizer_params** (*dict*) Arguments specific to the optimizer, passed as a dictionary.

Supported keys:

- 'lr' (float): learning rate (used by all the optimizers). Default: 0.1.
- 'momentum' (float): learning momentum (only used when optimizer=momentum). Default: 0.9.

Example: optimizer_params={'lr': 0.01}

- **loss** (*string*) The type of loss function to use during training.
 - pairwise the model will use pairwise margin-based loss function.
 - nll the model will use negative loss likelihood.
 - absolute_margin the model will use absolute margin likelihood.
 - self_adversarial the model will use adversarial sampling loss function.
 - multiclass_nll the model will use multiclass nll loss. Switch to multiclass loss defined in [aC15] by passing 'corrupt_side' as ['s','o'] to embedding_model_params. To use loss defined in [KBK17] pass 'corrupt_side' as 'o' to embedding_model_params.
- **loss_params** (*dict*) Dictionary of loss-specific hyperparameters. See *loss functions* documentation for additional details.

Example: optimizer_params={'lr': 0.01} if loss='pairwise'.

- **regularizer** (*string*) The regularization strategy to use with the loss function.
 - None: the model will not use any regularizer (default)
 - LP: the model will use L1, L2 or L3 based on the value of regularizer_params['p'] (see below).
- **regularizer_params** (*dict*) Dictionary of regularizer-specific hyperparameters. See the *regularizers* documentation for additional details.

```
Example: regularizer_params={'lambda': 1e-5, 'p': 2} if
regularizer='LP'.
```

- **initializer** (*string*) The type of initializer to use.
 - normal: The embeddings will be initialized from a normal distribution
 - uniform: The embeddings will be initialized from a uniform distribution
 - xavier: The embeddings will be initialized using xavier strategy (default)
- **initializer_params** (*dict*) Dictionary of initializer-specific hyperparameters. See the *initializer* documentation for additional details.

```
Example: initializer_params={'mean': 0, 'std': 0.001} if
initializer='normal'.
```

- **large_graphs** (*bool*) Avoid loading entire dataset onto GPU when dealing with large graphs.
- **verbose** (*bool*) Verbose mode.
- fit (X, early_stopping=False, early_stopping_params={}, focusE_numeric_edge_values=None, tensorboard_logs_path=None)

Train an EmbeddingModel (with optional early stopping).

The model is trained on a training set X using the training protocol described in [TWR+16].

Parameters

- X (ndarray (shape [n, 3]) or object of AmpligraphDatasetAdapter) - Numpy array of training triples OR handle of Dataset adapter which would help retrieve data.
- **early_stopping** (*bool*) Flag to enable early stopping (default:False)
- **early_stopping_params** (*dictionary*) Dictionary of hyperparameters for the early stopping heuristics.

The following string keys are supported:

- 'x_valid': ndarray (shape [n, 3]) or object of AmpligraphDatasetAdapter : Numpy array of validation triples OR handle of Dataset adapter which would help retrieve data.
- 'criteria': string : criteria for early stopping 'hits10', 'hits3', 'hits1' or 'mrr'(default).
- 'x_filter': ndarray, shape [n, 3] [Positive triples to use as filter if a 'filtered' early] stopping criteria is desired (i.e. filtered-MRR if 'criteria': 'mrr'). Note this will affect training time (no filter by default). If the filter has already been set in the adapter, pass True
- 'burn_in': int : Number of epochs to pass before kicking in early stopping (default: 100).

- check_interval': int : Early stopping interval after burn-in (default:10).
- 'stop_interval': int : Stop if criteria is performing worse over n consecutive checks (default: 3)
- 'corruption_entities': List of entities to be used for corruptions. If 'all', it uses all entities (default: 'all')
- 'corrupt_side': Specifies which side to corrupt. 's', 'o', 's+o', 's,o' (default)

```
Example: early_stopping_params={x_valid=X['valid'],
'criteria': 'mrr'}
```

- **focusE_numeric_edge_values** (*nd array (n, 1)*) Numeric values associated with links. Semantically, the numeric value can signify importance, uncertainity, significance, confidence, etc. If the numeric value is unknown pass a NaN weight. The model will uniformly randomly assign a numeric value. One can also think about assigning numeric values by looking at the distribution of it per predicate.
- tensorboard_logs_path(*str or None*) Path to store tensorboard logs, e.g. average training loss tracking per epoch (default: None indicating no logs will be collected). When provided it will create a folder under provided path and save tensorboard files there. To then view the loss in the terminal run: tensorboard --logdir <tensorboard_logs_path>.

get_embeddings (*entities*, *embedding_type='entity'*) Get the embeddings of entities or relations.

Note: Use *ampligraph.utils.create_tensorboard_visualizations()* to visualize the embeddings with TensorBoard.

Parameters

- **entities** (*array-like*, *dtype=int*, *shape=[n]*) The entities (or relations) of interest. Element of the vector must be the original string literals, and not internal IDs.
- **embedding_type** (*string*) If 'entity', entities argument will be considered as a list of knowledge graph entities (i.e. nodes). If set to 'relation', they will be treated as relation types instead (i.e. predicates).

Returns embeddings - An array of k-dimensional embeddings.

Return type ndarray, shape [n, k]

get_hyperparameter_dict()

Returns hyperparameters of the model.

Returns hyperparam_dict – Dictionary of hyperparameters that were used for training.

Return type dict

predict (X, from_idx=False)

Predict the scores of triples using a trained embedding model. The function returns raw scores generated by the model.

Note: To obtain probability estimates, calibrate the model with *calibrate()*, then call *predict_proba()*.

Parameters

- X (ndarray, shape [n, 3]) The triples to score.
- from_idx (bool) If True, will skip conversion to internal IDs. (default: False).

Returns scores_predict – The predicted scores for input triples X.

Return type ndarray, shape [n]

calibrate (*X_pos*, *X_neg=None*, *positive_base_rate=None*, *batches_count=100*, *epochs=50*) Calibrate predictions

The method implements the heuristics described in [TC20], using Platt scaling [P+99].

The calibrated predictions can be obtained with *predict_proba()* after calibration is done.

Ideally, calibration should be performed on a validation set that was not used to train the embeddings.

There are two modes of operation, depending on the availability of negative triples:

- 1. Both positive and negative triples are provided via X_pos and X_neg respectively. The optimization is done using a second-order method (limited-memory BFGS), therefore no hyperparameter needs to be specified.
- 2. Only positive triples are provided, and the negative triples are generated by corruptions just like it is done in training or evaluation. The optimization is done using a first-order method (ADAM), therefore batches_count and epochs must be specified.

Calibration is highly dependent on the base rate of positive triples. Therefore, for mode (2) of operation, the user is required to provide the positive_base_rate argument. For mode (1), that can be inferred automatically by the relative sizes of the positive and negative sets, but the user can override that by providing a value to positive_base_rate.

Defining the positive base rate is the biggest challenge when calibrating without negatives. That depends on the user choice of which triples will be evaluated during test time. Let's take WN11 as an example: it has around 50% positives triples on both the validation set and test set, so naturally the positive base rate is 50%. However, should the user resample it to have 75% positives and 25% negatives, its previous calibration will be degraded. The user must recalibrate the model now with a 75% positive base rate. Therefore, this parameter depends on how the user handles the dataset and cannot be determined automatically or a priori.

Note: Incompatible with large graph mode (i.e. if self.dealing_with_large_graphs=True).

Note: Experiments for the ICLR-21 calibration paper are available here [TC20].

Parameters

- **X_pos** (*ndarray* (*shape* [*n*, 3])) Numpy array of positive triples.
- X_neg (ndarray (shape [n, 3])) Numpy array of negative triples.

If *None*, the negative triples are generated via corruptions and the user must provide a positive base rate instead.

• **positive_base_rate** (float) – Base rate of positive statements.

For example, if we assume there is a fifty-fifty chance of any query to be true, the base rate would be 50%.

If X_neg is provided and this is *None*, the relative sizes of X_pos and X_neg will be used to determine the base rate. For example, if we have 50 positive triples and 200 negative triples, the positive base rate will be assumed to be 50/(50+200) = 1/5 = 0.2.

This must be a value between 0 and 1.

- **batches_count** (*int*) Number of batches to complete one epoch of the Platt scaling training. Only applies when X_neg is *None*.
- **epochs** (*int*) Number of epochs used to train the Platt scaling model. Only applies when X_neg is *None*.

Examples

```
>>> import numpy as np
>>> from sklearn.metrics import brier score loss, log loss
>>> from scipy.special import expit
>>>
>>> from ampligraph.datasets import load_wn11
>>> from ampligraph.latent_features.models import TransE
>>>
>>> X = load wn11()
>>> X_valid_pos = X['valid'][X['valid_labels']]
>>> X_valid_neg = X['valid'][~X['valid_labels']]
>>>
>>> model = TransE(batches_count=64, seed=0, epochs=500, k=100, eta=20,
                   optimizer='adam', optimizer_params={'lr':0.0001},
>>>
                   loss='pairwise', verbose=True)
>>>
>>>
>>> model.fit(X['train'])
>>>
>>> # Raw scores
>>> scores = model.predict(X['test'])
>>>
>>> # Calibrate with positives and negatives
>>> model.calibrate(X_valid_pos, X_valid_neq, positive_base_rate=None)
>>> probas_pos_neg = model.predict_proba(X['test'])
>>>
>>> # Calibrate with just positives and base rate of 50%
>>> model.calibrate(X_valid_pos, positive_base_rate=0.5)
>>> probas_pos = model.predict_proba(X['test'])
>>>
>>> # Calibration evaluation with the Brier score loss (the smaller, the
\rightarrow better)
>>> print("Brier scores")
>>> print("Raw scores:", brier_score_loss(X['test_labels'], expit(scores)))
>>> print("Positive and negative calibration:", brier_score_loss(X['test_
→labels'], probas pos neg))
>>> print("Positive only calibration:", brier_score_loss(X['test_labels'],...

→probas_pos))
```

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```
Brier scores
Raw scores: 0.4925058891371126
Positive and negative calibration: 0.20434617882733366
Positive only calibration: 0.22597599585144656
```

$predict_proba(X)$

Predicts probabilities using the Platt scaling model (after calibration).

Model must be calibrated beforehand with the calibrate method.

Parameters X (*ndarray* (*shape* [*n*, 3])) – Numpy array of triples to be evaluated.

Returns probas – Probability of each triple to be true according to the Platt scaling calibration.

Return type ndarray (shape [n])

_fn (*e_s*, *e_p*, *e_o*)

The scoring function of the model.

Assigns a score to a list of triples, with a model-specific strategy. Triples are passed as lists of subject, predicate, object embeddings. This function must be overridden by every model to return corresponding score.

Parameters

- **e_s** (*Tensor*, *shape* [*n*]) The embeddings of a list of subjects.
- **e_p**(*Tensor*, *shape* [*n*]) The embeddings of a list of predicates.
- **e_o** (*Tensor*, *shape* [*n*]) The embeddings of a list of objects.

Returns score – The operation corresponding to the scoring function.

Return type TensorFlow operation

_initialize_parameters()

Initialize parameters of the model.

This function creates and initializes entity and relation embeddings (with size k). If the graph is large, then it loads only the required entity embeddings (max:batch_size*2) and all relation embeddings. Overload this function if the parameters needs to be initialized differently.

_get_model_loss (dataset_iterator)

Get the current loss including loss due to regularization. This function must be overridden if the model uses combination of different losses(eg: VAE).

Parameters dataset_iterator(*tf.data.Iterator*) – Dataset iterator.

Returns loss – The loss value that must be minimized.

Return type tf.Tensor

get_embedding_model_params (*output_dict*) Save the model parameters in the dictionary.

Parameters output_dict (*dictionary*) – Dictionary of saved params. It's the duty of the model to save all the variables correctly, so that it can be used for restoring later.

restore_model_params(in_dict)

Load the model parameters from the input dictionary.

Parameters in_dict (*dictionary*) – Dictionary of saved params. It's the duty of the model to load the variables correctly.

_save_trained_params()

After model fitting, save all the trained parameters in trained_model_params in some order. The order would be useful for loading the model. This method must be overridden if the model has any other parameters (apart from entity-relation embeddings).

_load_model_from_trained_params()

Load the model from trained params. While restoring make sure that the order of loaded parameters match the saved order. It's the duty of the embedding model to load the variables correctly. This method must be overridden if the model has any other parameters (apart from entity-relation embeddings). This function also set's the evaluation mode to do lazy loading of variables based on the number of distinct entities present in the graph.

_initialize_early_stopping()

Initializes and creates evaluation graph for early stopping.

_perform_early_stopping_test (epoch)

Performs regular validation checks and stop early if the criteria is achieved.

Parameters epoch (*int*) – current training epoch.

Returns stopped – Flag to indicate if the early stopping criteria is achieved.

Return type bool

configure_evaluation_protocol(config=None)

Set the configuration for evaluation

- **Parameters config** (*dictionary*) Dictionary of parameters for evaluation configuration. Can contain following keys:
 - **corruption_entities**: List of entities to be used for corruptions. If all, it uses all entities (default: all)
 - **corrupt_side**: Specifies which side to corrupt. s, o, s+o, s, o (default) In 's,o' mode subject and object corruptions are generated at once but ranked separately for speed up (default: False).

set_filter_for_eval()

Configures to use filter

_initialize_eval_graph (mode='test')

Initialize the evaluation graph.

Parameters mode (*string*) – Indicates which data generator to use.

end_evaluation()

End the evaluation and close the Tensorflow session.

Loss

class ampligraph.latent_features.**Loss** (*eta*, *hyperparam_dict*, *verbose=False*) Abstract class for loss function.

Methods

init(eta, hyperparam_dict[, verbose])	Initialize Loss.
get_state(param_name)	Get the state value.

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init_hyperparams(hyperparam_dict)	Initializes the hyperparameters needed by the algo-	
	rithm.	
inputs_check(scores_pos, scores_neg)	Creates any dependencies that need to be checked	
	before performing loss computations	
<pre>apply(scores_pos, scores_neg)</pre>	Interface to external world.	
_apply(scores_pos, scores_neg)	Apply the loss function.	

Table 14 - continued from previous page

___init___(*eta*, *hyperparam_dict*, *verbose=False*) Initialize Loss.

Parameters

- **eta** (*int*) number of negatives
- **hyperparam_dict** (*dict*) dictionary of hyperparams. (Keys are described in the hyperparameters section)

get_state (param_name)

Get the state value.

Parameters param_name (*string*) – Name of the state for which one wants to query the value.

Returns The value of the corresponding state.

Return type param_value

_init_hyperparams (hyperparam_dict)

Initializes the hyperparameters needed by the algorithm.

Parameters hyperparam_dict (*dictionary*) – Consists of key value pairs. The Loss will check the keys to get the corresponding params.

_inputs_check (scores_pos, scores_neg)

Creates any dependencies that need to be checked before performing loss computations

Parameters

- **scores_pos** (*tf.Tensor*) A tensor of scores assigned to positive statements.
- **scores_neg** (*tf.Tensor*) A tensor of scores assigned to negative statements.

apply (scores_pos, scores_neg)

Interface to external world. This function does the input checks, preprocesses input and finally applies loss function.

Parameters

- **scores_pos** (*tf.Tensor*) A tensor of scores assigned to positive statements.
- **scores_neg** (*tf.Tensor*) A tensor of scores assigned to negative statements.

Returns loss – The loss value that must be minimized.

Return type tf.Tensor

_apply (scores_pos, scores_neg)

Apply the loss function. Every inherited class must implement this function. (All the TF code must go in this function.)

Parameters

- **scores_pos** (*tf.Tensor*) A tensor of scores assigned to positive statements.
- **scores_neg** (*tf.Tensor*) A tensor of scores assigned to negative statements.

Returns loss – The loss value that must be minimized.

Return type tf. Tensor

Regularizer

class ampligraph.latent_features.**Regularizer** (*hyperparam_dict*, *verbose=False*) Abstract class for Regularizer.

Methods

init(hyperparam_dict[, verbose])	Initialize the regularizer.
get_state(param_name)	Get the state value.
init_hyperparams(hyperparam_dict)	Initializes the hyperparameters needed by the algo-
	rithm.
apply(trainable_params)	Interface to external world.
_apply(trainable_params)	Apply the regularization function.

__init___(hyperparam_dict, verbose=False) Initialize the regularizer.

Parameters hyperparam_dict (*dict*)-dictionary of hyperparams (Keys are described in the hyperparameters section)

get_state (param_name)

Get the state value.

Parameters param_name (*string*) – name of the state for which one wants to query the value

Returns the value of the corresponding state

Return type param_value

_init_hyperparams (hyperparam_dict)

Initializes the hyperparameters needed by the algorithm.

Parameters hyperparam_dict (*dictionary*) – Consists of key value pairs. The regularizer will check the keys to get the corresponding params

apply (*trainable_params*)

Interface to external world. This function performs input checks, input pre-processing, and and applies the loss function.

Parameters trainable_params(*list*, *shape* [*n*])-List of trainable params that should be regularized

Returns loss – Regularization Loss

Return type tf.Tensor

__apply(*trainable_params*)

Apply the regularization function. Every inherited class must implement this function.

(All the TF code must go in this function.)

Parameters trainable_params (*list*, *shape* [*n*]) – List of trainable params that should be regularized

Returns loss - Regularization Loss

Return type tf. Tensor

Initializer

Abstract class for initializer .

Methods

	Initialize the Class
_init_hyperparams(hyperparam_dict)	Initializes the hyperparameters.
_get_tf_initializer([in_shape, out_shape,	Create a tensorflow node for initializer
])	
_get_np_initializer([in_shape, out_shape,	Create an initialized numpy array
])	
_display_params()	Display the parameter values
<pre>get_entity_initializer([in_shape,])</pre>	Initializer for entity embeddings
gee_enerey_interarizer([in_simple,])	8-

__init__ (initializer_params={}, verbose=True, seed=0) Initialize the Class

Parameters

- **initializer_params** (*dict*) dictionary of hyperparams that would be used by the initializer.
- **verbose** (*bool*) set/reset verbose mode
- **seed** (*int/np.random.RandomState*) random state for random number generator

_init_hyperparams (hyperparam_dict)

Initializes the hyperparameters.

Parameters hyperparam_dict (*dictionary*) – Consists of key value pairs. The initializer will check the keys to get the corresponding params

_get_tf_initializer (*in_shape=None*, *out_shape=None*, *concept='e'*) Create a tensorflow node for initializer

Parameters

- **in_shape** (*int*) number of inputs to the layer (fan in)
- **out_shape** (*int*) number of outputs of the layer (fan out)
- **concept** (*char*) **concept** type (e for entity, r for relation)

Returns initializer_instance

Return type An Initializer instance.

_get_np_initializer (in_shape=None, out_shape=None, concept='e')

Create an initialized numpy array

Parameters

- in_shape (int) number of inputs to the layer (fan in)
- **out_shape** (*int*) number of outputs of the layer (fan out)
- **concept** (*char*) **concept** type (e for entity, r for relation)

Returns initialized_values - Initialized weights

Return type n-d array

```
_display_params()
```

Display the parameter values

get_entity_initializer (in_shape=None, out_shape=None, init_type='tf')

Initializer for entity embeddings

Parameters

- **in_shape** (*int*) number of inputs to the layer (fan in)
- **out_shape** (*int*) number of outputs of the layer (fan out)
- **init_type** (*string*) Type of initializer ('tf' for tensorflow, 'np' for numpy)

Returns initialized_values - Weights initializer

Return type tf.Op or n-d array

get_relation_initializer (in_shape=None, out_shape=None, init_type='tf')
Initializer for relation embeddings

Parameters

- **in_shape** (*int*) number of inputs to the layer (fan in)
- **out_shape** (*int*) number of outputs of the layer (fan out)
- **init_type** (*string*) Type of initializer ('tf' for tensorflow, 'np' for numpy)

Returns initialized_values - Weights initializer

Return type tf.Op or n-d array

Scoring functions

Existing models propose scoring functions that combine the embeddings \mathbf{e}_s , \mathbf{r}_p , $\mathbf{e}_o \in \mathcal{R}^k$ of the subject, predicate, and object of a triple t = (s, p, o) according to different intuitions:

• *TransE* [BUGD+13] relies on distances. The scoring function computes a similarity between the embedding of the subject translated by the embedding of the predicate and the embedding of the object, using the L_1 or L_2 norm $|| \cdot ||$:

$$f_{TransE} = -||\mathbf{e}_s + \mathbf{r}_p - \mathbf{e}_o||_n$$

• DistMult [YYH+14] uses the trilinear dot product:

$$f_{DistMult} = \langle \mathbf{r}_p, \mathbf{e}_s, \mathbf{e}_o \rangle$$

• Complex [TWR+16] extends DistMult with the Hermitian dot product:

$$f_{ComplEx} = Re(\langle \mathbf{r}_p, \mathbf{e}_s, \overline{\mathbf{e}_o} \rangle)$$

• *Hole* [NRP+16] uses circular correlation (denoted by \otimes):

$$f_{HolE} = \mathbf{w}_r \cdot (\mathbf{e}_s \otimes \mathbf{e}_o) = \frac{1}{k} \mathcal{F}(\mathbf{w}_r) \cdot (\overline{\mathcal{F}(\mathbf{e}_s)} \odot \mathcal{F}(\mathbf{e}_o))$$

• ConvE [DMSR18] uses convolutional layers (g is a non-linear activation function, * is the linear convolution operator, vec indicates 2D reshaping):

$$f_{ConvE} = \langle \sigma \left(vec \left(g \left(\left[\overline{\mathbf{e}_s}; \overline{\mathbf{r}_p} \right] * \Omega \right) \right) \mathbf{W} \right) \right) \mathbf{e}_o \rangle$$

• ConvKB [NNNP18] uses convolutional layers and a dot product:

$$f_{ConvKB} = concat \left(g\left(\left[\mathbf{e}_s, \mathbf{r}_p, \mathbf{e}_o \right] \right) * \Omega \right) \right) \cdot W$$

Loss Functions

AmpliGraph includes a number of loss functions commonly used in literature. Each function can be used with any of the implemented models. Loss functions are passed to models as hyperparameter, and they can be thus used *during model selection*.

<pre>PairwiseLoss(eta[, loss_params, verbose])</pre>		Pairwise, max-margin loss.
AbsoluteMarginLoss(eta[, loss_params,	ver-	Absolute margin, max-margin loss.
bose])		
SelfAdversarialLoss(eta[, loss_params,	ver-	Self adversarial sampling loss.
bose])		
NLLLoss(eta[, loss_params, verbose])		Negative log-likelihood loss.
<pre>NLLMulticlass(eta[, loss_params, verbose])</pre>		Multiclass NLL Loss.
BCELoss(eta[, loss_params, verbose])		Binary Cross Entropy Loss.

PairwiseLoss

class ampligraph.latent_features.**PairwiseLoss**(*eta*, *loss_params=None*, *verbose=False*) Pairwise, max-margin loss.

Introduced in [BUGD+13].

$$\mathcal{L}(\Theta) = \sum_{t^+ \in \mathcal{G}} \sum_{t^- \in \mathcal{C}} max(0, [\gamma + f_{model}(t^-; \Theta) - f_{model}(t^+; \Theta)])$$

where γ is the margin, \mathcal{G} is the set of positives, \mathcal{C} is the set of corruptions, $f_{model}(t; \Theta)$ is the model-specific scoring function.

Methods

_____init___(eta[, loss_params, verbose]) Initialize Loss.

___init___(eta, loss_params=None, verbose=False) Initialize Loss.

Parameters

• **eta** (*int*) – Number of negatives.

• **loss_params** (*dict*) – Dictionary of loss-specific hyperparams:

- 'margin': (float). Margin to be used in pairwise loss computation (default: 1)

```
Example: loss_params={'margin': 1}
```

AbsoluteMarginLoss

class ampligraph.latent_features.AbsoluteMarginLoss(eta, loss_params=None, verbose=False)

Absolute margin , max-margin loss.

Introduced in [HOSM17].

$$\mathcal{L}(\Theta) = \sum_{t^+ \in \mathcal{G}} \sum_{t^- \in \mathcal{C}} f_{model}(t^-; \Theta) - max(0, [\gamma - f_{model}(t^+; \Theta)])$$

where γ is the margin, \mathcal{G} is the set of positives, \mathcal{C} is the set of corruptions, $f_{model}(t; \Theta)$ is the model-specific scoring function.

Methods

init(eta[, loss_params, verbose])	Initialize Loss	
-----------------------------------	-----------------	--

____init___(eta, loss_params=None, verbose=False) Initialize Loss

Parameters

• **eta** (*int*) – Number of negatives.

- **loss_params** (*dict*) Dictionary of loss-specific hyperparams:
 - 'margin': float. Margin to be used in pairwise loss computation (default:1)

Example: loss_params={ 'margin': 1}

SelfAdversarialLoss

class ampligraph.latent_features.SelfAdversarialLoss(eta, loss_params=None, ver-

Self adversarial sampling loss.

Introduced in [SDNT19].

$$\mathcal{L} = -\log\sigma(\gamma + f_{model}(\mathbf{s}, \mathbf{o})) - \sum_{i=1}^{n} p(h_{i}^{'}, r, t_{i}^{'}) \log\sigma(-f_{model}(\mathbf{s}_{i}^{'}, \mathbf{o}_{i}^{'}) - \gamma)$$

where $\mathbf{s}, \mathbf{o} \in \mathcal{R}^k$ are the embeddings of the subject and object of a triple $t = (s, r, o), \gamma$ is the margin, σ the sigmoid function, and $p(s'_i, r, o'_i)$ is the negatives sampling distribution which is defined as:

$$p(s'_j, r, o'_j | \{(s_i, r_i, o_i)\}) = \frac{\exp \alpha f_{model}(\mathbf{s}'_j, \mathbf{o}'_j)}{\sum_i \exp \alpha f_{model}(\mathbf{s}'_i, \mathbf{o}'_i)}$$

where α is the temperature of sampling, f_{model} is the scoring function of the desired embeddings model.

Methods

____init___(eta[, loss_params, verbose]) Initialize Loss

___init___(eta, loss_params=None, verbose=False) Initialize Loss

Parameters

- **eta** (*int*) number of negatives
- **loss_params** (*dict*) Dictionary of loss-specific hyperparams:
 - 'margin': (float). Margin to be used for loss computation (default: 1)
 - 'alpha': (float). Temperature of sampling (default:0.5)

Example: loss_params={'margin': 1, 'alpha': 0.5}

NLLLoss

class ampligraph.latent_features.NLLLoss(eta, loss_params=None, verbose=False)
 Negative log-likelihood loss.

As described in [TWR+16].

$$\mathcal{L}(\Theta) = \sum_{t \in \mathcal{G} \cup \mathcal{C}} \log(1 + exp(-y f_{model}(t; \Theta)))$$

where $y \in -1, 1$ is the label of the statement, \mathcal{G} is the set of positives, \mathcal{C} is the set of corruptions, $f_{model}(t; \Theta)$ is the model-specific scoring function.

Methods

init	_(eta[, loss_params, verbose])	Initialize Loss.	
------	--------------------------------	------------------	--

__init___(*eta*, *loss_params=None*, *verbose=False*) Initialize Loss.

Parameters

- **eta** (*int*) Number of negatives.
- **loss_params** (*dict*) Dictionary of hyperparams. No hyperparameters are required for this loss.

NLLMulticlass

```
class ampligraph.latent_features.NLLMulticlass(eta, loss_params=None, ver-
bose=False)
```

Multiclass NLL Loss.

Introduced in [aC15] where both the subject and objects are corrupted (to use it in this way pass corrupt_sides = ['s', 'o'] to embedding_model_params).

This loss was re-engineered in [KBK17] where only the object was corrupted to get improved performance (to use it in this way pass corrupt_sides = 'o' to embedding_model_params).

$$\mathcal{L}(\mathcal{X}) = -\sum_{x_{e_1, e_2, r_k} \in X} \log p(e_2|e_1, r_k) - \sum_{x_{e_1, e_2, r_k} \in X} \log p(e_1|r_k, e_2)$$

Examples

```
>>> from ampligraph.latent_features import TransE
>>> model = TransE(batches_count=1, seed=555, epochs=20, k=10,
>>> embedding_model_params={'corrupt_sides':['s', 'o']},
>>> loss='multiclass_nll', loss_params={})
```

Methods

___init___(eta, loss_params=None, verbose=False) Initialize Loss

Parameters

- **eta** (*int*) number of negatives
- **loss_params** (*dict*) Dictionary of loss-specific hyperparams:

BCELoss

class ampligraph.latent_features.BCELoss(eta, loss_params={}, verbose=False)
Binary Cross Entropy Loss.

$$\mathcal{L} = -\frac{1}{N} \sum_{i=1}^{N} y_i \cdot \log(p(y_i)) + (1 - y_i) \cdot \log(1 - p(y_i))$$

Examples

Methods

____init___(eta[, loss_params, verbose]) Initialize Loss

___init___(eta, loss_params={}, verbose=False) Initialize Loss

Parameters loss_params (*dict*) – Dictionary of loss-specific hyperparams:

Regularizers

AmpliGraph includes a number of regularizers that can be used with the *loss function*. *LPRegularizer* supports L1, L2, and L3.

LPRegularizer([regularizer_params, verbose]) Performs LP regularization

LPRegularizer

class ampligraph.latent_features.LPRegularizer(regularizer_params=None, verbose=False)

Performs LP regularization

$$\mathcal{L}(Reg) = \sum_{i=1}^{n} \lambda_i * \mid w_i \mid_p$$

where n is the number of model parameters, $p \in 1, 2, 3$ is the p-norm and λ is the regularization weight. For example, if p = 1 the function will perform L1 regularization. L2 regularization is obtained with p = 2. The nuclear 3-norm proposed in the ComplEx-N3 paper [LUO18] can be obtained with regularizer_params={'p': 3}.

Methods

 Initializes the hyperparameters needed by the algo-
rithm.

__init___(regularizer_params=None, verbose=False)

Initializes the hyperparameters needed by the algorithm.

Parameters regularizer_params (*dictionary*) – Consists of key-value pairs. The regularizer will check the keys to get the corresponding params:

- 'lambda': (float). Weight of regularization loss for each parameter (default: 1e-5)
- 'p': (int): norm (default: 2)

```
Example: regularizer_params={'lambda': 1e-5, 'p': 1}
```

Initializers

AmpliGraph includes a number of initializers that can be used to initialize the embeddings. They can be passed as hyperparameter, and they can be thus used *during model selection*.

<pre>RandomNormal([initializer_params, verbose, seed])</pre>	Initializes from a normal distribution with specified
	mean and std
RandomUniform([initializer_params, verbose,])	Initializes from a uniform distribution with specified
	low and high
Xavier([initializer_params, verbose, seed])	Follows the xavier strategy for initialization of layers
	[GB10].
Constant([initializer_params, verbose, seed])	Initializes with the constant values provided by the user

RandomNormal

 $\mathcal{N}(\mu, \sigma)$

Methods

init([initializer_params, verbose, seed])	Initialize the Random Normal initialization strategy
init(<i>initializer_params={}</i> , <i>verbose=True</i> Initialize the Random Normal initialization	

Parameters

- **initializer_params** (*dict*) Consists of key-value pairs. The initializer will check the keys to get the corresponding params:
 - mean: (float). Mean of the weights(default: 0)
 - std: (float): std of the weights (default: 0.05)

Example: initializer_params={'mean': 0, 'std': 0.01}

- **verbose** (*bool*) Enable/disable verbose mode
- **seed** (*int/np.random.RandomState*) random state for random number generator

RandomUniform

 $\mathcal{U}(low, high)$

Methods

[Initializer_parallis, verbose, seed]) Initialize the official initialization strategy	init_	_([initializer_params, verbose, seed])	Initialize the Uniform initialization strategy
--	-------	--	--

__init___(initializer_params={}, verbose=True, seed=0)
Initialize the Uniform initialization strategy

Parameters

- **initializer_params** (*dict*) Consists of key-value pairs. The initializer will check the keys to get the corresponding params:
 - low: (float). lower bound for uniform number (default: -0.05)
 - high: (float): upper bound for uniform number (default: 0.05)

Example: initializer_params={'low': 0, 'high': 0.01}

- **verbose** (*bool*) Enable/disable verbose mode
- **seed** (*int/np.random.RandomState*) random state for random number generator

Xavier

class ampligraph.latent_features.**Xavier** (*initializer_params={*}, *verbose=True*, *seed=0*) Follows the xavier strategy for initialization of layers [GB10].

If uniform is set to True, then it initializes the layer from the following uniform distribution:

$$\mathcal{U}(-\sqrt{\frac{6}{fan_{in}+fan_{out}}},\sqrt{\frac{6}{fan_{in}+fan_{out}}})$$

If uniform is False, then it initializes the layer from the following normal distribution:

$$\mathcal{N}(0, \sqrt{\frac{2}{fan_{in} + fan_{out}}})$$

where fan_{in} and fan_{out} are number of input units and output units of the layer respectively.

Methods

1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	/r· ·/· 1·	1 17\	T '.' 1' .1 XZ '
init	(linifializer narams	verbose seedly	Initialize the Xavier strategy
	([initializer_params,	verbose, seeup	minumize the Mayler strategy

__init__ (*initializer_params={}*, *verbose=True*, *seed=0*) Initialize the Xavier strategy

Parameters

- **initializer_params** (*dict*) Consists of key-value pairs. The initializer will check the keys to get the corresponding params:
 - **uniform**: (bool). indicates whether to use Xavier Uniform or Xavier Normal initializer.

Example: initializer_params={ 'uniform': False}

- **verbose** (*bool*) Enable/disable verbose mode
- **seed** (*int/np.random.RandomState*) random state for random number generator

Constant

class ampligraph.latent_features.Constant (initializer_params={}, verbose=True, seed=0)
Initializes with the constant values provided by the user

Methods

 Initialize the the constant values provided by the user
initialize the the constant values provided by the user

___init___(*initializer_params={}*, *verbose=True*, *seed=0*) Initialize the the constant values provided by the user

Parameters

- **initializer_params** (*dict*) Consists of key-value pairs. The initializer will check the keys to get the corresponding params:
 - entity: (np.ndarray.float32). Initial values for entity embeddings
 - relation: (np.ndarray.float32). Initial values for relation embeddings

```
Example: initializer_params={'entity': ent_init_value,
'relation': rel_init_value}
```

- **verbose** (bool) Enable/disable verbose mode
- **seed** (*int/np.random.RandomState*) random state for random number generator

Optimizers

The goal of the optimization procedure is learning optimal embeddings, such that the scoring function is able to assign high scores to positive statements and low scores to statements unlikely to be true.

We support SGD-based optimizers provided by TensorFlow, by setting the optimizer argument in a model initializer. Best results are currently obtained with Adam.

AdamOptimizer(optimizer_params, batches_count)	Wrapper around Adam Optimizer
AdagradOptimizer(optimizer_params,	Wrapper around adagrad optimizer
batches_count)	
SGDOptimizer(optimizer_params, batches_count)	Wrapper around SGD Optimizer
<pre>MomentumOptimizer(optimizer_params,[,])</pre>	Wrapper around Momentum Optimizer

AdamOptimizer

class ampligraph.latent_features.AdamOptimizer(optimizer_params, batches_count, verbose=False)

Wrapper around Adam Optimizer

Methods

		Initialize the Optimizer	
])			
minimize(loss)		Create an optimizer to minimize the model loss	
update_feed_dict(feed_dict,	batch_num,	Fills values of placeholders created by the optimiz-	
epoch_num)		ers.	

___init___(*optimizer_params*, *batches_count*, *verbose=False*) Initialize the Optimizer

Parameters

• **optimizer_params** (*dict*) – Consists of key-value pairs. The optimizer will check the keys to get the corresponding params:

- 'lr': (float). Learning Rate (default: 0.0005)

Example: optimizer_params={'lr': 0.001}

- **batches_count** (*int*) number of batches in an epoch
- **verbose** (*bool*) Enable/disable verbose mode

minimize (loss)

Create an optimizer to minimize the model loss

- **Parameters loss** (*tf.Tensor*) Node which needs to be evaluated for computing the model loss.
- Returns train Node that needs to be evaluated for minimizing the loss during training

Return type tf.Operation

update_feed_dict (*feed_dict*, *batch_num*, *epoch_num*) Fills values of placeholders created by the optimizers.

Parameters

- **feed_dict** (*dict*) Dictionary that would be passed while optimizing the model loss to sess.run.
- **batch_num** (*int*) current batch number
- **epoch_num** (*int*) current epoch number

AdagradOptimizer

Wrapper around adagrad optimizer

Methods

init(optimizer_params, batches_count[,]		Initialize the Optimizer	
])			
minimize(loss)		Create an optimizer to minimize the model loss	
update_feed_dict(feed_dict,	batch_num,	Fills values of placeholders created by the optimiz-	
epoch_num)		ers.	

___init___(optimizer_params, batches_count, verbose=False) Initialize the Optimizer

Parameters

• **optimizer_params** (*dict*) – Consists of key-value pairs. The optimizer will check the keys to get the corresponding params:

- 'lr': (float). Learning Rate (default: 0.0005)

Example: optimizer_params={'lr': 0.001}

• **batches_count** (*int*) – number of batches in an epoch

• **verbose** (*bool*) – Enable/disable verbose mode

minimize (loss)

Create an optimizer to minimize the model loss

Parameters loss (*tf.Tensor*) – Node which needs to be evaluated for computing the model loss.

Returns train – Node that needs to be evaluated for minimizing the loss during training

Return type tf.Operation

update_feed_dict (feed_dict, batch_num, epoch_num)
Fills values of placeholders created by the optimizers.

Parameters

- **feed_dict** (*dict*) Dictionary that would be passed while optimizing the model loss to sess.run.
- **batch_num** (*int*) current batch number
- **epoch_num** (*int*) current epoch number

SGDOptimizer

class ampligraph.latent_features.SGDOptimizer(optimizer_params, batches_count, verbose=False)

Wrapper around SGD Optimizer

Methods

init(optimizer_params, batches_count[,]		Initialize the Optimizer	
])			
minimize(loss)		Create an optimizer to minimize the model loss	
<pre>update_feed_dict(feed_dict,</pre>	batch_num,	Fills values of placeholders created by the optimiz-	
epoch_num)		ers.	

___init___(*optimizer_params*, *batches_count*, *verbose=False*) Initialize the Optimizer

Parameters

- **optimizer_params** (*dict*) Consists of key-value pairs. The optimizer will check the keys to get the corresponding params:
 - 'lr': (float). Learning Rate upper bound (default: 0.0005)
 - 'decay_cycle': (int). Cycle of epoch over which to decay (default: 0)
 - 'end_lr': (float). Learning Rate lower bound (default: 1e-8)
 - 'cosine_decay': (bool). Use cosine decay or to fixed rate decay (default: False)
 - 'expand_factor': (float). Expand the decay cycle length by this factor after each cycle (default: 1)
 - 'decay_lr_rate': (float). Decay factor to decay the start lr after each cycle (default: 2)

Example: optimizer_params={'lr': 0.01, 'decay_cycle':30, 'end_lr':0.0001, 'sine_decay':True}

- **batches_count** (*int*) number of batches in an epoch
- **verbose** (bool) Enable/disable verbose mode

minimize (loss)

Create an optimizer to minimize the model loss

Parameters loss (*tf.Tensor*) – Node which needs to be evaluated for computing the model loss.

Returns train - Node that needs to be evaluated for minimizing the loss during training

Return type tf.Operation

update_feed_dict (feed_dict, batch_num, epoch_num)

Fills values of placeholders created by the optimizers.

Parameters

- **feed_dict** (*dict*) Dictionary that would be passed while optimizing the model loss to sess.run.
- **batch_num** (*int*) current batch number
- **epoch_num** (*int*) current epoch number

MomentumOptimizer

Wrapper around Momentum Optimizer

Methods

init(optimizer_params, batches_count[,]		Initialize the Optimizer	
])			
minimize(loss)		Create an optimizer to minimize the model loss	
update_feed_dict(feed_dict,	batch_num,	Fills values of placeholders created by the optimiz-	
epoch_num)		ers.	

__init___(*optimizer_params*, *batches_count*, *verbose=False*) Initialize the Optimizer

Parameters

- **optimizer_params** (*dict*) Consists of key-value pairs. The optimizer will check the keys to get the corresponding params:
 - 'lr': (float). Learning Rate (default: 0.0005)
 - 'momentum': (float). Momentum (default: 0.9)

Example: optimizer_params={'lr': 0.001, 'momentum':0. 90}

• **batches_count** (*int*) – number of batches in an epoch

• **verbose** (*bool*) – Enable/disable verbose mode

minimize(loss)

Create an optimizer to minimize the model loss

Parameters loss (*tf.Tensor*) – Node which needs to be evaluated for computing the model loss.

Returns train – Node that needs to be evaluated for minimizing the loss during training

Return type tf.Operation

update_feed_dict (feed_dict, batch_num, epoch_num)
Fills values of placeholders created by the optimizers.

Parameters

- **feed_dict** (*dict*) Dictionary that would be passed while optimizing the model loss to sess.run.
- **batch_num** (*int*) current batch number
- **epoch_num** (*int*) current epoch number

Support for Numeric Literals on Edges

Numeric values associated to edges of a knowledge graph have been used to represent uncertainty, edge importance, and even out-of-band knowledge in a growing number of scenarios, ranging from genetic data to social networks.

Nevertheless, traditional knowledge graph embedding models (TransE, DistMult, ComplEx) are not designed to capture such information, to the detriment of predictive power.

AmpliGraph includes FocusE [PC21], a novel method to inject numeric edge attributes into the scoring layer of a traditional knowledge graph embedding architecture.

To process knowledge graphs with numeric-enriched edges, refer to the documentation and examples below:

- FocusE-TransE
- FocusE-DistMult
- FocusE-ComplEx

You can load publicly available benchmark knowledge graphs with numeric-enriched edges with the dataset loaders.

Note: Experiments for the IJCAI-21 FocusE paper are available here [PC21].

Saving/Restoring Models

Models can be saved and restored from disk. This is useful to avoid re-training a model.

More details in the *utils* module.

3.3.3 Evaluation

The module includes performance metrics for neural graph embeddings models, along with model selection routines, negatives generation, and an implementation of the learning-to-rank-based evaluation protocol used in literature.

Metrics

Learning-to-rank metrics to evaluate the performance of neural graph embedding models.

<pre>rank_score(y_true, y_pred[, pos_lab])</pre>	Rank of a triple
mrr_score(ranks)	Mean Reciprocal Rank (MRR)
mr_score(ranks)	Mean Rank (MR)
hits_at_n_score(ranks, n)	Hits@N

rank_score

ampligraph.evaluation.rank_score(y_true, y_pred, pos_lab=1)
Rank of a triple

The rank of a positive element against a list of negatives.

 $rank_{(s,p,o)_i}$

Parameters

- **y_true** (*ndarray*, *shape* [*n*]) An array of binary labels. The array only contains one positive.
- **y_pred** (*ndarray*, *shape* [*n*]) An array of scores, for the positive element and the n-1 negatives.
- **pos_lab** (*int*) The value of the positive label (default = 1).

Returns rank – The rank of the positive element against the negatives.

Return type int

Examples

```
>>> import numpy as np
>>> from ampligraph.evaluation.metrics import rank_score
>>> y_pred = np.array([.434, .65, .21, .84])
>>> y_true = np.array([0, 0, 1, 0])
>>> rank_score(y_true, y_pred)
4
```

mrr_score

```
ampligraph.evaluation.mrr_score(ranks)
    Mean Reciprocal Rank (MRR)
```

The function computes the mean of the reciprocal of elements of a vector of rankings ranks.

It is used in conjunction with the learning to rank evaluation protocol of *ampligraph.evaluation*. *evaluate_performance()*.

It is formally defined as follows:

$$MRR = \frac{1}{|Q|} \sum_{i=1}^{|Q|} \frac{1}{rank_{(s,p,o)_i}}$$

where Q is a set of triples and (s, p, o) is a triple $\in Q$.

Note: This metric is similar to mean rank (MR) *ampligraph.evaluation.mr_score()*. Instead of averaging ranks, it averages their reciprocals. This is done to obtain a metric which is more robust to outliers.

Consider the following example. Each of the two positive triples identified by \star are ranked against four corruptions each. When scored by an embedding model, the first triple ranks 2nd, and the other triple ranks first. The resulting MRR is:

S	р	C)	score	rank	
Jack	born_in	Ire	eland	0.789	1	
Jack	born_in	Ita	aly	0.753	2	*
Jack	born_in	Ger	rmany	0.695	3	
Jack	born_in	Chi	lna	0.456	4	
Jack	born_in	The	omas	0.234	5	
s	р	C)	score	rank	
Jack	friend_w:	lth	Thomas	0.901	1	*
Jack	friend_w:	lth	China	0.345	2	
Jack	friend_w:	lth	Italy	0.293	3	
Jack	friend_w:	lth	Ireland	0.201	4	
Jack	friend_w:	lth	Germany	0.156	5	
MRR=0.	75					

Parameters ranks (*ndarray or list*, *shape* [*n*] *or* [*n*, 2]) – Input ranks of n test statements.

Returns mrr_score – The MRR score

Return type float

Examples

```
>>> import numpy as np
>>> from ampligraph.evaluation.metrics import mrr_score
>>> rankings = np.array([1, 12, 6, 2])
>>> mrr_score(rankings)
0.4375
```

mr_score

The function computes the mean of of a vector of rankings ranks.

It can be used in conjunction with the learning to rank evaluation protocol of *ampligraph.evaluation*. *evaluate_performance()*.

It is formally defined as follows:

$$MR = \frac{1}{|Q|} \sum_{i=1}^{|Q|} rank_{(s,p,o)_i}$$

where Q is a set of triples and (s, p, o) is a triple $\in Q$.

Note: This metric is not robust to outliers. It is usually presented along the more reliable MRR *ampligraph*. *evaluation.mrr_score()*.

Consider the following example. Each of the two positive triples identified by * are ranked against four corruptions each. When scored by an embedding model, the first triple ranks 2nd, and the other triple ranks first. The resulting MR is:

S	р	C)	score	rank	
Jack	born_in	Ire	eland	0.789	1	
Jack	born_in	Ita	ly	0.753	2	*
Jack	born_in	Ger	many	0.695	3	
Jack	born_in	Chi	na	0.456	4	
Jack	born_in	The	omas	0.234	5	
S	р	C)	score	rank	
Jack	friend_wi	th	Thomas	0.901	1	*
Jack	friend_wi	th	China	0.345	2	
Jack	friend_wi	th	Italy	0.293	3	
Jack	friend_wi	th	Ireland	0.201	4	
Jack	friend_wi	th	Germany	0.156	5	
MR=1.5						

Parameters ranks (*ndarray or list*, *shape* [*n*] *or* [*n*, 2]) – Input ranks of n test statements.

Returns mr_score – The MR score

Return type float

Examples

```
>>> from ampligraph.evaluation import mr_score
>>> ranks= [5, 3, 4, 10, 1]
>>> mr_score(ranks)
4.6
```

hits_at_n_score

```
ampligraph.evaluation.hits_at_n_score(ranks, n)
    Hits@N
```

The function computes how many elements of a vector of rankings ranks make it to the top n positions.

It can be used in conjunction with the learning to rank evaluation protocol of *ampligraph.evaluation*. *evaluate_performance()*.

It is formally defined as follows:

$$Hits@N = \sum_{i=1}^{|Q|} 1 \text{ if } rank_{(s,p,o)_i} \leq N$$

where Q is a set of triples and (s, p, o) is a triple $\in Q$.

Consider the following example. Each of the two positive triples identified by \star are ranked against four corruptions each. When scored by an embedding model, the first triple ranks 2nd, and the other triple ranks first. Hits@1 and Hits@3 are:

```
s
       р
                0
                           score
                                  rank
Jack
     born_in Ireland
                          0.789
                                   1
Jack
     born_in Italy
                           0.753
                                     2
                           0.695
Jack born_in Germany
                                    3
    born_in China
                           0.456
Jack
                                     4
     born_in Thomas
                           0.234
                                     5
Jack
s
       р
               0
                           score
                                  rank
Jack friend_with Thomas
                                  1
                           0.901
    friend_with China
Jack
                           0.345
                                     2
                          0.293
Jack friend_with Italy
                                     3
Jack friend_with Ireland 0.201
                                     4
Jack friend_with Germany
                           0.156
                                     5
Hits@3=1.0
Hits@1=0.5
```

Parameters

- **ranks** (*ndarray or list*, *shape* [*n*] *or* [*n*,2]) Input ranks of n test statements.
- **n** (*int*) The maximum rank considered to accept a positive.

Returns hits_n_score – The Hits@n score

Return type float

Examples

```
>>> import numpy as np
>>> from ampligraph.evaluation.metrics import hits_at_n_score
>>> rankings = np.array([1, 12, 6, 2])
>>> hits_at_n_score(rankings, n=3)
0.5
```

Negatives Generation

Negatives generation routines. These are corruption strategies based on the Local Closed-World Assumption (LCWA).

generate_corruptions_for_eval(X,	[,	Generate corruptions for evaluation.
])		
generate_corruptions_for_fit(X[,	.])	Generate corruptions for training.

generate_corruptions_for_eval

```
ampligraph.evaluation.generate_corruptions_for_eval(X, entities_for_corruption, cor-
rupt_side='s, o')
Generate corruptions for evaluation.
```

Create corruptions (subject and object) for a given triple x, in compliance with the local closed

world assumption (LCWA), as described in [NMTG16].

Parameters

- X (Tensor, shape [1, 3]) Currently, a single positive triples that will be used to create corruptions.
- **entities_for_corruption** (*Tensor*) All the entity IDs which are to be used for generation of corruptions.
- **corrupt_side** (*string*) Specifies which side of the triple to corrupt:
 - 's': corrupt only subject.
 - 'o': corrupt only object
 - 's+o': corrupt both subject and object
 - 's,o': corrupt both subject and object but ranks are computed separately.

Returns out – An array of corruptions for the triples for x.

Return type Tensor, shape [n, 3]

generate_corruptions_for_fit

ampligraph.evaluation.generate_corruptions_for_fit (X, entities_list=None, eta=1, corrupt_side='s, o', entities_size=0, rnd=None)

Generate corruptions for training.

Creates corrupted triples for each statement in an array of statements, as described by [TWR+16].

Note: Collisions are not checked, as this will be computationally expensive [TWR+16]. That means that some corruptions *may* result in being positive statements (i.e. *unfiltered* settings).

Note: When processing large knowledge graphs, it may be useful to generate corruptions only using entities from a single batch. This also brings the benefit of creating more meaningful negatives, as entities used to corrupt are sourced locally. The function can be configured to generate corruptions *only* using the entities from the current batch. You can enable such behaviour be setting entities_size=0. In such case, if entities_list=None all entities from the *current batch* will be used to generate corruptions.

Parameters

- X (Tensor, shape [n, 3]) An array of positive triples that will be used to create corruptions.
- **entities_list** (*list*) List of entities to be used for generating corruptions. (default:None).

If entities_list=None and entities_size is the number of all entities, all entities will be used to generate corruptions (default behaviour).

If entities_list=None and entities_size=0, the batch entities will be used to generate corruptions.

- **eta** (*int*) The number of corruptions per triple that must be generated.
- **corrupt_side** (*string*) Specifies which side of the triple to corrupt:

- 's': corrupt only subject.
- 'o': corrupt only object
- 's+o': corrupt both subject and object
- 's,o': corrupt both subject and object
- **entities_size** (*int*) Size of entities to be used while generating corruptions. It assumes entity id's start from 0 and are continuous. (default: 0). When processing large knowledge graphs, it may be useful to generate corruptions only using entities from a single batch. This also brings the benefit of creating more meaningful negatives, as entities used to corrupt are sourced locally. The function can be configured to generate corruptions *only* using the entities from the current batch. You can enable such behaviour be setting entities_size=0. In such case, if entities_list=None all entities from the *current batch* will be used to generate corruptions.
- **rnd** (*numpy.random.RandomState*) A random number generator.
- **Returns out** An array of corruptions for a list of positive triples X. For each row in X the corresponding corruption indexes can be found at [index+i*n for i in range(eta)]

Return type Tensor, shape [n * eta, 3]

Evaluation & Model Selection

Functions to evaluate the predictive power of knowledge graph embedding models, and routines for model selection.

<pre>evaluate_performance(X, model[,])</pre>	Evaluate the performance of an embedding model.
<pre>select_best_model_ranking(model_class,</pre>	Model selection routine for embedding models via ei-
)	ther grid search or random search.

evaluate_performance

ampligraph.evaluation.evaluate_performance	(<i>X</i> ,	model,	filter_triple	es=None,	ver-
	bose	=False,	filter_unse	en=True,	en-
	tities	ities_subset=None, corrupt_		corrupt_si	de = s,
	<i>o'</i> ,		ranking_	_strategy='v	vorst',
	use_default_protocol=False)				

Evaluate the performance of an embedding model.

The evaluation protocol follows the procedure defined in [BUGD+13] and can be summarised as:

- 1. Artificially generate negative triples by corrupting first the subject and then the object.
- 2. Remove the positive triples from the set returned by (1) positive triples are usually the concatenation of training, validation and test sets.
- 3. Rank each test triple against all remaining triples returned by (2).

With the ranks of both object and subject corruptions, one may compute metrics such as the MRR by calculating them separately and then averaging them out. Note that the metrics implemented in Ampli-Graph's evaluate.metrics module will already work that way when provided with the input returned by evaluate_performance.

The artificially generated negatives are compliant with the local closed world assumption (LCWA), as described in [NMTG16]. In practice, that means only one side of the triple is corrupted at a time (i.e. either the subject or the object).

Note: The evaluation protocol assigns the worst rank to a positive test triple in case of a tie with negatives. This

is the agreed upon behaviour in literature.

Hint: When entities_subset=None, the method will use all distinct entities in the knowledge graph X to generate negatives to rank against. This might slow down the eval. Some of the corruptions may not even make sense for the task that one may be interested in.

For eg, consider the case <Actor, acted_in, ?>, where we are mainly interested in such movies that an actor has acted in. A sensible way to evaluate this would be to rank against all the movie entities and compute the desired metrics. In such cases, where focus us on particular task, it is recommended to pass the desired entities to use to generate corruptions to entities_subset. Besides, trying to rank a positive against an extremely large number of negatives may be overkilling.

As a reference, the popular FB15k-237 dataset has ~15k distinct entities. The evaluation protocol ranks each positives against 15k corruptions per side.

Parameters

- X (ndarray, shape [n, 3]) An array of test triples.
- model (EmbeddingModel) A knowledge graph embedding model
- filter_triples (ndarray of shape [n, 3] or None) The triples used to filter negatives.

Note: When *filtered* mode is enabled (i.e. *filtered_triples* is not None), to speed up the procedure, we use a database based filtering. This strategy is as described below:

- Store the filter_triples in the DB
- For each test triple, we generate corruptions for evaluation and score them.
- The corruptions may contain some False Negatives. We find such statements by quering the database.
- From the computed scores we retrieve the scores of the False Negatives.
- We compute the rank of the test triple by comparing against ALL the corruptions.
- We then compute the number of False negatives that are ranked higher than the test triple; and then subtract this value from the above computed rank to yield the final filtered rank.

Execution Time: This method takes ~4 minutes on FB15K using ComplEx (Intel Xeon Gold 6142, 64 GB Ubuntu 16.04 box, Tesla V100 16GB)

- **verbose** (bool) Verbose mode
- **filter_unseen** (*bool*) This can be set to False to skip filtering of unseen entities if train_test_split_unseen() was used to split the original dataset.
- **entities_subset** (*array-like*) List of entities to use for corruptions. If None, will generate corruptions using all distinct entities. Default is None.
- **corrupt_side** (*string*) Specifies which side of the triple to corrupt:
 - 's': corrupt only subject.
 - 'o': corrupt only object.
 - 's+o': corrupt both subject and object.

's,o': corrupt subject and object sides independently and return 2 ranks. This corresponds to the evaluation protocol used in literature, where head and tail corruptions are evaluated separately.

Note: When corrupt_side='s, o' the function will return 2*n ranks as a [n, 2] array. The first column of the array represents the subject corruptions. The second column of the array represents the object corruptions. Otherwise, the function returns n ranks as [n] array.

- **ranking_strategy** (*string*) Specifies the type of score comparison strategy to use while ranking:
 - 'worst': assigns the worst rank when scores are equal
 - 'best': assigns the best rank when scores are equal
 - 'middle': assigns the middle rank when scores are equal

Our recommendation is to use worst. Think of a model which assigns constant score to any triples. If you use the best strategy then the ranks will always be 1 (which is incorrect because the model has not learnt anything). If you choose this model and try to do knowledge discovery, you will not be able to deduce anything as all triples will get the same scores. So to be on safer side while choosing the model, we would recommend either worst or middle strategy.

- **use_default_protocol** (*bool*) Flag to indicate whether to use the standard protocol used in literature defined in [BUGD+13] (default: False). If set to *True*, corrupt_side will be set to '*s*,*o*'. This corresponds to the evaluation protocol used in literature, where head and tail corruptions are evaluated separately, i.e. in corrupt_side='s,o' mode
- **Returns ranks** An array of ranks of test triples. When corrupt_side='s, o' the function returns [n,2]. The first column represents the rank against subject corruptions and the second column represents the rank against object corruptions. In other cases, it returns [n] i.e. rank against the specified corruptions.

Return type ndarray, shape [n] or [n,2] depending on the value of corrupt_side.

Examples

```
>>> import numpy as np
>>> from ampligraph.datasets import load_wn18
>>> from ampligraph.latent features import ComplEx
>>> from ampligraph.evaluation import evaluate_performance, mrr_score, hits_at_n_
⇔score
>>>
>>> X = load_wn18()
>>> model = ComplEx(batches_count=10, seed=0, epochs=10, k=150, eta=1,
>>>
                    loss='nll', optimizer='adam')
>>> model.fit(np.concatenate((X['train'], X['valid'])))
>>>
>>> filter_triples = np.concatenate((X['train'], X['valid'], X['test']))
>>> ranks = evaluate_performance(X['test'][:5], model=model,
>>>
                                  filter_triples=filter_triples,
>>>
                                 corrupt_side='s+o',
>>>
                                 use_default_protocol=False)
```

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```
>>> ranks
array([ 1, 582, 543, 6, 31])
>>> mrr_score(ranks)
0.24049691297347323
>>> hits_at_n_score(ranks, n=10)
0.4
```

select_best_model_ranking

ampligraph.evaluation.select_best_model_ranking	$X_train,$				
	X_valid,	X_test ,	param_grid,		
	max_combinations=None,				
	param_grid_random_seed=0,				
	use_filter=True,				
	early_stopping=False,				
	early_stopping_params=None,				
use_test_for_selection=H					
	tities_subset=None, corrupt_s				
	o', use_default_protocol=1				
	retrain_be	st_model=Fa	lse, ver-		
	bose=Fals	e)			

Model selection routine for embedding models via either grid search or random search.

For grid search, pass a fixed param_grid and leave max_combinations as *None* so that all combinations will be explored.

For random search, delimit max_combinations to your computational budget and optionally set some parameters to be callables instead of a list (see the documentation for param_grid).

Note: Random search is more efficient than grid search as the number of parameters grows [BB12]. It is also a strong baseline against more advanced methods such as Bayesian optimization [LJ18].

The function also retrains the best performing model on the concatenation of training and validation sets.

Note we generate negatives at runtime according to the strategy described in [BUGD+13].

Note: By default, model selection is done with raw MRR for better runtime performance (use_filter=False).

Parameters

- model_class (*class*) The class of the EmbeddingModel to evaluate (TransE, DistMult, ComplEx, etc).
- X_train (ndarray, shape [n, 3]) An array of training triples.
- **X_valid** (*ndarray*, *shape* [*n*, 3]) An array of validation triples.
- X_test (ndarray, shape [n, 3]) An array of test triples.
- **param_grid** (*dict*) A grid of hyperparameters to use in model selection. The routine will train a model for each combination of these hyperparameters.

Parameters can be either callables or lists. If callable, it must take no parameters and return a constant value. If any parameter is a callable, max_combinations must be set to some value.

For example, the learning rate could either be "lr": [0.1, 0.01] or "lr": lambda: np.random.uniform(0.01, 0.1).

- **max_combinations** (*int*) Maximum number of combinations to explore. By default (None) all combinations will be explored, which makes it incompatible with random parameters for random search.
- **param_grid_random_seed** (*int*) Random seed for the parameters that are callables and random.
- **use_filter** (*bool*) If True, will use the entire input dataset X to compute filtered MRR (default: True).
- **early_stopping** (bool) Flag to enable early stopping (default:False).

If set to True, the training loop adopts the following early stopping heuristic:

- The model will be trained regardless of early stopping for burn_in epochs.
- Every check_interval epochs the method will compute the metric specified in criteria.

If such metric decreases for stop_interval checks, we stop training early.

Note the metric is computed on x_valid . This is usually a validation set that you held out.

Also, because criteria is a ranking metric, it requires generating negatives. Entities used to generate corruptions can be specified, as long as the side(s) of a triple to corrupt. The method supports filtered metrics, by passing an array of positives to x_filter . This will be used to filter the negatives generated on the fly (i.e. the corruptions).

Note: Keep in mind the early stopping criteria may introduce a certain overhead (caused by the metric computation). The goal is to strike a good trade-off between such overhead and saving training epochs.

A common approach is to use MRR unfiltered:

early_stopping_params={x_valid=X['valid'], 'criteria': 'mrr'}

Note the size of validation set also contributes to such overhead. In most cases a smaller validation set would be enough.

• **early_stopping_params** (*dict*) – Dictionary of parameters for early stopping.

The following keys are supported:

- x_valid: ndarray, shape [n, 3]: Validation set to be used for early stopping. Uses X['valid'] by default.
- criteria: criteria for early stopping hits10, hits3, hits1 or mrr. (default)
- x_filter: ndarray, shape [n, 3] : Filter to be used(no filter by default)
- burn_in: Number of epochs to pass before kicking in early stopping(default: 100)
- check_interval: Early stopping interval after burn-in(default:10)

- stop_interval: Stop if criteria is performing worse over n consecutive checks (default: 3)
- use_test_for_selection (bool) Use test set for model selection. If False, uses validation set (default: False).
- **entities_subset** (*array-like*) List of entities to use for corruptions. If None, will generate corruptions using all distinct entities (default: None).
- corrupt_side (string) Specifies which side to corrupt the entities: s is to corrupt only subject. o is to corrupt only object. s+o is to corrupt both subject and object. s, o is to corrupt both subject and object but ranks are computed separately (default).
- **use_default_protocol** (*bool*) Flag to indicate whether to evaluate head and tail corruptions separately(default:False). If this is set to true, it will ignore corrupt_side argument and corrupt both head and tail separately and rank triples i.e. corrupt_side='s,o' mode.
- **retrain_best_model** (*bool*) Flag to indicate whether best model should be re-trained at the end with the validation set used in the search. Default: False.
- **verbose** (*bool*) Verbose mode for the model selection procedure (which is independent of the verbose mode in the model fit).

Verbose mode includes display of the progress bar, logging info for each iteration, evaluation information, and exception details.

If you need verbosity inside the model training itself, change the verbose parameter within the param_grid.

Returns

- **best_model** (*EmbeddingModel*) The best trained embedding model obtained in model selection.
- **best_params** (*dict*) The hyperparameters of the best embedding model *best_model*.
- **best_mrr_train** (*float*) The MRR (unfiltered) of the best model computed over the validation set in the model selection loop.
- ranks_test (*ndarray, shape [n] or [n,2] depending on the value of corrupt_side.*) An array of ranks of test triples. When corrupt_side='s, o' the function returns [n,2]. The first column represents the rank against subject corruptions and the second column represents the rank against object corruptions. In other cases, it returns [n] i.e. rank against the specified corruptions.
- mrr_test (*float*) The MRR (filtered) of the best model, retrained on the concatenation of training and validation sets, computed over the test set.
- **experimental_history** (*list of dict*) A list containing all the intermediate experimental results: the model parameters and the corresponding validation metrics.

Examples

```
>>> from ampligraph.datasets import load_wn18
>>> from ampligraph.latent_features import ComplEx
>>> from ampligraph.evaluation import select_best_model_ranking
>>> import numpy as np
>>>
```

```
>>> X = load_wn18()
>>>
>>> model_class = ComplEx
>>> param_grid = {
        "batches_count": [50],
>>>
        "seed": 0,
>>>
        "epochs": [100],
>>>
>>>
        "k": [100, 200],
        "eta": [5, 10, 15],
>>>
        "loss": ["pairwise", "nll"],
>>>
        "loss_params": {
>>>
>>>
            "margin": [2]
>>>
        },
        "embedding_model_params": {
>>>
>>>
        },
        "regularizer": ["LP", None],
>>>
        "regularizer_params": {
>>>
            "p": [1, 3],
>>>
            "lambda": [1e-4, 1e-5]
>>>
>>>
        },
>>>
        "optimizer": ["adagrad", "adam"],
>>>
        "optimizer_params": {
            "lr": lambda: np.random.uniform(0.0001, 0.01)
>>>
>>>
        },
>>>
        "verbose": False
>>> }
>>> select_best_model_ranking(model_class, X['train'], X['valid'], X['test'],
>>>
                               param_grid,
>>>
                               max_combinations=100,
>>>
                               use_filter=True,
>>>
                               verbose=True,
                               early_stopping=True)
>>>
```

Helper Functions

Utilities and support functions for evaluation procedures.

<pre>train_test_split_no_unseen(X[,</pre>	test_size,	Split into train and test sets.
])		
create_mappings(X)		Create string-IDs mappings for entities and relations.
to_idx(X, ent_to_idx, rel_to_idx)		Convert statements (triples) into integer IDs.

train_test_split_no_unseen

```
ampligraph.evaluation.train_test_split_no_unseen (X, test_size=100, seed=0, al-
low_duplication=False, fil-
tered_test_predicates=None, back-
ward_compatible=False)
```

This function carves out a test set that contains only entities and relations which also occur in the training set.

Parameters

- **X** (*ndarray*, *size*[*n*, *3*]) The dataset to split.
- **test_size** (*int*, *float*) If int, the number of triples in the test set. If float, the percentage of total triples.
- seed (int) A random seed used to split the dataset.
- **allow_duplication** (*boolean*) Flag to indicate if the test set can contain duplicated triples.
- **filtered_test_predicates** (*None*, *list*) If None, all predicate types will be considered for the test set. If list, only the predicate types in the list will be considered for the test set.
- **backward_compatible** (*boolean*) Uses the old (slower) version of the API for reproducibility of splits in older pipelines(if any) Avoid setting this to True, unless necessary. Set this flag only if you want to use the train_test_split_no_unseen of Ampligraph versions 1.3.2 and below. The older version is slow and inefficient

Returns

- X_train (*ndarray*, *size*[*n*, 3]) The training set.
- X_test (*ndarray*, *size[n*, 3]) The test set.

Examples

```
>>> import numpy as np
>>> from ampligraph.evaluation import train_test_split_no_unseen
>>> # load your dataset to X
>>> X = np.array([['a', 'y', 'b'],
                  ['f',
                        'y', 'e'],
>>>
                  ['b', 'y', 'a'],
>>>
                  ['a', 'y', 'c'],
>>>
                  ['c', 'y', 'a'],
>>>
                  ['a', 'y', 'd'],
>>>
                  ['c', 'y', 'd'],
>>>
                  ['b', 'y', 'c'],
>>>
                  ['f', 'y', 'e']])
>>>
>>> # if you want to split into train/test datasets
>>> X_train, X_test = train_test_split_no_unseen(X, test_size=2)
>>> X_train
array([['a', 'y', 'd'],
   ['b', 'y', 'a'],
   ['a',
        'y', 'c'],
   ['f',
        'y', 'e'],
   ['a', 'y', 'b'],
   ['c', 'y', 'a'],
   ['b', 'y', 'c']], dtype='<U1')
>>> X_test
array([['f', 'y', 'e'],
   ['c', 'y', 'd']], dtype='<U1')</pre>
>>> # if you want to split into train/valid/test datasets, call it 2 times
>>> X_train_valid, X_test = train_test_split_no_unseen(X, test_size=2)
>>> X_train, X_valid = train_test_split_no_unseen(X_train_valid, test_size=2)
>>> X_train
array([['a', 'y', 'b'],
   ['a', 'y', 'd'],
   ['a', 'y', 'c'],
```

```
['c', 'y', 'a'],
 ['f', 'y', 'e']], dtype='<U1')
>>> X_valid
array([['c', 'y', 'd'],
 ['f', 'y', 'e']], dtype='<U1')
>>> X_test
array([['b', 'y', 'c'],
 ['b', 'y', 'a']], dtype='<U1')</pre>
```

create_mappings

```
ampligraph.evaluation.create_mappings(X)
```

Create string-IDs mappings for entities and relations.

Entities and relations are assigned incremental, unique integer IDs. Mappings are preserved in two distinct dictionaries, and counters are separated for entities and relations mappings.

Parameters X (*ndarray*, *shape* [*n*, 3]) – The triples to extract mappings.

Returns

- **rel_to_idx** (*dict*) The relation-to-internal-id associations.
- ent_to_idx (*dict*) The entity-to-internal-id associations.

to_idx

ampligraph.evaluation.to_idx (X, ent_to_idx, rel_to_idx)

Convert statements (triples) into integer IDs.

Parameters

- X (ndarray) The statements to be converted.
- ent_to_idx (dict) The mappings between entity strings and internal IDs.
- **rel_to_idx** (*dict*) The mappings between relation strings and internal IDs.

Returns X – The ndarray of converted statements.

Return type ndarray, shape [n, 3]

3.3.4 Discovery

This module includes a number of functions to perform knowledge discovery in graph embeddings.

Functions provided include discover_facts which will generate candidate statements using one of several defined strategies and return triples that perform well when evaluated against corruptions, find_clusters which will perform link-based cluster analysis on a knowledge graph, find_duplicates which will find duplicate entities in a graph based on their embeddings, and query_topn which when given two elements of a triple will return the top_n results of all possible completions ordered by predicted score.

<pre>discover_facts(X, model[, top_n, strategy,])</pre>	Discover new facts from an existing knowledge graph.
<pre>find_clusters(X, model[,])</pre>	Perform link-based cluster analysis on a knowledge
	graph.

Continued on next page

<pre>find_duplicates(X, model[, mode, metric,])</pre>	Find duplicate entities, relations or triples in a graph
	based on their embeddings.
<pre>query_topn(model[, top_n, head, relation,])</pre>	Queries the model with two elements of a triple and re-
	turns the top_n results of all possible completions or-
	dered by score predicted by the model.
find_nearest_neighbours(kge_model, enti-	Return the nearest neighbors of entities.
ties)	

Table 40 – continued from previous page

discover_facts

Discover new facts from an existing knowledge graph.

You should use this function when you already have a model trained on a knowledge graph and you want to discover potentially true statements in that knowledge graph.

The general procedure of this function is to generate a set of candidate statements C according to some sampling strategy strategy, then rank them against a set of corruptions using the *ampligraph.evaluation*. *evaluate_performance()* function. Candidates that appear in the top_n ranked statements of this procedure are returned as likely true statements.

The majority of the strategies are implemented with the same underlying principle of searching for candidate statements:

- from among the less frequent entities ('entity_frequency'),
- less connected entities ('graph_degree', cluster_coefficient'),
- less frequent local graph structures ('cluster_triangles', 'cluster_squares'), on the assumption that densely connected entities are less likely to have missing true statements.
- The remaining strategies ('random_uniform', 'exhaustive') generate candidate statements by a random sampling of entity and relations and exhaustively, respectively.

Warning: Due to the significant amount of computation required to evaluate all triples using the 'exhaustive' strategy, we do not recommend its use at this time.

The function will automatically filter entities that haven't been seen by the model, and operates on the assumption that the model provided has been fit on the data X (determined heuristically), although X may be a subset of the original data, in which case a warning is shown.

The target_rel argument indicates what relation to generate candidate statements for. If this is set to None then all target relations will be considered for sampling.

Parameters

- **X** (*ndarray*, *shape* [*n*, *3*]) The input knowledge graph used to train model, or a subset of it.
- **model** (EmbeddingModel) The trained model that will be used to score candidate facts.
- top_n (*int*) The cutoff position in ranking to consider a candidate triple as true positive.
- **strategy** (*string*) The candidates generation strategy:
 - 'random_uniform' : generates N candidates (N <= max_candidates) based on a uniform sampling of entities.

- 'entity_frequency' : generates candidates by weighted sampling of entities using entity frequency.
- 'graph_degree' : generates candidates by weighted sampling of entities with graph degree.
- 'cluster_coefficient' : generates candidates by weighted sampling entities with clustering coefficient.
- 'cluster_triangles' : generates candidates by weighted sampling entities with cluster triangles.
- 'cluster_squares' : generates candidates by weighted sampling entities with cluster squares.
- max_candidates (*int or float*) The maximum numbers of candidates generated by 'strategy'. Can be an absolute number or a percentage [0,1] of the size of the `X` parameter.
- **target_rel** (*str* or *list* (*str*)) Target relations to focus on. The function will discover facts only for that specific relation types. If None, the function attempts to discover new facts for all relation types in the graph.
- **seed** (*int*) Seed to use for reproducible results.

Returns X_pred – A list of new facts predicted to be true.

Return type ndarray, shape [n, 3]

Examples

```
>>> import requests
>>> from ampligraph.datasets import load_from_csv
>>> from ampligraph.latent_features import ComplEx
>>> from ampligraph.discovery import discover_facts
>>>
>>> # Game of Thrones relations dataset
>>> url = 'https://ampligraph.s3-eu-west-1.amazonaws.com/datasets/GoT.csv'
>>> open('GoT.csv', 'wb').write(requests.get(url).content)
>>> X = load_from_csv('.', 'GoT.csv', sep=',')
>>>
>>> model = ComplEx(batches_count=10, seed=0, epochs=200, k=150, eta=5,
                    optimizer='adam', optimizer_params={'lr':1e-3},
>>>
>>>
                    loss='multiclass_nll', regularizer='LP',
>>>
                    regularizer_params={'p':3, 'lambda':1e-5},
>>>
                    verbose=True)
>>> model.fit(X)
>>>
>>> discover_facts(X, model, top_n=3, max_candidates=20000, strategy='entity_
\hookrightarrow frequency',
>>>
                   target_rel='ALLIED_WITH', seed=42)
array([['House Reed of Greywater Watch', 'ALLIED_WITH', 'Sybelle Glover'],
       ['Hugo Wull', 'ALLIED_WITH', 'House Norrey'],
       ['House Grell', 'ALLIED_WITH', 'Delonne Allyrion'],
       ['Lorent Lorch', 'ALLIED_WITH', 'House Ruttiger']], dtype=object)
```

find clusters

ampligraph.discovery.find_clusters(X, model. *mode='entity'*)

clustering_algorithm=DBSCAN(),

Perform link-based cluster analysis on a knowledge graph.

The clustering happens on the embedding space of the entities and relations. For example, if we cluster some entities of a model that uses k=100 (i.e. embedding space of size 100), we will apply the chosen clustering algorithm on the 100-dimensional space of the provided input samples.

Clustering can be used to evaluate the quality of the knowledge embeddings, by comparing to natural clusters. For example, in the example below we cluster the embeddings of international football matches and end up finding geographical clusters very similar to the continents. This comparison can be subjective by inspecting a 2D projection of the embedding space or objective using a clustering metric.

The choice of the clustering algorithm and its corresponding tuning will greatly impact the results. Please see scikit-learn documentation for a list of algorithms, their parameters, and pros and cons.

Clustering is exclusive (i.e. a triple is assigned to one and only one cluster).

Parameters

- X (ndarray, shape [n, 3] or [n]) The input to be clustered. X can either be the triples of a knowledge graph, its entities, or its relations. The argument mode defines whether X is supposed an array of triples or an array of either entities or relations.
- model (EmbeddingModel) The fitted model that will be used to generate the embeddings. This model must have been fully trained already, be it directly with fit() or from a helper function such as ampligraph.evaluation. select_best_model_ranking().
- clustering_algorithm (object) The initialized object of the clustering algorithm. It should be ready to apply the *fit_predict* method. Please see: scikit-learn documentation to understand the clustering API provided by scikit-learn. The default clustering model is sklearn's DBSCAN with its default parameters.
- mode (*string*) Clustering mode. Choose from:
 - 'entity' (default): the algorithm will cluster the embeddings of the provided entities.
 - 'relation': the algorithm will cluster the embeddings of the provided relations.
 - 'triple' : the algorithm will cluster the concatenation of the embeddings of the subject, predicate and object for each triple.

Returns labels - Index of the cluster each triple belongs to.

Return type ndarray, shape [n]

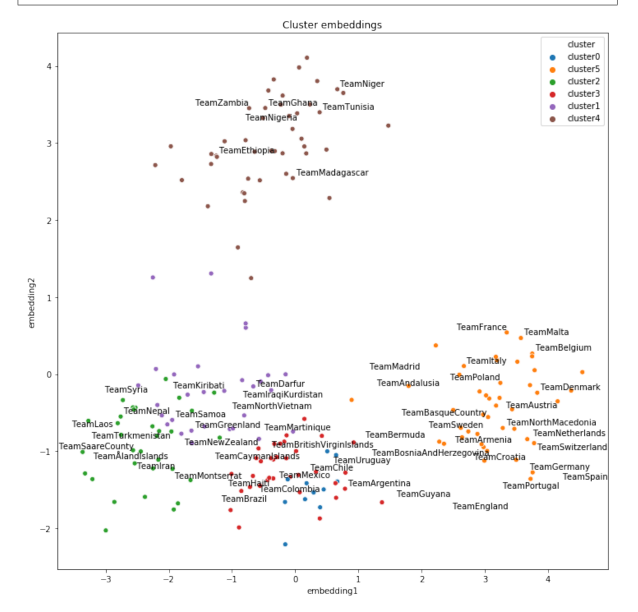
Examples

```
>>> # Note seaborn, matplotlib, adjustText are not AmpliGraph dependencies.
>>> # and must therefore be installed manually as:
>>> #
>>> # $ pip install seaborn matplotlib adjustText
>>>
```

```
>>> import requests
>>> import pandas as pd
>>> import numpy as np
>>> from sklearn.decomposition import PCA
>>> from sklearn.cluster import KMeans
>>> import matplotlib.pyplot as plt
>>> import seaborn as sns
>>>
>>> # adjustText lib: https://github.com/Phlya/adjustText
>>> from adjustText import adjust_text
>>>
>>> from ampligraph.datasets import load_from_csv
>>> from ampligraph.latent_features import ComplEx
>>> from ampligraph.discovery import find_clusters
>>>
>>> # International football matches triples
>>> # See tutorial here to understand how the triples are created from a tabular.
\rightarrow dataset:
>>> # https://github.com/Accenture/AmpliGraph/blob/master/docs/tutorials/
→ClusteringAndClassificationWithEmbeddings.ipynb
>>> url = 'https://ampligraph.s3-eu-west-1.amazonaws.com/datasets/football.csv'
>>> open('football.csv', 'wb').write(requests.get(url).content)
>>> X = load_from_csv('.', 'football.csv', sep=',')[:, 1:]
>>>
>>> model = ComplEx (batches_count=50,
>>>
                    epochs=300,
>>>
                    k=100,
>>>
                    eta=20,
>>>
                    optimizer='adam',
                    optimizer_params={'lr':1e-4},
>>>
                    loss='multiclass_nll',
>>>
>>>
                    regularizer='LP',
>>>
                    regularizer_params={'p':3, 'lambda':1e-5},
>>>
                    seed=0.
>>>
                    verbose=True)
>>> model.fit(X)
>>>
>>> df = pd.DataFrame(X, columns=["s", "p", "o"])
>>>
>>> teams = np.unique(np.concatenate((df.s[df.s.str.startswith("Team")],
>>>
                                       df.o[df.o.str.startswith("Team")])))
>>> team_embeddings = model.get_embeddings(teams, embedding_type='entity')
>>>
>>> embeddings_2d = PCA(n_components=2).fit_transform(np.array([i for i in team_
→embeddings]))
>>>
>>> # Find clusters of embeddings using KMeans
>>> kmeans = KMeans(n_clusters=6, n_init=100, max_iter=500)
>>> clusters = find_clusters(teams, model, kmeans, mode='entity')
>>>
>>> # Plot results
>>> df = pd.DataFrame({"teams": teams, "clusters": "cluster" + pd.
→Series(clusters).astype(str),
                       "embedding1": embeddings_2d[:, 0], "embedding2":...
>>>

→embeddings_2d[:, 1]})

>>>
>>> plt.figure(figsize=(10, 10))
```



Chapter 3. How to Cite

find_duplicates

```
ampligraph.discovery.find_duplicates(X, model, mode='entity', metric='l2', toler-
ance='auto', expected_fraction_duplicates=0.1,
verbose=False)
```

Find duplicate entities, relations or triples in a graph based on their embeddings.

For example, say you have a movie dataset that was scraped off the web with possible duplicate movies. The movies in this case are the entities. Therefore, you would use the 'entity' mode to find all the movies that could de duplicates of each other.

Duplicates are defined as points whose distance in the embedding space are smaller than some given threshold (called the tolerance).

The tolerance can be defined a priori or be found via an optimisation procedure given an expected fraction of duplicates. The optimisation algorithm applies a root-finding routine to find the tolerance that gets to the closest expected fraction. The routine always converges.

Distance is defined by the chosen metric, which by default is the Euclidean distance (L2 norm).

As the distances are calculated on the embedding space, the embeddings must be meaningful for this routine to work properly. Therefore, it is suggested to evaluate the embeddings first using a metric such as MRR before considering applying this method.

Parameters

- X (*ndarray*, *shape* [*n*, 3] or [*n*]) The input to be clustered. X can either be the triples of a knowledge graph, its entities, or its relations. The argument *mode* defines whether X is supposed an array of triples or an array of either entities or relations.
- model (EmbeddingModel) The fitted model that will be used to generate the embeddings. This model must have been fully trained already, be it directly with fit() or from a helper function such as *ampligraph.evaluation*. select_best_model_ranking().
- **mode** (*string*) Choose from:
 - 'entity' (default): the algorithm will find duplicates of the provided entities based on their embeddings.
 - 'relation': the algorithm will find duplicates of the provided relations based on their embeddings.
 - 'triple' : the algorithm will find duplicates of the concatenation of the embeddings of the subject, predicate and object for each provided triple.
- **metric** (*str*) A distance metric used to compare entity distance in the embedding space. See options here.
- **tolerance** (*int* or *str*) Minimum distance (depending on the chosen metric) to define one entity as the duplicate of another. If 'auto', it will be determined automatically in a way that you get the expected_fraction_duplicates. The 'auto' option can be much slower than the regular one, as the finding duplicate internal procedure will be repeated multiple times.
- **expected_fraction_duplicates** (*float*) Expected fraction of duplicates to be found. It is used only when tolerance is 'auto'. Should be between 0 and 1 (default: 0.1).
- **verbose** (*bool*) Whether to print evaluation messages during optimisation (if tolerance is 'auto'). Default: False.

Returns

- **duplicates** (*set of frozensets*) Each entry in the duplicates set is a frozenset containing all entities that were found to be duplicates according to the metric and tolerance. Each frozenset will contain at least two entities.
- **tolerance** (*float*) Tolerance used to find the duplicates (useful in the case of the automatic tolerance option).

Examples

```
>>> import pandas as pd
>>> import numpy as np
>>> import re
>>>
>>> # The IMDB dataset used here is part of the Movies5 dataset found on:
>>> # The Magellan Data Repository (https://sites.google.com/site/anhaidgroup/
⇔projects/data)
>>> import requests
>>> url = 'http://pages.cs.wisc.edu/~anhai/data/784_data/movies5.tar.gz'
>>> open('movies5.tar.gz', 'wb').write(requests.get(url).content)
>>> import tarfile
>>> tar = tarfile.open('movies5.tar.gz', "r:gz")
>>> tar.extractall()
>>> tar.close()
>>>
>>> # Reading tabular dataset of IMDB movies and filling the missing values
>>> imdb = pd.read_csv("movies5/csv_files/imdb.csv")
>>> imdb["directors"] = imdb["directors"].fillna("UnknownDirector")
>>> imdb["actors"] = imdb["actors"].fillna("UnknownActor")
>>> imdb["genre"] = imdb["genre"].fillna("UnknownGenre")
>>> imdb["duration"] = imdb["duration"].fillna("0")
>>>
>>> # Creating knowledge graph triples from tabular dataset
>>> imdb_triples = []
>>>
>>> for _, row in imdb.iterrows():
>>>
        movie_id = "ID" + str(row["id"])
        directors = row["directors"].split(",")
>>>
        actors = row["actors"].split(",")
>>>
        genres = row["genre"].split(",")
>>>
>>>
        duration = "Duration" + str(int(re.sub("\D", "", row["duration"])) // 30)
>>>
>>>
        directors_triples = [(movie_id, "hasDirector", d) for d in directors]
        actors_triples = [(movie_id, "hasActor", a) for a in actors]
>>>
        genres_triples = [(movie_id, "hasGenre", g) for g in genres]
>>>
        duration_triple = (movie_id, "hasDuration", duration)
>>>
>>>
        imdb_triples.extend(directors_triples)
>>>
>>>
        imdb_triples.extend(actors_triples)
>>>
        imdb_triples.extend(genres_triples)
>>>
        imdb_triples.append(duration_triple)
>>>
>>> # Training knowledge graph embedding with ComplEx model
>>> from ampligraph.latent_features import ComplEx
>>>
>>> model = ComplEx (batches_count=10,
```

```
>>>
                    seed=0,
>>>
                    epochs=200,
                    k=150,
>>>
>>>
                    eta=5,
>>>
                    optimizer='adam',
                    optimizer_params={'lr':1e-3},
>>>
                    loss='multiclass_nll',
>>>
>>>
                    regularizer='LP',
                    regularizer_params={'p':3, 'lambda':1e-5},
>>>
                    verbose=True)
>>>
>>>
>>> imdb_triples = np.array(imdb_triples)
>>> model.fit(imdb_triples)
>>>
>>> # Finding duplicates movies (entities)
>>> from ampligraph.discovery import find_duplicates
>>>
>>> entities = np.unique(imdb_triples[:, 0])
>>> dups, _ = find_duplicates(entities, model, mode='entity', tolerance=0.4)
>>> print(list(dups)[:3])
[frozenset({'ID4048', 'ID4049'}), frozenset({'ID5994', 'ID5993'}), frozenset({
→'ID6447', 'ID6448'})]
>>> print(imdb[imdb.id.isin((4048, 4049, 5994, 5993, 6447, 6448))][['movie_name',
movie_name year
4048
             Ulterior Motives 1993
4049
             Ulterior Motives 1993
5993
             Chinese Hercules 1973
5994
             Chinese Hercules 1973
6447 The Stranglers of Bombay 1959
6448 The Stranglers of Bombay 1959
```

query_topn

ampligraph.discovery.query_topn(model, top_n=10, head=None, relation=None, tail=None,

ents_to_consider=None, *rels_to_consider=None*)

Queries the model with two elements of a triple and returns the top_n results of all possible completions ordered by score predicted by the model.

For example, given a <subject, predicate> pair in the arguments, the model will score all possible triples <subject, predicate, ?>, filling in the missing element with known entities, and return the top_n triples ordered by score. If given a <subject, object> pair it will fill in the missing element with known relations.

Note: This function does not filter out true statements - triples returned can include those the model was trained on.

Parameters

- **model** (EmbeddingModel) The trained model that will be used to score triple completions.
- top_n (*int*) The number of completed triples to returned.
- head (string) An entity string to query.
- **relation** (*string*) A relation string to query.

- tail An object string to query.
- ents_to_consider (*array-like*) List of entities to use for triple completions. If None, will generate completions using all distinct entities. (Default: None.)
- **rels_to_consider** (*array-like*) List of relations to use for triple completions. If None, will generate completions using all distinct relations. (Default: None.)

Returns

- X (*ndarray, shape* [n, 3]) A list of triples ordered by score.
- **S** (*ndarray*, *shape* [*n*]) A list of scores.

Examples

```
>>> import requests
>>> from ampligraph.datasets import load_from_csv
>>> from ampligraph.latent_features import ComplEx
>>> from ampligraph.discovery import discover_facts
>>> from ampligraph.discovery import query_topn
>>>
>>> # Game of Thrones relations dataset
>>> url = 'https://ampligraph.s3-eu-west-1.amazonaws.com/datasets/GoT.csv'
>>> open('GoT.csv', 'wb').write(requests.get(url).content)
>>> X = load_from_csv('.', 'GoT.csv', sep=',')
>>>
>>> model = ComplEx (batches_count=10, seed=0, epochs=200, k=150, eta=5,
                    optimizer='adam', optimizer_params={'lr':1e-3}, loss=
>>>
→'multiclass_nll',
>>>
                    regularizer='LP', regularizer_params={'p':3, 'lambda':1e-5},
>>>
                    verbose=True)
>>> model.fit(X)
>>>
>>> query_topn(model, top_n=5,
               head='Catelyn Stark', relation='ALLIED_WITH', tail=None,
>>>
               ents_to_consider=None, rels_to_consider=None)
>>>
>>>
(array([['Catelyn Stark', 'ALLIED_WITH', 'House Tully of Riverrun'],
        ['Catelyn Stark', 'ALLIED_WITH', 'House Stark of Winterfell'],
        ['Catelyn Stark', 'ALLIED_WITH', 'House Wayn'],
        ['Catelyn Stark', 'ALLIED_WITH', 'House Mollen'],
        ['Catelyn Stark', 'ALLIED_WITH', 'Orton Merryweather']],
       dtype='<U44'), array([[10.261374]],
        [ 8.84298 ],
        [ 2.78139 ],
        [ 1.9809164],
        [ 1.833096 ]], dtype=float32))
```

find_nearest_neighbours

```
ampligraph.discovery.find_nearest_neighbours(kge_model, entities, n_neighbors=10, enti-
```

ties_subset=None, metric='euclidean')

Return the nearest neighbors of entities.

The method works in the embedding space and finds a desired number of neighboring embeddings. It can operate from all the entities in the graph or from a subset of interest.

Parameters

- kge_model (ampligraph.latent_features.EmbeddingModel) Trained kge model
- **entities** (*list or np.array*) List of entities whose neighbors need to be found
- n_neighbors (int) number of neighbors to be computed
- entities_subset (list or np.array) List of entities from which neighbors need to be computed. If this list is not passed, all the entities in the graph would be used
- **metric** (*string or callable*) distance metric to be used with NearestNeighbors algorithm For values that can be passed, refer sklearn NearestNeighbors

Returns

- **neighbors** (*np.array of size (len(entities), n_neighbors)*) Each row contains the n_neighbors neighbours of corresponding concepts in entities
- distance (*np.array of size (len(entities), n_neighbors)*) Each row contains distances of corresponding neighbours

Examples

```
>>> model = DistMult(batches_count=2, seed=555, epochs=1, k=10,
                     loss='pairwise', loss_params={'margin': 5},
>>>
                     optimizer='adagrad', optimizer_params={'lr': 0.1})
>>>
>>> X = np.array([['a', 'y', 'b'],
                  ['b', 'y', 'a'],
>>>
                  ['e', 'y', 'c'],
>>>
                  ['c', 'z', 'a'],
>>>
                  ['a', 'z', 'd'],
>>>
                  ['f', 'z', 'g'],
>>>
                  ['c', 'z', 'g']])
>>>
>>> model.fit(X)
>>> neighbors, dist = find_nearest_neighbours(model,
>>>
                                               entities=['b'],
>>>
                                               n_neighbors=3,
                                               entities_subset=['a', 'c', 'd', 'e',
>>>
↔ 'f'])
>>> print(neighbors, dist)
[['e' 'd' 'c']] [[0.97474706 0.979108
                                         1.2323136 ]]
```

3.3.5 Utils

This module contains utility functions for neural knowledge graph embedding models.

Saving/Restoring Models

Models can be saved and restored from disk. This is useful to avoid re-training a model.

<pre>save_model(model[, model_name_path, protocol])</pre>	Save a trained model to disk.
<pre>restore_model([model_name_path])</pre>	Restore a saved model from disk.

save_model

```
ampligraph.utils.save_model (model, model_name_path=None, protocol=4)
Save a trained model to disk.
```

Examples

```
>>> import numpy as np
>>> from ampligraph.latent_features import ComplEx
>>> from ampligraph.utils import save_model
>>> model = Complex(batches_count=2, seed=555, epochs=20, k=10)
>>> X = np.array([['a', 'y', 'b'],
                  ['b', 'y', 'a'],
>>>
                  ['a', 'y', 'c'],
>>>
                  ['c', 'y', 'a'],
>>>
                  ['a', 'y', 'd'],
>>>
                  ['c', 'y', 'd'],
>>>
                  ['b', 'y', 'c'],
>>>
                  ['f', 'y', 'e']])
>>>
>>> model.fit(X)
>>> y_pred_before = model.predict(np.array([['f', 'y', 'e'], ['b', 'y', 'd']]))
>>> example_name = 'helloworld.pkl'
>>> save_model(model, model_name_path = example_name)
>>> print(y_pred_before)
[-0.29721245, 0.07865551]
```

Parameters

- model (EmbeddingModel) A trained neural knowledge graph embedding model, the model must be an instance of TransE, DistMult, ComplEx, or HolE.
- model_name_path (*string*) The name of the model to be saved. If not specified, a default name model with current datetime is named and saved to the working directory

restore_model

See also save_model().

Examples

```
>>> from ampligraph.utils import restore_model
>>> import numpy as np
>>> example_name = 'helloworld.pkl'
>>> restored_model = restore_model(model_name_path = example_name)
>>> y_pred_after = restored_model.predict(np.array([['f', 'y', 'e'], ['b', 'y', 'd
___']]))
>>> print(y_pred_after)
[-0.29721245, 0.07865551]
```

Parameters model_name_path (*string*) – The name of saved model to be restored. If not specified, the library will try to find the default model in the working directory.

Returns model – the neural knowledge graph embedding model restored from disk.

Return type *EmbeddingModel*

Visualization

Functions to visualize embeddings.

```
create_tensorboard_visualizations(model, Export embeddings to Tensorboard.
loc)
```

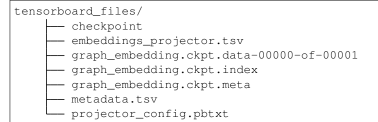
create_tensorboard_visualizations

Export embeddings to Tensorboard.

This function exports embeddings to disk in a format used by TensorBoard and TensorBoard Embedding Projector. The function exports:

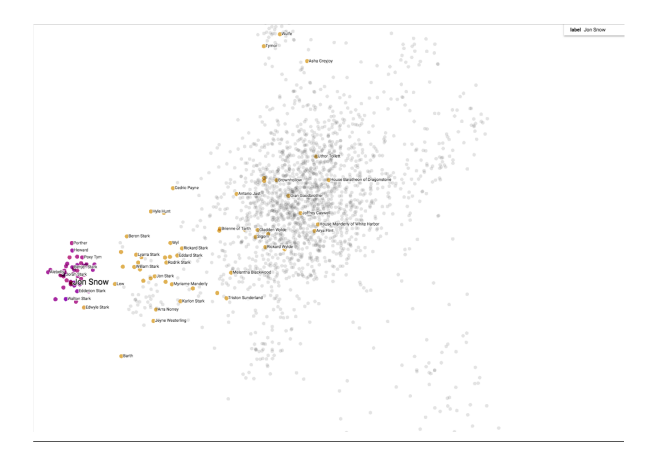
- A number of checkpoint and graph embedding files in the provided location that will allow you to visualize embeddings using Tensorboard. This is generally for use with a local Tensorboard instance.
- a tab-separated file of embeddings embeddings_projector.tsv. This is generally used to visualize embeddings by uploading to TensorBoard Embedding Projector.
- embeddings metadata (i.e. the embeddings labels from the original knowledge graph), saved to metadata.tsv. Such file can be used in TensorBoard or uploaded to TensorBoard Embedding Projector.

The content of loc will look like:



Note: A TensorBoard guide is available at this address.

Note: Uploading embeddings_projector.tsv and metadata.tsv to TensorBoard Embedding Projector will give a result similar to the picture below:



Examples

```
>>> import numpy as np
>>> from ampligraph.latent_features import TransE
>>> from ampligraph.utils import create_tensorboard_visualizations
>>>
>>> X = np.array([['a', 'y', 'b'],
                         'y', 'a'],
>>>
                   ['b',
                         'Y',
                   ['a',
                              'c'],
>>>
                         'Y',
                              'a'],
                   ['c',
>>>
>>>
                   ['a',
                         'y',
                              'd'],
>>>
                   ['c',
                         'v',
                              'd'],
>>>
                   ['b',
                         'y',
                              'c'],
>>>
                   ['f', 'y', 'e']])
>>>
>>> model = TransE(batches_count=1, seed=555, epochs=20, k=10, loss='pairwise',
>>>
                    loss_params={'margin':5})
>>> model.fit(X)
>>>
>>> create_tensorboard_visualizations(model, 'tensorboard_files')
```

Parameters

- model (EmbeddingModel) A trained neural knowledge graph embedding model, the model must be an instance of TransE, DistMult, ComplEx, or HolE.
- **loc** (*string*) Directory where the files are written.

- **labels** (*pd.DataFrame*) Label(s) for each embedding point in the Tensorboard visualization. Default behaviour is to use the embeddings labels included in the model.
- **export_tsv_embeddings** (*bool* (*Default: True*) If True, will generate a tab-separated file of embeddings at the given path. This is generally used to visualize embeddings by uploading to TensorBoard Embedding Projector.
- write_metadata (bool (Default: True)) If True will write a file named 'metadata.tsv' in the same directory as path.

Others

Function to convert a pandas DataFrame with headers into triples.

```
dataframe_to_triples(X, schema) Convert DataFrame into triple format.
```

dataframe_to_triples

Parameters

- X (pandas DataFrame with headers) -
- **schema** (List of (subject, relation_name, object) tuples) where subject and object are in the headers of the data frame

Examples

3.4 How to Contribute

3.4.1 Git Repo and Issue Tracking

AmpliGraph repository is available on GitHub.

A list of open issues is available here.



3.4.2 How to Contribute

We welcome community contributions, whether they are new models, tests, or documentation.

You can contribute to AmpliGraph in many ways:

- · Raise a bug report
- · File a feature request
- Help other users by commenting on the issue tracking system
- Add unit tests
- Improve the documentation
- Add a new graph embedding model (see below)

3.4.3 Adding Your Own Model

The landscape of knowledge graph embeddings evolves rapidly. We welcome new models as a contribution to Ampli-Graph, which has been built to provide a shared codebase to guarantee a fair evaluation and comparison acros models.

You can add your own model by raising a pull request.

To get started, read the documentation on how current models have been implemented.

3.4.4 Developer Notes

Additional documentation on data adapters, AmpliGraph support for large graphs, and further technical details is available here.

3.4.5 Clone and Install in editable mode

Clone the repository and checkout the develop branch. Install from source with pip. use the -e flag to enable editable mode:

```
git clone https://github.com/Accenture/AmpliGraph.git
git checkout develop
cd AmpliGraph
pip install -e .
```

3.4.6 Unit Tests

To run all the unit tests:

\$ pytest tests

See pytest documentation for additional arguments.

3.4.7 Documentation

The project documentation is based on Sphinx and can be built on your local working copy as follows:

```
cd docs
make clean autogen html
```

The above generates an HTML version of the documentation under docs/_built/html.

3.4.8 Packaging

To build an AmpliGraph custom wheel, do the following:

```
pip wheel --wheel-dir dist --no-deps .
```

3.5 Examples

These examples show how to get started with AmpliGraph APIs, and cover a range of useful tasks. Note that additional tutorials are also *available*.

3.5.1 Train and evaluate an embedding model

```
import numpy as np
from ampligraph.datasets import load_wn18
from ampligraph.latent_features import ComplEx
from ampligraph.evaluation import evaluate_performance, mrr_score, hits_at_n_score
def main():
    # load Wordnet18 dataset:
   X = load_wn18()
    # Initialize a Complex neural embedding model with pairwise loss function:
    # The model will be trained for 300 epochs.
   model = Complex(batches_count=10, seed=0, epochs=20, k=150, eta=10,
                    # Use adam optimizer with learning rate 1e-3
                    optimizer='adam', optimizer_params={'lr':1e-3},
                    # Use pairwise loss with margin 0.5
                    loss='pairwise', loss_params={'margin':0.5},
                    # Use L2 regularizer with regularizer weight 1e-5
                    regularizer='LP', regularizer_params={'p':2, 'lambda':1e-5},
                    # Enable stdout messages (set to false if you don't want to
→display)
                    verbose=True)
    # For evaluation, we can use a filter which would be used to filter out
    # positives statements created by the corruption procedure.
    # Here we define the filter set by concatenating all the positives
    filter = np.concatenate((X['train'], X['valid'], X['test']))
    # Fit the model on training and validation set
   model.fit(X['train'],
```

```
early_stopping = True,
              early_stopping_params = \
                      {
                          'x_valid': X['valid'],
                                                     # validation set
                          'criteria':'hits10',
                                                      # Uses hits10 criteria for
→early stopping
                          'burn_in': 100,
                                                       # early stopping kicks in
→after 100 epochs
                          'check_interval':20,
                                                     # validates every 20th epoch
                          'stop_interval':5,
                                                       # stops if 5 successive_
⇔validation checks are bad.
                          'x_filter': filter,
                                                      # Use filter for filtering out.
⇔positives
                          'corruption_entities':'all', # corrupt using all entities
                          'corrupt_side':'s+o' # corrupt subject and object_
\hookrightarrow (but not at once)
                     }
              )
    # Run the evaluation procedure on the test set (with filtering).
    # To disable filtering: filter_triples=None
    # Usually, we corrupt subject and object sides separately and compute ranks
   ranks = evaluate_performance(X['test'],
                                model=model,
                                 filter_triples=filter,
                                use_default_protocol=True, # corrupt subj and obj_
→ separately while evaluating
                                verbose=True)
    # compute and print metrics:
   mrr = mrr_score(ranks)
   hits_10 = hits_at_n_score(ranks, n=10)
   print("MRR: %f, Hits@10: %f" % (mrr, hits_10))
    # Output: MRR: 0.886406, Hits@10: 0.935000
if __name__ == "__main__":
   main()
```

3.5.2 Model selection

```
from ampligraph.datasets import load_wn18
from ampligraph.latent_features import ComplEx
from ampligraph.evaluation import select_best_model_ranking
def main():
    # load Wordnet18 dataset:
    X_dict = load_wn18()
    model_class = ComplEx
    # Use the template given below for doing grid search.
    param_grid = {
```

```
"batches_count": [10],
                     "seed": 0,
                     "epochs": [4000],
                     "k": [100, 50],
                     "eta": [5,10],
                     "loss": ["pairwise", "nll", "self_adversarial"],
                     # We take care of mapping the params to corresponding classes
                     "loss_params": {
                          #margin corresponding to both pairwise and adverserial loss
                          "margin": [0.5, 20],
                          #alpha corresponding to adverserial loss
                         "alpha": [0.5]
                     },
                     "embedding_model_params": {
                          # generate corruption using all entities during training
                          "negative_corruption_entities":"all"
                     },
                     "regularizer": [None, "LP"],
                     "regularizer_params": {
                          "p": [2],
                          "lambda": [1e-4, 1e-5]
                     },
                     "optimizer": ["adam"],
                     "optimizer_params":{
                         "lr": [0.01, 0.0001]
                     },
                     "verbose": True
                 }
    # Train the model on all possibile combinations of hyperparameters.
    # Models are validated on the validation set.
    # It returnes a model re-trained on training and validation sets.
   best_model, best_params, best_mrr_train, \
    ranks_test, mrr_test = select_best_model_ranking(model_class, # Class handle of...
\rightarrow the model to be used
                                                       # Dataset
                                                      X_dict['train'],
                                                      X_dict['valid'],
                                                      X_dict['test'],
                                                       # Parameter grid
                                                      param_grid,
                                                       # Use filtered set for eval
                                                      use_filter=True,
                                                       # corrupt subject and objects
→ separately during eval
                                                      use_default_protocol=True,
                                                       # Log all the model hyperparams.
→and evaluation stats
                                                      verbose=True)
    print(type(best_model).__name__, best_params, best_mrr_train, mrr_test)
if __name__ == "__main__":
   main()
```

3.5.3 Get the embeddings

3.5.4 Save and restore a model

```
import numpy as np
from ampligraph.latent_features import ComplEx
from ampligraph.utils import save_model, restore_model
model = ComplEx(batches_count=2, seed=555, epochs=20, k=10)
X = np.array([['a', 'y', 'b']],
            ['b', 'y', 'a'],
            ['a', 'y', 'c'],
            ['c', 'y', 'a'],
            ['a', 'y', 'd'],
            ['c', 'y', 'd'],
            ['b', 'y', 'c'],
            ['f', 'y', 'e']])
model.fit(X)
# Use the trained model to predict
y_pred_before = model.predict(np.array([['f', 'y', 'e'], ['b', 'y', 'd']]))
print(y_pred_before)
#[-0.29721245, 0.07865551]
# Save the model
example_name = "helloworld.pkl"
save_model(model, model_name_path = example_name)
# Restore the model
restored_model = restore_model(model_name_path = example_name)
# Use the restored model to predict
y_pred_after = restored_model.predict(np.array([['f', 'y', 'e'], ['b', 'y', 'd']]))
print(y_pred_after)
# [-0.29721245, 0.07865551]
```

3.5.5 Split dataset into train/test or train/valid/test

```
import numpy as np
from ampligraph.evaluation import train_test_split_no_unseen
from ampligraph.datasets import load_from_csv
. . .
Assume we have a knowledge graph stored in my_folder/my_graph.csv,
and that the content of such file is:
a,y,b
f,y,e
b,y,a
a,y,c
c,y,a
a, y, d
c,y,d
b,y,c
f,y,e
# Load the graph in memory
X = load_from_csv('my_folder', 'my_graph.csv', sep=',')
# To split the graph in train and test sets:
# (In this toy example the test set will include only two triples)
X_train, X_test = train_test_split_no_unseen(X, test_size=2)
print(X_train)
. . .
X_train:[['a' 'y' 'b']
         ['f' 'y' 'e']
         ['b' 'y' 'a']
         ['c' 'y' 'a']
         ['c' 'y' 'd']
         ['b' 'y' 'c']
         ['f' 'y' 'e']]
. . .
print(X_test)
. . .
X_test: [['a' 'y' 'c']
        ['a' 'y' 'd']]
. . .
# To split the graph in train, validation, and test the method must be called twice:
X_train_valid, X_test = train_test_split_no_unseen(X, test_size=2)
X_train, X_valid = train_test_split_no_unseen(X_train_valid, test_size=2)
print(X_train)
. . .
X_train: [['a' 'y' 'b']
           ['b' 'y' 'a']
           ['c' 'y' 'd']
```

```
['b' 'y' 'c']
['f' 'y' 'e']]
'''
print(X_valid)
'''
X_valid: [['f' 'y' 'e']
['c' 'y' 'a']]
'''
print(X_test)
'''
X_test: [['a' 'y' 'c']
['a' 'y' 'd']]
''''
```

3.5.6 Clustering and projectings embeddings into 2D space

Embedding training

```
import numpy as np
import pandas as pd
import requests
from ampligraph.datasets import load_from_csv
from ampligraph.latent_features import ComplEx
from ampligraph.evaluation import evaluate_performance
from ampligraph.evaluation import mr_score, mrr_score, hits_at_n_score
from ampligraph.evaluation import train_test_split_no_unseen
# International football matches triples
url = 'https://ampligraph.s3-eu-west-1.amazonaws.com/datasets/football.csv'
open('football.csv', 'wb').write(requests.get(url).content)
X = load_from_csv('.', 'football.csv', sep=',')[:, 1:]
# Train test split
X_train, X_test = train_test_split_no_unseen(X, test_size=10000)
# ComplEx model
model = ComplEx(batches_count=50,
               epochs=300,
                k=100,
                eta=20,
                optimizer='adam',
                optimizer_params={'lr':1e-4},
                loss='multiclass_nll',
                regularizer='LP',
                regularizer_params={'p':3, 'lambda':1e-5},
                seed=0,
                verbose=True)
model.fit(X train)
```

Embedding evaluation

```
filter_triples = np.concatenate((X_train, X_test))
ranks = evaluate_performance(X_test,
                              model=model,
                              filter_triples=filter_triples,
                              use_default_protocol=True,
                              verbose=True)
mr = mr_score(ranks)
mrr = mrr_score(ranks)
print("MRR: %.2f" % (mrr))
print("MR: %.2f" % (mr))
hits_10 = hits_at_n_score(ranks, n=10)
print("Hits@10: %.2f" % (hits_10))
hits_3 = hits_at_n_score(ranks, n=3)
print("Hits@3: %.2f" % (hits_3))
hits_1 = hits_at_n_score(ranks, n=1)
print("Hits@1: %.2f" % (hits_1))
. . .
MRR: 0.25
MR: 4927.33
Hits@10: 0.35
Hits@3: 0.28
Hits@1: 0.19
. . .
```

Clustering and 2D projections

Please install lib adjustText first with pip install adjustText. For incf.countryutils, do the following steps:

```
git clone https://github.com/wyldebeast-wunderliebe/incf.countryutils.git
cd incf.countryutils
pip install .
```

incf.countryutils is used to map countries to the corresponding continents.

```
import re
from sklearn.decomposition import PCA
from sklearn.cluster import KMeans
import matplotlib.pyplot as plt
import seaborn as sns
from adjustText import adjust_text
from incf.countryutils import transformations
from ampligraph.discovery import find_clusters
# Get the teams entities and their corresponding embeddings
triples_df = pd.DataFrame(X, columns=['s', 'p', 'o'])
teams = triples_df.s[triples_df.s.str.startswith('Team')].unique()
team_embeddings = dict(zip(teams, model.get_embeddings(teams)))
team_embeddings_array = np.array([i for i in team_embeddings.values()])
# Project embeddings into 2D space via PCA
embeddings_2d = PCA(n_components=2).fit_transform(team_embeddings_array)
```

```
# Cluster embeddings (on the original space)
clustering_algorithm = KMeans(n_clusters=6, n_init=100, max_iter=500, random_state=0)
clusters = find_clusters(teams, model, clustering_algorithm, mode='entity')
# This function maps country to continent
def cn_to_ctn(country):
   try:
       original_name = ' '.join(re.findall('[A-Z][^A-Z]*', country[4:]))
        return transformations.cn_to_ctn(original_name)
   except KeyError:
       return "unk"
plot_df = pd.DataFrame({"teams": teams,
                        "embedding1": embeddings_2d[:, 0],
                        "embedding2": embeddings_2d[:, 1],
                        "continent": pd.Series(teams).apply(cn_to_ctn),
                        "cluster": "cluster" + pd.Series(clusters).astype(str)})
# Top 20 teams in 2019 according to FIFA rankings
top20teams = ["TeamBelgium", "TeamFrance", "TeamBrazil", "TeamEngland", "TeamPortugal
∽",
              "TeamCroatia", "TeamSpain", "TeamUruguay", "TeamSwitzerland",

→ "TeamDenmark",

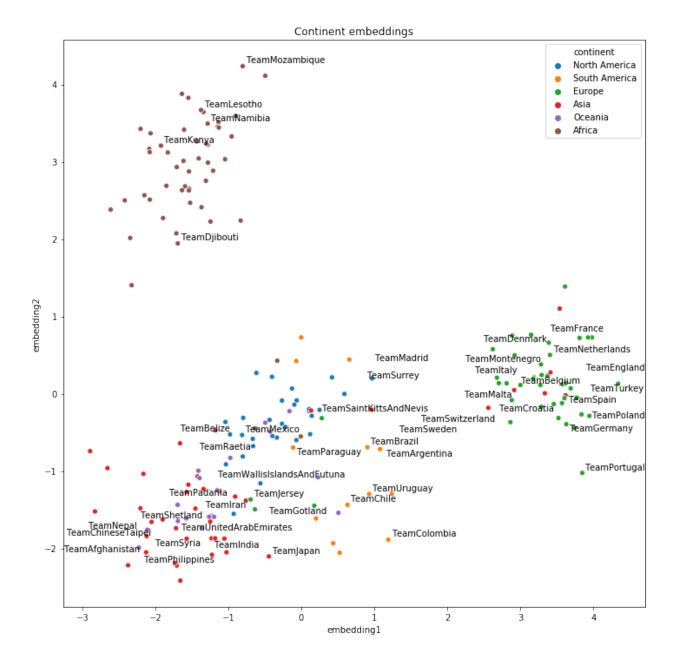
              "TeamArgentina", "TeamGermany", "TeamColombia", "TeamItaly",

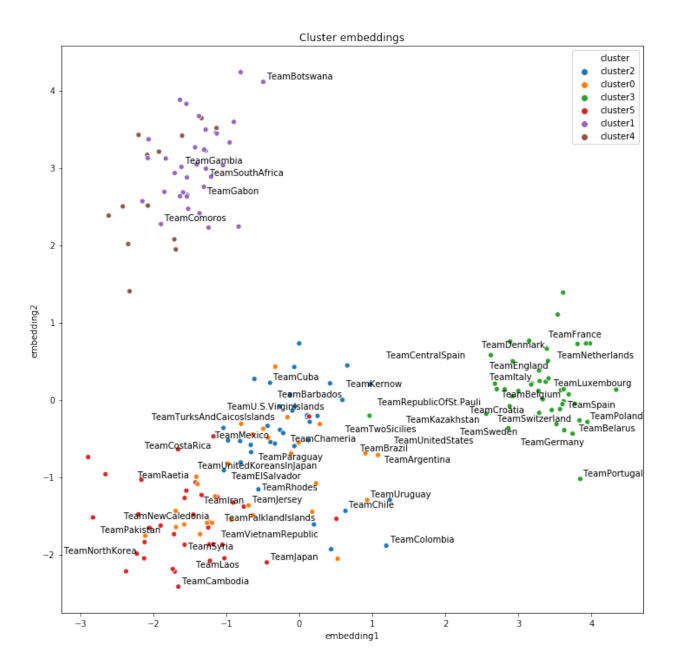
→ "TeamNetherlands",

              "TeamChile", "TeamSweden", "TeamMexico", "TeamPoland", "TeamIran"]
np.random.seed(0)
# Plot 2D embeddings with country labels
def plot_clusters(hue):
   plt.figure(figsize=(12, 12))
   plt.title("{} embeddings".format(hue).capitalize())
   ax = sns.scatterplot(data=plot_df[plot_df.continent!="unk"],
                         x="embedding1", y="embedding2", hue=hue)
   texts = []
   for i, point in plot_df.iterrows():
        if point["teams"] in top20teams or np.random.random() < 0.1:
            texts.append(plt.text(point['embedding1']+0.02,
                         point['embedding2']+0.01,
                         str(point["teams"])))
   adjust_text (texts)
```

Results visualization

```
plot_clusters("continent")
plot_clusters("cluster")
```





3.6 Tutorials

For a comprehensive theoretical and hands-on overview of KGE models and hands-on AmpliGraph, check out our ECAI-20 Tutorial (Slides + Recording + Colab Notebook).

The following Jupyter notebooks will guide you through the most important features of AmpliGraph:

- AmpliGraph basics: training, saving and restoring a model, evaluating a model, discover new links, visualize embeddings. [Jupyter notebook] [Colab notebook]
- Link-based clustering and classification: how to use the knowledge embeddings generated by a graph of international football matches in clustering and classification tasks. [Jupyter notebook] [Colab notebook]

Additional examples and code snippets are available here.

3.7 Performance

3.7.1 Predictive Performance

We report the filtered MR, MRR, Hits@1,3,10 for the most common datasets used in literature.

Results are computed assigning the worst rank to a positive test triple in case of tie with its synthetic negatives. Although this is the most conservative approach, some published literature may adopt an evaluation protocol that assigns the best rank instead. Check out the documentation of *ampligraph.evaluation*. *evaluate_performance()* for details.

Note: On ConvE Evaluation. Results reported in the literature for ConvE are based on the alternative *1-N* evaluation protocol which requires that reciprocal relations are added to the dataset [DMSR18]:

$$D \leftarrow (D, D_{recip})$$
$$D_{recip} \leftarrow \{ (o, p_r, s) | \forall x \in D, x = (s, p, o) \}$$

During training each unique pair of subject and predicate can predict all possible object scores for that pairs, and therefore object corruptions evaluation can be performed with a single forward pass:

ConvE(s, p, o)

In the standard corruption procedure the subject entity is replaced by corruptions:

$$ConvE(s_{corr}, p, o),$$

However in the *1-N* protocol subject corruptions are interpreted as object corruptions of the reciprocal relation:

$$ConvE(o, p_r, s_{corr})$$

To reproduce the results reported in the literature using the *1-N* evaluation protocol, add reciprocal relations by specifying add_reciprocal_rels in the dataset loader function, e.g. load_fb15k(add_reciprocal_rels=True), and run the evaluation protocol with object corruptions by specifying corrupt_sides='o'.

Results obtained with the standard evaluation protocol are labeled ConvE, while those obtained with the *1-N* protocol are marked ConvE(1-N).

3.7.2 FB15K-237

Mode	elMR	MRF	MRR Hits@Hits@Hyperparameters					
Trans	E208	0.31	0.22	0.35	0.50	k: 400; epochs: 4000; eta: 30; loss: multiclass_nll; regularizer: LP;		
						regularizer_params: lambda: 0.0001; p: 2; optimizer: adam; opti-		
						mizer_params: lr: 0.0001; embedding_model_params: norm: 1; normal-		
						ize_ent_emb: false; seed: 0; batches_count: 64;		
Dist-	199	0.31	0.22	0.35	0.49	k: 300; epochs: 4000; eta: 50; loss: multiclass_nll; regularizer: LP;		
Mult						regularizer_params: lambda: 0.0001; p: 3; optimizer: adam; op-		
						timizer_params: lr: 0.00005; seed: 0; batches_count: 50; normal-		
						ize_ent_emb: false;		
Com	· 184	0.32	0.23	0.35	0.50	k: 350; epochs: 4000; eta: 30; loss: multiclass_nll; optimizer: adam; opti-		
plEx						mizer_params: lr: 0.00005; seed: 0; regularizer: LP; regularizer_params:		
						lambda: 0.0001; p: 3; batches_count: 64;		
HolE	184	0.31	0.22	0.34	0.49	k: 350; epochs: 4000; eta: 50; loss: multiclass_nll; regularizer: LP;		
						regularizer_params: lambda: 0.0001; p: 2; optimizer: adam; opti-		
						mizer_params: lr: 0.0001; seed: 0; batches_count: 64;		
Con-	327	0.23	0.15	0.25	0.40	k: 200; epochs: 500; eta: 10; loss: multiclass_nll; loss_params:		
vKB						{} optimizer: adam; optimizer_params: lr: 0.0001; embed-		
						ding_model_params:{ num_filters: 32, filter_sizes: 1, dropout: 0.1};		
						seed: 0; batches_count: 300;		
Conv	E1060	0.26	0.19	0.28	0.38	k: 200; epochs: 4000; loss: bce; loss_params: {label_smoothing=0.1}		
						optimizer: adam; optimizer_params: lr: 0.0001; embed-		
						ding_model_params:{ conv_filters: 32, conv_kernel_size: 3,		
						dropout_embed: 0.2, dropout_conv: 0.1, dropout_dense: 0.3,		
						use_batchnorm: True, use_bias: True}; seed: 0; batches_count:		
						100;		
Conv	EQB4	0.32	0.23	0.35	0.50	k: 200; epochs: 4000; loss: bce; loss_params: {label_smoothing=0.1}		
N)						optimizer: adam; optimizer_params: lr: 0.0001; embed-		
						ding_model_params:{ conv_filters: 32, conv_kernel_size: 3,		
						dropout_embed: 0.2, dropout_conv: 0.1, dropout_dense: 0.3,		
						use_batchnorm: True, use_bias: True}; seed: 0; batches_count:		
						100;		

Note: FB15K-237 validation and test sets include triples with entities that do not occur in the training set. We found 8 unseen entities in the validation set and 29 in the test set. In the experiments we excluded the triples where such entities appear (9 triples in from the validation set and 28 from the test set).

3.7.3 WN18RR

Mod	elMR	MRR	MRR Hits@Hits@Blits@Hyperparameters					
Trans	E2692	0.22	0.03	0.37	0.54	k: 350; epochs: 4000; eta: 30; loss: multiclass_nll; optimizer: adam;		
						optimizer_params: lr: 0.0001; regularizer: LP; regularizer_params:		
						lambda: 0.0001; p: 2; seed: 0; normalize_ent_emb: false; embed-		
						ding_model_params: norm: 1; batches_count: 150;		
Dist-	5531	0.47	0.43	0.48	0.53	k: 350; epochs: 4000; eta: 30; loss: multiclass_nll; optimizer: adam; op-		
Mult						timizer_params: lr: 0.0001; regularizer: LP; regularizer_params: lambda:		
						0.0001; p: 2; seed: 0; normalize_ent_emb: false; batches_count: 100;		
Com	4177	0.51	0.46	0.53	0.58	k: 200; epochs: 4000; eta: 20; loss: multiclass_nll; loss_params: margin:		
plEx						1; optimizer: adam; optimizer_params: lr: 0.0005; seed: 0; regularizer:		
						LP; regularizer_params: lambda: 0.05; p: 3; batches_count: 10;		
HolE	7028	0.47	0.44	0.48	0.53	k: 200; epochs: 4000; eta: 20; loss: self_adversarial; loss_params:		
						margin: 1; optimizer: adam; optimizer_params: lr: 0.0005; seed: 0;		
						batches_count: 50;		
Con-	3652	0.39	0.33	0.42	0.48	k: 200; epochs: 500; eta: 10; loss: multiclass_nll; loss_params:		
vKB						{} optimizer: adam; optimizer_params: lr: 0.0001; embed-		
						ding_model_params:{ num_filters: 32, filter_sizes: 1, dropout: 0.1};		
						seed: 0; batches_count: 300;		
Conv	E5346	0.45	0.42	0.47	0.52	k: 200; epochs: 4000; loss: bce; loss_params: {label_smoothing=0.1}		
						optimizer: adam; optimizer_params: lr: 0.0001; embed-		
						ding_model_params:{ conv_filters: 32, conv_kernel_size: 3,		
						dropout_embed: 0.2, dropout_conv: 0.1, dropout_dense: 0.3,		
						use_batchnorm: True, use_bias: True}; seed: 0; batches_count:		
						100;		
	E (18 42	0.48	0.45	0.49	0.54	k: 200; epochs: 4000; loss: bce; loss_params: {label_smoothing=0.1}		
N)						optimizer: adam; optimizer_params: lr: 0.0001; embed-		
						ding_model_params:{ conv_filters: 32, conv_kernel_size: 3,		
						dropout_embed: 0.2, dropout_conv: 0.1, dropout_dense: 0.3,		
						use_batchnorm: True, use_bias: True}; seed: 0; batches_count:		
						100;		

Note: WN18RR validation and test sets include triples with entities that do not occur in the training set. We found 198 unseen entities in the validation set and 209 in the test set. In the experiments we excluded the triples where such entities appear (210 triples in from the validation set and 210 from the test set).

3.7.4 YAGO3-10

Mod	elMR	MRR	MRR Hits@Hits@Blits@Hyperparameters					
Trans	E1264	0.51	0.41	0.57	0.67	k: 350; epochs: 4000; eta: 30; loss: multiclass_nll; optimizer: adam; op-		
						timizer_params: lr: 0.0001; regularizer: LP; regularizer_params: lambda:		
						0.0001; p: 2; embedding_model_params: norm: 1; normalize_ent_emb:		
						false; seed: 0; batches_count: 100;		
Dist-	1107	0.50	0.41	0.55	0.66	k: 350; epochs: 4000; eta: 50; loss: multiclass_nll; optimizer: adam; op-		
Mult						timizer_params: lr: 5e-05; regularizer: LP; regularizer_params: lambda:		
						0.0001; p: 3; seed: 0; normalize_ent_emb: false; batches_count: 100;		
Com	· 1227	0.49	0.40	0.54	0.66	k: 350; epochs: 4000; eta: 30; loss: multiclass_nll; optimizer: adam; op-		
plEx						timizer_params: lr: 5e-05; regularizer: LP; regularizer_params: lambda:		
						0.0001; p: 3; seed: 0; batches_count: 100		
HolE	6776	0.50	0.42	0.56	0.65	k: 350; epochs: 4000; eta: 30; loss: self_adversarial; loss_params: alpha:		
						1; margin: 0.5; optimizer: adam; optimizer_params: lr: 0.0001; seed: 0;		
						batches_count: 100		
Con-	2820	0.30	0.21	0.34	0.50	k: 200; epochs: 500; eta: 10; loss: multiclass_nll; loss_params:		
vKB						{} optimizer: adam; optimizer_params: lr: 0.0001; embed-		
						ding_model_params:{ num_filters: 32, filter_sizes: 1, dropout: 0.1};		
						seed: 0; batches_count: 3000;		
Conv	E6063	0.40	0.33	0.42	0.53	k: 300; epochs: 4000; loss: bce; loss_params: {label_smoothing=0.1}		
						optimizer: adam; optimizer_params: lr: 0.0001; embed-		
						ding_model_params:{ conv_filters: 32, conv_kernel_size: 3,		
						dropout_embed: 0.2, dropout_conv: 0.1, dropout_dense: 0.3,		
						use_batchnorm: True, use_bias: True}; seed: 0; batches_count:		
-		0.55	0.40	0.60	0.60	300;		
	EQ1741	0.55	0.48	0.60	0.69	k: 300; epochs: 4000; loss: bce; loss_params: {label_smoothing=0.1}		
N)						optimizer: adam; optimizer_params: lr: 0.0001; embed-		
						ding_model_params:{ conv_filters: 32, conv_kernel_size: 3,		
						dropout_embed: 0.2, dropout_conv: 0.1, dropout_dense: 0.3,		
						use_batchnorm: True, use_bias: True}; seed: 0; batches_count:		
						300;		

Note: YAGO3-10 validation and test sets include triples with entities that do not occur in the training set. We found 22 unseen entities in the validation set and 18 in the test set. In the experiments we excluded the triples where such entities appear (22 triples in from the validation set and 18 from the test set).

3.7.5 FB15K

Warning: The dataset includes a large number of inverse relations, and its use in experiments has been deprecated. Use FB15k-237 instead.

Mode	elMR	MRR	MRR Hits@Hits@Hits@Higperparameters					
Trans	E44	0.63	0.50	0.73	0.85			
						optimizer_params: lr: 5e-5; regularizer: LP; regularizer_params: lambda:		
						0.0001; p: 3; embedding_model_params: norm: 1; normalize_ent_emb:		
						false; seed: 0; batches_count: 100;		
Dist-	179	0.78	0.74	0.82	0.86	k: 200; epochs: 4000; eta: 20; loss: self_adversarial; loss_params: mar-		
Mult						gin: 1; optimizer: adam; optimizer_params: lr: 0.0005; seed: 0; normal-		
						ize_ent_emb: false; batches_count: 50;		
Com-	184	0.80	0.76	0.82	0.86	k: 200; epochs: 4000; eta: 20; loss: self_adversarial; loss_params:		
plEx						margin: 1; optimizer: adam; optimizer_params: lr: 0.0005; seed: 0;		
						batches_count: 100;		
HolE	216	0.80	0.76	0.83	0.87	k: 200; epochs: 4000; eta: 20; loss: self_adversarial; loss_params:		
						margin: 1; optimizer: adam; optimizer_params: lr: 0.0005; seed: 0;		
-		0.65				batches_count: 50;		
Con-	331	0.65	0.55	0.71	0.82	k: 200; epochs: 500; eta: 10; loss: multiclass_nll; loss_params:		
vKB						{} optimizer: adam; optimizer_params: lr: 0.0001; embed-		
						ding_model_params:{ num_filters: 32, filter_sizes: 1, dropout: 0.1};		
	D2 0 <i>5</i>	0.50	0.42	0.50	0.66	seed: 0; batches_count: 300;		
Conv	E385	0.50	0.42	0.52	0.66	k: 300; epochs: 4000; loss: bce; loss_params: {label_smoothing=0.1}		
						optimizer: adam; optimizer_params: lr: 0.0001; embed-		
						ding_model_params:{ conv_filters: 32, conv_kernel_size: 3,		
						dropout_embed: 0.2, dropout_conv: 0.1, dropout_dense: 0.3,		
						use_batchnorm: True, use_bias: True}; seed: 0; batches_count:		
Conv	F65	0.80	0.74	0.84	0.89	100; k: 300; epochs: 4000; loss: bce; loss_params: {label_smoothing=0.1}		
N)	сбр-	0.80	0.74	0.84	0.89	optimizer: adam; optimizer_params: lr: 0.0001; embed-		
11)						ding_model_params: { conv_filters: 32, conv_kernel_size: 3,		
						dropout_embed: 0.2, dropout_conv: 0.1, dropout_dense: 0.3,		
						use_batchnorm: True, use_bias: True}; seed: 0; batches_count:		
						100;		
						100,		

3.7.6 WN18

Warning: The dataset includes a large number of inverse relations, and its use in experiments has been deprecated. Use WN18RR instead.

Mode	IMR	MRR	MRR Hits@Hits@Hyperparameters					
Transl	E260	0.66	0.44	0.88	0.95	k: 150; epochs: 4000; eta: 10; loss: multiclass_nll; optimizer: adam;		
						optimizer_params: lr: 5e-5; regularizer: LP; regularizer_params: lambda:		
						0.0001; p: 3; embedding_model_params: norm: 1; normalize_ent_emb:		
						false; seed: 0; batches_count: 100;		
	675	0.82	0.73	0.92	0.95	k: 200; epochs: 4000; eta: 20; loss: nll; loss_params: margin: 1; opti-		
Mult						mizer: adam; optimizer_params: lr: 0.0005; seed: 0; normalize_ent_emb:		
						false; batches_count: 50;		
Com	726	0.94	0.94	0.95	0.95	k: 200; epochs: 4000; eta: 20; loss: nll; loss_params: margin: 1; op-		
plEx						timizer: adam; optimizer_params: lr: 0.0005; seed: 0; batches_count:		
						50;		
HolE	665	0.94	0.93	0.94	0.95	k: 200; epochs: 4000; eta: 20; loss: self_adversarial; loss_params:		
						margin: 1; optimizer: adam; optimizer_params: lr: 0.0005; seed: 0;		
						batches_count: 50;		
Con-	331	0.80	0.69	0.90	0.94	k: 200; epochs: 500; eta: 10; loss: multiclass_nll; loss_params:		
vKB						{} optimizer: adam; optimizer_params: lr: 0.0001; embed-		
						ding_model_params:{ num_filters: 32, filter_sizes: 1, dropout: 0.1};		
	7.40.2	0.02	0.01	0.04	0.05	seed: 0; batches_count: 300;		
Conv	5492	0.93	0.91	0.94	0.95	k: 300; epochs: 4000; loss: bce; loss_params: {label_smoothing=0.1}		
						optimizer: adam; optimizer_params: lr: 0.0001; embed-		
						ding_model_params:{ conv_filters: 32, conv_kernel_size: 3,		
						dropout_embed: 0.2, dropout_conv: 0.1, dropout_dense: 0.3,		
						use_batchnorm: True, use_bias: True}; seed: 0; batches_count:		
ConvE	74176	0.95	0.93	0.95	0.95	100;		
	0d#C	0.93	0.93	0.93	0.93	k: 300; epochs: 4000; loss: bce; loss_params: {label_smoothing=0.1} optimizer: adam; optimizer params: lr: 0.0001; embed-		
N)						optimizer: adam; optimizer_params: lr: 0.0001; embed- ding_model_params:{ conv_filters: 32, conv_kernel_size: 3,		
						dropout_embed: 0.2, dropout_conv: 0.1, dropout_dense: 0.3,		
						use_batchnorm: True, use_bias: True}; seed: 0; batches_count:		
						100;		
						100,		

To reproduce the above results:

```
$ cd experiments
$ python predictive_performance.py
```

Note: Running predictive_performance.py on all datasets, for all models takes ~115 hours on an Intel Xeon Gold 6142, 64 GB Ubuntu 16.04 box equipped with a Tesla V100 16GB. The long running time is mostly due to the early stopping configuration (see section below).

Note: All of the experiments above were conducted with early stopping on half the validation set. Typically, the validation set can be found in X['valid']. We only used half the validation set so the other half is available for hyperparameter tuning.

The exact early stopping configuration is as follows:

- x_valid: validation[::2]
- criteria: mrr
- x_filter: train + validation + test
- stop_interval: 4

- burn_in: 0
- check_interval: 50

Note that early stopping adds a significant computational burden to the learning procedure. To lessen it, you may either decrease the validation set, the stop interval, the check interval, or increase the burn in.

Note: Due to a combination of model and dataset size it is not possible to evaluate Yago3-10 with ConvKB on the GPU. The fastest way to replicate the results above is to train ConvKB with Yago3-10 on a GPU using the hyperparameters described above (~15hrs on GTX 1080Ti), and then evaluate the model in CPU only mode (~15 hours on Intel(R) Xeon(R) CPU E5-2620 v4 @ 2.10GHz).

Note: ConvKB with early-stopping evaluation protocol does not fit into GPU memory, so instead is just trained for a set number of epochs.

Experiments can be limited to specific models-dataset combinations as follows:

3.7.7 Runtime Performance

Training the models on FB15K-237 (k=100, eta=10, batches_count=100, loss=multiclass_nll), on an Intel Xeon Gold 6142, 64 GB Ubuntu 16.04 box equipped with a Tesla V100 16GB gives the following runtime report:

model	seconds/epoch
ComplEx	1.33
TransE	1.22
DistMult	1.20
HolE	1.30
ConvKB	2.83
ConvE	1.13

Note: ConvE is trained with bce loss instead of multiclass_nll.

3.8 Bibliography

3.9 Changelog

3.9.1 1.4.0

26 May 2021

- Added support for numerical attributes on edges (FocusE) (#235)
- Added loaders for benchmark datasets with numeric values on edges (O*NET20K, PPI5K, NL27K, CN15K)
- Added discovery API to find nearest neighbors in embedding space (#240)
- Change of optimizer (from BFSG to Adam) to calibrate models with ground truth negatives (#239)
- 10x speed improvement on train_test_split_unseen API (#242)
- Added support to visualize training progression via tensorboard (#230)
- Bug fix in large graph mode (when evaluate_performance with entities_subset is used) (#231)
- Updated save model api to save embedding matrix > 6GB (#233)
- Doc updates (#247, #221)
- Fixed ntriples loader spurious trailing dot.
- Add tensorboard_logs_path to model.fit() for tracking training loss and early stopping criteria.

3.9.2 1.3.2

25 Aug 2020

- Added constant initializer (#205)
- Ranking strategies for breaking ties (#212)
- ConvE Bug Fixes (#210, #194)
- Efficient batch sampling (#202)
- Added pointer to documentation for large graph mode and Docs for Optimizer (#216)

3.9.3 1.3.1

18 Mar 2020

• Minor bug fix in ConvE (#189)

3.9.4 1.3.0

9 Mar 2020

- ConvE model Implementation (#178)
- Changes to evaluate_performance API (#183)
- Option to add reciprocal relations (#181)

- Minor fixes to ConvKB (#168, #167)
- Minor fixes in large graph mode (#174, #172, #169)
- Option to skip unseen entities checks when train_test_split is used (#163)
- Stability of NLL losses (#170)
- ICLR-20 calibration paper experiments added in branch paper/ICLR-20

3.9.5 1.2.0

22 Oct 2019

- Probability calibration using Platt scaling, both with provided negatives or synthetic negative statements (#131)
- Added ConvKB model
- Added WN11, FB13 loaders (datasets with ground truth positive and negative triples) (#138)
- Continuous integration with CircleCI, integrated on GitHub (#127)
- Refactoring of models into separate files (#104)
- Fixed bug where the number of epochs did not exactly match the provided number by the user (#135)
- Fixed some bugs on RandomBaseline model (#133, #134)
- Fixed some bugs on discover_facts with strategies "exhaustive" and "graph_degree"
- Fixed bug on subsequent calls of model.predict on the GPU (#137)

3.9.6 1.1.0

16 Aug 2019

- Support for large number of entities (#61, #113)
- Faster evaluation protocol (#74)
- New Knowledge discovery APIs: discover facts, clustering, near-duplicates detection, topn query (#118)
- API change: model.predict() does not return ranks anymore
- API change: friendlier ranking API output (#101)
- Implemented nuclear-3 norm (#23)
- Jupyter notebook tutorials: AmpliGraph basics (#67) and Link-based clustering
- Random search for hyper-parameter tuning (#106)
- Additional initializers (#112)
- Experiment campaign with multiclass-loss
- · System-wide bugfixes and minor improvements

3.9.7 1.0.3

7 Jun 2019

- Fixed regression in RandomBaseline (#94)
- Added TensorBoard Embedding Projector support (#86)
- Minor bugfixing (#87, #47)

3.9.8 1.0.2

19 Apr 2019

- Added multiclass loss (#24 and #22)
- Updated the negative generation to speed up evaluation for default protocol.(#74)
- Support for visualization of embeddings using Tensorboard (#16)
- Save models with custom names. (#71)
- Quick fix for the overflow issue for datasets with a million entities. (#61)
- Fixed issues in train_test_split_no_unseen API and updated api (#68)
- Added unit test cases for better coverage of the code(#75)
- Corrupt_sides : can now generate corruptions for training on both sides, or only on subject or object
- Better error messages
- Reduced logging verbosity
- Added YAGO3-10 experiments
- Added MD5 checksum for datasets (#47)
- Addressed issue of ambiguous dataset loaders (#59)
- Renamed 'type' parameter in models.get_embeddings to fix masking built-in function
- Updated String comparison to use equality instead of identity.
- Moved save_model and restore_model to ampligraph.utils (but existing API will remain for several releases).
- Other minor issues (#63, #64, #65, #66)

3.9.9 1.0.1

22 Mar 2019

- · evaluation protocol now ranks object and subjects corruptions separately
- Corruption generation can now use entities from current batch only
- FB15k-237, WN18RR loaders filter out unseen triples by default
- Removed some unused arguments
- Improved documentation
- Minor bugfixing

3.9.10 1.0.0

16 Mar 2019

- TransE
- DistMult
- ComplEx
- FB15k, WN18, FB15k-237, WN18RR, YAGO3-10 loaders
- generic loader for csv files
- RDF, ntriples loaders
- Learning to rank evaluation protocol
- Tensorflow-based negatives generation
- save/restore capabilities for models
- pairwise loss
- nll loss
- self-adversarial loss
- absolute margin loss
- Model selection routine
- LCWA corruption strategy for training and eval
- rank, Hits@N, MRR scores functions

3.10 About

AmpliGraph is developed and maintained by Accenture Labs Dublin.

3.10.1 Contact us

You can contact us by email at about@ampligraph.org.



3.10.2 How to Cite

If you like AmpliGraph and you use it in your project, why not starring the project on GitHub!

If you instead use AmpliGraph in an academic publication, cite as:

```
@misc{ampligraph,
author= {Luca Costabello and
        Sumit Pai and
        Chan Le Van and
        Rory McGrath and
        Nicholas McCarthy and
        Pedro Tabacof},
title = {{AmpliGraph: a Library for Representation Learning on Knowledge Graphs}},
month = mar,
year = 2019,
doi = {10.5281/zenodo.2595043},
url = {https://doi.org/10.5281/zenodo.2595043}
}
```

3.10.3 Contributors

Active contributors (in alphabetical order)

- Luca Costabello
- Adrianna Janik
- Chan Le Van
- Nicholas McCarthy
- Rory McGrath
- Sumit Pai

Past contributors

• Pedro Tabacof

3.10.4 License

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