Classifying Malware into families Based on File Content

Patrick Rand, Reynier Ortiz Florida International University Miami, Florida, USA

Abstract

We present a mechanism to classify a malware file into one of nine families: Rammit, Gatak, Tracur, Vundo, Simda, Kelihos_ver1, Obfuscator.ACY, Lollipop and Kelihos_ver3. The system uses a large training set of disassembled malware executables, both in their binary and assembly forms. We ran two separate experimental processes: the first involved extracting n-grams from the binary files using the kfNgram tool, and the second used a shell script to parse the assembly files for method calls to external API libraries. In both cases, the attributes are merged to form a master list and later reduced by selecting the top 500 with the highest information gain (IG). These 500 attributes are the boolean attributes to determine if they are present or not in each of the malware executables. The training set, using the selected attributes, was transformed into an .arff file required by Weka [17] to run several classifiers, i.e. nave bayes, decision trees and support vector machines (SVM). We then compared the different algorithms using some common measures: accuracy, error rate, true and false positive error rates.

1. Introduction

The malware industry continues to evolve in complexity, causing increased personal and commercial damage. Traditional methods of malware detection usually involve matching a piece of malware to a signature that has been predefined in the system. However, malware programmers have learned to evade signature detection by polymorphically obfuscating their code. This means, that, for any given piece of malware, the program itself will be recompiled with randomized variable and string names, rendering standard signature detection near useless in identifying an instance of malware [15]. This has given rise to another goal of both research and industry: given a malware file, can we classify it into a common malware family, such that each instance of that family was morphed from the same original malware program? The ability to categorize a malicious program into a known family provides great importance because it allows for a more focused recovery and repair process. Researchers have developed methods for identifying malware by only considering static portions of the malicious code that are unaffected by polymorphism. In particular, these attributes are the method calls of the external libraries used in the code, and the byte sequence of executable. While the research behind these evaluation processes have shown to have good results, much of this research was performed on small sample size of data [12], while others have low accuracy of prediction [15]. The small data sizes is most likely attributed to the difficulty involved with building a data set of malware. First, a program will need to be trapped and identified as malware. From there, researchers can examine the code either dynamically, or statically. The former involves tracing the program's execution in secure, virtual sandbox environment, while the latter involves running the program through a disassembler, and outputting its source code. This paper and its referenced work focus on static analysis of programs, which can be generalized into a form of text/document classification, of which there is extensive research in the machine learning community. The purpose of this paper will be to survey and examine previous research in the field, and evaluate how these methods scale and perform on a very large data set. Microsoft has provided a sample data set of 10,838 disassembled malware executables, of which each program is given in both its hexadecimal byte form, and its assembly form [7]. Each of these training examples have been labeled as belonging of one of the aforementioned malware families. The results of our experiment showed similar, and at times, improved classification accuracy compared to previous research. Also, our modified system, designed to handle our 400 GB training performed well and showed its ability to scale with the large data set. Given more time, we would have liked to explore how are system handles classifying both programs that do not belong to one of the 9 origin families, as well as those that are not malware.

1.1. Paper length

For CVPR 2015, the rules about paper length have changed, so please read this section carefully. Papers, ex-

cluding the references section, must be no longer than eight pages in length. The references section will not be included in the page count, and there is no limit on the length of the references section. For example, a paper of eight pages with two pages of references would have a total length of 10 pages. Unlike previous years, there will be no extra page charges for CVPR 2015.

Overlength papers will simply not be reviewed. This includes papers where the margins and formatting are deemed to have been significantly altered from those laid down by this style guide. Note that this LaTeX guide already sets figure captions and references in a smaller font. The reason such papers will not be reviewed is that there is no provision for supervised revisions of manuscripts. The reviewing process cannot determine the suitability of the paper for presentation in eight pages if it is reviewed in eleven.

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Please number all of your sections and displayed equations. It is important for readers to be able to refer to any particular equation. Just because you didn't refer to it in the text doesn't mean some future reader might not need to refer to it. It is cumbersome to have to use circumlocutions like "the equation second from the top of page 3 column 1". (Note that the ruler will not be present in the final copy, so is not an alternative to equation numbers). All authors will benefit from reading Mermin's description of how to write mathematics: http://www.pamitc.org/documents/mermin.pdf.

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Many authors misunderstand the concept of anonymizing for blind review. Blind review does not mean that one must remove citations to one's own work—in fact it is often impossible to review a paper unless the previous citations are known and available.

Blind review means that you do not use the words "my" or "our" when citing previous work. That is all. (But see below for techreports.)

Saying "this builds on the work of Lucy Smith [1]" does not say that you are Lucy Smith; it says that you are building on her work. If you are Smith and Jones, do not say "as we show in [7]", say "as Smith and Jones show in [7]" and at the end of the paper, include reference 7 as you would any other cited work.

An example of a bad paper just asking to be rejected:

An analysis of the frobnicatable foo filter.

In this paper we present a performance analysis of our previous paper [1], and show it to be inferior to all previously known methods. Why the previous paper was accepted without this analysis is beyond me.

[1] Removed for blind review

An example of an acceptable paper:

An analysis of the frobnicatable foo filter.

In this paper we present a performance analysis of the paper of Smith *et al.* [1], and show it to be inferior to all previously known methods. Why the previous paper was accepted without this analysis is beyond me.

[1] Smith, L and Jones, C. "The frobnicatable foo filter, a fundamental contribution to human knowledge". Nature 381(12), 1-213.

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[1] Authors. "The frobnicatable foo filter", F&G 2014 Submission ID 324, Supplied as additional material fg324.pdf.

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You can handle this paper like any other. Don't write "We show how to improve our previous work [Anonymous, 1968]. This time we tested the algorithm on a lunar lander [name of lander removed for blind review]". That would be silly, and would immediately identify the authors. Instead write the following:

We describe a system for zero-g frobnication. This system is new because it handles the following cases: A, B. Previous systems [Zeus et al. 1968] didn't handle case B properly. Ours handles it by including a foo term in the bar integral.

The proposed system was integrated with the Apollo lunar lander, and went all the way to the moon, don't you know. It displayed the following behaviours which show how well we solved cases A and B: ...

As you can see, the above text follows standard scientific convention, reads better than the first version, and does not explicitly name you as the authors. A reviewer might think it likely that the new paper was written by Zeus *et al.*, but cannot make any decision based on that guess. He or she would have to be sure that no other authors could have been contracted to solve problem B.

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1.5. Miscellaneous

Compare the following:

$$\begin{array}{lll} & & conf_a \\ & & \\ &$$

The space after e.g., meaning "for example", should not be a sentence-ending space. So e.g. is correct, e.g. is not. The provided \eq macro takes care of this.

When citing a multi-author paper, you may save space by using "et alia", shortened to "et al." (not "et. al." as "et" is a complete word.) However, use it only when there are three or more authors. Thus, the following is correct: "Frobnication has been trendy lately. It was introduced by Alpher [1], and subsequently developed by Alpher and Fotheringham-Smythe [2], and Alpher et al. [3]."

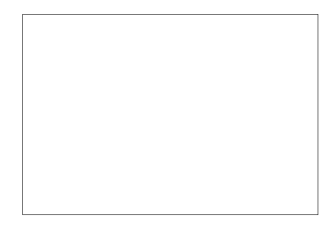


Figure 1. Example of caption. It is set in Roman so that mathematics (always set in Roman: $B\sin A = A\sin B$) may be included without an ugly clash.

This is incorrect: "... subsequently developed by Alpher *et al.* [2] ..." because reference [2] has just two authors. If you use the \etal macro provided, then you need not worry about double periods when used at the end of a sentence as in Alpher *et al.*

For this citation style, keep multiple citations in numerical (not chronological) order, so prefer [2, 1, 4] to [1, 2, 4].

2. Problem Definition and Methods

2.1. Task Definition

Our approach to feature extraction and classification is based on a combination of the methods from the common text classification techniques developed in previous research [12, 15]. Where our process differed from these methods is our focus on the design of the complete system architecture, in particular, creating a system that was not only able to replicate the successful results of the referenced research, but to do so on a data set of a large magnitude of size, and by trying to classify malware into families. The size of our training set required us to design the feature extraction process in a MapReduce-styled pattern [10], but also to only consider machine learning classifiers that were able to scale with our large input. Thus, classifiers such as k-nearest-neighbors and Hidden Markov Models [6] were not considered due to their performance on large data-sets and the impracticality of scaling them in regards to our limited computing resources. For both types of feature-sets, we evaluated the classification performance of three classifiers: Naive Bayes, Decision Trees, and Support Vector Machines. Of these classifiers, Decision Trees and Support Vector Machines performed best, and from there we extended our analysis into the boosted forms of these two algorithms.

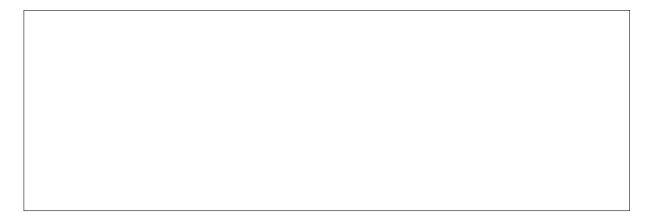


Figure 2. Example of a short caption, which should be centered.

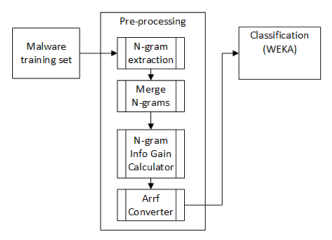


Figure 3. System architecture.

2.2. Algorithms and Methods

2.2.1 System Architecture

As shown in figure Fig 3, the architecture of the system is rather simple. The main tasks were the preprocessing of the data which involved extracting the N-grams and transform the training set into a format that Weka [17] accepts as input. And then apply several learning methods.

2.2.2 Data Preprocessing

The preprocessing part was a process that required a considerable effort because of the size of the training set, 10,868 files. The first step was to extract the hexadecimal N-grams from the binary files using the kfNgram tool [11], we used n=4.

It was then necessary to merge the resulting sorted N-gram files to effectively calculate the *information gain (IG)* of each N-gram. Because these files were extremely long, we could not use the merge functionality built in the kfN-gram tool, and we have to halt the application after running

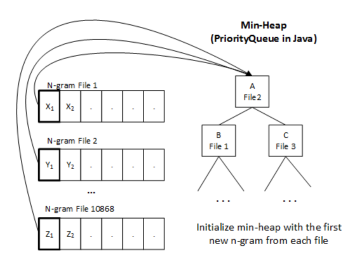


Figure 4. Initializing PriorityQueue to merge N-grams.

for three days. To optimize this process we merged the files maintaining a min-heap (or *PriorityQueue* in Java) in memory with the first distinct N-gram from each file, as shown in Fig 4. We were also able to keep 10,878 Java scanners in memory to speed up I/O. Then, to merging process would consist in each iteration to perform a pop() operation from the PriorityQueue, append the element to merged file, and insert the next N-gram from the same file of the element that was popped to the PriorityQueue. This process is represented in Fig 5.

Once merging was complete, we obtained a total of N-grams was 255,942,370, therefore we had to reduce them by calculating the *information gain (IG)* of each N-gram and selecting the top 500 (Fig. 7), the same approach used in [12]. The *average mutual information* [18] was used to select the most relevant attributes:

$$IG(j) = \sum_{v_j \in \{0,1\}} \sum_{C_i} P(v_j, C_i) \log \frac{P(v_j, C_i)}{P(v_j)P(C_i)}$$
 (1)

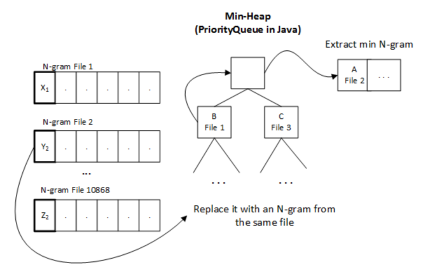


Figure 5. Process of merging N-grams. The minimum element is extracted and replaced with another N-gram from the same file

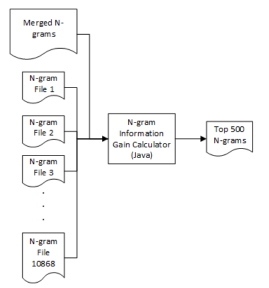


Figure 6. Process of calculating the information gain of each N-gram.

where C_i is the ith class, v_j is the value of the jth attribute, $P(v_j, C_i)$ is the proportion that the jth attribute has the value v_j in the class C_i , $P(v_j)$ is the proportion that the jth n-gram takes the value v_j in the training data, and $P(C_i)$ is the proportion of the training data belonging to the class C_i [12].

The final step, as shown in Fig ??, in the data preprocessing was to transform the training set in .arff format where each example was represented by the presence (or absence) of the top 500 N-grams. This .arff file is one of the formats supported by Weka [17], which facilitated to experiment with different machine learning algorithms.

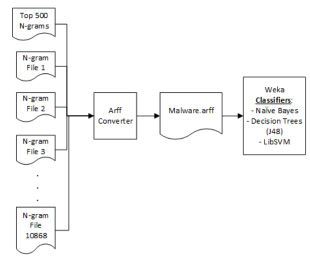


Figure 7. Transforming training set into .arff format.

Our system design for extracting the external API strings followed a near identical design, with the only difference being in the method use to parse the files (.asm) for the method calls. We used a simple shell-script that was designed to parse the files for any API methods using regular expressions. From there the we followed the same feature-set building pattern as for the n-grams.

2.2.3 Classification algorithms

The Nave Bayes classifier uses Bayes' Theorem to label examples based on the assumption of conditional independence between features. Given a data set consisting of word counts, the training of this algorithm is very simple, and only requires the computation of the prior probabil-

ity of each class, $P(C_i)$, and the conditional probability of each word given a class, $P(v_j|C_i)$. For a given instance \mathbf{v} , we aim to classify it into some class, C_i , thus computing $P(C_i|v)$. Using Bayes' Theorem, this can be decomposed into:

$$P(C_i|\mathbf{v}) = \frac{P(C_i) P(\mathbf{v}|C_i)}{P(\mathbf{v})}$$
(2)

We can ignore the $P(\mathbf{v})$ term, and estimate the remaining feature probabilities as:

$$P(C_i) = \frac{\text{# of occurences of class } i}{\text{# of total attributes}}$$
(3)

$$P(v_j|C_i) = \frac{\text{\# of occurences of attribute } v_j \text{ with class } C_i}{\text{\# of occurences of class } i}$$

From here, we can define the classification of training example **v** as:

$$\hat{y} = \operatorname{argmax} P(C_k) \prod_{i=1}^{n} p(v_i|C_k)$$
 (5)

Decision trees are a method of classification that use tree-like graphs to model the classification of an example as a path of probabilities from the root node attribute, to one of the leaf (class) attributes. The beauty of decision trees lies in their simplicity and easy interpretation. Likewise, decision trees can be thought of as a "white-box" classification model, because once the tree has been built, the researchers can visualize and analyze each sub-path of an example's classification path. The basic implementation of a decision tree works as follows: for each node of tree, we select the attribute that best splits the our training examples into their respective classes using the entropy gain for choosing that attribute. For our system, we used the J48 decision tree algorithm in Weka, which is an implementation of the ID3 decision tree algorithm [14]. We define the entropy for a data set S, and its set of attributes V, as:

$$H(S) = -\sum_{v \in V} P(v) \log_2 P(x) \tag{6}$$

Once we obtain the corresponding decision tree, we prune the tree to help minimize over-fitting that may have occurred as a result of tree's greedy construction.

Support Vector Machines (SVMs) produced the best classification results of our tested algorithms. The goal of an SVM is to find an optimal hyperplane that separates the training examples by their respective classes, while maximizing the margin between theses classes. The hyperplane is represented by a vector w and a scalar value b. Thus, the solution of the SVM can be represented by the minimization of:

$$P(\mathbf{w}, b, \boldsymbol{\xi}) = \frac{1}{2} \mathbf{w} \cdot \mathbf{w} + C \sum_{i=1}^{n} \xi_{i}$$
 (7)

such that:

$$\forall_{i=1}^n , \, \xi_i \ge 0 : \, y_i(\mathbf{w} \cdot \mathbf{x}_i + b) \ge 1 - \xi_i$$
 (8)

where ξ_i is the upper bound on the number of training errors and C is the parameter that controls the trade-off between the margin of the hyperplane and the training error. We use the LibSVM [8] implementation of this algorithm.

3. Experimental Evaluation

3.1. Methodology

based on previous methods based on the coulter paper and the reiesk paper. Our paper is a blend of these two papers, in coulter they are classifying if the malware is benign or no and they have of a smaller magnitude data set of magnitude smaller. the riesk paper data set it comparable in size compare malware families like we are but our results seem to be better. we achieved better accuracy and a better AUC measure in our results

All of our samples were collected from this was run on our 10,000 files in both their binary and assembly formats which we acquired from the Microsoft Malware Data Set [] they are real viruses the Microsoft caught, the data set is huge

the performance evaluation data was our test error rate using 10-Fold Cross-Validation, area under the curve (AUC), and precision and recall rates. and how it relates to our ability to classify a Malware file into one of the nine origin families. useful because a Malware family describes the way a group a malicious files do and the damage they cause begin to diagnose and repair, undo whatever is going on or to prevent and narrows doewn yur search for corrupted files and gives you a roadmap for where your problems are the data we analyzed the system byu running three machine learning classifiers on each of the file formats the classification algorithms that we ran are the naiive base classifer the tdecision tree ID3 algorithm and the linear support vector machines

coulter given a file, is it malware. to build our feature set we use similar feature extraction form them n-gram extraciton process taken directly from their paper. feature set is what you pump into your algorithm they are trying to classify things into malware families we are too very large data set but

theyre doing more of a behavior analysis of the API class Its more dynmaic our is more static monitoring the behavior we are just looking at the code they have a data set the same size as us on averge only clasify 88dynamic behavioral modeling approach. what it does to the computer. how are you ever goingto run that on a consumers computer? its not very applicable transferable whereas our system detection system has real world app in the just needs the file run through disassembler and such as IDA [] from there the file

just needs to be parsed for the n-grams or the api streams and can be run through the support vector machine classifier this can all be done in real time and does not require a virtual sandbox environment that is used by the reisk experiment

Our What are the criteria you are using to evaluate? How does this evaluation (e.g. experiment) relate to the questions you are trying to answer? Describe the experimental methodology that you used. What is the training/test data that was used, and why is it realistic or interesting? Exactly what performance data did you collect and how are you presenting and analyzing it? Comparisons to competing methods that address the same problem or to variations of your own algorithm are particularly useful. Give enough detail about your experiment setup so that others could reproduce the work.

3.2. Results

Present the quantitative results of your experiments. Graphical data presentation such as graphs and histograms are frequently better than tables. Explain the basic findings by explaining the results contained in the tables and graphs.

3.3. Discussion

Our results showed the importance of selecting the N-grams that would provide more information about the underlying family by calculating the IG, instead of choosing the most frequent N-grams. With this method we intuitively consider that irrelevant N-grams are avoided, which are those that are product of mutation engines and other techniques used by hackers, and also we assume that by calculating the information gain we somewhat extracted the most common instructions per malware family. This, we think, is the reason why we obtained a high accuracy with Decision Trees and Support Vector Machines, about 99% in both. The results are also inspiring in the sense that, at difference with other papers, our data set is considerable large, and using several evaluation methods as cross-fold validation we presume we are not incurring in overfitting.

This method is far from perfection, as there is still a considerable change of selecting irrelevant N-grams, which could have been mitigated by using sequential pattern extraction or Markov Models. The reasoning is that the mutation engines, we believe, would not be capable of generating sequences of instructions that would mislead the classification if we apply Markov Models or sequence pattern extraction correctly.

4. Related Work

Malware detection and classification is a problem being addressed through several angles. In [9], the goal is to detect if a program exhibits a specified malicious behavior by determining if a set of templates of sequence of instructions are present in the executable files. This approach requires to have knowledge on semantics of each of the malware families. Although the results are promising, there are several reasons we could not follow this approach, for example, as inferred by the title, it would be necessary to create a set of sequence of instructions for each of the nine families we need to classify the malwares into. This is infeasible due to the length of the course, but also it is outside of the scope of the course. As exhibited in the results, this strategy is resilient to obfuscation and showed improvements when compared to McAfee VirusScan.

Several malware classification algorithms are based on n-grams extracted from the executable files. In [13], instead of byte sequences, the n-grams extracted are formed by machine codes. After obtaining the n-grams from the malwares, a centroid for each family is created by selecting the most frequent n-grams. Then, the strategy to classify a malware into one of the families, is to determine the centroid which the malware is more similar to by counting the number of matching n-grams. When considering this approach, one of the possible limitations analyzed was that selecting the most frequent n-gram could implicate choosing an n-gram irrelevant to the malware family.

We also explored sequential pattern extraction of ngrams. The proposed methodology in [16] outlines a procedure to use the n-grams patterns to classify the malware by family. The kfNgram tool [11] was used to extract the ngrams from the disassembled files with n=1, n=2, n=3 and n=4, obtaining the best results with n=4. In [12] the accuracy achieved was higher with n=4, therefore we skipped this evaluation and use only n-grams with n=4. The sequential pattern extraction technique in [19] was used to generate frequently occurred sequences of n-grams to represent the data [16]. Then the patterns significance was calculated using the term frequency-inverse document frequency (TF-IDF) where the term refers to the n-gram pattern and a document to the malware file. Since the number of patterns was too large, the sequential floating forward selection (SFFS) procedure was applied to reduce the number of features, in this case, n-gram sequential patterns. With all the features extracted, three classification algorithms were used, C4.5, multilayer perceptron and support vector machine. The training set was randomly split into two partitions using 80% for training and 20% for testing achieving a 96.64% of accuracy [16]. Because of the duration of the course and the complexity of sequential pattern extraction, we were not able to experiment with this approach.

Following Occam's razor, suggesting that the simplest hypothesis is the best, we applied an approach similar to the one described in [12]. The n-grams extracted from the executable files represented boolean features, present (i.e., 1) or absent (i.e., 0). Since the n-grams list was too large, it was necessary to select the most relevant attributes (i.e.,

n-grams) by computing the *information gain (IG)* described in [18] for each, also called *average mutual information*. Through pilot studies, it was determined to use the top 500 n-grams, and then applied classifiers implemented in the Wakaito Environment for Knowledge Acquisition (WEKA) [17]: IBk, Nave Bayes, SVM, and J48 (decision tree), and also *boosted* the last three of these learners [12]. The results indicated 98% the highest accuracy using boosted decision trees.

5. Future Work

Malware classification is a topic that is always evolving because viruses' authors are constantly developing techniques to avoid been detected, for example by obfuscating the code or by designing polymorphic behaviours. The proposed approach is agnostic of these techniques, therefore, we assume intuitively that a set of the extracted N-grams are product of this ingenious methods. Mutation engines are capable of generating millions of variations of the same virus, therefore, to overcome this problem, it would be necessary to have knowledge of the malware behaviors to develop heuristics or detect specific sequence of instructions to find in the malicious files.

For time reasons, we could not research on the possibilities of removing the irrelevant computations in the malware files created to evade anti-viruses. This would help, if using sequential N-gram patterns extraction, to detect patterns that are truly correlated to the malware family. These patterns could, somewhat, describe the common semantics present in each class of malwares.

6. Conclusion

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AUTHOR NAME(s) and AFFILIATION(s) are to be centered beneath the title and printed in Times 12-point,

Method	Frobnability
Theirs	Frumpy
Yours	Frobbly
Ours	Makes one's heart Frob

Table 1. Results. Ours is better.

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List and number all bibliographical references in 9-point Times, single-spaced, at the end of your paper. When referenced in the text, enclose the citation number in square brackets, for example [4]. Where appropriate, include the name(s) of editors of referenced books.

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All graphics should be centered. Please ensure that any point you wish to make is resolvable in a printed copy of the paper. Resize fonts in figures to match the font in the body text, and choose line widths which render effectively in print. Many readers (and reviewers), even of an electronic copy, will choose to print your paper in order to read it. You cannot insist that they do otherwise, and therefore must not assume that they can zoom in to see tiny details on a graphic.

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