# Predicting The Energy Consumption Level of Java Classes in Android Apps: An Exploratory Analysis

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#### **ABSTRACT**

Mobile applications usage has considerably increased since the last decade. Successful apps need to make the users feel comfortable while using them, thus demanding high-quality design and implementation. One of the most influencing factors for user experience is battery consumption, which should have the minimum possible impact on the battery. The current body of knowledge on energy consumption measurement only reports approaches relying on complex instrumentation or stressing the application with many test scenarios, thus making it hard to measure energy consumption in practice. In this work, we explore the performance of machine learning to predict the energy consumption level of JAVA classes in Android apps, leveraging only on a set of structural properties extracted via source code analysis, without requiring any hardware measurements tools or executing the app at all. The preliminary results show the poor performance of learning-based estimation models, likely caused by (1) an insufficient amount of training data, (2) a limited feature set, and (3) an inappropriate way to label the dependent variable. The paper concludes by presenting the limitations of the experimented models and the possible strategies to address them.

## **CCS CONCEPTS**

• Software and its engineering → Software maintenance tools.

## **KEYWORDS**

Energy Consumption, Machine Learning, Android Apps

#### **ACM Reference Format:**

Emanuele Iannone, Manuel De Stefano, Fabiano Pecorelli, and Andrea De Lucia. 2018. Predicting The Energy Consumption Level of Java Classes in Android Apps: An Exploratory Analysis. In Woodstock '18: ACM Symposium on Neural Gaze Detection, June 03–05, 2018, Woodstock, NY. ACM, New York, NY, USA, 5 pages. https://doi.org/10.1145/1122445.1122456

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MobileSoft'22, May 22–23, 2022, Pittsburgh, PA, USA © 2018 Association for Computing Machinery. ACM ISBN 978-1-4503-XXXX-X/18/06...\$15.00 https://doi.org/10.1145/1122445.1122456 Manuel De Stefano SeSa Lab - University of Salerno, Italy madestefano@unisa.it

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#### 1 INTRODUCTION

Mobile applications (a.k.a. apps) have changed our lives and habits, and their use has been increasing at a very high pace [7, 32]. Users need to be comfortable with them in terms of functional and nonfunctional aspects, and one of the main aspects influencing the user's satisfaction is energy consumption [40], that is one of the main reasons a user uninstalls an application [25, 43].

In this regard, previous studies have shown that this is one of the main motivations that could lead users to uninstall an application Therefore, measuring the consumption of mobile apps is a key factor for their success. In the last decade, several techniques have been proposed in this respect. On the one hand, there exists purely hardware-based solutions [11, 18], which rely on special instrumentation to directly measure the amount of energy drained from the battery when the app is exercised with specific test scenarios. On the other hand, model- or software-based solutions [9, 14, 28, 33, 42] leverages on the APIs offered by the device, which compute an estimate of the energy consumed using the sensors installed.

Although these solutions have proven reliable and precise, they need to run realistic test cases that stress the different parts of the code, which are not easy to define. Moreover, hardware-based solutions require costly equipment, which can also be difficult to set up and run. In this respect, the current state of the practice lacks lightweight and rapid measurements techniques, which can be run without any particular requirements and in a short amount of time. Such techniques may address both the developers' and users' need to have early feedback on the app energy consumption level.

In this paper, we investigate the effectiveness of predicting the energy consumption level of classes in mobile apps by the use of machine learning algorithms that leverage structural properties i.e., code metrics and smells-extracted via source code analysis. In this respect, previous works have shown the impact that specific design and implementation decisions have on the consumption of applications [10, 17, 31], motivating their use as features for a prediction model. This investigation is carried on the dataset by Palomba et al. [31], made of 620 JAVA classes belonging to 60 open-source Android apps. The preliminary results show that our machine learning-based estimation models failed, in most cases, to assign the right energy consumption level. Future work should provide additional insights on the efficiency of such energy estimation models, such as by (1) building a dataset with additional training data, (2) identifying new features affecting the energy consumption, and (3) exploring new strategies for making more meaningful and interpretable predictions.

To summarize, this paper (1) presents an empirical comparison of four different machine learners having the goal to predict the energy consumption of JAVA classes in Android apps by leveraging code smells and code metrics, and (2) provides an online appendix [2] containing all the data and scripts employed to foster replications and further experiments built upon this study.

# 2 RELATED WORK

Energy consumption measurement of mobile apps has been the subject of several studies. In recent years the research has been focused on envisioning tools and mechanisms to measure the battery drain reliably and efficiently. On the one hand, there are hardware-based solutions such as POWERSCOPE [11] and GREENMINER [18], which rely on circuits and physical measurement tools (e.g., a multimeter); on the other hand, there exists software solutions that require little or no equipment at all, e.g., PowerBooter [42], eLens [14], or PETRA [9]. Almost all approaches calculate the consumption with direct measurements on the device's battery or via sensor data APIs. However, there exist alternative ways to this purpose, such as the one proposed by Gupta et al. [13], who built a linear regression model to estimate the battery consumption of Windows Phone apps based on the .dll files called. Similarly, Aggarwal et al. [1] relied on the number of system calls and the number of times they were changed to predict the energy consumption of two Android apps.

Several empirical studies have shown other factors that negatively or positively affect energy consumption. The way design patterns are implemented [37], the choice of certain data structures [15], and the presence of specific types of code smells [6, 16, 31] may largely decrease the energy efficiency of mobile apps. Conversely, there is also evidence of the beneficial effects of good programming practices, such as query optimization [35], the use of lock-free data structures [19], and an appropriate management of resources [22]. Violations to such practices have been branded as Android-specific code smells by Reimann et al. [36]. Other choices, such as calls to specific Android API methods [23], screen color palettes [24], and sorting algorithms [5] have a variable impact, depending on how they are applied.

**Our contribution.** This work increases the granularity level of such predictions by providing estimates at the class-level of Android apps. We rely on software metrics and characteristics that can be computed via the sole static code analysis of the Java classes that constitute an app. We also compare the performance of different machine learning algorithms.

# 3 PRELIMINARY EVALUATION

The *goal* of the study is to predict the energy consumption level of Java classes in the *context* of Android apps with the *purpose* of making rapid and lightweight estimates by using structural properties of the source code. Specifically, we analyzed the performance of different machine learning algorithms on making multi-class classification, i.e., predicting which is the overall energy consumption level, at the *class* granularity level, irrespective of the belonging Android app. We opted for a classification as its predictions provide more *meaningful* and *interpretable* results than a regression model, which would estimate the consumption using a continuous value (measured in Joule). Based on these considerations, we asked:

**RQ.** How good is a machine-learning-based classification model in predicting the energy consumption level of JAVA class in ANDROID apps?

# 3.1 Context

The *context* of the study consists of 60 real-world open-source Android apps whose energy consumption was measured in the work by Palomba et al. [31] These apps have differs in terms of size and scope and come from the F-Droid repository. The dataset contains the estimated energy consumption of methods of 620 public classes belonging to the 60 apps. The estimates were done with PETRA [9], a fully software-based energy measurement tool that stress each method with multiple runs of randomly generated test cases to profile the energy consumption, measured in Joule, according to the device's battery sensors. The method-level consumption values of all the repeated runs had been aggregated using the mean since no outlier was found in the measurements distributions.

In addition, the dataset reports the number of occurrences of 19 types of code smells affecting each class. Specifically, five of which are regular object-oriented smells that the authors detected using the DECOR detection rules [26], while the remaining 14 are specific to the Android context—as defined in the catalog<sup>4</sup> of Reimann et al. [36]—and detected using the tool ADOCTOR [30].

The described dataset was used to train and test several prediction models, built using three learning algorithms: *Multinomial Logistic Regression* (MLR) [3], *Support Vector Machine* (SVM) [8], and *Decision Tree* (DT) [4], as they are widely adopted in classification tasks. We involved a *random multi-class classifier*—which assigns a random energy consumption class to a given data point—as comparison baseline for the models' performances. We exploited the implementation of these models provided by the TidyModels<sup>5</sup> package for the R programming language.

### 3.2 Study Design

In the following we describe how we arranged the experimental process, starting from the selection of the dependent variable and the independent variables, to the definition of the entire training and testing pipeline, alongside the metrics used to evaluate the models' performances.

**Dependent Variable.** Since our study focuses on predicting the energy consumption at the *class* granularity level, we brought the method-level energy measurements—contained in the input dataset—to the class level by aggregating the consumption of all the methods belonging to a given class. This choice was driven by the fact that our goal was to investigate whether static factors can effectively predict energy consumption. Unfortunately, most of the static factors that have been associated with energy consumption in the past are all defined at a class-level granularity, hence we had to aggregate method-level observations to be able to build our

<sup>&</sup>lt;sup>1</sup>F-Droid website: https://f-droid.org/

<sup>&</sup>lt;sup>2</sup>Generated with Monkey: https://developer.android.com/studio/test/monkey

<sup>&</sup>lt;sup>3</sup>Collected with BATTERYSTATS: https://tinyurl.com/batterystats

<sup>&</sup>lt;sup>4</sup>The smells catalog:https://martinbrylski.github.io/android\_smells/

<sup>&</sup>lt;sup>5</sup>TidyModels website: http://tidymodels.org

models. More details are reported in Section 3.4. We took into consideration three ways of aggregation: mean, sum, and maximum of the methods of a class. We applied a set of pre-processing steps to prepare the data for the classification task. We first normalized the dependent variable (for each aggregation) using the log transformation [20] to make its distribution more similar to a Gaussian one. Then, we mapped the normalized values to their percentiles and assign an energy consumption level based on the quartiles, so that each bin contains the same amount of classes. Specifically:  $[min..Q_1[\rightarrow `Negligible', [Q_1..Q_2[\rightarrow `Low', [Q_2..Q_3[\rightarrow `Moderate', and [Q_3..max] \rightarrow `High'.]]$ 

**Independent Variables.** The occurrences of the 19 code smells appearing in the dataset were treated as *independent variables* for our models, as some of them have been shown to be positively correlated with the energy consumption—i.e., the higher the number of instances, the higher the battery drain. This is the case of *Internal Setter, Leaking Thread*, and *Member Ignoring Method* Android-specific code smells, which can affect the consumption up to 87 times more than non-smelly classes.

Once we had collected the smell data, we mined additional metrics concerning structural code properties—such as size, cohesion, coupling, and complexity. These metrics were computed via inspection of the Abstract Syntax Trees (ASTs) of all the 620 Java classes found in the dataset. Specifically, we used an our own script (available in our online appendix [2]), to extract 22 different metrics, which we considered as additional *independent variables* of the models. We suspect that these kind of metrics may give additional guidance to our prediction models, as hinted by Hindle [17]. For example, long or complex classes may affect energy consumption since they requires executing more code or more complex programming constructs. To summarize, we involved 41 different independent variables, whose descriptions are all reported in our online appendix due to space limitations [2].

Machine Learning Pipeline. After collecting the dependent and independent variables, we configured the machine learners for predicting the energy consumption. We evaluated the four learning algorithms on the same dataset using a 10-fold cross-validation strategy [39], in which the dataset is divided into 10 random subsets of approximately equal size (folds). The training and testing phases were repeated 10 times. Iteratively, nine folds were used to train the models, while the remaining one was used as the test set, ensuring that in each iteration a different permutation of folds is used. Before running the training phase of each iteration, we applied a feature selection strategy to remove those predictors (1) showing nearzero variance (i.e., predictors having the same value in almost all data points), and (2) having large absolute correlations (in this case 0.6)—using the Spearman's rank correlation coefficient [38]—with other predictors. An example of near-zero variance predictor is one that, for 1000 samples, has two distinct values and 999 of them are a single value. These steps were accomplished to remove those variables that would have given poor or no information to the models, and mitigate the effect of multi-collinearity [29]. Then, we took into account the tuning of models' hyper-parameters by running the GRID SEARCH algorithm. Internally, it relies on another cross validation to systematically search the best combination of

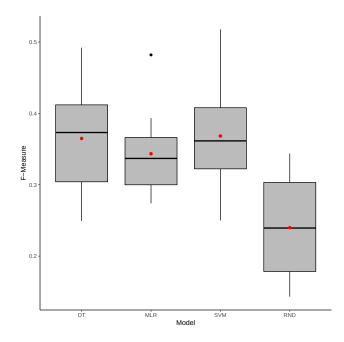


Figure 1: Box plots representing the F-measure values obtained during the 10-fold cross validation of the four classifiers selected, in the aggregation by sum case. The straight horizontal lines in the boxes indicate the median, while the red dots the mean.

hyper-parameters, so that a given error measure is minimized—here we selected the F-measure [34]. Finally, we trained the models on the test set using the optimal hyper-parameter configuration, and run the predictions on the test set—without considering the predictors dropped after the feature selection done previously.

Evaluation Metrics. The prediction results were used to obtain the confusion matrix, reporting the outcomes for each energy consumption level (i.e.,  $4 \times 4$  matrix). For each class c, we computed Precision, Recall, and F-measure performance metrics [34] for the multi-class classification task, obtaining their one-vs-all version (a.k.a., macro measures), i.e., when treating c as the positive class and the remaining ones as a single negative class. Section 3.3 discusses the achieved results only in terms of F-measure distributions obtained by the 10-fold cross validation on the sum aggregation, leaving the other results in our online appendix [2]. The F-measure distributions are represented via box plots and compared using the Friedman test [12] with the Nemenyi post-hoc test [27], and reported the results using MCB plots (Multiple comparisons with the best) [21]. The Friedman test is a non-parametric equivalent to ANOVA which does not assume any distribution of the data. It allows us to assess the significance of the different outcomes achieved across multiple test attempts (in our case, datasets) when using different treatments (in our case, the various models). We employed the confidence level  $\alpha$  = 0.95 [12]. All the study materials can be found on the online appendix[2].

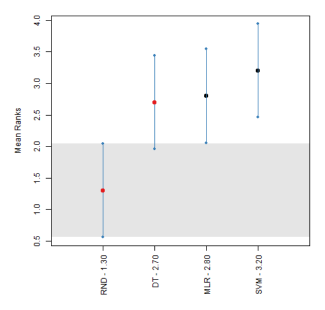


Figure 2: Nemenyi test results for statistical significance between the F-measure values obtained during the 10-fold cross validation of the four classifiers selected in the case of aggregation by usum. The results are presented by mean of MCB plots [21].

### 3.3 Results

Figure 1 shows boxplots reporting the F-measure values achieved by Decision Tree (DT), Multinomial Logistic Regression (MLR), Support Vector Machine (SVM), and the Random Classifier (RND), respectively. We report only the case of aggregation by sum, since it is the one performing better than the others, however similar. According to the figure, the classification performance appears to be pretty low, regardless of the classifier, with an F-measure ranging between 0.30 and 0.40. When turning to the comparison between the classifiers, SVM appears to perform slightly better than the other two while the Random Classifier achieves a clearly lower f-measure, thus indicating that, poor as it is, the performance of the proposed classification technique is still better than a simple random classification. These observations are also confirmed by the results of Nemenyi test, as reported in Figure 2, in which SVM and MLR appear to perform significantly better than both DT and the Random Classifier. However, given the preliminary nature of the study, we can not draw certain conclusions.

The poor performances might be the consequence of several choices. Firstly, the dataset we exploited is made of only a scarce amount of data points (the JAVA classes), limiting what the models learned from them. This urges the need for richer datasets having a large amount of energy measurements, to enable machine learning-based energy estimation. Secondly, we considered a limited set of features, motivated by the availability of tools able to compute them in short time and without running or building the apps. As a matter of fact, the literature showed the existence of other static

properties correlated with the energy consumption of apps, that should be involved in future work. Finally, the performance may be influenced by the splitting criterion adopted to assign each class to the corresponding energy consumption level. The splitting was performed by dividing the variable in quartiles according to its distribution, hence being strongly related to other observation in the dataset. More details about this aspect are reported in Section 3.4.

#### 3.4 Limitations

One of the study's major limitations lies in the calculation of our dependent variable. While the experimental dataset provides method-level measurements for the energy consumption, we aggregated such measures by computing, for each class, the mean, the sum, and the maximum of the energy consumption for all the methods in it contained. This choice could have biased our results since an aggregated measure may not represent the class energy consumption effectively. In the future, we plan to replicate the experiment by directly relying on a dataset containing class-level information about the dependent variable.

Other possible imitations are related to the data set used to train the classifiers, since it did not include secondary features that might impact both dependent and some of the independent variables. In this preliminary work, we did not consider any confounding factors, restricting the interpretation of the selected structural properties. We plan to consider them in a complete evaluation.

Finally, the splitting criterion we applied to label the energy consumption values into the four levels strongly depends on the corresponding variable distribution in the dataset. Such a criterion may threaten the generalizability of the presented classification approach. Indeed, the labeling phase should be performed on a larger and more representative dataset to be applicable in practice. We suspect that using a different and more robust labeling criteria, may solve this limitation and provide "fairer" energy consumption levels. For instance, an ideal schema could be based on the *EU energy labeling schema*, <sup>6</sup>resambling Wilke et al's idea [41], where appliances are assigned to a letter (A, B, C, etc.) depending on their energy consumption (the closer to A, the lower).

## 4 CONCLUSION

This paper presents a mobile app's energy consumption estimation technique based on static properties (i.e., code metrics and smells). The proposed approach predicts the energy consumption level for Java class in Android applications. Results of a preliminary evaluation indicate an average F-measure of  $\approx 0.4$ , suggesting that the technique still needs to be further improved to be considered for real applications. Our future research directions aim to replicate the study on a larger dataset and consider a broader set of predictors.

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<sup>&</sup>lt;sup>6</sup>https://eur-lex.europa.eu/legal-content/EN/TXT/?uri=CELEX:32017R1369

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