Detecting Code Smells using Machine Learning Techniques: Are We There Yet?

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Abstract—Code smells are symptoms of poor design and implementation choices weighing heavily on the quality of produced source code. During the last decades several code smell detection tools have been proposed. However, the literature shows that the results of these tools can be subjective and are intrinsically tied to the nature and approach of the detection. In a recent work Arcelli Fontana et al. \cite{1} proposed the use of Machine-Learning (ML) techniques for code smell detection, possibly solving the issue of tool subjectivity giving to a learner the ability to discern between smelly and non-smelly source code elements. While this work opened a new perspective for code smell detection, in the context of our research we found a number of possible limitations that might threaten the results of this study. The most important issue is related to the metric distribution of smelly instances in the used dataset, which is strongly different than the one of non-smelly instances. In this work, we investigate this issue and our findings show that the high performance achieved in the study by Arcelli Fontana et al. was in fact due to the specific dataset employed rather than the actual capabilities of machine-learning techniques for code smell detection.

Index Terms—Code Smells; Machine Learning; Empirical Studies; Replication Study;

I. INTRODUCTION

Nowadays, the complexity of software systems is growing fast and software companies are required to continuously update their source code \cite{2}. Those continuous changes frequently occur under time pressure and lead developers to set aside good programming practices and principles in order to deliver the most appropriate but still immature product in the shortest time possible \cite{3}--\cite{5}. This process can often result in the introduction of so-called technical debt \cite{6}, design problems likely to have negative consequences during the system maintenance and evolution.

One of the symptoms of the technical debt are code smells \cite{7}, suboptimal design decisions applied by developers that can negatively affect the overall maintainability of a software system. Over the last decade, the research community heavily investigated (i) how code smells are introduced \cite{8}, \cite{9}, (ii) how they evolve \cite{10}--\cite{13}, (iii) what is their effect on program comprehension \cite{14}, \cite{15} as well as on the change- and bug-proneness of the affected source code elements \cite{16}, \cite{17}, and (iv) the perception and ability of developers to deal with them \cite{18}--\cite{20}.

Moreover, several code smell detectors have been proposed \cite{21}, \cite{22}: the detectors mainly differ in the underlying algorithm (e.g., metric-based \cite{23}--\cite{26} vs search-based techniques \cite{27}, \cite{28}) and for the specific metric types considered (e.g., product metrics \cite{23}, \cite{24} vs process metrics \cite{26}).

Despite the good performance shown by the detectors, recent studies highlight a number of important limitations threatening adoption of the detectors in practice \cite{21}, \cite{29}. In the first place, code smells detected by existing approaches can be subjectively perceived and interpreted by developers \cite{30}, \cite{31}. Secondly, the agreement between the detectors is low \cite{32}, meaning that different tools can identify the smelliness of different code elements. Last, but not least, most of the current detectors require the specification of thresholds that allow them to distinguish smelly and non-smelly instances \cite{21}: as a consequence, the selection of thresholds strongly influence the detectors’ performance.

To overcome these limitations, machine-learning (ML) techniques are being adopted to detect code smells \cite{1}. Usually a supervised method is exploited, i.e., a set of independent variables (a.k.a. predictors) are used to determine the value of a dependent variable (i.e., presence of a smell or degree of the smelliness of a code element) using a machine-learning classifier (e.g., Logistic Regression \cite{33}).

In order to empirically assess the actual capabilities of ML techniques for code smell detection, Arcelli Fontana et al. \cite{1} conducted a large-scale study where 32 different ML algorithms were applied to detect four code smell types, i.e., Data Class, Large Class, Feature Envy and Long Method. The authors reported that most of the classifiers exceeded 95\% both in terms of accuracy and of F-Measure, with J48 and RANDOM FOREST obtaining the best performance. The authors see in these results an indication that “using machine learning algorithms for code smell detection is an appropriate approach” and that “performances are already so good that we think it does not really matter in practice what machine learning algorithm one chooses for code smell detection” \cite{1}. In our research, we have observed important limitations of the work by Arcelli Fontana et al. \cite{1} that might affect the generalizability of their findings. Specifically, the high performance reported might be due to the way the dataset was constructed: for each type of code smell analyzed, the dataset contains only instances affected by this type of smell or non-smelly instances, with a non-realistic balance of smelly and non-smelly instances \cite{8}, \cite{30} and a strongly different distribution of the metrics between the two groups of instances, which is far from reality.

In this paper, we propose a replicated study on the usage of
ML techniques for code smell detection that aims at addressing the issue related to the metric distribution of smelly and non-smelly elements exploited in the work by Arcelli Fontana et al. [1]. For this reason, we first statistically analyze the metric distribution of smelly and non-smelly code elements; then we replicate the study conducted by Arcelli Fontana et al. [1] on a different dataset containing code elements affected by different types of code smells, with a less balanced distribution of smelly and non-smelly instances and with a more smoothed boundary between the metrics distributions of the two groups of instances, thus depicting a more realistic scenario.

Our results show that the high performance achieved by Arcelli Fontana et al. [1] can be attributed to the dataset exploited, rather than to the real capabilities of prediction models. Indeed, we found the metric distributions of smelly and non-smelly elements in the dataset used by Arcelli Fontana et al. to be very different. When testing code smell prediction models on the revised dataset, we noticed that they are up to 90% less accurate in terms of F-Measure than those reported by Arcelli Fontana et al. [1].

Our findings have important implications for the research community: the problem of detecting code smells through the adoption of ML techniques is far from being solved, and therefore more research is needed to devise proper tools for software engineers.

Structure of the paper. Section II discusses the related literature. Section III describes the reference work and its limitations. Section IV reports the overall methodology adopted. In Section V we report the design and results of the study aimed at analyzing the metric distribution of smelly and non-smelly elements in the dataset used in the reference work, while Section VI discusses how we replicated the reference study and what are the achieved results. Section VII reports possible threats affecting our findings and how we mitigated them. Finally, Section VIII concludes the paper and outlines our future research agenda.

II. RELATED WORK

The problem of detecting code smells [7] in source code has attracted the attention of several researchers over the recent years [21]. The research literature can be roughly divided in two main groups: on the one hand, empirical studies have been performed with the aim of understanding code smell evolution [8]–[13], [34], their perception [18], [19], [35]–[37], as well as their impact on non-functional properties of source code [14]–[17], [38]–[40]. On the other hand, several detection approaches have been devised: most of them rely on the analysis of structural information extractable from the source code [23]–[25], [41], while a recent trend is concerned with the analysis of alternative sources of information [26], [42], [43] or the usage of search-based software engineering methods [27], [28], [44], [45]. In the context of this paper we mainly focused on supervised methods for the detection of design flaws. Thus, in this section we discuss papers leveraging machine-learning models to identify design flaws.

Kreimer [46] originally proposed the use of decision trees for the detection of the Blob and Long Method code smells on two small-scale software systems, finding that such a prediction model can lead to high values of accuracy. The findings were then confirmed by Amorin et al. [47], who tested decision trees over four medium-scale open-source projects. Later on, Vaucher et al. [48] relied on a Naive Bayes technique to track the evolution of the Blob smell, while Maiga et al. [49], [50] devised a SVM approach for the incremental detection of the same smell which is able to reach a F-Measure of ≈80%.

Khomh et al. [51], [52] proposed the use of Bayesian Belief Networks to detect Blob, Functional Decomposition, and Spaghetti Code instances on open-source programs, finding an overall F-Measure close to 60%. Following this direction, Hassaine et al. [53] defined an immune-inspired approach for the detection of Blob smells, while Oliveto et al. [54] relied on B-Splines for understanding the “signatures” of code smells and training a machine learner for detecting them. More recently, machine-learning techniques have been also adapted for the detection of a specific type of code smell, i.e., the Duplicated Code (a.k.a., code clones) [55]–[57].

Arcelli Fontana et al. [1], [58], [59] provided the most relevant advances in this field: in the first place, they theorized that the use of machine-learning might have lead to a more objective evaluation of the harmfulness of code smells [58]. Furthermore, they provided a machine-learning method for the assessment of code smell intensity, i.e., the severity of a code smell instance perceived by developers [59]. Finally, they empirically benchmarked a set of 16 machine-learning techniques for the detection of four code smell types [1]; they performed their analyses over 74 software systems belonging to the Qualitas Corpus dataset [60]. This is clearly the reference work for our study. In their study, they found that all the machine learners experimented achieved high performance in a cross-project scenario, with the J48 and RANDOM FOREST classifiers obtaining the highest accuracy. Perhaps more importantly, they discovered that a hundred training examples are enough for reaching at least 95% accuracy.

Our study aims at addressing one of the limitations of this work, to understand whether the problem of detecting code smells using ML techniques can be actually considered solved.

III. THE REFERENCE WORK

The reference work of our replication is the one by Arcelli Fontana et al. [1]. The authors analyze three main aspects related to the use of machine-learning algorithms for code smell detection: (i) performance of a set of classifiers over a sample of the total instances contained in the dataset, (ii) analysis of the minimum training set size needed to accurately detect code smells, and (iii) analysis of the number of code smells detected by different classifiers over the entire dataset.

In this paper, we focus on the first research question of the reference work. As discussed later in this section, we have identified important limitations that might have led to biased results. In the following subsections we detail the methodological process adopted by Arcelli Fontana et al. [1].
A. Context Selection

The context of the study by Arcelli Fontana et al. was composed of software systems and code smells.

The authors have analyzed systems from the Qualitas Corpus [60], release 20120401r, one of the largest curated benchmark datasets to date, specially designed for empirical software engineering research. Among 111 Java systems of the corpus, 37 were discarded because they could not be compiled and therefore code smell detection could be applied. Hence, the authors focused on the remaining 74 systems.

For each system 61 source code metrics were computed at class level and 82—at method level. The former were used by Arcelli Fontana et al. as independent variables for predicting class-level smells Data Class and God Class, the latter for predicting method-level smells Feature Envy and Long Method:

1) **God Class.** It arises when a source code class implements more than one responsibility; it is usually characterized by a large number of attributes and methods, and has several dependencies with other classes of the system;

2) **Data Class.** This smell refers to classes that store data without providing complex functionalities;

3) **Feature Envy.** This is a method-level code smell that appears when a method uses much more data than another class with respect to the one it is actually in;

4) **Long Method.** It represents a large method that implements more than one function;

The choice of these smells is due to the fact that they capture different design issues, e.g., large classes or misplaced methods.

B. Machine-learning Techniques Experimented

Arcelli Fontana et al. [1] evaluated six basic ML techniques: J48 [61], JRIP [62], RANDOM FOREST [63], NAIVE BAYES [64], SMO [65], and LibSVM [66]. As for J48, the three types of pruning techniques available in WEKA [67] were used, SMO was based on two kernels (e.g., POLYNOMIAL and RBF), while for LibSVM eight different configurations, using C-SVC and v-SVC, were used. Thus, in total Arcelli Fontana et al. [1] have evaluated 16 different ML techniques. Moreover, the eight ML techniques were also combined with the ADABOOSTM1 boosting technique [68], i.e., a method that iteratively uses a set of models built in previous iterations to manipulate the training set and make it more suitable for the classification problem [69], leading to 32 different variants.

An important step for an effective construction of machine-learning models consists in the identification of the best configuration of parameters [70]: the authors applied to each classifier the Grid-search algorithm [71], capable of exploring the parameter space to find an optimal configuration.

C. Dataset Building

To establish the dependent variable for code smell prediction models, the authors applied for each code smell the set of automatic detectors shown in Table I. However, code smell detectors cannot usually achieve 100% recall, meaning that an automatic detection process might not identify actual code smell instances (i.e., false positives) even in the case that multiple detectors are combined. To cope with false positives and to increase their confidence in validity of the dependent variable, Arcelli Fontana et al. [1] applied a stratified random sampling of the classes/methods of the considered systems: this sampling produced 1,986 instances (826 smelly elements and 1,160 non-smelly ones), which were manually validated by the authors in order to verify the results of the detectors.

As a final step, the sampled dataset was normalized for size: the authors randomly removed smelly and non-smelly elements building four disjoint datasets, i.e., one for each code smell type, composed of 140 smelly instances and 280 non-smelly ones (for a total of 420 elements). These four datasets represented the training set for the ML techniques above.

D. Validation Methodology

To test the performance of the different code smell prediction models built, Arcelli Fontana et al. [1] applied ten-fold cross validation [72]: each of the four datasets was randomly partitioned in ten folds of equal size, such that each fold has the same proportion of smelly elements. A single fold was retained as test set, while the remaining ones were used to train the ML models. The process was then repeated ten times, using each time a different fold as the test set. Finally, the performance of the models was assessed using mean accuracy, F-Measure, and AUC-ROC [73] over the ten runs.

E. Limitations and Replication Problem Statement

The results achieved by Arcelli Fontana et al. [1] reported that most of the classifiers have accuracy and F-Measure higher than 95%, with J48 and RANDOM FOREST being the most powerful ML techniques. These results seem to suggest that the problem of code smell detection can be solved almost perfectly through ML approaches, while other unsupervised techniques (e.g., the ones based on detection rules [24]) only provide suboptimal recommendations.

However, we identified possible limitations of the work by Arcelli Fontana et al. [1] that might have threatened this view:

1) **Selection of the instances in the dataset.** A first factor possibly affecting the results might be represented by the characteristics of smelly and non-smelly instances present in the four datasets exploited (one for each smell type): in particular, if the metric distribution of smelly elements is strongly different than the metric distribution of non-smelly instances, then any ML technique might easily distinguish the two classes. Clearly, this does not properly represent a real-case scenario, where the boundary between the structural characteristics of smelly and non-smelly
code components is not always clear [8], [30]. In addition, the authors built four datasets, one for each smell. Each dataset contained code components affected by that type of smell or non-smelly components. This also makes the datasets unrealistic (a software system contains different types of smells) and might have made easier for the classifiers to discriminate smelly components.

2) **Unrealistically balanced dataset.** In the reference work, one third of the instances in the dataset was composed of smelly elements. According to recent findings on the diffuseness of code smells [17], software systems are usually affected by a small percentage of code smells. For instance, Palomba et al. [17] found that God Classes represent less than 1% of the total classes in a software system.

3) **Biased validation methods.** The authors applied the ten-fold cross validation on a balanced dataset. The problem raises because in this way both the training and the test sets are balanced, thus leading the model to be exercised and evaluated on a test set having much more smelly instances than in reality.

4) **Missing analysis of relevant features.** As reported in literature [74], building a model containing independent variables that are highly correlated with each other might lead to model over-fitting, leading to biased results.

Because of the points above, we argue that the capabilities of ML techniques for code smell detection should be reevaluated. In this paper, we start addressing this challenge by focusing on the first two issues, i.e., instance selection in the dataset and unrealistic dataset balancing.

### IV. Empirical Study Definition

The goal of the empirical study reported in this paper was to analyze the sensitivity of the results achieved by our reference work, i.e., the study by Arcelli Fontana et al. [1] with respect to the metric distribution of smelly and non-smelly instances, with the purpose of understanding the real capabilities of existing prediction models in the detection of code smells. The perspective is of both researchers and practitioners: the former are interested in understanding possible limitations of current approaches in order to devise better ones; the latter are interested in evaluating the actual applicability of code smell prediction in practice.

We pose the following research questions:

- **RQ1.** What is the difference in the metric distribution of smelly and non-smelly instances in the four datasets exploited in the reference work?
- **RQ2.** What is the performance of ML techniques when trained on a more realistic dataset, containing different types of smells, with a reduced proportion of smelly components and with a smoothed boundary between the metric distribution of smelly and non-smelly components?

**RQ1** can be considered as a preliminary analysis aimed at assessing whether and to what extent the metric distribution of smelly and non-smelly instances in the datasets used by Arcelli Fontana et al. [1] is different. With **RQ2** we aim at assessing the performance of code smell prediction techniques in a more realistic setting where the differences between smelly and non-smelly instances are less prominent.

To enable a proper replication, the context of our study was composed of the same dataset and code smells used by the reference work. Thus, we took into account the 74 software systems and the four code smells discussed in Section III.

### V. RQ1—Metrics Analysis

We start by comparing the distributions of metrics over smelly and non-smelly code elements (**RQ1**).

#### A. Design

To answer this research question, we compared the distribution of the metrics representing smelly and non-smelly instances in the dataset exploited by Arcelli Fontana et al. [1]. Given the amount of metrics composing class- and method-level code smells, i.e., 61 and 82, respectively, an extensive comparison would have been prohibitively expensive, other than being not practically useful since not all the metrics actually have an impact on the prediction of the dependent variable [74]. For this reason, we first reduce the number of independent variables by means of feature selection, i.e., we consider only the metrics impacting more the prediction of code smells. To this aim, we employed the widely-adopted *Gain Ratio Feature Evaluation* algorithm [75], identifying the features having more weight in the code smell detection. Let \( M \) be a code smell prediction model, \( P = \{m_1, \ldots, m_n\} \) be the set of independent variables composing \( M \), then the *Gain Ratio Feature Evaluation* algorithm [75] computes the difference in entropy before and after the set \( M \) is split on the metrics \( m_i \):

\[
\text{GainRatio}(M, m_i) = H(M) - H(M|m_i),
\]

where \( H(M) \) indicates the entropy of the model that includes the metric \( m_i \), \( H(M|m_i) \) the entropy of the model that does not include \( m_i \), and the entropy is computed as

\[
H(M) = - \sum_{i=1}^{n} \text{prob}(p_i) \log_2 \text{prob}(p_i).
\]

The algorithm quantifies the extent of uncertainty in \( M \) that was reduced after considering \( M \) without \( m_i \). The output of the algorithm is represented by a ranked list—ranks ranging between 0 and 1—in which the metrics having the higher expected reduction in entropy are placed at the top, i.e., the metrics giving more weight to the prediction are ranked first. As cut-off point of the ranked list we selected 0.1, as recommended by Quinlan [75]. The output of this process consisted of the set of metrics really relevant for the prediction.

Afterwards, we compared the distribution of the metrics representing smelly and non-smelly instances among them using the Wilcoxon Rank Sum statistical test [76] with \( p \)-value = 0.05 as significance threshold. Since we performed
multiple tests, we adjusted \( p \)-values using the Bonferroni-Holm's correction procedure [77]. The procedure firstly sorts the \( p \)-values resulting from \( n \) tests in ascending order of values, multiplying the smallest \( p \)-value by \( n \), the next by \( n - 1 \), and so on. Then, each resulting \( p \)-value is then compared with the desired significance level (e.g., 0.05) to determine whether or not it is statistically significant. Furthermore, we estimated the magnitude of the observed differences using Cliff’s Delta (or \( d \)), a non-parametric effect size measure [78] for ordinal data. To interpret the effect size values, we follow established guidelines [78]: small for \( d < 0.33 \) (positive as well as negative values), medium for \( 0.33 \leq d < 0.474 \) and large for \( d \geq 0.474 \).

If the differences in the metric distributions of smelly and non-smelly instances are statistically significant and with a large effect size, then the two classes can be too easily classified for any machine-learning technique.

### B. Results

#### Table II

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<td>&lt; 0.01 Medium</td>
<td>&lt; 0.01 Large</td>
<td>&lt; 0.01 Medium</td>
<td>&lt; 0.01 Large</td>
</tr>
<tr>
<td>36</td>
<td>- -</td>
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</tbody>
</table>

Table II reports the comparison of the metric distribution of smelly and non-smelly instances. For sake of readability, we avoid adding the complete metric names in the table. However, our online appendix contains the information about the relevant metrics for each dataset. It is important to note that the feature selection process detected approximately one third of the total features as relevant: this actually means that most of the metrics present in the original dataset did not impact the prediction of code smells, but rather might have caused overfitting of the model.

It is possible to notice that in most cases the differences between the distributions are statistically significant. Analyzing \( p \)-values, we can notice that in 26 of 34 considered metrics there is a significant difference (14 with Large effect size). Moreover looking at the first 10 features in terms of \( \text{GainRatio} \), we can notice that in 8 out 10 case there is a Large effect size.

More evident results are observable when analyzing \( \text{God Class} \). In this case, 35 out of 36 metrics distributions are statistically different (\( p \)-value < 0.05). In particular, in 29 cases the distributions are different with a Large effect size, while in 3 cases the effect size is medium.

Analyzing the method-level code smells (e.g., Feature Envy and Long Method), we can notice similar results. Indeed, in the first case, 96% of the metrics distributions filtered with \( \text{GainRatio} \) are statistically different (73% with Large effect size), while in the second case, 94% of metrics distributions are statistically different (74% with Large effect size).

These results demonstrate that smelly and non-smelly instances selected in the original dataset are clearly different. Hence, the selected instances could lead to overestimate the performances of machine learning techniques in the context of bad smell prediction.

#### Summary for \( \text{RQ1} \).

Based on the results of our \( \text{RQ1} \), we decided to modify the dataset in order to have (i) a less strong difference in the metrics distribution, (ii) a less balanced dataset between smelly and non-smelly instances, and (iii) different types of smells in the same dataset so that we can model a more realistic scenario [17].

#### A. Design

To answer our second research question, we replicated the reference study after setting up the original dataset. Specifically, we designed our replication in two steps, described below.

**Dataset Setup.** In \( \text{RQ1} \), we found that the two sets of elements to predict, i.e., smelly and non-smelly ones, are almost totally disjoint from a statistical point of view and hence any machine-learning approach can be expected to easily discriminate them. To verify the actual capabilities of the code smell prediction models experimented in the reference study, we built a dataset closer to reality by merging the instances contained in the four original datasets and replicated the study by Arcelli Fontana et al. [1] on such a new dataset. In this way, we created a new dataset including source code elements affected by different smells, thus generating a more realistic scenario in which different code elements having a similar metric profile are affected by different design issues.

To this aim, we firstly merged the datasets regarding \( \text{God Class} \) and \( \text{Data Class} \) for class-level code smells and secondly the ones related to \( \text{Feature Envy} \) and \( \text{Long Method} \) for method-level ones. Afterwards, we duplicated the datasets so that we
could have (i) two class-level datasets in which the dependent variable was represented by the presence of God Class or Data Class, respectively, and (ii) two method-level datasets having as dependent variables the presence of Feature Envy or Long Method, respectively.

**Replicating the study.** Once we had created the four datasets, we performed exactly the same experiment as done in the reference work, thus using the same machine-learners configured in the same way. Finally, to measure the performance of the different machine-learning algorithms and compare them with the one achieved by Arcelli Fontana et al. [1], we computed the mean accuracy, F-Measure, and AUC-ROC achieved over 10 runs, as done in the reference work.

![Figure 1. Data Class: Bar charts of Accuracy, F-Measure, and AUC-ROC achieved by the methods under study on the dataset we created.](image1)

**Figure 1.** Data Class: Bar charts of Accuracy, F-Measure, and AUC-ROC achieved by the methods under study on the dataset we created.

**B. Results**

Before discussing the results of this research question, it is important to note that we replicated the statistical analyses made in RQ1 on the merged dataset we created. We reported the detailed results in our online appendix [79], however we found that in our datasets smelly and non-smelly elements were much less different in terms of metric distribution than the original dataset: most of the differences between the distributions of smelly and non-smelly elements are not statistically significant or have a small/negligible effect size.

![Figure 2. God Class: Bar charts of Accuracy, F-Measure, and AUC-ROC achieved by the methods under study on the dataset we created.](image2)

**Figure 2.** God Class: Bar charts of Accuracy, F-Measure, and AUC-ROC achieved by the methods under study on the dataset we created.
created. For sake of space limitations, we made available the fine-grained results in our online appendix [79]. Note that due to computational reasons, in our experiment we limited the execution time to 10 minutes: N/A values in the figures indicate that a certain classifier did not complete the computation in the given time slot. Moreover, we could not run the nu-based VSM models because (i) the default parameter set in Weka was too high to allow the model to complete the execution in the considered time slot and (ii) the parameter values assigned by Arcelli Fontana et al. [1] was not specified.

As a first point of discussion, we noticed that the accuracy of all the models is still noticeable high when compared to the results of the reference study (on average, 76% vs 96%): this is mainly due to the characteristics of the accuracy metric, which takes into account the ability of a model to classify true negative instances, possibly leading to a misinterpretation of the performance a prediction model [80]. It is worth noting that non-smelly elements in each dataset are 5/6 of the total instances (whereas in the datasets by Arcelli Fontana et al. they were 2/3 of the instances).

However, the F-Measure tells a different story. Indeed, the results are 90% lower than in the reference work (e.g., Random Forest), indicating that the models were not actually able to properly classify the smelliness of the analyzed code elements.
This result holds for all the code smell types considered, thus confirming that the high performance achieved by Arcelli Fontana et al. [1] was due to the dataset selection rather than to the real capabilities of the experimented models.

The best performance (for all the smells) is achieved by the tree-based classifiers, i.e., RANDOM FOREST and J48: this confirms the results of the reference study, which highlighted how this type of classifiers perform better than the others.

Finally, the results for AUC-ROC contrast with the ones achieved in the reference study. While Arcelli Fontana et al. [1] reported that basically all the classifiers had an AUC-ROC ranging from 95% to 99%, we found that instead the choice of the ML approach might be highly relevant for effectively detecting code smells, thus paving the way for a more sophisticated way to combine machine-learning approaches for an effective detection of code smells.

Summary for RQ2. The performance of code smell prediction models is up to 90% lower than the one reported in the reference study. High performance reported in the reference study can be therefore mainly attributed to the specific dataset employed rather than to the capabilities of ML techniques for code smell detection.

VII. Threats to Validity

In this section we discuss the threats that might have affected our empirical study and how we mitigated them.

Threats to construct validity. As for potential issues related to the relationship between theory and observation, a first discussion point regards the dataset used in the study. Specifically, we exploited the same dataset used by Arcelli Fontana et al. [1] in order to rely on their classification of the dependent variable (i.e., the smelliness of source code elements), thus reducing the bias of a different manual classification. Of course, we cannot exclude possible imprecisions contained in the Qualitas Corpus dataset [60], e.g., imprecisions in the computations of the metrics for the source code elements exploited in this study.

In the context of RQ1, before comparing the distribution of the metrics in the dataset we applied a feature selection algorithm named Gain Ratio Feature Evaluation. Doing this, we did not compare all the metrics distributions, but we only limited the analysis to the relevant ones, i.e., approximatively one third of the total features. However, the missing extensive comparison does not represent an issue for our results. Indeed, the unselected features do not have an impact on the dependent variable: even in case of negligible differences between non-relevant metrics distributions of smelly and non-smelly instances, this would have not affected the performance of the experimented prediction models.

As for the experimented prediction models, we exploited the implementation provided by the WEKA framework [67], which is widely considered as a reliable tool. Moreover, to faithfully replicate the empirical study by Arcelli Fontana et al. [1] we adopted the same classifiers and their best configurations.

Threats to conclusion validity. Threats in this category impact the relation between treatment and outcome. A first discussion point is related to the validation methodology: in particular, we adopted the 10-fold cross validation. We are aware of the existence of other validation methodologies that might possibly provide a better interpretation of the real performance of code smell prediction models [81], however we choose a multiple 10-fold cross validation in order to directly compare our results with those achieved in the reference work by Arcelli Fontana et al. [1]. Future effort will be devoted to establish the impact of the validation technique on the results.

We are also aware of other possible confounding effects like (i) data unbalance [82] and (ii) wrong data scaling [83]. However, we preferred replicate the reference study using the same methodology. As part of our future work, we plan to assess the role of such preprocessing techniques on the results achieved.

As for the evaluation metrics adopted to interpret the performance of the experimented models, we adopted the same metrics as Arcelli Fontana et al. [1]. We are aware that measures like AUC-ROC and MCC have been highly recommended by Hall et al. [80] since they are threshold-independent. Also in this case, we aimed at replicating as closer as possible the reference study to demonstrate its limitation, but we plan to investigate more in depth the performance of code smell prediction models.

Threats to external validity. With respect to the generalizability of the findings, we took into account one of the largest datasets publicly available, and containing 74 software projects coming from different application domains and with different characteristics. We are aware that our findings may and may not be directly applicable to industrial environments, however the replication of our study on closed-source and industrial projects is part of our future research agenda.

VIII. Conclusion and Future Directions

In this work, we presented a replicated study of the work by Arcelli Fontana et al. [1]. We highlighted some limitations of this study and started addressing the issue related to the construction of the dataset, which contained smelly and non-smelly elements clearly distinguishable for any machine learning approach. As a result, we found that the problem of detecting code smells using machine learning models is still far from being solved, and therefore more research is needed toward this direction.

As future work, we firstly plan to assess the impact of (i) dataset size, (ii) feature selection, and (iii) validation methodology on the results of our study. At the same, we aim at addressing these issues, thus defining new prediction models for code smell detection.

REFERENCES
