Source-Code Embedding-Based Software Defect Prediction

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Abstract: Software defect prediction is an essential software development activity, a highly researched topic and yet a still difficult problem. One of the difficulties is that the most prevalent software metrics are insufficiently relevant for predicting defects. In this paper we are proposing the use of Graph2Vec embeddings unsupervisedly learnt from the source code as basis for prediction of defects. The reliability of the Graph2Vec embeddings is compared to that of the alternative embeddings based on Doc2Vec and LSI through a study performed on 16 versions of Calcite and using three classification models: FastAI, as a deep learning model, Multilayer Perceptron, as an untuned conventional model, and Random Forests with hyperparameter tuning, as a tuned conventional model. The experimental results suggest a complementarity of the Graph2Vec, Doc2Vec and LSI-based embeddings, their combination leading to the best performance for most software versions. When comparing the three classifiers, the empirical results highlight the superiority of the tuned Random Forests over FastAI and Multilayer Perceptron, which confirms the power of hyperparameter optimization.

1 INTRODUCTION

Software defect prediction (SDP) is the task of identifying defective software components, so that testing effort can be focused on them. Since the resources for testing are always limited, having accurate SDP models can lead to the best results with the available resources. Given its importance, it is not surprising that SDP is an actively researched topic.

In order to build a Machine Learning (ML) model for SDP it is necessary to have a common representation (feature vector) for all the software entities. The earliest and prevalent representation is based on software metrics. Initially, the metrics considered have been procedural code metrics exclusively, but they have been replaced (or complemented) later by object-oriented and code change metrics. More recently, however, several approaches for constructing software features by directly or indirectly considering the source code, through the Abstract Syntax Tree (AST), have been proposed.

In a recent study (Miholca et al., 2022), an extensive series of traditional software metrics have been compared to conceptual software features extracted directly from the source code using Doc2Vec and LSI. The study’s conclusion is that, on average, the Doc2Vec and LSI-based software features outperform the traditional ones in terms of their soundness in predicting defect-proneness. In extracting the Doc2Vec and LSI-based features, the source code is seen as a piece of text. So, its actual structure, which is captured by the AST, is not taken into consideration.

Consequently, in the current paper, we want to build on the work of Miholca et al. (Miholca et al., 2022), while also capitalizing on the structure underlying the source code. To this end, we are proposing an additional representation unsupervisedly learnt from the AST of the source code using Graph2Vec (Narayanan et al., 2017). Graph2Vec is a neural embedding framework which can learn a fixed-length feature vector (called embedding) for an entire graph. The Graph2Vec-based embeddings will be evaluated in terms of their discriminative power when it comes to classifying software entities as defective or non-defective. Having the intuition of a possible complementarity between the Graph2Vec-based embedding, that capitalizes on the software structure, and the Doc2Vec and LSI-based embeddings, that capitalize on the semantics of the source code (especially of the comments and identifiers), we will also investigate if combining them leads to a superior SDP performance. Accordingly, we define the following research questions:

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• **RQ1.** What is the relative relevance of the embedding learnt using Graph2Vec for SDP?
• **RQ2.** Does the combination of the Graph2Vec, Doc2Vec and LSI embeddings improve the performance of SDP?

Miholca et al. (Miholca et al., 2022) have experimentally compared multiple classifiers of which FastAI proved to be the best performer, being followed by a feed-forward artificial neural network that is, in fact, a Multilayer Perceptron (MLP). Since we build on their results and conclusions, we will also use FastAI, which is a deep learning model. However, recently, some studies (for example (Fu and Menzies, 2017) and (Majumder et al., 2018)) have shown that there are many tasks for which there is no need for deep learning, which in general needs more training data and takes a longer time to run, since the same performance can be achieved with a simpler ML algorithm if proper hyperparameter tuning is performed. While these studies considered other tasks, not SDP, Tantithamthavorn et al. in (Tantithamthavorn et al., 2019) investigated the impact of automated parameter optimization for SDP and concluded that for most classification techniques a non-negligible performance improvement can be achieved by hyperparameter optimization. Starting from this conclusion, we have decided to use the Random Forest (RF) classifier with optimized parameters as well. Consequently, we add a third research question:

• **RQ3.** How does the SDP performance of FastAI compare to the one of RF with optimized parameters for the considered representations?

The main original contributions brought by this paper are as follows. To the best of our knowledge, the Graph2Vec-based embedding learnt from the source code AST has not yet been used as representation in the SDP literature. This being the case, we are proposing a new software representation for the benefit of SDP. Moreover, we are proposing combining the Graph2Vec-based embedding with the Doc2Vec-based and the LSI-based embeddings proposed in (Miholca et al., 2022) to unite the power of structural and conceptual information underlying the source code and thus to boost the SDP performance. In addition to this contributions, in this paper we are also assessing the potency of parameter optimization in the context of SDP by comparing a tuned Random Forest to the FastAI deep classifier, while feeding them with the proposed software embeddings.

The remainder of this paper is organized in the following manner. Section 2 shortly presents a selection of existing SDP approaches that are relevant for the current paper. Section 3 describes in detail the experimental methodology. Section 4 presents the results of the performed analysis and formulates the answers to our research questions. Finally, in Section 5, the conclusions are drawn and some directions to continue this work are outlined.

## 2 RELATED WORK

There is a huge amount of literature related to the problem of SDP. Throughout the years, researchers proposed many approaches using supervised and unsupervised machine learning, deep learning, ensemble approaches, etc. In the following we will focus only on a selection of papers that are most related to the main topic of our paper: different software representations to be used in SDP.

Most existing studies use for experimental evaluation the SDP data sets available in Promise Software Engineering Repository (Sayyad and Menzies, 2015), that is currently known as SeaCraft (Sea, 2017). Accordingly, many of the existing SDP approaches are based on the Promise metrics, that are either static OO metrics or traditional metrics associated with the quality of the procedural source code. Literature reviews reveals that about 87% (Malhotra, 2015) of the case studies used such procedural or object-oriented metrics while being focused on proposing accurate classifiers.

However, relatively recent studies have opened up a new, highly active, research direction in the field of SDP - the one of designing new relevant features for enabling the discrimination between defective and non-defective software entities. This research direction is motivated by the fact that the existing software features are insufficient or insufficiently relevant for SDP, making it still a difficult problem. In the following, we will focus on presenting state-of-the art approaches that are the most related to our approach.

Many approaches proposing new features consider the AST of the source code. One such approach is presented by Wang et al. in (Wang et al., 2016) and (Wang et al., 2020). They automatically learn semantic features starting from the AST and using Deep Belief Networks (DBNs). The resulted features have been comparatively evaluated against traditional metrics in terms of their relevance for SDP, on open source projects from the Promise repository as case studies. The evaluation results have confirmed that the proposed semantic features outperform traditional SDP features.

Li et al (Li et al., 2017) have proposed a similar process, but using Convolutional Neural Networks (CNNs) instead of DBNs, and combining the ex-
tracted features with some traditional ones. The combined representation has been evaluated in terms of its soundness for SDP by using the Logistic Regression (LR) classifier. The results of the experimental evaluation confirmed that the AST-based features outperform traditional features, while combining the two of them leads to even better performance.

Another study using AST-based features for SDP is the one performed by Dam et al. (Dam et al., 2018). They have introduced a tree-structured network of Long-Short Term Memory (LSTM) units as a SDP model starting from AST embeddings. Using the same set of 10 case studies as Wang et al. (Wang et al., 2016), they train two traditional classifiers: LR and Random Forest on the features generated by LSTM.

Aladics et al. (Aladics et al., 2021) have proposed an approach, where the AST of the source code was parsed in a depth-first order, and the sequence of nodes was recorded and considered as a document, on which Doc2Vec was applied. Their experimental evaluation considering several ML algorithms showed that these embedding might not outperform traditional source code metrics, but combining the two of them will improve the performance.

A SDP model based on a Convolutional Graph Neural Network has been proposed by Sikic et al. in (Sikic et al., 2022). The neural network architecture employed is specifically tailored for graph data so that AST data can be fed into it. For the experimental evaluation 7 SDP data sets from the Promise repository were considered. The results showed that the proposed model outperforms standard SDP models and is comparable to the state-of-the-art AST-based SDP models, including (Wang et al., 2020).

Another direction of defining features derived from the source code, but not involving AST, is the one of Miholca et al. (Miholca et al., 2020), who have proposed the COMET metrics suite. This suite starts from the representations of the source code learnt using LSI and Doc2Vec, but these are not used directly for SDP. Rather, for every entity, different descriptive statistical measures are computed between its representation and the other (all or just defective or non-defective) representations. These statistical measures together provide the representation of an entity. The experimental evaluation was performed on 7 Promise data sets and the COMET metrics suite outperformed the traditional Promise metrics.

Doc2Vec and LSI vectorial representations of the source code have been directly used for SDP in a study by Miholca et al. (Miholca et al., 2022). The authors have compared them to an extensive set of 4189 metrics containing static code metrics, clone metrics, warning-based metrics, changes-based and refactoring-based metrics, AST node counts and code churn metrics. As experimental case study, multiple versions of a software system have been considered. Different combinations of the metrics subsets have been evaluated in terms of their relevance for SDP. The experimental results led to the conclusion that combining Doc2Vec and LSI produces a predominantly superior performance of SDP when compared to the 4189 software metrics, as well as to the separate use of Doc2Vec and LSI representations.

Subsequent to the release of BERT, an advanced language representation model, its feasibility for predicting software defects has begun to be investigated. Cong et al. (Fan et al., 2021) have employed CodeBERT, a BERT model pre-trained on open source repositories, while Uddin et al. (Uddin et al., 2022) have pre-trained themselves a BERT model on source code. In both approaches the code comments have been eliminated in the pre-processing phase, whereas in our approach, presented in the following, we intentionally keep them, considering that they are potential carriers of semantic information that can benefit SDP.

3 APPROACH

3.1 Case Study

As a case study, we have selected 16 releases of Apache Calcite, an open-source dynamic data management framework (Begoli et al., 2018). Details about the considered versions of Calcite are presented in Table 1.

We started from the Calcite data sets provided by Herbold et al. (Herbold et al., 2022), that have been produced by an extended version of the SZZ-RA (Neto et al., 2018) algorithm. These data sets contain the names of the classes, the value of 4189 software metrics and the class label. Since in this study we use as features different embeddings extracted from the source code, we have only used the labels provided by Herbold et al. and added them to our representations, using the class name to match the instances.

3.2 Proposed Vectorial Representations

Instead of considering static structural or code churn metrics like the vast majority of SDP approaches, in this paper we consider three different embeddings constructed (directly or indirectly) from the source code of a software system.

The first embedding is unsupervisedly learnt by Doc2Vec (Le and Mikolov, 2014), a prediction-based model for representing texts (in our case, source code)
Table 1: Number of non-defective and defective instances, total number of instances and rate of defective instances for all Calcte versions.

<table>
<thead>
<tr>
<th>Version</th>
<th>Non-defective</th>
<th>Defective</th>
<th>Total</th>
<th>Defective rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.0</td>
<td>897</td>
<td>178</td>
<td>1075</td>
<td>0.166</td>
</tr>
<tr>
<td>1.1</td>
<td>990</td>
<td>113</td>
<td>1103</td>
<td>0.102</td>
</tr>
<tr>
<td>1.2</td>
<td>982</td>
<td>126</td>
<td>1108</td>
<td>0.114</td>
</tr>
<tr>
<td>1.3</td>
<td>1003</td>
<td>112</td>
<td>1115</td>
<td>0.100</td>
</tr>
<tr>
<td>1.4</td>
<td>1004</td>
<td>123</td>
<td>1127</td>
<td>0.109</td>
</tr>
<tr>
<td>1.5</td>
<td>1073</td>
<td>103</td>
<td>1176</td>
<td>0.088</td>
</tr>
<tr>
<td>1.6</td>
<td>1086</td>
<td>107</td>
<td>1193</td>
<td>0.090</td>
</tr>
<tr>
<td>1.7</td>
<td>1124</td>
<td>128</td>
<td>1252</td>
<td>0.102</td>
</tr>
<tr>
<td>1.8</td>
<td>1200</td>
<td>101</td>
<td>1301</td>
<td>0.078</td>
</tr>
<tr>
<td>1.9</td>
<td>1220</td>
<td>90</td>
<td>1310</td>
<td>0.069</td>
</tr>
<tr>
<td>1.10</td>
<td>1226</td>
<td>84</td>
<td>1310</td>
<td>0.064</td>
</tr>
<tr>
<td>1.11</td>
<td>1251</td>
<td>80</td>
<td>1331</td>
<td>0.060</td>
</tr>
<tr>
<td>1.12</td>
<td>1334</td>
<td>81</td>
<td>1415</td>
<td>0.057</td>
</tr>
<tr>
<td>1.13</td>
<td>1222</td>
<td>53</td>
<td>1275</td>
<td>0.042</td>
</tr>
<tr>
<td>1.14</td>
<td>1255</td>
<td>53</td>
<td>1308</td>
<td>0.041</td>
</tr>
<tr>
<td>1.15</td>
<td>1307</td>
<td>45</td>
<td>1352</td>
<td>0.033</td>
</tr>
</tbody>
</table>

As a fixed-length numeric vector. It is a MLP based model that extends Word2Vec and an alternative to traditional models such as bag-of-words and bag-of-n-grams. The main advantage of Doc2Vec over traditional models is that it considers the semantic distance between words (Le and Mikolov, 2014).

The second embedding is extracted by LSI (Deerwester et al., 1990), a count-based model for representing texts (in our case, source code). LSI builds a matrix of occurrences of words in documents and then uses singular value decomposition to reduce the number of words while keeping the similarity structure between documents.

The third embedding is based on Graph2Vec, a model which can create a fixed-length vector of an entire graph, using unsupervised learning (Narayanan et al., 2017). The basic idea behind the algorithm is to view an entire graph as a document and its rooted subgraphs around the nodes as the words of the document and apply document embedding models (more exactly, Doc2Vec) to learn graph embeddings. Source code can easily be parsed into AST which is in fact a graph, for which an embedding can be generated using Graph2Vec.

For any embedding, the software entities are represented as numeric vectors. These are composed of numerical values corresponding to a set $\mathcal{F} = \{f_1, f_2, \ldots, f_s\}$ of features learned from the source code directly (in case of Doc2Vec and LSI) or indirectly, through the AST (in case of Graph2Vec).

Therefore, a software entity $se$ is represented as an $s$-dimensional vector in an emb space:

- $se^{emb} = (se_1^{emb}, \ldots, se_s^{emb})$, where $se_i^{emb}$ (forall $1 \leq i \leq s$) denotes the value of the $i$-th feature computed for the entity $se$ by using embedding $emb$, $emb \in \{\text{Doc2Vec}, \text{LSI}, \text{Graph2Vec}\}$.

For extracting the conceptual vectors, we have opted for $s = 30$ as the length of the embedding. In case of Doc2Vec and LSI, the source code (including comments) afferent to each class was filtered so as to keep only the tokens presumably carrying semantic meaning. So, operators, special symbols, English stop words or Java keywords have been eliminated. For both Doc2Vec and LSI, we have used the implementation offered by Gensim (Rehurek and Sojka, 2010). For Graph2Vec we have used the implementation from Karateclub (Rozemberczki et al., 2020), with the number of epochs set to 100 and the flag to consider the labels of the AST-nodes set to True.

The experimental results of previous studies (Miholca and Cizbulia, 2019) (Miholca and Onet-Marian, 2020) (Miholca et al., 2022) revealed that combining Doc2Vec and LSI is appropriate and increases the performance of SDP, while, to the best of our knowledge, Graph2Vec was not used previously for SDP. Consequently, we have decided to use the following three representation in our experimental evaluation:

- **Doc2Vec + LSI** - each software entity $se$ is represented as a $2 \times s$-dimensional vector: $se_{\text{Doc2Vec} + \text{LSI}} = (se_1^{\text{Doc2Vec}}, \ldots, se_s^{\text{Doc2Vec}}, se_1^{\text{LSI}}, \ldots, se_s^{\text{LSI}})$, where $se_i^{\text{Doc2Vec}}$ (forall $1 \leq i \leq s$) denotes the value of the $i$-th feature computed for the entity $se$ by using Doc2Vec and $se_i^{\text{LSI}}$ (forall $1 \leq i \leq s$) denotes the value of the $i$-th feature computed for the entity $se$ by using LSI.

- **Graph2Vec** - each software entity $se$ is represented as the embedding provided by Graph2Vec for it: $se_{\text{Graph2Vec}} = (se_1^{\text{Graph2Vec}}, \ldots, se_s^{\text{Graph2Vec}})$, where $se_i^{\text{Graph2Vec}}$ (forall $1 \leq i \leq s$) denotes the value of the $i$-th feature computed for the entity $se$ by using Graph2Vec.

- **Doc2Vec + LSI + Graph2Vec** - each software entity $se$ is represented as a $3 \times s$-dimensional vector, the concatenation of the three embeddings for it: $se_{\text{Doc2Vec} + \text{LSI} + \text{Graph2Vec}} = (se_1^{\text{Doc2Vec}}, \ldots, se_s^{\text{Doc2Vec}}, se_1^{\text{LSI}}, \ldots, se_s^{\text{LSI}}, se_1^{\text{Graph2Vec}}, \ldots, se_s^{\text{Graph2Vec}})$.

The third representation is inspired by the intuition that the Doc2Vec + LSI embedding captures semantic (or conceptual) information contained by the source code, including the comments, while overlooking the structural information, whereas the Graph2Vec embedding also captures data related to the syntactic structure of the source code. Consequently, we intend to experimentally verify whether or not joining them enhances the SDP performance.
3.3 Vectorial Representations Relevance Analysis

Given that the representation obtained by joining the Doc2Vec and LSI conceptual vectors have been experimentally proven to be prevalently superior to the representation based on traditional software metrics in terms of their effectiveness for SDP (Miholca et al., 2022), for assessing the relative relevance of the Graph2Vec-based representation, we will compare it to the combined representation Doc2Vec + LSI.

3.3.1 Difficulty Analysis

To comparatively analyse how the different embeddings facilitate the discrimination between defective and non-defective software entities, but also to outline the complexity of the SDP task, we have computed for each version of Calcite and for each of the three representations (Doc2Vec + LSI, Graph2Vec and Doc2Vec + LSI + Graph2Vec) two difficulty measures.

As defined by Zhang et al. (Zhang et al., 2007), the difficulty of a class \( c \), in a binary classification context, is the proportion of data instances belonging to class \( c \) for which the nearest neighbor (computed using the Euclidean distance, when ignoring the class label) belongs to the opposite class. In our case, the two difficulty measures are computed for the defective (positive) and for the non-defective (negative) classes. Intuitively, the difficulty of a class indicates how hard it is to distinguish the instances belonging to that class from the others, considering a given vectorial representation for the software entities.

3.3.2 Supervised Analysis

The difficulty analysis is supplemented by a supervised analysis which involves three different classification models: FastAI, a deep learning model that proved to be the best-performing one in (Miholca et al., 2022), Multilayer Perceptron, an untuned conventional model that proved to be the second best-performing classifier in (Miholca et al., 2022) and Random Forest with hyperparameter optimization, as a tuned conventional classifier.

FastAI. FastAI, the first classification model employed in the supervised analysis, is a deep learning classifier implemented in the FastAI machine learning library (Howard et al., 2018). It is composed of an artificial neural network with embeddings of the input layer. The architecture consists of 1 input, 1 output and 2 hidden layers. Compared to other deep learning models, especially Convolutional Neural Networks, FastAI is very small and fast, which makes it suitable for real-time usage scenarios. The model is trained using the FastAI fit one cycle method, which uses a learning rate that varies according to a specific pattern: first it increases, then it decreases and the process is repeated for each epoch.

Multilayer Perceptron. The second classification model employed is a Multilayer Perceptron. We used the scikit-learn implementation for this model, while opting for one single hidden layer and a rectified linear unit activation function. The model has been trained using a stochastic gradient-based optimizer for maximum 2000 epochs and with a constant learning rate of 0.001.

Random Forest. The third classification model used in this analysis is Random Forest (Breiman, 2001), an ensemble learning method, which builds a set of decision trees and uses a majority voting mechanism to make a prediction for an unseen instance. Each decision tree is built considering only a random selection of features from the data set and, in general, using only a subset of the training instances, sampled randomly with replacement. RF have previously been used extensively for SDP (Matloob et al., 2021).

RF was one of the classifiers used in a study about the effect of hyperparameter tuning for SDP models (Tantithamthavorn et al., 2019), but only one parameter, the number of classification trees, was tuned. The conclusions of that study were that RF is not that sensitive to hyperparameter values when AUC is used as a performance measure. Since we use different data sets and different representations than the ones used in (Tantithamthavorn et al., 2019), we have decided to see if their conclusions are valid for our data as well and to try and find the best set of parameters for the RF classifier.

We have used the implementation for RF from the scikit learn library (Pedregosa et al., 2011), where there are a set of parameters related to building the trees that can be used to find the best model. We have decided to tune 8 of them, that are presented in Table 2 together with the considered values.

Out of the parameter values from Table 2 not all possible combinations are valid: if the value of bootstrap is set to False, the value of max_samples has to be set to None. Considering this restriction, the total number of valid parameter combinations for the RF algorithm is 32,400. Checking such a large number of parameter combinations using grid search is not possible, especially since some of the parameters (for example bootstrap and max_features) introduce randomness in the process, so in order to get a more accurate view of the performance of a point from the hy-
### Table 2: Hyperparameters of the Random Forest algorithm and the values used for tuning.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>Values considered for tuning</th>
</tr>
</thead>
<tbody>
<tr>
<td>n_estimators</td>
<td>number of trees to build</td>
<td>50, 75, 100, 150, 200</td>
</tr>
<tr>
<td>criterion</td>
<td>function used to measure the quality of the split</td>
<td>gini, entropy, log_loss</td>
</tr>
<tr>
<td>max_depth</td>
<td>maximum depth of a tree</td>
<td>3, 5, 7, 9</td>
</tr>
<tr>
<td>max_features</td>
<td>number (fraction) of features to check for the best split</td>
<td>log2, sqrt, 0.25, 0.5, 0.75, 0.95</td>
</tr>
<tr>
<td>bootstrap</td>
<td>whether bootstrap samples are used for building the trees</td>
<td>True, False</td>
</tr>
<tr>
<td>class_weight</td>
<td>weights assigned for the classes</td>
<td>None, balanced, balanced_subsample</td>
</tr>
<tr>
<td>ccp_alpha</td>
<td>parameter used for determining which subtree to prune</td>
<td>0, 0.01, 0.02, 0.03, 0.04</td>
</tr>
<tr>
<td>max_samples</td>
<td>the maximum fraction of samples to use if bootstrap is True</td>
<td>0.5, 0.6, 0.7, 0.8, 0.9</td>
</tr>
</tbody>
</table>

Figure 1: Steps of the hyperparameter tuning process.

perparameter space, several runs for that point should be executed. In order to balance the total run time and the exploration of the hyperparameter space, we have decided to use a two-step parameter tuning process, which is presented on Figure 1.

For each of the 16 data sets and each of the three representations, first we have selected 1000 random hyperparameter points and evaluated them using a single run of a 3-fold cross validation on the current data set using the AUC measure. This is called random search-based hyperparameter tuning and is an alternative for grid search which was shown in (Tantithamthavorn et al., 2019) to perform just as well as grid search for SDP.

In the second step of the process, the 20 points with the highest AUC from the first step were selected and re-evaluated. The 3-fold cross validation was repeated 20 times and for each repetition 5 different random splittings into the 3-folds were considered. Consequently, for every point, we had the value of AUC for 100 evaluations. The average of these values was considered to be the evaluation value of that hyperparameter point. Finally, the point with the highest AUC was selected as the best hyperparameter for the given data set and representation.

### 3.3.3 Evaluation Methodology

In order to evaluate the performance of the three supervised models, when fed with different vectorial representations, scaled in [0,1] using Min-Max normalization, we employed the following evaluation methodology. For FastAI and MLP the data was split into 70% train, 10% validation (for early stopping) and 20% test sets. For RF there was no early stopping, so the data was split into 80% train and 20% test set. In order to get consistent results, 30 experiments with different splits had been performed.

During this evaluation process, the confusion matrix for the binary classification task has been computed for each of the 30 testing subsets. Based on the confusion matrix (TP - number of true positives, FN - number of false negatives), the area under the ROC curve (AUC) has been computed as a performance indicator. The reported values have been averaged over the 30 experiments.

We considered AUC as a performance evaluation measure because the SDP literature reveals that AUC is particularly suitable for evaluating the performance of the software defect classifiers (Fawcett, 2006). In general, the AUC measure is employed for approaches that yield a single value, which is then converted into a class label using a threshold. Thus, for each threshold value, the point \((1 - \text{Spec}, \text{Sens})\) is represented on a plot and the AUC is computed as the area under this curve. For the approaches where the direct output of the defect classifier is the class label, there is only one \((1 - \text{Spec}, \text{Sens})\) point, which is linked to the \((0, 0)\) and \((1, 1)\) points. The AUC measure represents the area under the trapezoid and is computed as the mean of sensitivity (Sens) and specificity (Spec): \(\text{AUC} = \frac{\text{Sens} + \text{Spec}}{2}, \text{where} \ \text{Sens} = \frac{TP}{TP + FN}\).
and $Spec = \frac{T_N}{T_N + FP}$. AUC ranges from 0 to 1. Higher values indicate better classification performance.

4 RESULTS AND DISCUSSION

4.1 Difficulty Analysis

In order to answer our first research question presented in Section 1, we have first computed the positive and negative difficulty of the data sets generated by using each of the three embeddings for each of the 16 versions. The computation of the difficulty values was performed according to Section 3.3.1.

The difficulty values computed for the defective class can be seen on Figure 2, while the ones for the non-defective class are represented on Figure 3.

By analyzing Figure 2, where, for a clearer presentation of the differences, the y-axis labels start from 0.5, we can observe that the difficulty for the positive class is quite high. Most values are around 0.6-0.7, the minimum is 0.551 and for some versions and representations it can be as high as 0.83. This means that, on average, around two thirds of the defective instances are closer to a non-defective instance than to a defective one, which makes the correct prediction of defective entities a very difficult task.

If we compare the values for different embeddings, we can see that for 12 versions out of 16, the Graph2Vec representation leads to a lower difficulty than the other two. This suggests that the Graph2Vec embedding captures structural information which could be relevant for identifying defects.

However, analyzing Figure 3 that represents the difficulty for the negative class, besides observing that the values are a lot smaller, most of them below 0.1, we can also observe that this time the Doc2Vec + LSI embedding is the one that produces lower difficulties. This suggests that the Doc2Vec + LSI embedding manages to better capture the characteristics of the non-defective entities.

The fact that one representation produces lower difficulty values for the positive class while the other produces lower difficulty values for the negative class suggests that the two representations capture different aspects of the software entities and thus confirm our intuition that they complement each other by catching together both semantic and syntactic information underlying the source code.

4.2 Supervised Analysis

For performing the comparative analysis from the perspective of supervised learning, we have run the three classification algorithms presented in Section 3.3.2 on all 16 versions of Calcite fed with the three embeddings proposed in Section 3.2.

The AUC values for FastAI, MLP and RF are depicted on Figures 4, 5 and 6, respectively. All the AUC values are the averages for the 30 runs, as presented in Section 3.3.3. While we did not put them on the figures to avoid visual overcrowding, we have computed the 95% confidence intervals (CI) for all AUC values. For FastAI and RF the margin of error is at most 0.03 (although there is one single case for RF where it is 0.08), while for MLP it is at most 0.02.

Even if, due to the lack of space, we do not detail all the numeric results in the current paper, a document with the complete values for all algorithms and all runs is available on Figshare (fig, 2023).

4.2.1 RQ1: The Relative Relevance of Graph2Vec-Based Embedding

As regards RQ1, considering the FastAI algorithm (Figure 4), we can see that the Graph2Vec and Doc2Vec + LSI embeddings lead to quite similar SDP performance; but almost always (15 cases out of 16) Doc2Vec + LSI outperforms Graph2Vec. The same stands if we look at the results of the MLP classifier (Figure 5): the values are close, but Doc2Vec + LSI performs better for 14 Calcite versions. When considering the results of RF (Figure 6), we can notice that Doc2Vec + LSI is better in only 8 cases. Moreover, there are 4 cases (versions 1.5, 1.6, 1.11 and 1.15) where the AUC values for Doc2Vec + LSI are a lot lower than the ones for Graph2Vec. This aspect will be discussed in Section 4.2.3. In conclusion, our answer for RQ1 is that the Graph2Vec embedding performs similarly to Doc2Vec + LSI, but in most cases, irrespectively of the classification algorithm used, it is outperformed by Doc2Vec + LSI.

4.2.2 RQ2: The Potency of Combining Doc2Vec, LSI and Graph2Vec-Based Embeddings

In order to answer RQ2, we look at Figures 4, 5, and 6 again. We can see that, for all three algorithms, in 12 out of 16 cases, the Doc2Vec + LSI + Graph2Vec is the best-performing one. This is summarized on Figure 7. To further verify if the Doc2Vec + LSI + Graph2Vec representation indeed captures information about the characteristics of defective and non-defective entities, we have performed an additional experiment. We run the RF classifier with the best hyperparameter selected for each version of the Calcite software system, but we have randomly shuffled the class labels. In this way, we try to learn from a random data set. If the classifier has, on the random data,
results comparable to the ones on the original one, it suggests that only random noise has been learnt. We have used the same methodology as for the other experiments. 20% of data being randomly selected for testing (the labels being randomly shuffled) and for each version the experiment being repeated 30 times to account for randomness in the classifier. The average AUC values obtained for the 16 versions are between 0.478 and 0.521, so a lot below the AUC values for the original data set, which are above 0.71. This confirms that the \textit{Doc2Vec + LSI + Graph2Vec} embedding captures aspects carrying discriminative information that enables differentiating between defective and defect-free software entities.

Consequently, our answer for RQ2 is that the \textit{Graph2Vec} and \textit{Doc2Vec + LSI} embeddings improves the performance of the SDP model.
4.2.3 RQ3: FastAI Versus Tuned Random Forest as Defect Prediction Models

Our third research question is related to the performance of the hyperparameter-tuned RF compared to FastAI, the deep learning classifier.

To answer this question, we have looked at our results from a different perspective. Instead of comparing the performance obtained by the same classifier for different embeddings, we have compared...
As illustrated in Figure 8, the tuned RF classifier outperforms FastAI for all representations. This confirms the conclusions presented in (Fu and Menzies, 2017) and (Majumder et al., 2018), claiming that properly tuned simpler models can outperform deep learning models. In order to check that the parameter tuning indeed matters, we have repeated the experiments for the Doc2Vec + LSI + Graph2Vec representation (since, according to the conclusions of RQ2, this is the best-performing embedding) considering the default parameters of RF from scikit-learn. The results were all less than 0.55, which is significantly lower than the AUC values achieved for the tuned RF, that are all above 0.71.

As presented in Section 3.3.2 we have used the random search-based parameter tuning approach for RF. The previous experiment demonstrated that default parameters perform quite poor on the Doc2Vec + LSI + Graph2Vec embedding, so we have decided to look at the initially generated 1000 hyperparameter points, to see how varied their AUC values are. For all data sets and representations there were hyperparameter points with an AUC value of 0 (at least 97, at most 462) and a lot of points with an AUC value below 0.5 (at least 162 and at most 995). Actually, there were 4 versions of the Calcite data set, for which there was an exceptionally high number of hyperparameter points with an initial AUC value below 0.5: version 1.5 (995 points), 1.6 (903), 1.11 (844) and 1.15 (929). On Figure 6 we can see that these are exactly the versions with an AUC value a lot less than the other versions and representations. This is probably the result of a random search which produced a lot of hyperparameter points with very poor performance. This suggests that in some cases, random search might not generate good enough points, and probably the number of points should be increased in such cases.

Our findings about how much parameter tuning
matters for RF are different from the findings of Tantithamthavorn (Tantithamthavorn et al., 2019), who concluded that RF is not sensitive to hyperparameter tuning (although they only tuned the parameter denoting the number of trees). The reason for this discrepancy might be given by the difference in the considered features: we have used embeddings learned from the source code, while in (Tantithamthavorn et al., 2019) structural, complexity and size metrics are used. There is another observation regarding RFs in (Tantithamthavorn et al., 2019): they are not among the top-performing classifiers. In our study, RFs had the best performance, but we compared only 3 approaches and only the parameters of RF were tuned, while Tantithamthavorn et al. compared 26 classifiers. Consequently, it is possible that even better performance can be achieved by considering other classifiers.

Consequently, our answer for RQ3 is that, for our data sets, the RF classifier with the best hyperparameter setting outperforms FastAI in most cases. Nevertheless, there were 4 identified cases when the random search parameter tuning approach could not identify good enough parameters and in these cases the performance of FastAI was better than that of RF.

4.2.4 Statistical Tests

In order to test if the conclusions of the three research questions are statistically significant, we have performed some statistical tests. RQ1 and RQ2 focus on the comparison of different source code embeddings, so we have first considered the 48 AUC values for each source code embedding (16 values for each classifier). According to the Kolmogorov-Smirnov test (with Lilliefors’ method) all three variables (i.e. set of AUC values for code embeddings) are normally distributed so we have used a one-way repeated measures ANOVA test. Mauchly’s test indicated that the assumption of sphericity had been violated, therefore degrees of freedom were corrected using Greenhouse-Geisser estimates of sphericity. The results of ANOVA show that the AUC values for the three embeddings differ significantly. Post hoc tests revealed that the Doc2Vec + LSI + Graph2Vec representation has significantly higher AUC values than both other representations. However, there is no significant difference between Doc2Vec + LSI and Graph2Vec embeddings.

For RQ3, we have compared the 16 AUC values for the FastAI and RF classifier for the Doc2Vec + LSI + Graph2Vec embedding. Since the Kolmogorov-Smirnov test showed that both variables are normally distributed, we used a one-way repeated measures ANOVA again (even if we had only two variables) whose results show that the AUC values for RF are significantly higher than those for FastAI.

Consequently, we can conclude that the results for RQ2 and RQ3 are statistically significant.

5 CONCLUSIONS

In this paper, we proposed using Graph2Vec embedding as a new representation of software entities in defect prediction models. Results from several experimental analyses have confirmed the relevance of the proposed representation in identifying defect-prone code. Additionally, when using Graph2Vec embedding in conjunction with Doc2Vec and LSI embeddings, their complementarity boosts the defect prediction performance. In our comparative analysis, we have employed three classification models, including the FastAI deep learning model and a hyperparameter tuned Random Forest, the latter outperforming the former, which reveals the potency of hyperparameter optimisation in the case of software defect predictors.

To reinforce the conclusions of our present study we aim to further extend the experimental analyses by considering additional open-source software systems and machine learning models. We also envision investigating the potency of CodeBERT and graph-CodeBERT for SDP and to reliably compare it to that of our current approach.

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