

# GALE: Active Adversarial Learning for Erroneous Node Detection in Graphs

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**Abstract**—We introduce GALE, an active adversarial learning framework to detect nodes with erroneous information in attributed graphs. GALE is empowered by a new adversarial active error detection framework, which interacts active learning with a graph generative adversarial model to best exploit limited labeled examples of erroneous nodes. It dynamically determines diversified query nodes in batches with bounded size in terms of node typically to enrich a pool of examples, which in turn provide representative examples to best train an adversarial classifier to capture different types of errors. Moreover, GALE provides an annotation algorithm to suggest a context of possible correct attribute values and error types, to facilitate the labeling of query nodes. We show that using limited queries and examples, GALE significantly improves competing methods such as constraint-based detection, outlier detection, and Graph Neural Networks (e.g. GCNs), with 32%, 31%, and 17% gain in F-1 score on average, and is feasible in learning cost for large graphs.

## I. INTRODUCTION

Error detection in attributed graphs is critical for ensuring high-quality graph data and downstream analysis in e.g., knowledge graphs and social networks [37]. In practice, there often exist multiple types of errors in the node attributes. Several methods have been studied to capture erroneous nodes (with incorrect attribute values).

(1) Errors can be defined as violations of data constraints [13], anomalies [31], or inferred with statistical models [34]. These methods can be accurate for specific types of errors, yet may not achieve good recall to capture multiple types of errors [17].  
(2) Node classifiers [17], [36] can be trained from examples of errors, by solving a node classification problem. Another approach is to train individual “base detectors” with ensemble learning [47] to improve the recall. This often requires sufficient training examples from matching class of errors [46]. High-quality labeled examples require heavy manual effort, and remain a luxury for error detection in real-world scenarios.

Consider the example below from a real knowledge graph. Erroneous nodes are chosen from their real editing history<sup>1</sup>.

**Example 1:** Fig. 1 illustrates a fraction of a knowledge graph about films (series) from Wikipedia. A node carries a type (e.g., film) and a set of attributes (e.g., “name”) with values (e.g., “Avengers: Infinity War”). An edge denotes a relationship between nodes (e.g., “subsequent”). There are four erroneous nodes with the type “film” (with the errors marked in red).

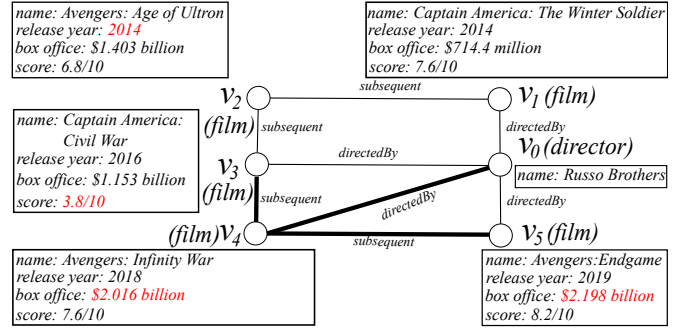


Fig. 1: Detecting Erroneous Nodes in Knowledge Graphs

- Case 1: The Film node  $v_2$  has an incorrect release year “2014”, which should be “2015”;
- Case 2: The Film node  $v_3$  has a wrong rate score “3.8/10” that should be “7.7/10”;
- Case 3: The Film node  $v_4$  has an inaccurate record of \$2.048 billion at the worldwide box office; instead, a correct value should be \$2.016 billion;
- Case 4: Film  $v_5$  makes \$2.798 billion at the worldwide box office instead of \$2.198 billion, which is inaccurate.

Consider the following options.

**Data constraints.** Rules and dependencies that enforce equivalence on node attribute values [13] may not capture Case 1, as the true value “2015” cannot be inferred via node equivalence or is hard to be specified by rules for an individual node. A “negative” rule [6] may specify that “if two films are connected by ‘subsequent’ relation, then their release years must be different”. Nevertheless, either  $v_1$  or  $v_2$  can have a wrong release year “2014”, which cannot be distinguished by the rule due to the vagueness of undirected “subsequent” relation.

**Outlier detection** [31], [38] may capture Case 2 error at node  $v_3$ , given its significantly low value (an outlier) over the attribute “score”. Nevertheless, the errors of the box office (Case 3 and Case 4) at node  $v_4$  and node  $v_5$  cannot be easily captured by an outlier detection process, since both values are in a “normal” range of value distribution.

A single approach may not capture all the errors. Stacking them to a sequential process, or using simple ensemble strategies (e.g., union or voting) may improve the recall but not the overall accuracy due to overlap or false positives [1], [17].

**Train a classifier** [9], [17]. One may train a node classifier to distinguish correct and erroneous nodes given their learned

<sup>1</sup><https://www.wikidata.org/w/index.php?title=Q1765358&action=history>

representation (“node embeddings”). For example, films  $v_5$  and  $v_4$  are “close” in terms of similar embeddings from “score”, “release year” and “box office gross”. A classifier may infer that  $v_4$  is likely to contain errors given  $v_5$  as an example of an erroneous node [9], [17]. Learning-based methods, on the other hand, require a sufficient set of labeled examples. This can be hard to build from scratch.  $\square$

It has been verified that obtaining additional labels with data exploration and active learning [11], [35] can improve error detection especially when training examples are sparse. For example, one may iteratively choose a few nodes to query an oracle (e.g., a data analyst) for a label (“correct” or “error”), and update the classifier to detect errors given the enriched training examples. Although desirable, challenges remain for active learning-based error detection in graphs.

*Query Selection.* Effective active learning often requires proper choices of queries [28], [41], [41]. This is in particular nontrivial for active learning in graph data, as traditional sampling strategies may fail [32]. Moreover, there are often more than one type of errors in real-world graphs [1], [17], [37], [42]. *How to choose a representative set of queries* for the diversified error types to best exploit the oracle, especially with limited or even no labels to start with?

*Labeling Effort.* Most active learning assumes an oracle with a full capability to provide correct labels. In practice, it is not easy to obtain a true label by inspecting the query node and its attribute values alone. The low-quality labels remain a major issue in error detection by crowd-sourcing [10]. In addition, new unknown types of errors may arise in evolving graphs. These limit the application of active learning in leveraging human knowledge for large-scale error detection in graphs.

*Limited Examples.* Even with the new labels obtained from the oracles, the pool of labeled nodes (“training examples”) may still be sparse and even biased. Generative adversarial learning has been verified to be effective for error detection by augmenting training examples with synthetic ones [17]. We are interested in combining active learning and adversarial detection to best exploit the limited amount of examples.

We illustrate a more desirable error detection scenario.

**Example 2:** A desirable approach allows us to “cold start” error detection even when no example is available. One may iteratively choose unlabeled nodes and query an oracle (a validated detection method, or a human expert) and obtain examples to *improve* a node classifier. (1) An initial round captures  $v_3$  as an erroneous node (with e.g., outlier detection). (2) Observe that (a)  $\{v_1, v_3, v_4, v_5\}$  forms a cluster of films directed by the same director, and (b)  $v_4$  is closer to the “center” of the cluster in terms of node similarity. Choosing  $\{v_2, v_4\}$  as a query set improves the decision boundary of the classifier, which in turn distinguish  $v_5$  and  $v_1$  as erroneous and correct nodes, respectively. (3) Moreover, the neighbors of  $v_4$  contain rich information (highlighted in Fig. 1) and can be annotated with error types and statistics to reduce the labeling effort and optimize the choice of follow up queries.  $\square$

These call for an effective detection framework with the following desirable properties. (1) It automatically selects representative query nodes that, if labeled, can best improve a classifier in distinguishing erroneous nodes from correct ones. (2) The query set properly covers diversified classes of errors. As suggested by the above example, the choice should balance both the diversity and the coverage of different types of errors. (3) The oracle is consulted by a *bounded* number of queries. This requires effective exploitation of limited examples via e.g., data augmentation [43]. (4) Moreover, it actively provides useful auxiliary information to help the (human) oracle to derive the labels for “hard” queries. *Can we enable an iterative error detection framework with all these desirable properties?*

**Contribution.** This paper introduces **GALE**, a **Graph Active Learning** framework for **E**rroneous node detection. Our approach is to *improve* a node classifier  $\mathcal{M}$  for error detection, by building an iterative framework that interacts active learning, data augmentation, with adversarial learning that can maximally exploit the labeled examples obtained from an oracle. GALE has the following novel features.

*An Active Adversarial Framework* (Section III and IV). GALE interacts active learning with adversarial learning in a single framework, where a node classifier  $\mathcal{M}$  is continuously improved by examples of erroneous nodes that are obtained and enriched from a query generator. By querying high-quality nodes for training  $\mathcal{M}$ , GALE improves error detection with gained new knowledge. By casting error detection as a “two-players game”, GALE further improves the classifier with augmented examples. These allow GALE to cold start error detection without initial examples, achieve desirable performance in both precision and recall using a small amount of examples, and for capturing multiple types of errors.

*Query Generation with Diversified Typicality.* (Section V). GALE admits high-quality query nodes that are both representative and cover diversified error types. Following the intuition in Example 2, it dynamically generates query nodes in small batches, to continuously improve the decision boundary of the node classifier  $\mathcal{M}$ . We model the query generation as a bi-criteria optimization problem, and introduce a 2-approximation algorithm to approach the best choices of query nodes.

*Query Annotation* (Section VI). GALE also extracts useful auxiliary information and annotates the query nodes and their neighbors. We introduce an annotation algorithm to extract and track such information. These information help users to verify and label the query nodes, understand possible error types, and facilitate further inspection for e.g., data repairing.

*System Evaluation* (Section VII and Section VIII). We have developed a prototype system GALE that nontrivially enhances a pilot prototype [17] with the active adversarial framework. Our experiments using real-world graphs verified the effectiveness of GALE. It improves GEDet with a gain, on average, of 4.98 percentage points and 2.2 percentage points in recall and F-1 score, respectively, with a small batch of queries in only 5 iterations and additional training cost less than 300 seconds,

see details in Fig. 7. Our case study also verified that GALE could suggest useful annotation to facilitate graph data repair.

**Related Work.** We categorize the related work as follows.

*Error detection in graphs.* While error detection for relational data has been extensively studied [21], effective error detection in emerging graph data is much less addressed. Several methods have been proposed to detect errors in graphs. (1) Data dependencies are extended to graphs [13], [23], [30] to capture errors as violations of value constraints, contextualized by graph patterns. (2) Embedding-based methods [9], [17] exploit node and topological features to learn node classifiers to detect errors in graphs. For example, GEDet [17] utilizes semi-supervised generative adversarial networks (GAN) to detect heterogeneous errors and to improve the recall of error detection. PGE [9] jointly leverages textual and graph structures to learn embeddings of knowledge graphs for error detection. Unlike prior methods, GALE is empowered by a generative active adversarial framework, which continuously exploits new knowledge from oracles to improve error detection.

*Graph Representation Learning.* Graph representation learning aims to properly map nodes to a low-dimensional vectors in the embedding space [8], [45]. The embeddings can be used to capture the node similarity that is hard to capture in the original irregular space. Graph neural networks (GNNs) [16] have been verified to be effective for graph representation learning. GNNs take the topological structure and nodal attributes as input and learn encoder functions that aggregates the node’s local neighborhood, with variants such as GCN [25] (which applies a localized first-order approximation of spectral graph convolutions), Graph autoencoders (GAE) [26], [44] (that learn meaningful node embeddings to minimize reconstruction error). Generative adversarial networks (GANs) are used to generate synthetic data that approximates real data distribution by jointly learning a generator  $\mathcal{G}$  and a discriminator  $\mathcal{D}$ . GALE nontrivially interacts GAE to learn node representation for error detection and GAN to enrich examples for few-shot learning. Beyond these enabling techniques, GALE (1) provides new query generation strategy to sample typical and representative nodes for multiple types of errors; (2) exploits annotation component that can help human oracles label the nodes during the active learning process, and (3) suggests correct attribute values for further verification.

*Active Learning.* Active learning has demonstrated its success in error detection for relational data [20]. A main goal is to select small amount of queries to (iteratively) improve model accuracy. Pool-based sampling [28] chooses the best query sample based on evaluation and ranking. Several examples include Random sampling [41] (selects instances uniformly in random), Entropy-based sampling (favors instances with large information entropy), and Margin-based sampling [14] (samples instances that have lowest margin between the two highest softmax outputs of embeddings). It has been observed that traditional strategies may not work well for graph data [32], due to that the choice tends to be biased to “informative” yet

suboptimal queries that are not reflecting true distribution. This is also verified in our tests with multiple type of errors.

The recent Clustering-based sampling has gained attention in the low-budget active learning regime, and has been experimentally verified to outperform random- and uncertainty-based methods (e.g., margin-based sampling and entropy-based sampling) [39]. It picks unlabeled examples that are nearest to the clustering centroids. We proposes a new query selection strategy that maximizes typicality and diversity in terms of both node similarity and topological features, and provide a greedy algorithm that optimizes the query selection with 2-approximation. These are not addressed by prior work.

Recent effort exploits active learning for graph analytics. GraphUCB [11] issues a limited number of queries in the unsupervised setting to detect node anomalies. SEAL [29] proposes a semi-supervised adversarial learning structure for general classification tasks and shows that the graph embedding network and the discriminator can work together to improve the classifiers. These methods are general and cannot be directly applied for error detection, and are not designed to cope with multiple error scenarios in the graphs.

To the best of our knowledge, GALE is the first framework that integrates and interacts GAN, graph representation learning and active learning under a limited query budget for error detection in the graph. In addition, GALE provides auxiliary information to help the human inspect the query node and explore the examples, which is not addressed in prior work.

## II. ERROR DETECTION IN GRAPHS

We start with several notations used by GALE.

**Attributed Graphs.** A graph  $G = (V, E)$  consists of a set of nodes  $V$  and a set of edges  $E \subseteq V \times V$ . Each node  $v \in V$  is a tuple  $(v.A_1, \dots, v.A_n)$  defined on  $n$  attributes, where  $v.A_i = a_i$  ( $i \in [1, n]$ ) denotes that the attribute  $A_i$  of  $v$  has value  $a_i$ .

GALE processes attributed graphs in their feature representation. A *feature representation* of  $G$  is a pair  $G = (X_G, A_G)$ . (1)  $X_G$  is a matrix of node features, where each row  $X_v$  is a vector encoding of a node tuple  $v \in V$ . The encoding can be obtained by e.g., word embedding or one-hot encoding [15]. (2)  $A_G$  is the adjacency matrix of  $G$ .

**Erroneous Nodes.** We assume the existence of a “ground truth” node  $v^*$  for each node  $v \in V$  that carries correct values of all the attributes of  $v$ . A node  $v$  is *erroneous* if there exists an attribute  $A$ , such that  $v.A \neq v^*.A$  ( $v.A$  can be ‘null’ that denotes a missing value). Otherwise,  $v$  is correct.

An *example* denotes a labeled node in  $G$ , which is a pair  $(v, l)$ , where  $l$  is either ‘correct’ for a correct node  $v$ , or ‘error’ if  $v$  is an erroneous node.  $v$  is unlabeled if  $l$  is unknown.

**Error Detection.** We formulate error detection in a graph  $G=(V, E)$  as a node classification problem. The *training nodes*  $V_{\mathcal{T}} = V^e \cup V^c$  ( $V_{\mathcal{T}} \subseteq V$ ) is a set of examples, where  $V^e$  (resp.  $V^c$ ) refers to a set of examples of erroneous (resp. correct) nodes. Given  $G$  and  $V_{\mathcal{T}}$ , the error detection is to train

Symbol	Notation
$G = (V, E)$	Heterogeneous graph with nodes $V$ and edges $E$
$V_T = V^e \cup V^c$	$V_T$ : Labeled training nodes; $V^e$ : erroneous nodes; $V^c$ : correct nodes
$(X_G, A_G)$	feature representation of $G$ : ( $X_G$ : feature matrix; $A_G$ : adjacency matrix)
$\mathbf{h}_n(\mathbf{x}_v)$	the embedding of node $v$ (at layer $n$ )
$H_n$	graph embedding matrix at layer $n$
$\mathcal{G}, \mathcal{D}$	generator and discriminator of SGAN
$\mathcal{L}(\mathcal{G}), \mathcal{L}(\mathcal{D})$	loss function of $\mathcal{G}$ and $\mathcal{D}$
$\mathcal{L}_s, \mathcal{L}_u$	supervised and unsupervised loss of $\mathcal{L}(\mathcal{D})$
$S$	query selector
$\mathcal{A}$	query annotator

TABLE I: Table of Notations

a classifier  $\mathcal{M}$  that accesses  $G$  to infer the labels of a set of unlabeled nodes from  $V \setminus V_T$ .

Given a node classifier model  $\mathcal{M}$ , GALE aims to exploit active learning with an oracle  $\mathcal{O}$  to improve the performance of  $\mathcal{M}$ . To this end, it also uses the following notations.

**Queries.** A *query*  $q$  is an unlabeled node  $(v, \emptyset)$  that requests for the real label of a node  $v$ . Once assigned a label,  $q$  is converted to an example  $(v, \text{'correct'})$  or  $(v, \text{'error'})$ .

**Oracles.** An oracle  $\mathcal{O}$  can be queried and returns the true label  $\mathcal{O}(q)$  of a query  $q$ . In practice, an oracle can be a human expert, or be simulated by invoking and ensembling (with *e.g.*, majority voting) a set of user-defined classifiers called *base detectors*. GALE admits a library of base detectors  $\Psi$ .

### III. FRAMEWORK OVERVIEW

We next provide an overview of GALE framework. We first introduce a principled design that enables the interaction of active learning and adversarial learning for error detection in graphs. We then specify the principles to major modules and models, followed by a general workflow of GALE.

#### A. An Adversarial Active Learning Framework

**Error Detection as “Two-players Game”.** GALE casts error detection into a two-players game between a generator  $\mathcal{G}$  and a discriminator  $\mathcal{D}$ , and derives the node classifier  $\mathcal{M}$  from  $\mathcal{D}$ . It exploits adversarial detection, with the following principle.

- A generator  $\mathcal{G}$  aims to “fool” the discriminator  $\mathcal{D}$  by representing  $G$  with synthetically generated representations that as close to their true counterparts as possible.
- The discriminator  $\mathcal{D}$  meanwhile learns to distinguish nodes with real labels from the synthetic ones from  $\mathcal{G}$ .

A “competition” between  $\mathcal{G}$  and  $\mathcal{D}$  *should* improve the performance of both:  $\mathcal{G}$  learns how to better simulate the distribution of the real labels with synthetic ones, and  $\mathcal{D}$  learns the node embeddings that better discriminates the synthetic errors, real errors, and correct nodes.

**Intuition.** Let us consider a labeled set  $\mathbb{L} = \{(x, y)\}$ , where  $x$  represents a node and  $y \in \{\text{'error'} (y = 1), \text{'correct'} (y = 2)\}$ . GALE introduces a *third type of label* called ‘synthetic error’ ( $y = 3$ ) to annotate nodes with synthetic errors from  $\mathcal{G}$ . We then enforce a learning objective of  $\mathcal{D}$  in the form of:

$$\max_{\mathcal{D}} \mathbb{E}_{x \sim \mathbb{L}} \log P_{\mathcal{D}}(y|x, y \leq 2) + \mathbb{E}_{x \sim p} \log P_{\mathcal{D}}(y \leq 2|x) + \mathbb{E}_{x \sim p_{\mathcal{G}}} \log P_{\mathcal{D}}(3|x) \quad (1)$$

where  $p$  and  $p_{\mathcal{G}}$  refers to the distribution of the real data and its synthetic counterpart from  $\mathcal{G}$ , respectively. The first, second, and third terms are to maximize the log conditional probability for (1) the labeled nodes with true labels (examples), (2) the unlabeled nodes (as either ‘error’ or ‘correct’), and (3) the generated synthetic examples (‘synthetic error’). As such,  $\mathcal{D}$  is able to better distinguish synthetic errors, real errors, and real correct nodes. This yields more accurate classifiers  $\mathcal{M}$ .

**Joint learning scheme.** In a nutshell, GALE improves  $\mathcal{G}$  and  $\mathcal{D}$  in a joint learning process computes a node embedding matrix (derived from the classification layer of  $\mathcal{D}$ ) that minimizes a weighted combination of supervised loss (from examples  $V_T$ ) and an unsupervised (feature matching) loss. A node classifier  $\mathcal{M}$  is yielded by converting the matrix (via a softmax function) to a matrix of class probabilities for the unlabeled nodes in  $G$ . The label with largest probability is chosen for each node (with “error” or “correct” as labels of interests for error detection).

**Graph Augmentation.** This scheme has been verified to be effective for error detection in graphs [17], [18]. The specific scheme is enabled by a *graph augmentation* process [17], which (1) leverages a library  $\Psi$  of base detectors (rules, constraints or string transformations) to inject synthetic errors, and (2) convert the augmented graph to a compact featurized encoding (Section II) by learning a graph autoencoder GAE of  $G$ , yielding representations of real data  $X_R$  and synthetic data  $X_S$ , respectively. The generator and discriminator then play a competitive game over the compact encodings to improve the decision boundary by specifying and optimizing the above objective. GALE adapts the above scheme to iteratively update the discriminator  $\mathcal{D}$  and the classifier  $\mathcal{M}$  upon receiving new examples from the oracle  $\mathcal{O}$  (see Section IV).

**Active Adversarial Detection.** Despite its effectiveness [17], [18], the above scheme requires sufficient high-quality examples. Moreover, the learning of  $\mathcal{M}$  is a “one-shot” process, and cannot evolve upon new error types. To adapt  $\mathcal{M}$  to sparse, biased and possibly new examples, GALE interacts active learning and adversarial learning in an *iterative* framework.

(1) Given an oracle  $\mathcal{O}$ , graph  $G$ , and a current classifier  $\mathcal{M}$ , it chooses from unlabeled nodes a set of queries and consults  $\mathcal{O}$  to obtain their labels. The selection of the queries follows a general strategy that favors nodes that properly represents multiple error types in  $G$ . This enriches a pool of examples that are in turn used by the adversarial detection scheme. GALE then updates the discriminator  $\mathcal{D}$  in the “two-players game”, and derives the classifier  $\mathcal{M}$  accordingly from  $\mathcal{D}$ .

(2) The learned classifier  $\mathcal{M}$  is applied to detect new erroneous nodes in  $G$ . This in turn improves the quality of future query selection, mitigating the bias of active learning [32].

The above steps runs in multiple iterations. As such, GALE allows us to query the oracle and gain new knowledge that

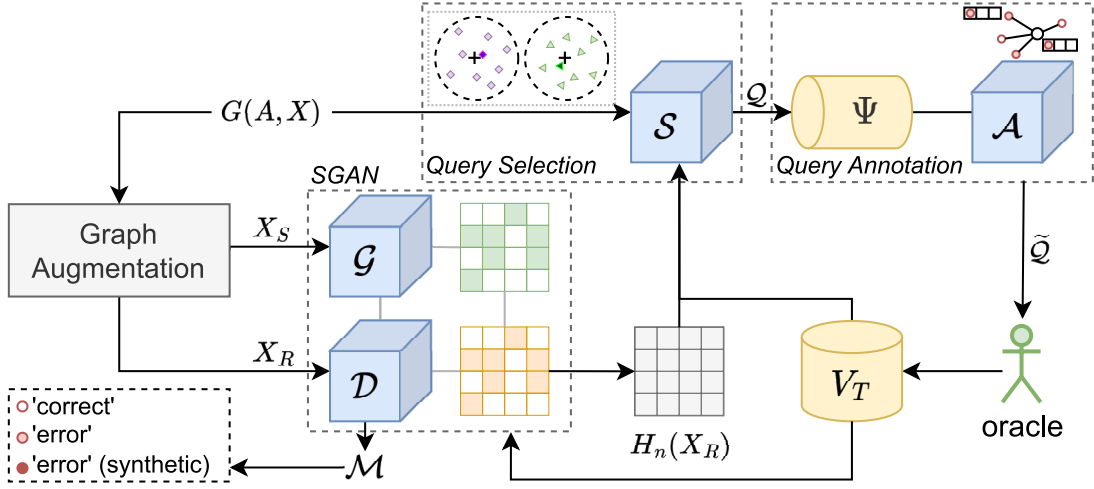


Fig. 2: Overview of GALE Framework

approach the real distribution of unknown erroneous nodes, thus improving the accuracy of error detection by continuously refining the decision boundary of  $\mathcal{M}$ .

### B. Modules and Models

We next introduce the major modules in GALE. These modules and associated models specify the principled adversarial active framework into an *iterative* error detection process.

**Semi-supervised Adversarial Module.** *How to maintain and improve a node classifier  $\mathcal{M}$  for error detection as “two-players game”?* GALE specifies the adversarial training of  $\mathcal{M}$  in an *iterative semi-supervised* generative adversarial module (SGAN). This module is responsible for dynamically maintaining  $\mathcal{M}$  upon new examples from the oracle  $\mathcal{O}$  for error detection. Given a set of new examples, it works in two stages.

**Initialization.** SGAN initializes the joint learning of  $\mathcal{G}$  and  $\mathcal{D}$  from scratch, and derives the classifier  $\mathcal{M}$  from  $\mathcal{D}$ . SGAN takes as input the node representation from graph  $G$  that includes synthetic erroneous node features  $X_S$  and real example features  $X_R$  via the graph augmentation process (see Section IV). The classification layer of  $\mathcal{D}$  predicts the class label for unlabeled nodes (“real” or “error”).

**Update  $\mathcal{D}$ .** Upon receiving changes in  $X_R$  due to the new examples obtained from the oracle  $\mathcal{O}$ , SGAN incrementally updates  $\mathcal{D}$  by updating its parameters in response to the changed examples  $V_T^i$ .  $\mathcal{M}^i$  is derived accordingly.

That is,  $\mathcal{D}^i$  is maintained at iterations  $i$  as follows:

$$\begin{aligned} (\mathcal{G}, \mathcal{D}^0) &= \text{SGAN}(G, V_T^0, X_R, X_S); \\ \mathcal{D}^i &= \text{SGAN}(G, V_T^i, X_R, X_S) \end{aligned}$$

The module also forwards  $H_n^i(X_R)$ , the node embeddings learned on an intermediate  $n$ -th layer of the discriminator  $\mathcal{D}$  to Query Selection module  $\mathcal{Q}$  to facilitate the selection of queries from unlabeled nodes in future iterations.

**Query Selection Module.** *How to select a small set of queries to improve error detection?* GALE aims to improve  $\mathcal{M}$  by

obtaining additional labels from oracle  $\mathcal{O}$ . The query selection module utilizes a Query Selector  $\mathcal{S}$  to generate promising queries from the unlabeled training node set, and query  $\mathcal{Q}$  for additional true labels. The model  $\mathcal{S}$  initializes  $k$  queries with clustering-based sampling [39] from unlabeled nodes. Here  $k$  is a *local budget* that specifies the number of queries allowed per iteration. In the follow-up iteration  $i$ , it exploits the node embeddings  $H_n^{i-1}(X_R)$  learned by  $\mathcal{D}^{i-1}$ , and examples  $V_T^{i-1}$  from the last iteration, and returns query set  $\mathcal{Q}^i$  with bounded size  $k$ , with a goal to represent diversified errors.

$$\begin{aligned} \mathcal{Q}^0 &= \mathcal{S}(\emptyset, \emptyset, G, k); \\ \mathcal{Q}^i &= \mathcal{S}(H_n^{i-1}(X_R), V_T^{i-1}, G, k) \end{aligned}$$

The queries are sent to be enriched with available auxiliary information (see Query Annotation Module), assigned with true labels (thus converted to examples  $\Delta V_T$ ), and featurized to enrich the example features  $X_R$  to improve  $\mathcal{M}$ .

**Query Annotation Module.** *How to adapt active adversarial framework for iterative error detection?* To escape from locally biased choices [32], GALE dynamically chooses promising queries by extracting and aggregating useful annotated information of the queries  $\mathcal{Q}$ , and enrich  $\mathcal{Q}$  to an *annotated* counterpart  $\tilde{\mathcal{Q}}$ . By default, GALE collects and encodes the following information, whenever applicable: 1-hop neighborhood induced subgraph of  $\mathcal{Q}$ , most influential labeled nodes to  $\mathcal{Q}$ , degree assortativity [32], erroneous attribute values detected by base detectors from  $\Psi$  [17], among others (see Section VI). The benefit is twofold: it provides useful local and global structural information of query nodes in  $G$  that facilitate the oracle  $\mathcal{O}$  for labeling, and allow query selector  $\mathcal{S}$  to re-estimate the node importance for dynamic selection in future iterations.

Specifically, a query annotator  $\mathcal{A}$  provides  $\mathcal{O}$  and  $\mathcal{S}$  with annotated  $\tilde{\mathcal{Q}}$  in each iteration as follows:

$$\begin{aligned} \tilde{\mathcal{Q}}^0 &= \mathcal{A}(\mathcal{Q}^0, \Psi, G); \\ \tilde{\mathcal{Q}}^i &= \mathcal{A}(\mathcal{Q}^i, \Psi, G) \end{aligned}$$

The module also naturally augments the examples with the new ones by querying the oracle  $\mathcal{O}$  with the queries  $\mathcal{Q}$ :

$$\begin{aligned} V_T^0 &= \mathcal{O}(\tilde{\mathcal{Q}}^0); \\ V_T^i &= V_T^{i-1} \cup \mathcal{O}(\tilde{\mathcal{Q}}^i) \end{aligned}$$

### C. Workflow of GALE

Putting the modules together, we next describe the workflow of GALE (illustrated in Fig. 2, with an algorithmic description of the learning process in Fig. 3). It works with a graph  $G$ , an oracle  $\mathcal{O}$ , and (optionally) a library of base detectors  $\Psi$ .

**Initialization.** GALE “cold starts” the framework by initializing a set of queries  $\mathcal{Q}$  with annotated information from  $\Psi$ , which provides an initial set of detected errors in attributes, whenever applicable (lines 1-2). It then consults an initial verification of  $\mathcal{Q}$  with oracle  $\mathcal{O}$  (line 3) to initialize the examples. It also invokes a procedure `GAugment` to preprocess  $G$  and generates the feature representation of real (resp. synthetic) examples  $X_R$  (resp.  $X_S$ ) following [17], to enable the adversarial learning of the generator  $\mathcal{G}$  and discriminator  $\mathcal{D}$ . This yields an initial node classifier  $\mathcal{M}$  (lines 5-6).

**Iterative Improvement.** At iteration  $i$ , GALE interacts the modules as follows. (1) With the learned embeddings  $H_n(X_R)$  of the real examples from  $\mathcal{D}$  in the last round, it selects a representative query set  $\mathcal{Q}^i$  (line 8), and enrich the queries with their auxiliary annotated information (line 9). (2) It consults the oracle  $\mathcal{O}$  with the batch of new queries and augment the examples  $V_T$ . (line 10). (3) The adversarial learning is then activated to update  $\mathcal{D}$  with newly updated examples; The classifier  $\mathcal{M}$  along with the learned embeddings are then updated accordingly (lines 11-12). Here the procedure `SGAND` is a variant of `SGAN`. Instead of retraining both  $\mathcal{G}$  and  $\mathcal{D}$  from scratch, it incrementally updates  $\mathcal{D}$  in response to the necessary change of the examples  $V_T$  (see Section IV).

The above process repeats up to (user-defined)  $T$  times, and issues at most  $T \cdot k$  queries. The improved classifier  $\mathcal{M}$  is then returned (line 13).

We next introduce the details of iterative semi-supervised GAN module, query selection module and annotation module, in Sections IV, V, and VI, respectively.

## IV. ITERATIVE ADVERSARIAL DETECTION

SGAN enforce  $\mathcal{G}$  and  $\mathcal{D}$  to compete in a two player’s game, yet assumes fixed examples. We formulate an optimization problem to minimize a bi-criteria loss in an *iterative* semi-supervised setting. We use the following construction.

**Loss Function of  $\mathcal{D}$ .** To adapt the discriminator  $\mathcal{D}$  to recognize new erroneous nodes and differentiate real and synthetic labels, we quantify the loss of  $\mathcal{D}$  as  $\mathcal{L}^i(\mathcal{D}) = \mathcal{L}_s^i + \lambda \mathcal{L}_u$ , where (1) a *supervised loss*  $\mathcal{L}_s^i$  quantifies the loss of accuracy on node label classification (‘error’ or ‘correct’) for a *current* set of examples  $V_T^i$ ; and (2) an *unsupervised loss*  $\mathcal{L}_u$  quantifies the loss of accuracy on distinguishing synthetic or real labels.

### Algorithm GALE

*Input:*  $G(V, E)$ , library  $\Psi$ , integers  $k$  and  $T$ ;

*Output:* (improved) SGAN classifier  $\mathcal{M}$ .

1. set  $V_T := \emptyset$ ; set  $\mathcal{Q} := \emptyset$ ; integer  $i := 1$ ;  
matrix  $X_S := \emptyset$ , matrix  $X_R := \emptyset$ ; matrix  $H_n := \emptyset$ ;  
/\* preprocess  $G$  and initialize models \*/
2.  $\mathcal{Q} := \mathcal{S}(\emptyset, \emptyset, G, k)$ ;  $\tilde{\mathcal{Q}} := \mathcal{A}(\mathcal{Q}, \Psi, G)$ ;
3.  $V_T := \mathcal{O}(\tilde{\mathcal{Q}})$ ;
4.  $(X_R, X_S) := \text{GAugment}(G, \Psi)$ ;
5.  $(\mathcal{G}, \mathcal{D}) := \text{SGAN}(G, V_T, X_R, X_S)$ ;
6. derive an initial classifier  $\mathcal{M}$  and  $H_n$  from discriminator  $\mathcal{D}$ ;  
/\* iteratively improve the classifier \*/
7. **while**  $i < T$  **do**
8. set  $\mathcal{Q}^i := \mathcal{S}(H_n(X_R), V_T, G, k)$ ;
9. set  $\tilde{\mathcal{Q}}^i := \mathcal{A}(\mathcal{Q}^i, \Psi, G)$ ;
10. set  $V_T := V_T \cup \mathcal{O}(\tilde{\mathcal{Q}}^i)$ ;
11.  $\mathcal{D}^i := \text{SGAND}(G, V_T, X_R, X_S)$ ;
12. update  $\mathcal{M}$  and  $H_n$  from  $\mathcal{D}^i$ ;
13. **return** updated  $\mathcal{M}$ ;

Fig. 3: GALE: Learning Framework

Unlike [17] that assume a fixed learning objective, we guide the learning of  $\mathcal{D}$  to minimize the supervised and unsupervised loss given the current examples and latest error distribution.

**Supervised Loss.** We define the supervised loss as:

$$\mathcal{L}_s^i = \sum_{v \in V_T^i} \ell(f_\theta(x_v), l_v)$$

where  $V_T^i$  is the  $i$ -th batch of the examples obtained from the oracle  $\mathcal{O}$ , with  $l_v$  the ground truth label of query  $v$ .  $\theta$  is the learnable parameter and  $f_\theta(x_v)$  is the prediction of node  $v$ .  $\ell(\cdot, \cdot)$  measures the difference between prediction and ground truth label using e.g., cross entropy.

**Unsupervised Loss.** The discriminator  $\mathcal{D}$  also aims to classify the real or synthetic examples as accurately as possible by minimizing the unsupervised loss  $\mathcal{L}_u$ . We define  $\mathcal{L}_u$  as:

$$\mathcal{L}_u = -\mathbb{E}_{x_v \sim X_R}[\log(1 - \mathcal{D}(x_v))] - \mathbb{E}_{x_v \sim \mathcal{G}(X_S)}[\log \mathcal{D}(x_v)]$$

where  $\mathcal{D}(x_v)$  is the predicted synthetic probability of  $v$ .

**Loss Function of  $\mathcal{G}$ .** The generator  $\mathcal{G}$  aims to minimize the difference between the synthetic graph embeddings  $X_S$  and the real counterpart from  $X_R$ . We define the loss function  $L(\mathcal{G})$  of  $\mathcal{G}$  to measure the feature matching loss [40].

$$L(\mathcal{G}) = \|\mathbb{E}_{\mathbf{x}_v \in X_R} \mathbf{h}(\mathbf{x}_v) - \mathbb{E}_{\mathbf{x}'_v \in X_S} \mathbf{h}(\mathcal{G}(\mathbf{x}'_v))\|_2^2$$

where  $\mathbb{E}(\cdot)$  computes the expected value of a graph feature matrix. The generator is trained to match the expected value of the embeddings on an intermediate layer of  $\mathcal{D}$  between the features of real examples  $\mathbf{h}(\mathbf{x}_v)$  and synthetic erroneous counterpart  $\mathbf{h}(\mathcal{G}(\mathbf{x}'_v))$ . Intuitively,  $\mathcal{G}$  has higher chance to fool the discriminator  $\mathcal{D}$  by faking node embeddings closer to the real counterpart, by minimizing  $L(\mathcal{G})$ .

$\mathcal{G}$  and  $\mathcal{D}$  consist of a sequence of transpose convolution and batch normalization layers, adding regularization layers e.g., dropout layers to prevent overfitting.  $\mathcal{D}$  takes real feature



---

**Procedure: SGAN** ( $G, V_T, X_R, X_S$ )

1. **initialize** the parameters of  $\mathcal{G}$  and  $\mathcal{D}$ ;
2. update  $\mathcal{L}(\mathcal{D})$  with  $V_T, X_R$ , and  $X_S$ ;
3. **while**  $\mathcal{G}$  and  $\mathcal{D}$  do not reaches a Nash equilibrium **do**
4.   update  $\mathcal{G}(\cdot)$  by descending gradients of losses:
5.    $\nabla_{\theta_{\mathcal{G}}} L(\mathcal{G})$
6.   update  $\mathcal{D}(\cdot)$  by descending gradients of losses:
7.    $\nabla_{\theta_{\mathcal{D}}} L(\mathcal{D})$
8.   reduce learning rate  $\beta$
9. **return**  $\mathcal{D}$ ;

**Procedure: SGAND** ( $G, V_T, X_R, X_S$ )

1. update  $\mathcal{L}^i(\mathcal{D})$  with  $V_T, X_R$ , and  $X_S$ ;
  2. Update  $\mathcal{D}(\cdot)$  by descending gradients of losses:
  3.    $\nabla_{\theta_{\mathcal{D}}} L(\mathcal{D})$
  4. **return**  $\mathcal{D}$ ;
- 

Fig. 4: Procedure SGAN and SGAND

matrix  $X_R$  and its synthetic counterpart  $X_G$  that is generated from  $\mathcal{G}$  as inputs. Letting  $\mathbf{h}_n(\mathbf{x}_v)$  be the embeddings on an intermediate layer of  $\mathcal{D}$ .  $H_n^i(X_R)$  refers to the learned real node embeddings for real feature matrix  $X_R$  at iteration  $i$ . The matrix  $H_n^i(X_R)$  is then passed to Query Selection module.

**Procedures SGAN and SGAND.** GALE trains generator  $\mathcal{G}$  and discriminator  $\mathcal{D}$  by iteratively minimizing their corresponding losses in procedure GANDet, as shown as Fig. 4. The main training loop (lines 3-8) jointly optimize the generator loss  $\mathcal{L}(\mathcal{G})$  and discriminator loss  $\mathcal{L}(\mathcal{D})$ . We optimize the gradients by applying Adam [24] and update  $\mathcal{D}$  and  $\mathcal{G}$ .

We distinguish SGAND with SGAN, a variant that follows the joint learning scheme as in SGAN but only maintains the discriminator  $\mathcal{D}$ , by iteratively minimizing the corresponding supervised loss. It *incrementalizes* the learning of SGAN, by performing necessary updates to  $\mathcal{D}$  in response to the change of the *supervised loss*  $\mathcal{L}^i(\mathcal{D})$  (line 1) due to enriched  $V_T$ .

## V. QUERY SELECTION MODULE

We next describe the query selection strategy of GALE. We first introduce a notion of *diversified typically* to measure the quality of the queries in terms of “typically” and diversity. We then study the query selection problem, and provide an approximation algorithm to implement the query selector  $\mathcal{S}$ .

### A. Diversified Typicality

**Clustering Typicality.** The *typically* of a query in active learning strategies [19], [39] measures the quality of a query in terms of its closeness to the centroid of a proper clustering. An intuition is that when the clustering is treated as a density function mapping, the data point that is closer to the centroid indicates that it falls into a high density region that is easier to be classified. It has been observed that such queries [19], [39] achieves good performance especially in the low-budget regime (where only limited queries are allowed). Example 2 consistently illustrates this intuition. Based on this intuition, we define the *clustering typicality* of a node  $v$  as follows:

$$\text{clusT}(v) = (\|\mathbf{h}(v) - c(v)\|_2)^{-1}$$

where (1)  $c(v)$  is the centroid of the cluster that  $v$  belongs to, where the clustering is performed on the node embedding space, and (2)  $\mathbf{h}(v)$  refers to the embedding of  $v$ . The embedding is readily obtained from  $H_n^i(X_R)$  at the  $i$ -th round of execution of SGAND (line 11 of Fig. 3). Intuitively, it computes the inverse of the Euclidean distance of the node embeddings between  $v$  and its cluster centroid.

**Topological Typicality.** As observed in [7], [32], the topology influence also plays an important role for query selection especially for graph learning. One intuition is that the “locations” of a labeled node can influence how information spreads over the graph. We want to select nodes that whose relative topological position is “close” to the class center.

We introduce a second measure of *topological typicality* (denoted as  $\text{topoT}(v)$ ) of a node  $v$ . We quantify this with the probability that a node  $v$  *does not* cause a node influence conflict, for which Personalized PageRank matrix is used to encode the distribution of influence outward from nodes [7]:

$$\text{topoT}(v) = 1 - \mathbb{E}_{x \sim \mathbf{P}_v, \cdot} \left[ \frac{1}{|c|} \sum_{i \in c, c \neq L_s(v)} \mathbf{P}_{i,x} \right]$$

where  $L_s(v)$  denotes the estimated class type of node  $v$  with soft-label obtained in Label Propagation algorithm [48] following  $\mathbf{Y} = \mathbf{P}\mathbf{Y}^0$ ;  $\mathbf{Y}^0$  is the initial label distribution of  $X_R$ ,  $\mathbf{P} = \alpha(\mathbf{I} - (1 - \alpha)\mathbf{D}^{-\frac{1}{2}}\mathbf{A}\mathbf{D}^{-\frac{1}{2}})^{-1}$ .  $\mathbf{I}$  is the identity matrix,  $\alpha$  is the random walk restart probability,  $\mathbf{A}$  is the adjacency matrix, and  $\mathbf{D}$  is the diagonal degree matrix.  $\mathbf{P}_v$  is the Personalized PageRank probability vector for node  $v$ , which indicates a distribution of influence exerted outward to all the nodes in  $G$  from node  $v$ . Here  $\frac{1}{|c|}$  is a normalization term.

We take the expectation of the influence conflict that node  $v$  is classified to be the other “conflict” class (the opposite class of  $L_s(v)$ ), following a random walks across  $G$  as the probability measurement. As such,  $\text{topoT}(v)$  denotes the probability that node  $v$  *does not* lead to a influence conflict due to its topological location. The smaller influence conflict a node  $v$  has as in message passing-based GNN learning, the closer  $v$  to its topological class center is.

We define the *typicality* of a node  $v$  as

$$\mathcal{T}(v) = \text{clusT}(v) \cdot \text{topoT}(v)$$

Given a set of queries  $\mathcal{Q}$ , the typicality of  $\mathcal{Q}$  is defined as  $\mathcal{T}(\mathcal{Q}) = \sum_{v \in \mathcal{Q}} \mathcal{T}(v)$ . **cite the pagerank centrality strategy and make a quick comparison - justify the benefit of considering both embedding and topological features for deciding centrality.**

**Diversified Typicality Selection.** Given a graph  $G$  with unlabeled training node set  $U$ , a graph node embedding  $H^n(X_R)$ , and query budget  $k$ , the *diversified typicality selection* problem is to compute a set of queries  $\mathcal{Q} \subseteq U$  such that

$$\mathcal{Q} = \arg \max_{|\mathcal{Q}'|=k} (\mathcal{T}(\mathcal{Q}') + \lambda \sum_{v, v' \in \mathcal{Q}'} d(\mathbf{h}(v), \mathbf{h}(v'))) \quad (2)$$

---

**Procedure:** QSelect ( $H_n(X_R)$ ,  $V_T$ ,  $G$ ,  $k$ )

1. set  $\mathcal{Q} := \emptyset$ ; set  $U = V \setminus V_T$ ;
2.  $\mathcal{C}(U) := \text{ClusterU}(U, H_n(X_R))$ ;
3. **while**  $|\mathcal{Q}| < k$  **do**
4.     compute  $v \in U \setminus \mathcal{Q}$  that maximizes  $\mathcal{B}'_v(\mathcal{Q}, \mathcal{C})$ ;
5.      $\mathcal{Q} = \mathcal{Q} \cup \{v\}$ ;  $U = U \setminus \mathcal{Q}$ ;
6. **return**  $\mathcal{Q}$ ;

---

Fig. 5: Algorithm QSelect

Here  $d(\cdot)$  computes the Euclidean distance of the node embeddings of  $v$  and  $v'$ .

While the diversified typicality selection problem is NP-hard, we next present an efficient query selection algorithm.

### B. Query Selection Algorithm

We present an algorithm, denoted as QSelect, to implement the query selector  $\mathcal{S}$  (line 8 in Fig. 3). We focus on the query selection per iteration.

**Lemma 1:** *Given a graph  $G$  with unlabeled node set  $U$ , a graph node embedding  $H_n(X_R)$ , a query budget  $k$ , there exists a 2-approximation algorithm that selects a set of query  $\mathcal{Q}$  for the diversified typicality selection problem.*  $\square$

**Proof sketch:** We observe that diversified typicality selection can be reduced to the max-sum  $p$ -dispersion problem [4]. The latter maximizes the relevance of a given subset  $S$  to a search query and the distance between any pair in the subset  $S$ , with the cardinality constraint that  $|S| = p$ . Specifically, we show that (1)  $\mathcal{T}(\mathcal{Q})$  is a monotone (submodular) set function defined on the unlabeled node set. and (2) there is an approximation preserving reduction from the diversified typicality selection problem to the max-sum diversification problem.  $\square$

**Algorithm.** We next outline the algorithm QSelect (Fig. 5). It first performs a clustering algorithm ClusterU over the node features  $H_n(X_R)$ , and induces the clusters  $\mathcal{C}(U)$  for the unlabeled nodes  $U$  (line 2). It then adopts a greedy strategy to prioritize the selection of an unlabeled node  $v$  that has the largest marginal gain  $\mathcal{B}'_v(\mathcal{Q}, \mathcal{C}) = F(\mathcal{Q} \cup \{v\}) - F(\mathcal{Q})$ , where  $F(\mathcal{Q}) = \mathcal{T}(\mathcal{Q}) + \lambda \sum_{v, v' \in \mathcal{Q}}$ . By default, GALE implements ClusterU with  $k'$ -means clustering with  $k'$  between  $k$  and  $3k$ .

The 2-approximability can be verified by the approximation preserving reduction to the max-sum  $p$ -dispersion problem.

**For the time cost, the clustering can be implemented in time  $O(|\mathcal{C}|)$ , and the cost of computing  $\mathcal{B}'_v(\mathcal{Q}, \mathcal{C})$  can be optimized to only cost time  $O(1)$  [3], [4] (see more details in Section VII). Hence, the total time cost is  $O(k|V| + |\mathcal{C}(U)|)$ .**

## VI. QUERY ANNOTATION

We finally describe the query annotation process  $\mathcal{A}$  in GALE. The query annotator  $\mathcal{A}$  (implemented by the algorithm QAnnotate) enables  $\mathcal{O}$  to inspect the to-be-labelled

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**Procedure:** QAnnotate ( $\mathcal{Q}$ ,  $\Psi$ ,  $G$ )

1. **for each** base detector  $f_i \in \Psi$  **do**
2.     execute  $f_i \in \Psi$  to generate error distribution  $\Pi_{\psi_i}$ ;
3. **for each**  $v \in \mathcal{Q}$  **do**
4.     derive correct values for detected erroneous values;
5.     GetAnnotate ( $v$ ,  $G$ ,  $\Psi$ );
6. **return**  $\tilde{\mathcal{Q}}$ ;

---

Fig. 6: Algorithm QAnnotate

nodes with informative auxiliary structures, and obtain suggested correct attribute values of the detected erroneous nodes for further verification.

**Auxiliary structures.** GALE initializes a map structure for unlabeled nodes that are selected as queries. Given a node  $v \in \mathcal{Q}$ , the auxiliary data structures  $f(v)$  includes the following information, whenever applicable: (1) a “local” subgraph that is induced by the  $p$ -hop neighbors of  $v$ , (2) a list of erroneous attribute values  $v.A = 'a'$  that are captured by base detectors in  $\Psi$ , (3) for each erroneous attribute value  $v.A = 'a'$ , a set of suggested attribute values that are likely to be correct, (4) a probabilistic distribution of the errors learned by  $\Psi$ , and (5) statistical information such as degree centrality. The structures are shared with both the oracle  $\mathcal{O}$  and the query selector  $\mathcal{S}$  to facilitate label generation and query selection. We remark that these auxiliary structures can be user-defined to match the need given the different choices of the typicality functions and the type of oracles.

For each nodes in  $\mathcal{Q}$ ,  $\mathcal{A}$  also extracts their surrounding local graph information, e.g.,  $p$ -hop neighborhood induced subgraph and suggested “correct” attribute values as support information to the oracle. It invokes the procedure the method GetAnnotate( $\cdot$ ) to generate and aggregate the structural statistics (Fig. 6).

**Annotation Generation.** We focus on how the query annotator  $\mathcal{A}$  generates the auxiliary information of error distribution (case (2) - (4)) with base detectors  $\Psi$ .  $\mathcal{A}$  utilizes  $\Psi$  to generate error probability distribution of each training node as follows. (1) For each base detector  $f_i \in \Psi$  of a particular class  $C_i$ , it runs  $f_i$  over the nodes  $V$  to evaluate the number of erroneous nodes that are captured by  $f_i$  (denoted as  $|\Psi_i|$ ), and computes the counterpart  $|\Psi_{C_i}|$  that counts all the erroneous nodes captured by the same class of base detectors. This yields a normalized confidence score for each base detector  $f_i$  as  $\frac{|\Psi_i|}{|\Psi_{C_i}|}$ . (2) The overall error distribution is then estimated as a weighted sum of the scores to represent the error probability that an erroneous node  $v$  is likely to be “caused” by some particular error type. (3) When possible, a list of correct values for a detected wrong value is also suggested by “reversing” the detection process.

## VII. IMPLEMENTATION AND OPTIMIZATION

GALE optimizes the iterative learning of the active adversarial framework (lines 7-12 in Fig. 3) with the following



strategy. The general idea is to record previously computed model parameters and reduce unnecessary calculation, and avoid unnecessary update of the model parameters if the changes to the node embeddings are small. We make a case of the optimization for the query selector  $\mathcal{S}$  (algorithm QSelect), a major source of redundant computation. Indeed, it requires pairwise comparison of the large amount of unlabeled nodes. On the other hand, the distance measure satisfies triangle inequality. GALE maintains the following structures: (a) a matrix to store the node pair embedding distance  $d(u, v)$  for unlabeled node pair  $(v, v')$  whenever  $u, v \in \{U \cup C\}$ ; (b) a vector of flags to indicate whether the learned embedding of unselected nodes changes or not in two consecutive iterations, and (c) a dictionary that records the typicality of  $\mathcal{Q}$ , where the key is the size of the  $\mathcal{Q}$ . GALE simply retrieves an “approximate” distance  $d'(u, v)$  that is stored before if the embeddings of node  $v$  and  $v'$  are not significantly changed even  $\mathcal{D}^i$  is updated. If the embedding of  $u$  and  $v$  are element-wise equal within a tolerance, then  $d(u, v)$  is not updated.

GALE outperforms the state-of-the-art methods, including unsupervised constraint-based graph error detection and graph neural-network-based error detection with a cost of small overhead as shown in Section VIII.

**System Implementation.** We have developed a prototype system GALE, built on our pilot system GEDet [17].

**Feature Engineering.** GALE adopts word embedding [15] that maps attribute-level tokens (e.g., words) to vectors of real numbers. We firstly feed the node attribute embedding to a graph autoencoder GAE, which exploits graph structure information to learn node representations. GALE concatenates the attribute-level representation and node-level representation as the input of the SGAN. Furthermore, dimensional reduction techniques e.g., Principal Component Analysis (PCA) [27] is used to facilitates the error detection with less training cost.

**Built-in Library.** By default, GALE has three classes of built-in base detectors: (1) “constraints-based detectors”, which detects errors as violations of data constraints, e.g., graph functional dependencies [13], (2) “outlier detectors”, which encodes the algorithm in e.g., [5] to detect errors. (3) “string error detectors”, including the detection of string noises such as spelling errors. GALE keeps track of a set of “invertable” base detectors  $f$ , such that their consequence can be used to suggest possibly correct values, such as the equivalent literals in graph functional dependencies.

**Deployment.** GALE builders and servers are deployed in Google Colab environment with Tensorflow libraries and NVIDIA TESLA P100 with 16GB GPU memory. The base detector library, graph augmentation module, and an initial GUI has been well supported by the established implementation in GEDet [18]. The code and resources are made available <sup>2</sup>.

Dataset	$ V $	$ E $	# node types	# edge types	avg. # attr
DBP	2.2M	7.4M	73	584	4
OAG	0.6M	1.7M	5	6	2
Yelp	1.5M	1.6M	42	20	5

TABLE II: Overview of Real-world Graphs

Dataset	$ V_T $	$ E_T $	avg.# attrs	$ V^e $	$ V^e $
Species(SP:DBP)	17.7K	20K	4	1062	134
Data Mining(DM:OAG)	11.2K	12.9K	3	670	158
Machine Learning(ML:OAG)	3.4K	3.3K	3	203	54
UserGroup1(UG1:Yelp)	3.4K	2.6K	3	202	57
UserGroup2(UG2:Yelp)	3.3K	2.5K	3	196	45

TABLE III: Examples of Processed Graphs

## VIII. EXPERIMENT

We experimentally verify the effectiveness and the training cost of GALE for error detection in real-world graphs.

### A. Experiment settings

**Datasets.** We used the following real-world graph data. (1) DBP<sup>3</sup> is a knowledge graph extracted from Wikipedia, includes entities such as “plant” or “animal” and relationships among entities such as “studiedBy”; (2) OAG, a fraction of the open academic graph<sup>4</sup>, contains nodes such as e.g., articles, authors, organizations, and edges such as “cite” or “affiliatedTo”; (3) Yelp<sup>5</sup> is a graph that contains entities such as users and services (e.g., plumbers, restaurants), and edges such as “friendWith” and “reviews”. We also show the sizes of graphs that are induced by specific types (e.g., ‘Machine Learning’ from OAG denotes an induced subgraph with nodes that belong to the topic of ‘Machine Learning’). Tables II and III provide the details of the original and several examples of processed graphs, respectively.

**Error Generation.** We injected erroneous values into the real graphs with a configurable error generator. The generator enhances BART [2], a tool that pollute attributes to evaluate error detection. Three error classes were introduced for node attribute values with the help of stand-alone base detectors in a built-in library  $\Psi$ .

(1) *Constraint violations* are violations of a set of data constraints  $\Sigma$  [13]. We discovered data constraints  $\Sigma$  by invoking the algorithm in [12]. Matching “If” condition in the data constraints, the generator perturbed attribute values to violate the consequent value constraints, which leads to erroneous nodes. To discover meaningful  $\Sigma$ , we ensure constraints with high metrics such as minimum support (the minimum number of node matches) and minimum confidence (the ratio of nodes that also satisfy the consequent of a data constraint). Setting minimum support as 1000, 10, and 20 and minimum confidence as 0.9, 0.8, and 0.85 for DBP, OAG, and Yelp, we discover 89, 21, and 112 data constraints, respectively.

(2) *Outliers* are injected disturbed values that are significantly

<sup>2</sup><https://github.com/sxgcase/GALE>

<sup>3</sup><https://wiki.dbpedia.org/develop/datasets>

<sup>4</sup><https://www.aminer.org/open-academic-graph>

<sup>5</sup><https://www.yelp.com/dataset>

different from the distribution of peer attribute values. Some can be captured by outlier detectors in  $\Psi$ .

(3) *String noises* are generated from multiple scenarios including misspelling, missing values ('null'), and random string disturbance. We ensured that injecting these errors alone are not leading to violations of  $\Sigma$  or as detectable outliers.

We set the node error rate, which denotes the probability that a node in the graph is erroneous; attribute error rate, which denotes the probability that an attribute is perturbed as an erroneous attribute; and detectable rate, which denotes that an error is detectable by base detectors in the configurable error generator. The node error rate, attribute error rate and detectable rate are set as 0.01, 0.33, and 0.5 by default, respectively. For each graph, we randomly partition the nodes to obtain 6 folds for training examples, 1 fold for the validation set, and 3 folds for testing nodes (see Table III). The constraint-based detectors and outlier detectors are integrated into the annotation module for attribute value suggestion if consulted, to test how well GALE exploits external knowledge.

**Oracle.** For controlled tests, we simulated the oracle with the set of base detectors  $\Psi$ . An 'error' label is assigned if a base detector identified an erroneous attribute values of the query. For our case study, we have invited students who are experienced in verifying knowledge graphs and Wikipedia editing history to manually verify the errors and correct nodes.

**Algorithms.** We implemented GALE, along with 3 variants that interacts with the adversarial learning with different query selection strategies, and compared their performances with 5 baselines, all categorized as follows.

(1) Three variants of GALE, that replaces query selector  $\mathcal{S}$  with different strategies, including (a) GALE (-Ran.), a uniformly sampling of the unlabeled nodes; (b) GALE (-Ent.), entropy-based uncertainty sampling using the softmax outputs; and (c) GALE (-Kme.), sampling the unlabeled nearest nodes to K-means clustering centroids [22].

(2) Alad [31], a state-of-the-art anomaly node detection framework that measures normality of the nodes by considering both the topological structures of the graph and attribute distribution estimation within local context of nodes.

(3) VioDet, a constraint-based error detection that detects errors as the union of the violations of a set of data constraints  $\Sigma$  mined from the original datasets.

(4) Raha [33] is a state-of-the-art method to detect errors in relational data. It configures a library of built-in detectors *e.g.*, outlier detection, to generate error detection strategies.

(5) Two Graph Neural Network-based error detectors: (a) GCN [25] applies a graph convolutional architecture to encode local graph structure and features of nodes as a semi-supervised node classifier; (b) GEDet [17], a state-of-the-art few-shot learning based error detection framework. GEDet detects erroneous nodes with a graph augmentation module to enhance examples with synthetic ones, and adopts adversarial learning to train the node classifier.

**Configuration.** We use consistent settings for fair comparison.

(1) As Alad ranks nodes and is evaluated by AUC-PR curve [31], we applied the default setting to learn Alad, selected the thresholds that enable its best performance in terms of AUC-PR curve, and derived anomalies as erroneous nodes. (2) We used the same set of data constraints  $\Sigma$  for GALE variants, GEDet, and VioDet. We used the same settings for shared hyperparameters in variants of GALE on each dataset. GALE and its variants are consistently trained with the same rounds of iterations  $T$  and same local budget  $k$ , within ranges [7, 17] and [40, 100], respectively. (3) As Raha is designed for relational data, we applied Raha to node tables with one table per node type.

**Evaluation metrics.** We evaluate the performance in effectiveness and efficiency. For effectiveness, we report precision, recall, and  $F_1$ -score. Denote  $\text{Err}_d$  as the set of erroneous nodes detected from the graph, and  $\text{Err}$  as the set of nodes that are erroneous in the graph. The precision, recall and  $F_1$ -score are defined as  $P = \frac{|\text{Err}_d \cap \text{Err}|}{|\text{Err}_d|}$ ,  $R = \frac{|\text{Err}_d \cap \text{Err}|}{|\text{Err}|}$ , and  $F = \frac{2PR}{P+R}$ , respectively. For efficiency, we report the training time of GALE (-Ent.), GALE (-Ran.), GALE (-Kme.), and GALE.

All Experiments were executed on a Unix environment with Intel 2.6GHz CPUs, and 16GB memory. All the algorithms were implemented in Python on Tensorflow. Each experiment was run 5 times and the median results were reported. Our code and datasets of GALE are made available<sup>6</sup>.

## B. Experiment results

We first evaluate the effectiveness of GALE and the baseline algorithms, and the impact of several factors. We then evaluate the training cost of GALE. In addition, we conduct case studies to evaluate the usability of query annotation.

**Exp-1: Accuracy of GALE.** We report the accuracy of the methods over all the datasets in Table IV. GALE variants are initialized by using 10% of the training nodes  $V_T$  (summarized in Table III). The total budget size for Species(DBP), Data Mining (OAG), Machine Learning (OAG), UserGroup1 (Yelp), and UserGroup2 (Yelp) are 800, 1190, 800, 600, and 700, respectively. We report our findings below.

(1) We first inspect the 5 competing methods (all excluding GALE variants). The low recall of VioDet suggests the errors are quite diversified. GCN (graph neural network learning) and Raha (assembles multiple learning methods) are able to improve  $F_1$ -scores, but may come with a cost of significant precision drop (*e.g.*, GCN on 'UG2' dataset). Among the competing methods, GEDet achieves the best  $F_1$ -scores since its few-shot learning module and graph augmentation module ensure the coverage of heterogeneous errors.

(2) Active learning can effectively improve the performance of error detection. Despite the diversified errors, GALE variants achieve either the top or the second best results in precision, recall or  $F_1$ -score in almost all the datasets. This is not recognized in individual competing methods. On average,

<sup>6</sup><https://anonymou.s.4open.science/r/galedet-7400>

Data	Met.	VioDet	Alad	Raha	GCN	GEDet	GALE (-Ent.)	GALE (-Ran.)	GALE (-Kme.)	GALE
SP	P	0.85	0.26	0.40	0.57	<b>0.9085</b>	0.8237	0.8880	0.8530	0.8173
	R	0.24	0.80	0.60	0.35	0.6009	0.6801	0.6398	0.6859	<b>0.7219</b>
	F <sub>1</sub>	0.38	0.39	0.48	0.43	0.7233	0.7451	0.7437	0.7604	<b>0.7666</b>
DM	P	0.26	0.23	0.50	0.35	0.9812	0.9813	<b>0.9814</b>	0.9813	<b>0.9814</b>
	R	0.30	<b>0.77</b>	0.43	0.74	0.4541	0.4566	0.4578	0.4566	0.4578
	F <sub>1</sub>	0.28	0.35	0.47	0.47	0.6209	0.6232	<b>0.6244</b>	0.6232	<b>0.6244</b>
ML	P	0.24	0.23	0.62	0.63	0.9725	<b>0.9730</b>	0.9339	0.9412	0.9417
	R	0.27	0.40	0.45	0.43	0.4569	0.4655	<b>0.4871</b>	0.4828	<b>0.4871</b>
	F <sub>1</sub>	0.25	0.30	0.52	0.51	0.6217	0.6297	0.6402	0.6382	<b>0.6420</b>
UG1	P	0.33	0.27	0.63	0.51	<b>0.7764</b>	0.7578	0.7538	0.7652	0.7491
	R	0.55	0.55	0.60	0.52	0.6389	0.6736	0.6806	0.6563	<b>0.7049</b>
	F <sub>1</sub>	0.41	0.36	0.62	0.52	0.7010	0.7132	0.7153	0.7065	<b>0.7263</b>
UG2	P	0.31	0.27	0.59	0.66	<b>0.9576</b>	0.9440	0.9030	0.9160	0.9231
	R	0.54	<b>0.73</b>	0.56	0.33	0.4502	0.4701	<b>0.4821</b>	0.4781	0.4781
	F <sub>1</sub>	0.39	0.39	0.57	0.44	0.6125	<b>0.6277</b>	0.6286	0.6283	<b>0.6299</b>

TABLE IV: Performance of Error Detection. **Bold**: best result; Underlined: second best.

GALE has improved the F<sub>1</sub>-score of VioDet, Alad, Raha, GCN and GEDet, by 100%, 94%, 28%, 44%, and 5%, respectively.

(3) In general, GALE variants with active adversarial framework achieve desirable, and robust performance in accuracy despite of the presence of multiple types of errors and datasets. For example, in all cases, GALE and the three variants achieve at least **TBF** in precision, **TBF** in recall and **TBF** in F<sub>1</sub>. Other methods demonstrate higher variance given different datasets. For example, VioDet has a high precision of 0.85 on SP, and only achieves 0.24 precision on ML; similarly for GCN.

(3) For *all* the datasets, GALE achieves the best performance among all the GALE variants. This verifies the effectiveness of the typicality-based query selection strategy. We found that typical nodes from diversified clusters should be recommended to be queried in the low budget regime, instead of simply choosing queries with highest uncertainty, or simple strategies such as random selection (GALE (-Ran.)).

**Exp-2: Impact of factors.** We next investigate the impact of the following factors: (1) the data imbalance  $p_e = \frac{|V^e|}{|V^e|}$ , and (2) the training data ratio  $p_t = \frac{|V_T|}{|V^e|}$ , and (3) a cumulative query size  $K$  (the total # queries  $T \cdot k$  with  $T$  grows and  $k$  fixed).

*Impact of Data Imbalance.* Fixing  $p_t = \text{TBF}$  and  $K = \text{TBF}$ , we vary the imbalance  $p_e$  from 0.1 to 0.9 over Machine Learning (OAG). Fig. 7(a) tells us that while all methods achieve better performance over more balanced data, GALE (-Ent.), GALE (-Ran.), GALE (-Kme.), GALE, GEDet, are more stable than GCN, due to the graph augmentation module can counteract imbalanced training examples. Table IV consistently justifies this observation as the fraction of  $|V^e|$  varies over different datasets (summarized in Table III): compared with GCN, the methods GALE (-Ent.), GALE (-Ran.), GALE (-Kme.), and GALE are less sensitive.

*Varying example size.* Fixing  $K = \text{TBF}$  and  $p_e = \text{TBF}$ , we vary  $p_t$  over dataset UserGroup1(Yelp) from 15% to 1% and report the result in Fig. 7(b) (the result of VioDet and Alad are insensitive and constantly 0.41 and 0.36, respectively; not shown). While the accuracy decreases for all models as less training data is available, GALE (-Ent.), GALE (-Ran.),

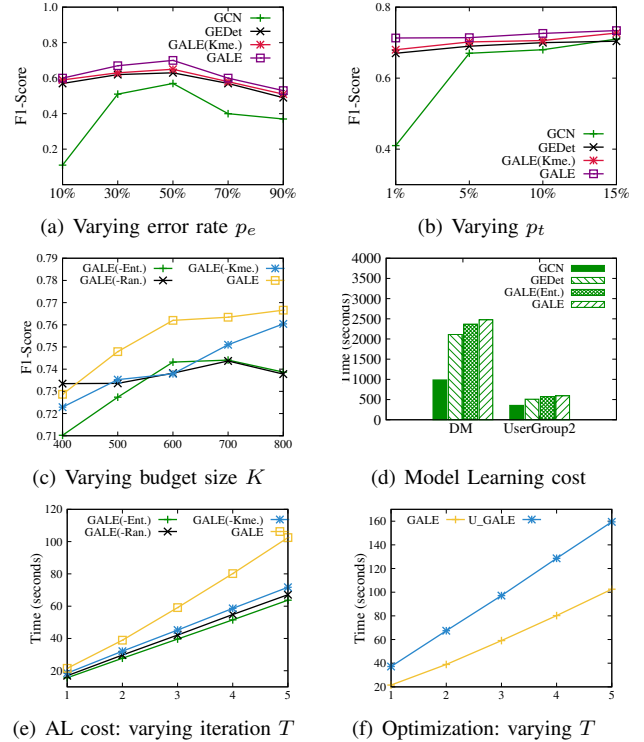


Fig. 7: Impact of factors to model performance

GALE (-Kme.), GEDet is the least sensitive (remains a recall above 0.6; not shown). Indeed, active learning variants in GALE effectively counteract the impact of lack of labels, with the graph augmentation model that improves recall, and the adversarial module that improves the accuracy.

*Varying cumulative budget  $K$ .* Active learning aims to label as few samples as possible while obtaining as many performance gains as possible. We study the effectiveness of different sampling strategies given a low budget regime. We vary cumulative budget  $T$  from 400 to 700 and report the result in Fig. 7(c). While the F<sub>1</sub>-scores increase as more unlabeled nodes are queried and get labeled by oracles for

all active learning sampling strategies in general, GALE and GALE (-Kme.) achieves the better  $F_1$ -scores compared with the other two methods, which affirms our claim that the typical and diversified nodes should be preferred to get selected over atypical (uncertain) nodes when the query budget is low. Hence, K-means clustering based sampling strategies (including GALE and GALE (-Kme.)) can help active learning strategies select typical graph nodes that are suited for low budgets. Furthermore, GALE outperforms GALE (-Kme.), which indicates the diversity that makes selected nodes be far apart benefits the active learning in low-budget regime.

**Exp-3: Learning cost.** We first compare the learning cost between GALE variants and other baselines. We found that GALE and variants do not incur significantly more time to learn a graph node error classifier. Then, we compare the learning cost with different sampling strategies in the low-budget regime for GALE variants. In the last, we show the optimization strategy applied on the GALE reduces the learning cost by 64%.

*Model Learning cost.* We evaluate the learning cost of GALE-based methods. We set the number of epochs as 220 for all the methods (200 epochs for the GAN in GALE variants to reach a Nash equilibrium and 20 epochs for the generative adversarial active learning module to query unlabeled nodes in the graph) and apply an “early-stop” strategy based on validation performance and make GEDet and GALE variants terminate early if no improvement is observed within consecutive 40 epochs. As shown in Fig. 7(d), (1) it is quite feasible to learn GALE. For example, it takes 520 seconds to learn GALE to achieve a recall at 0.48 over UG2; (2) GALE although is with the most sophisticated optimization goal, it still has a comparable running time with other GALE variants and improves the accuracy of error detection at a cost of small overhead. For example, it introduces on average 33%, 45%, 15% and 62% additional cost compared with GALE (-Kme.), GALE (-Ent.) (the running time of GALE (-Ran.) is close to GALE (-Ent.), not shown), GEDet and GCN, respectively.

*Active Learning cost.* We fix the queried nodes as 10 in each epoch on the Data Mining (OAG) of all GALE variants. After the labels of these queried nodes are provided by the oracle, we keep updating the model parameters of GALE variants. In the low-budget regime of Fig. 7(e), GALE introduces on average 53.8%, 42.9%, and 33.3% additional cost compared with GALE (-Ent.), GALE (-Ran.) and GALE (-Kme.). As we can observe that the learning cost of GALE is not significantly increased since we apply upper bound analysis for pruning during the queried node selection process and model memorization to skip unnecessary computation in the consecutive model updating.

*Optimization on GALE.* We apply upper bound analysis for node pruning when querying nodes to be labeled in the sampling strategy by leveraging the triangle inequality for distance computation. Furthermore, we store an embedding matrix and do not update distance of any node-pair embedding

if both node embedding values of the node-pair in consecutive model updating are close within a tolerance threshold. As Fig. 7(f) shows, GALE significantly reduces the running time by 40% compared to the naive implementation as  $U\_GALE$  on the Data Mining (OAG) when the number of queried nodes per epoch is 10.

**Exp-4: Case study: Usability of Query Annotation.** GALE can show clear advantages in identifying erroneous nodes when the given training examples are sparse. It can successfully detect erroneous nodes given additional labels, where base detectors fail to detect such errors. In a real-world knowledge graph, a node  $v$  with “name”: cavanillesia and its attribute “order” is associated with an erroneous attribute value “Lepidoptera”, which should be “Marvales”. There is no graph constraint that covers node  $v$  and the outlier detector detects node  $v$  with a normalized outlier score as 0.35 (cannot be classified as an error). In the query selection process, GALE considers the *typicality*, queries an unlabelled node  $u$ , and feeds node  $u$  to the *oracle*. Oracle can leverage the auxiliary information associated with  $u$  to correctly label node  $u$ . The identified node  $u$  is semantically similar to node  $v$  (sharing an common attribute “kingdom” with value “plantae”). The iterative updating of SGAN helps GALE detect the errors on attribute value “Lepidoptera” of node  $v$ .

## IX. CONCLUSION

We have introduced GALE, a graph error detection framework empowered by an active adversarial learning framework. GALE exploits active learning to best exploit the new knowledge from oracles by issuing a bounded number of queries. Moreover, we introduce new quality measures for selecting queries in graph data in terms of diversified typicality, and introduced approximation algorithms for query selection, query annotation schemes to facilitate oracle and query selection, and optimization strategies. Our experimental study verifies that the active learning and adversarial error detection of GALE achieve significant gain on accuracy compared with state-of-the-art baselines. A future topic is to enhance GALE for large-scale error detection with distributed learning techniques.

## REFERENCES

- [1] Z. Abedjan, X. Chu, D. Deng, R. C. Fernandez, I. F. Ilyas, M. Ouzzani, P. Papotti, M. Stonebraker, and N. Tang. Detecting data errors: Where are we and what needs to be done? *VLDB*, 2016.
- [2] P. C. Arocena, B. Glavic, G. Mecca, R. J. Miller, P. Papotti, and D. Santoro. Messing up with bart: error generation for evaluating data-cleaning algorithms. *VLDB*, 2015.
- [3] B. Birnbaum and K. J. Goldman. An improved analysis for a greedy remote-clique algorithm using factor-revealing lps. *Algorithmica*, 55(1):42–59, 2009.
- [4] A. Borodin, H. C. Lee, and Y. Ye. Max-sum diversification, monotone submodular functions and dynamic updates. In *Proceedings of the 31st ACM SIGMOD-SIGACT-SIGAI symposium on Principles of Database Systems*, pages 155–166, 2012.
- [5] M. M. Breunig, H.-P. Kriegel, R. T. Ng, and J. Sander. Lof: Identifying density-based local outliers. In *SIGMOD*, 2000.
- [6] G. Castagna, D. Colazzo, and A. Frisch. Error mining for regular expression patterns. In *Italian conference on Theoretical Computer Science*, pages 160–172. Springer, 2005.

- [7] D. Chen, Y. Lin, G. Zhao, X. Ren, P. Li, J. Zhou, and X. Sun. Topology-imbalance learning for semi-supervised node classification. *Advances in Neural Information Processing Systems*, 34:29885–29897, 2021.
- [8] F. Chen, Y.-C. Wang, B. Wang, and C.-C. J. Kuo. Graph representation learning: a survey. *APSIPA Transactions on Signal and Information Processing*, 9, 2020.
- [9] K. Cheng, X. Li, Y. E. Xu, X. L. Dong, and Y. Sun. Pge: Robust product graph embedding learning for error detection. *arXiv preprint arXiv:2202.09747*, 2022.
- [10] X. Chu, I. F. Ilyas, S. Krishnan, and J. Wang. Data cleaning: Overview and emerging challenges. In *SIGMOD*, 2016.
- [11] K. Ding, J. Li, and H. Liu. Interactive anomaly detection on attributed networks. In *Proceedings of the twelfth ACM international conference on web search and data mining*, 2019.
- [12] W. Fan, C. Hu, X. Liu, and P. Lu. Discovering graph functional dependencies. In *SIGMOD*, 2018.
- [13] W. Fan, Y. Wu, and J. Xu. Functional dependencies for graphs. In *SIGMOD*, 2016.
- [14] Y. Fu, X. Zhu, and B. Li. A survey on instance selection for active learning. *Knowledge and information systems*, 35(2):249–283, 2013.
- [15] M. Gardner, J. Grus, M. Neumann, O. Tafjord, P. Dasigi, N. Liu, M. Peters, M. Schmitz, and L. Zettlemoyer. Allennlp: A deep semantic natural language processing platform. *arXiv preprint arXiv:1803.07640*, 2018.
- [16] M. Gori, G. Monfardini, and F. Scarselli. A new model for learning in graph domains. In *Proceedings. 2005 IEEE International Joint Conference on Neural Networks, 2005.*, 2005.
- [17] S. Guan, P. Lin, H. Ma, and Y. Wu. Gedet: Adversarially learned few-shot detection of erroneous nodes in graphs. In *IEEE Big Data*, 2020.
- [18] S. Guan, H. Ma, S. Choudhury, and Y. Wu. Gedet: detecting erroneous nodes with a few examples. *Proceedings of the VLDB Endowment*, 14(12):2875–2878, 2021.
- [19] G. Hacohen, A. Dekel, and D. Weinshall. Active learning on a budget: Opposite strategies suit high and low budgets. *arXiv preprint arXiv:2202.02794*, 2022.
- [20] A. Heidari, J. McGrath, I. F. Ilyas, and T. Rekatsinas. Holodetect: Few-shot learning for error detection. In *SIGMOD*, 2019.
- [21] I. F. Ilyas, X. Chu, et al. Trends in cleaning relational data: Consistency and deduplication. *Foundations and Trends® in Databases*, 2015.
- [22] X. Jin and J. Han. K-means clustering. In *Encyclopedia of Machine Learning*, 2010.
- [23] Z. Khayyat, I. F. Ilyas, A. Jindal, S. Madden, M. Ouzzani, P. Papotti, J.-A. Quiané-Ruiz, N. Tang, and S. Yin. Bigdansing: A system for big data cleansing. In *Proceedings of the 2015 ACM SIGMOD international conference on management of data*, pages 1215–1230, 2015.
- [24] D. P. Kingma and J. Ba. Adam: A method for stochastic optimization. *arXiv preprint arXiv:1412.6980*, 2014.
- [25] T. N. Kipf and M. Welling. Semi-supervised classification with graph convolutional networks. *arXiv preprint arXiv:1609.02907*, 2016.
- [26] T. N. Kipf and M. Welling. Variational graph auto-encoders. *arXiv preprint arXiv:1611.07308*, 2016.
- [27] R. Lebrete and R. Collobert. Word emdeddings through hellinger pca. *arXiv preprint arXiv:1312.5542*, 2013.
- [28] D. D. Lewis and W. A. Gale. A sequential algorithm for training text classifiers. In *SIGIR'94*, 1994.
- [29] Y. Li, J. Yin, and L. Chen. Seal: Semisupervised adversarial active learning on attributed graphs. *IEEE Transactions on Neural Networks and Learning Systems*, 32(7):3136–3147, 2020.
- [30] P. Lin, Q. Song, Y. Wu, and J. Pi. Repairing entities using star constraints in multirelational graphs. In *ICDE*, 2020.
- [31] N. Liu, X. Huang, and X. Hu. Accelerated local anomaly detection via resolving attributed networks. In *IJCAI*, 2017.
- [32] K. Madhawa and T. Murata. Active learning for node classification: an evaluation. *Entropy*, 22(10):1164, 2020.
- [33] M. Mahdavi, Z. Abedjan, R. Castro Fernandez, S. Madden, M. Ouzzani, M. Stonebraker, and N. Tang. Raha: A configuration-free error detection system. In *SIGMOD*, 2019.
- [34] C. Mayfield, J. Neville, and S. Prabhakar. Eracer: a database approach for statistical inference and data cleaning. In *SIGMOD*, 2010.
- [35] F. Neutatz, M. Mahdavi, and Z. Abedjan. Ed2: A case for active learning in error detection. In *Proceedings of the 28th ACM international conference on information and knowledge management*, pages 2249–2252, 2019.
- [36] M. Nickel, K. Murphy, V. Tresp, and E. Gabrilovich. A review of relational machine learning for knowledge graphs. *Proceedings of the IEEE*, 104(1):11–33, 2015.
- [37] H. Paulheim. Knowledge graph refinement: A survey of approaches and evaluation methods. *Semantic web*, 2017.
- [38] Z. Peng, M. Luo, J. Li, H. Liu, and Q. Zheng. Anomalous: A joint modeling approach for anomaly detection on attributed networks. In *IJCAI*, 2018.
- [39] K. Pourahmadi, P. Nooralinejad, and H. Pirsiavash. A simple baseline for low-budget active learning. *arXiv preprint arXiv:2110.12033*, 2021.
- [40] T. Salimans, I. Goodfellow, W. Zaremba, V. Cheung, A. Radford, and X. Chen. Improved techniques for training gans. In *NeurIPS*, 2016.
- [41] B. Settles. Active learning literature survey. 2009.
- [42] D. J. Wang, X. Shi, D. A. McFarland, and J. Leskovec. Measurement error in network data: A re-classification. *Social Networks*, 2012.
- [43] Y. Wang, Q. Yao, J. Kwok, and L. M. Ni. Generalizing from a few examples: A survey on few-shot learning. In *arXiv: 1904.05046*. 2019.
- [44] Z. Wu, S. Pan, F. Chen, G. Long, C. Zhang, and S. Y. Philip. A comprehensive survey on graph neural networks. *IEEE Transactions on Neural Networks and Learning Systems*, 2020.
- [45] M. Xu. Understanding graph embedding methods and their applications. *SIAM Review*, 63(4):825–853, 2021.
- [46] C. Zhao, Y. Xin, X. Li, Y. Yang, and Y. Chen. A heterogeneous ensemble learning framework for spam detection in social networks with imbalanced data. *Applied Sciences*, 10(3):936, 2020.
- [47] Y. Zhong, W. Chen, Z. Wang, Y. Chen, K. Wang, Y. Li, X. Yin, X. Shi, J. Yang, and K. Li. Helad: A novel network anomaly detection model based on heterogeneous ensemble learning. *Computer Networks*, 169:107049, 2020.
- [48] D. Zhou and C. J. Burges. Spectral clustering and transductive learning with multiple views. In *Proceedings of the 24th international conference on Machine learning*, pages 1159–1166, 2007.