Needles in a Haystack: Mining Information from Public Dynamic Analysis Sandboxes for Malware Intelligence

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Abstract

Malware sandboxes are automated dynamic analysis systems that execute programs in a controlled environment. Within the large volumes of samples submitted every day to these services, some submissions appear to be different from others, and show interesting characteristics. For example, we observed that malware samples involved in famous targeted attacks – like the Regin APT framework or the recently disclosed malwares from the Equation Group – were submitted to our sandbox months or even years before they were detected in the wild. In other cases, the malware developers themselves interact with public sandboxes to test their creations or to develop a new evasion technique. We refer to similar cases as malware developments.

In this paper, we propose a novel methodology to automatically identify malware development cases from the samples submitted to a malware analysis sandbox. The results of our experiments show that, by combining dynamic and static analysis with features based on the file submission, it is possible to achieve a good accuracy in automatically identifying cases of malware development. Our goal is to raise awareness on this problem and on the importance of looking at these samples from an intelligence and threat prevention point of view.

1 Introduction

Malware sandboxes are automated dynamic analysis tools that execute samples in an isolated and instrumented environment. Security researchers use them to quickly collect information about the behavior of suspicious samples, typically in terms of their execution traces and API calls. While customized sandboxes are often installed in the premises of security companies, some sandboxes are available as public online services, as it is the case for Malwr [13], Anubis [10], ThreatExpert [14], VirusTotal [16], and many others [5, 18, 4, 6, 15, 1, 3].

The main advantage of these systems is the fact that the analysis is completely automated and easily parallelizable, thus providing a way to cope with the overwhelming number of new samples that are collected every day. However, due to this extreme parallelization, an incredible amount of reports are generated every day. This makes the task of distinguishing new and important malware from the background noise of polymorphic and uninteresting samples very challenging.

In particular, two important and distinct observations motivate our work. First, it is relatively common that malware samples used to carry out famous targeted attacks were collected by antivirus companies or public sandboxes long before the attacks were publicly discovered [25]. For instance, the binaries responsible for operation Aurora, Red October, Regin, and even some of the new one part of the Equation Group were submitted to the sandbox we used in our experiments several months before the respective attacks appeared in the news [11, 40, 17, 50, 45, 35]. The reasons behind this phenomenon are not always clear. It is possible that the files were automatically collected as part of an automated network or host-based protection system. Or maybe a security analyst noticed something anomalous on a computer and wanted to double-check if a suspicious file exhibited a potentially malicious behavior. It is even possible that the malware developers themselves submitted an early copy of their work to verify whether it triggered any alert on the sandbox system. Whatever the reason, the important point is that no one paid attention to those files until it was too late.

The second observation motivating our study is the constant arm race between the researchers that put continuous effort to randomize their analysis environments, and the criminals that try to fingerprint those systems to avoid being detected. As a consequence of this hidden battle, malware and packers often include evasion techniques for popular sandboxes [19] and updated information about the internal sandbox details are regu-
larly posted on public websites [2]. These examples prove that there must be a constant interaction between malware developers and popular public malware analysis services. This interaction is driven by the need to collect updated information as well as to make sure that new malware creation would go undetected. Even though detecting this interaction might be very difficult, we believe it would provide valuable information for malware triage.

Up to the present, malware analysis services have collected large volumes of data. This data has been used both to enhance analysis techniques [23, 46] and to extrapolate trends and statistics about the evolution of malware families [24]. Unfortunately, to the best of our knowledge, these datasets have never been used to systematically study malware development and support malware intelligence on a large scale. The only public exception is a research recently conducted by looking at VirusTotal to track the activity of specific high-profile hacking groups involved in APT campaigns [52, 27].

In this paper, we approach this objective by applying data-mining and machine learning techniques to study the data collected by Anubis Sandbox [10], a popular malware dynamic analysis service. At the time we performed our analysis, the dataset contained the analysis reports for over 30 millions unique samples. Our main goal is to automatically detect if miscreants submit their samples during the malware development phase and, if this is the case, to acquire more insights about the dynamics of malware development. By analyzing the metadata associated to the sample submissions, it might be possible to determine the software provenance and implement an early-warning system to flag suspicious submission behaviors.

It is important to understand that our objective is not to develop a full-fledged system, but instead to explore a new direction and to show that by combining metadata with static and dynamic features it is possible to successfully detect many examples of malware development submitted to public sandboxes. In fact, our simple prototype was able to automatically identify thousands of development cases, including botnets, keyloggers, backdoors, and over a thousand unique trojan applications.

2 Overview and Terminology

There are several reasons why criminals may want to interact with an online malware sandbox. It could be just for curiosity, in order to better understand the analysis environment and estimate its capabilities. Another reason could be to try to escape from the sandbox isolation to perform some malicious activity, such as scanning a network or attacking another machine. Finally, criminals may also want to submit samples for testing purposes, to make sure that a certain evasion technique works as expected in the sandbox environment, or that a certain malware prototype does not raise any alarm.

In this paper, we focus on the detection of what we call malware development. We use the term “development” in a broad sense, to include anything that is submitted by the author of the file itself. In many cases the author has access to the source code of the program – either because she wrote it herself or because she acquired it from someone else. However, this is not always the case, e.g., when the author of a sample uses a builder tool to automatically generate a binary according to a number of optional configurations (see Section 6 for a practical example of this scenario). Moreover, to keep things simple, we also use the word “malware” as a generic term to model any suspicious program. This definition includes traditional malicious samples, but also attack tools, packers, and small probes written with the only goal of exfiltrating information about the sandbox internals.

Our main goal is to automatically detect suspicious submissions that are likely related to malware development or to a misuse of the public sandbox. We also want to use the collected information for malware intelligence. In this context, intelligence means a process, supported by data analysis, that helps an analyst to infer the motivation, intent, and possibly the identity of the attacker.

Our analysis consists of five different phases. In the first phase, we filter out the samples that are not interesting for our analysis. Since the rest of the analysis is quite time-consuming, any sample that cannot be related to malware development or that we cannot process with our current prototype is discarded at this phase. In the second phase, we cluster the remaining samples based on their binary similarity. Samples in each cluster are then compared using a more fine-grained static analysis technique. Afterwards, we collect six sets of features, based respectively on static characteristics of the submitted files, on the results of the dynamic execution of the samples in the cluster, and on the metadata associated to the samples submissions. This features are finally provided to a classifier that we previously trained to identify the malware development clusters.

3 Data reduction

The first phase of our study has the objective of reducing the amount of data by filtering out all the samples that are not relevant for our analysis. We assume that a certain file could be a candidate for malware development only if two conditions are met. First, the sample must have been submitted to the public sandbox before it was observed in the wild. Second, it has to be part of a manual submission done by an individual user – and not, for example, originating from a batch submission of a secu-
We started by filtering out the large number of batch submissions Anubis Sandbox receives from several researchers, security labs, companies, universities and registered users that regularly submit large bulks of binaries. As summarized in Table 1, with this step we managed to reduce the data from 32 million to around 6.6 million binaries. These samples have been collected by Anubis Sandbox from 2006 to 2013.

Then, to isolate the new files that were never observed in the wild, we applied a two-step approach. First, we removed those submissions that, while performed by single users, were already part of a previous batch submission. This reduced the size of the dataset to half a million samples. In the second step, we removed the files that were uploaded to the sandbox after they were observed by two very large external data sources: Symantec’s Worldwide Intelligence Network (WINE), and VirusTotal.

After removing corrupted or not executable files (e.g., Linux binaries submitted to the Microsoft Windows sandbox), we remained with 184,548 files that match our initial definition of candidates for malware development. Before sending them to the following stages of our analysis, we applied one more filter to remove the packed applications. The rationale behind this choice is very simple. As explained in Section 4, the majority of our features work also on packed binaries, and, therefore, some potential malware development can be identified also in this category. However, it would be very hard for us to verify our results without having access to the decompiled code of the application. Therefore, in this paper we decided to focus on unpacked binaries, for which it is possible to double-check the findings of our system. The packed executables were identified by leveraging the SigBuster [37] signatures.

Table 1 summarizes the number of binaries that are filtered out after each step. The filtering phase reduced the data to be analyzed from over 32 millions to just above 121,000 candidate files, submitted by a total of 68,250 users, were already part of a previous batch submission. This reduced the size of the dataset to half a million samples. In the second step, we removed the files that were uploaded to the sandbox after they were observed by two very large external data sources: Symantec’s Worldwide Intelligence Network (WINE), and VirusTotal.

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### Symantec Filter

Symantec Worldwide Intelligence Network Environment (WINE) is a platform that allows researchers to perform data intensive analysis on a wide range of cyber security relevant datasets, collected from over a hundred million hosts [28]. The data provided by WINE is very valuable for the research community, because these hosts are computers that are actively used by real users which are potential victims of various cyber threats. WINE adopts a 1:16 sampling on this large-scale data such that all types of complex experiments can be held at scale.

To filter out from our analysis the binaries that are not good candidates to belong to malware development, we used two WINE datasets: the binary reputation and the AntiVirus telemetry datasets. The binary reputation dataset contains information about all of the executables (both malicious and benign) downloaded by Symantec customers over a period of approximately 5 years. To preserve the user privacy, this data is collected only from the users that gave explicit consent for it. At the time we performed our study, the binary reputation dataset included reports for over 400 millions of distinct binaries. On the other hand, the AntiVirus telemetry dataset records only the detections of known files that triggered the Norton Antivirus Engine on the users’ machines.

The use of binary reputation helps us locating the exact point in time in which a binary was first disseminated in the wild. The AntiVirus telemetry data provided instead the first time the security company deployed a signature to detect the malware. We combined these datasets to remove those files that had already been observed by Symantec either before the submission to Anubis Sandbox, or within 24 hours from the time they were first submitted to the sandbox.

### VirusTotal Filter

VirusTotal is a public service that provides virus scan results and additional information about hundreds of millions of analyzed files. In particular, it incorporates the detection results of over 50 different AntiVirus engines – thus providing a reliable estimation of whether a file is benign or malicious. Please note that we fetched the VirusTotal results for each file in our dataset several months (and in some cases even years) after the file was first submitted. This ensures that the AV signatures were up to date, and files were not misclassified just because they belonged to a new or emerging malware family.

Among all the information VirusTotal provides about
binaries, the most important piece of information we incorporate in our study is the first submission time of a certain file to the service. We believe that by combining the timestamps obtained from the VirusTotal and Symantec datasets, we achieved an acceptable approximation of the first time a certain malicious file was observed in the wild.

4 Sample Analysis

If a sample survived the data reduction phase, it means that (with a certain approximation due to the coverage of Symantec and VirusTotal datasets) it had never been observed in the wild before it was submitted to the online malware analysis sandbox. Although this might be a good indicator, it is still not sufficient to flag the submission as part of a potential malware development. In fact, there could be other possible explanations for this phenomenon, such as the fact that the binary was just a new metamorphic variation of an already known malware family.

Therefore, to reduce the risk of mis-classification, in this paper we consider a candidate for possible development only when we can observe at least two samples that clearly show the changes introduced by the author in the software. In the rest of this section we describe how we find these groups of samples by clustering similar submissions together based on the sample similarity.

4.1 Sample Clustering

In the last decade, the problem of malware clustering has been widely studied and various solutions have been proposed [31, 33, 51, 32]. Existing approaches typically use behavioral features to group together samples that likely belong to the same family, even when the binaries are quite different. Our work does not aim at proposing a new clustering method for malware. In fact, our goal is quite different and requires to group files together only when they are very similar (we are looking for small changes between two versions of the same sample) and not when they just belong to the same family. Therefore, we leverage a clustering algorithm that simply groups samples together based on their binary similarity (as computed by ssdeep [38]) and on a set of features we extract from the submission metadata.

Moreover, we decided to put together similar binaries into the same cluster only if they were submitted to our sandbox in a well defined time window. Again, the assumption is that when a malware author is working on a new program, the different samples would be submitted to the online sandbox in a short timeframe. Therefore, to cluster similar binaries we compute the binary similarities among all the samples submitted in a sliding window of seven days. We then shift the sliding window ahead of one day and repeat this step. We employ this sliding window approach in order (1) to limit the complexity of the computation and the total number of binary comparisons, and (2) to ensure that only the binaries that are similar and have been submitted within one week from each other are clustered together. We also experimented with other window sizes (between 2 and 15 days) but while we noticed a significant reduction of clusters for shorter thresholds, we did not observe any advantage in increasing it over one week.

Similarities among binaries are computed using the ssdeep [38] tool which is designed to detect similarities on binary data. ssdeep provides a light-weight solution for comparing a large-number of files by relying solely on similarity digests that can be easily stored in a database. As we already discarded packed binaries in the data reduction phase, we are confident that the similarity score computed by ssdeep is a very reliable way to group together binaries that share similar code snippets. After computing the similarity metrics, we executed a simple agglomerative clustering algorithm to group the binaries for which the similarity score is greater than 70%. Note that this step is executed separately for each time window, but it preserves transitivity between binaries in different sliding windows. For example, if file $A$ is similar to $B$ inside window 1, and $B$ is similar to file $C$ inside the next sliding window, at the end of the process $A$, $B$, and $C$ will be grouped into the same cluster. As a result, a single cluster can model a malware development spanning also several months.

Starting from the initial number of binaries, we identified 5972 clusters containing an average of 4.5 elements each.

Inter-Cluster Relationships

The ssdeep algorithm summarizes the similarity using an index between 0 (completely different) and 100 (perfect match). Our clustering algorithm groups together samples for which the difference between the fuzzy hashes is greater than the 70% threshold. This threshold was chosen according to previous experiments [38], which concluded that 70% similarity is enough to guarantee a probability of misclassification close to zero.

However, if the malware author makes very large changes on a new version of his program, our approach may not be able to find the association between the two versions. Moreover, the final version of a malware development could be compiled with different options, making a byte-level similarity too imprecise. To mitigate these side effects, after the initial clustering step, we perform a refinement on its output by adding inter-clusters edges whenever two samples in the same time window
share the same submission origin (i.e., either from the same IP address or using the same email address for the registration). These are “weak” connections that do not model a real similarity between samples, and therefore they are more prone to false positives. As a consequence, our system does not use them when performing its automated analysis to report suspicious clusters. However, as explained in Section 6, these extra connections can be very useful during the analysis of a suspicious cluster to gain a more complete picture of a malware development.

After executing this refinement step, we were able to link to our clusters an additional 10,811 previously isolated binaries. This procedure also connected several clusters together, to form 225 macro groups of clusters.

4.2 Intra-cluster Analysis

Once our system had clustered the binaries that likely belong to the same malware development, we investigate each cluster to extract more information about its characteristics. In particular, we perform a number of code-based analysis routines to understand if the samples in the same cluster share similar code-based features.

Code Normalization

Code normalization is a technique that is widely used to transform binary code to a canonical form [26]. In our study, we normalize the assembly code such that the differences between two binaries can be determined more accurately. Under the assumption that two consecutive variations of the same program are likely compiled with the same tool chain and the same options, code normalization can be very useful to remove the noise introduced by small variations between two binaries.

There are several approaches that have been proposed to normalize assembly code [36, 49, 34]. Some of them normalize just the operands, some the mnemonics, and some normalize both. In this paper, we chose to normalize only the operands so that we can preserve the semantics of the instructions. In particular, we implemented a set of IDA Pro plugins to identify all the functions in the code and then replace, for each instruction, each operand with a corresponding placeholder tag: reg for registers, mem for memory locations, val for constant values, near for near call offsets, and ref for references to memory locations. These IDA scripts were run in batch mode to pre-process all the samples in our clusters.

Programming Languages

The second step in our intra-cluster analysis phase consists in trying to identify the programming language used to develop the samples. The programming language can provide some hints about the type of development. For example, scripting languages are often used to develop tools or probes designed to exfiltrate information from the sandbox. Moreover, it is likely that a malware author would use the same programming language for all the intermediate versions of the same malware. Therefore, if a cluster includes samples of a malware development, all samples should typically share the same programming language. Exceptions, as the one explained in Section 6, may point to interesting cases.

To detect the programming language of a binary we implemented a simple set of heuristics that incorporate the information extracted by three tools: PEiD, the pefile python library, and the Linux strings command. First, we use pefile to parse the Import Address Table (IAT) and obtain the list of libraries that are linked to the binary. Then, we search for programming language specific keywords on the extracted list. For example, the “VB” keyword in the library name is a good indicator of using Visual Basic, and including mscoree.dll in the code can be linked to the usage of Microsoft .NET. In the second step of our analysis, we analyze the strings and the output of PEiD to detect compiler specific keywords (e.g., type_info and RTTI produced by C++ compilers, or “Delphi” strings generated by the homonymous language).

With these simple heuristics, we identified the programming language of 14,022 samples. The most represented languages are Visual Basic (49%), C (21%), Delphi (18%), Visual Basic .Net (7%), and C++ (3%). The large number of Visual Basic .Net could be a consequence of the fact that a large number of available tools that automatically create generic malware programs adopt this language.

Fine-grained Sample Similarity

In this last phase, we look in more detail at the similarity among the samples in the same cluster. In particular, we are interested to know why two binaries show a certain similarity: Did the author add a new function to the code? Did she modify a branch condition, or remove a basic block? Or maybe the code is exactly the same, and the difference is limited to some data items (such as a domain name, or a file path).

To answer these questions, we first extract the timeline of each cluster, i.e., the sequence in which each sample was submitted to the sandbox in chronological order. Moving along the timeline, we compare each couple of samples using a number of static analysis plugins we developed for IDA Pro.

The analysis starts by computing and comparing the call graph of the two samples. In this phase we compare
the normalized code of each function, to check which functions of the second binary were present unchanged in the first binary. The output is a list of additional function that were not present in the original file, plus a list of functions that were likely modified by the author — i.e., those function that share the same position in the call graph but whose code does not perfectly match. However, at this level of granularity it is hard to say if something was modified in the function or if the author just removed the function and added another with the same callee.

Therefore, in these cases, we “zoom” into the function and repeat our analysis, this time comparing their control flow graphs (CFGs). Using a similar graph-based approach, this time we look for differences at the basic block level. If the two CFGs are too different, we conclude that the two functions are not one the evolution of the other. Otherwise, we automatically locate the different basic blocks and we generate a similarity measure that summarize the percentage of basic blocks that are shared by the two functions.

4.3 Feature Extraction

Based on the analysis described in the previous sections, our system automatically extracts a set of 48 attributes that we believe are relevant to study the dynamics of malware development.

This was done in two phases. First, we enriched each sample with 25 individual features, divided in six categories (see the Appendix for a complete list of individual features). The first class includes self-explanatory file features (such as its name and size). The Timestamps features identify when the sample was likely created, when it was submitted to Anubis Sandbox, and when it was later observed in the wild. While the creation time of the binary (extracted from the PE headers) could be manually faked by the author, we observed that this is seldom the case in practice, in particular when the author submits a probe or an intermediate version of a program. In fact, in these cases we often observed samples in which the compilation time precedes the submission time by only a few minutes.

The third category of features contains the output of the VirusTotal analysis on the sample, including the set of labels associated by all AntiVirus software and the number of AVs that flag the sample as malicious. We then collect a number of features related to the user who submitted the sample. Since the samples are submitted using a web browser, we were able to extract information regarding the browser name and version, the language accepted by the system (sometime useful to identify the nationality of the user) and the IP from which the client was connecting from. Two features in this set require more explanation. The email address is an optional field that can be specified when submitting a sample to the sandbox web interface. The proxy flag is instead an attempt to identify if the submitter is using an anonymization service. We created a list of IP addresses related to these services and we flagged the submissions in which the IP address of the submitter appears in the blacklist. In the Binary features set we record the output of the fine-grained binary analysis scripts, including the number of sections and functions, the function coverage, and the metadata extracted by the PE files. Finally, in the last feature category we summarize the results of the sandbox behavioral report, such as the execution time, potential runtime errors, use of evasion techniques, and a number of boolean flags that represent which behavior was observed at runtime (e.g., HTTP traffic, TCP scans, etc.)

In the second phase of our analysis we extended the previous features from a single sample to the cluster that contains it. Table 2 shows the final list of aggregated attributes, most of which are obvious extensions of the values of each sample in the cluster. Some deserve instead a better explanation. For instance, the cluster shape (A3) describes how the samples are connected in the cluster: in a tightly connected group, in a chain in which each node is only similar to the next one, or in a mixed shape including a core group and a small tail. The Functions diff (B13) summarized how many functions have been modified in average between one sample and the next one. Dev time (B25) tells us how far apart in time each samples were submitted to the sandbox, and Connect back (B24) counts how many samples in the cluster open a TCP connection toward the same /24 subnetwork from which the sample was submitted. This is a very common behavior for probes, as well as for testing the data exfiltration component of a malicious program.

Finally, some features such as the number of crashes (C8) and the average VT detection (D4) are not very interesting per se, but they become more relevant when compared with the number of samples in the cluster. For example, imagine a cluster containing three very similar files. Two of them run without errors, while the third one crashes. Or two of them are not detected by AV signatures, but one is flagged as malware by most of the existing antivirus software.

While we are aware of the fact that each feature could be easily evaded by a motivated attacker, as described in Section 6 the combinations of all them is usually sufficient to identify a large number of development clusters. Again, our goal is to show the feasibility of this approach and draw attention to a new problem, and not to propose its definitive solution.
5 Machine Learning

Machine learning provides a very powerful set of techniques to conduct automated data analysis. As the goal of this paper is to automatically distinguishing malware developments from other submissions, we tested with a number of machine learning techniques applied to the set of features we presented in detail in the previous section.

Among the large number of machine learning algorithms we have tested our training data with, we have obtained the best results by using the logistic model tree (LMT). LMT combines the logistic regression and decision tree classifiers by building a decision tree whose leaves have linear regression models [41].

### Training Set

The most essential phase of machine learning is the training phase where the algorithm learns how to distinguish the characteristics of different classes. The success of the training phase strictly depends on a carefully prepared labeled data. If the labeled data is not prepared
carefully, the outcome of machine learning can be misleading. To avoid this problem, we manually labeled a number of clusters that were randomly chosen between the ones created at the end of our analysis phase. Manual labeling was carried out by an expert that performed a manual static analysis of the binaries to identify the type and objective of each modification. With this manual effort, we flagged 91 clusters as non-development and 66 as development. To estimate the accuracy of the LMT classifier, we conducted a 10-fold cross validation and a 70% percentage split evaluation on the training data.

Feature Selection

In the previous section, we have presented a comprehensive set of features that we believe can be related to the evolution of samples and to distinguish malware developments from ordinary malware samples. However, not all the features contribute in the same way to the final classification, and some works well only when used in combination with other classes.

To find the subset of features that achieves the optimal classification accuracy while helping us to obtain the list of features that contribute the most to it, we leveraged a number of features selection algorithms that are widely used in machine learning literature: Chi-Square, Gain Ratio and Relief-F attribute evaluation. Chi-square attribute evaluation computes the chi-square statistics of each feature with respect to the class, which in our case is the fact of being a malware development or not. The Gain Ratio evaluation, on the other hand, evaluates the effect of the feature by measuring its gain ratio. Finally, the Relief-F attribute evaluation methodology assigns particular weights to each feature according to how much they are successful to distinguish the classes from each other. This weight computation is based on the comparison of the probabilities of two nearest neighbors having the same class and the same feature value.

While the order slightly differs, the ten most effective features for the accuracy of the classifier for all three feature selection algorithms are the same. As also the common sense suggests, the features we extract from the binary similarity and the analysis of the samples are the most successful. For example, the connect back feature that checks if the sample connects back to the same IP address of the submitter, the average edit distance of the filenames of the samples, the binary function similarity, and the sample compile time features are constantly ranked on the top of the list. The submitter features and the sandbox features are following the sample features in the list. All of the features except the number of sandbox evasions, the VirusTotal results, and the features we extracted from the differences on the file sizes in the clusters had a contribution to the accuracy. After removing those features, we performed a number of experiments on the training set to visualize the contribution of the different feature sub-sets to the classification accuracy. Figure 1 shows (in log-scale) the impact of each class and combination of classes. Among all the classes the samples-based features produced the best combination of detection and false positive rates (i.e. 88.2% detection rate with 7.4% false positives). In particular, the ones based on the static and dynamic analysis of the binaries seem to be the core of the detection ability of the system. Interestingly, the cluster-based features alone are the worst between all sets, but they increase the accuracy of the final results when combined with other features.

The results of the final classifier are reported in Table 3: 97.4% detection with of 3.7% false positives, according to 10-folds cross validation experiment. Note that we decided to tune the classifier to favor detection over false positives, since the goal of our system is only to tag suspicious submissions that would still need to be manually verified by a malware analyst.

6 Results

Our prototype implementation was able to collect substantial evidences related to a large number of malware developments.

In total, our system flagged as potential development 3038 clusters over a six years period. While this number was too large for us to perform a manual verification of each case, if such a system would be deployed we estimate between two and three alerts generated per day. Therefore, we believe our tool could be used as part of an early warning mechanism to automatically collect information about suspicious submissions and report them to human experts for further investigation.

In addition to the 157 clusters already manually labeled to prepare the training set for the machine learning component, we also manually verified 20 random clusters automatically flagged as suspicious by our system. Although according to the 10-fold cross validation experiments the false positive rate is 3.7%, we have not found any false positives on the clusters we randomly selected for our manual validation.

Our system automatically detected the development of a diversified group of real-world malware, ranging from generic trojans to advanced rootkits. To better understand the distribution of the different malware families, we verified the AV labels assigned to each reported cluster. According to them, 1474 clusters were classified as malicious, out of which our system detected the development of 45 botnets, 1082 trojans, 83 backdoors, 4 keyloggers, 65 worms, and 21 malware development tools (note that each development contained several different samples modeling intermediate steps). A large fraction
of the clusters that were not identified by the AV signatures contained the development of probes, i.e., small programs whose goal is only to collect and transmit information about the system where they run. Finally, some clusters also contained the development or testing of offensive tools, such as packers and binders.

6.1 Targeted Attacks Campaigns

Before looking at some of the malware development cases detected by our system, we wanted to verify our initial hypothesis that even very sophisticated malware used in targeted attacks are often submitted to public sandboxes months before the real attacks are discovered. For this reason, we created a list of hashes of known and famous APT campaigns, such as the ones used in operation Aurora and Red October. In total, we collected 1271 MD5s belonging to twelve different campaigns. As summarized in Table 4, in all cases we found at least one sample in our database before the campaign was publicly discovered (Early Submission column). For example, for Red October the first sample was submitted in February 2012, while the campaign was later detected in October 2012.
2012. The sample of Regin was collected a record 44 months before the public discovery.

Finally, we checked from whom those samples were submitted to the system. Interestingly, several samples were first submitted by large US universities. A possible explanation is that those samples were automatically collected as part of a network-based monitoring infrastructure maintained by security researchers. Other were instead first submitted by individual users (for whom we do not have much information) from several different countries, including US, France, Germany, UK, and Romania. Even more interesting, some were first submitted from DSL home Internet connections. However, we cannot claim that we observed the development phase of these large and popular targeted attacks campaigns as in all cases the samples were already observed in the wild (even though undetected and no one was publicly aware of their existence) before they were submitted to our sandbox. It is important to note that for this experiment we considered the entire dataset, without applying any filtering and clustering strategy. In fact, in this case we did not want to spot the development of the APT samples, but simply the fact that those samples were submitted and available to researchers long before they were publicly discovered.

We believe the sad message to take away from this experiment is that all those samples went unnoticed. As a community, there is a need for some kind of early warning system to report suspicious samples to security researches. This could prevent these threats from flying under the radar and could save months (or even years) of damage to the companies targeted by these attacks.

6.2 Case studies

In the rest of this section we describe in more details three development scenarios. While our system identified many more interesting cases, due to space limitation we believe the following brief overview provides a valuable insight on the different ways in which attackers use (and misuse) public sandboxes. Moreover, it also shows how a security analyst can use the information collected by our system to investigate each case, and reconstruct both the author behavior and his final goal.

In the first example, the malware author introduced an anti-sandbox functionality to a Trojan application. In this case the analyst gathers intelligence information about the modus operandi of the attacker and about all the development phases.

In the second scenario, we describe a step by step development in which the attacker tries to collect information from the sandbox. This information is later used to detect the environment and prevent the execution of a future malware in the sandbox. In the last example, we show how an attacker uses the sandbox as a testbed to verify the behavior of the malware. In this case, the author generated the binary using one of the many dedicated builder applications that can be downloaded from the Internet or bought on the black market.

Example I: Anti-sandbox Malware

The cluster related to this example contains three samples. The timeline (summarized in Figure 2) already suggests a possible development. In fact, the difference between the submission time and the compile time is very small.

A quick look at the static features of the cluster shows that the three samples are very similar, and share the same strings as well as the same imphash (the import hash [20, 21] recently introduced also by VirusTotal). However, the first sample is composed of 21 functions, while the last two samples have 22 functions. Our report also shows how the first and the second samples differ for two functions: the author modified the function start, and introduced a new function CloseHandle. This information (so far extracted completely automatically by our system) is a good starting point for a closer analysis.

We opened the two executables in IDA Pro, and quickly identified the two aforementioned functions (snippet in Figure 3). It was immediately clear that the start function was modified to add an additional basic block and a call to the new CloseHandle function. The new basic block uses the rdtsc x86 instruction to read the value of the Timestamp Counter Register (TSC), which contains the number of CPU cycles since the last reset. The same snippet of assembly is called two times to check the time difference. After the first rdtsc instruction there is a call to CloseHandle, using the timestamp as handler (probably an invalid handler). These two well known tricks are here combined to detect the Anubis Sandbox environment – due to the delay introduced by its checks during program execution. The Anubis Sandbox’s core is slower in looking up the handlers table, and this time discrepancy is the key to detect the analysis environment. In this case the difference has to be less than 0x0000h, or the program would immediately terminate by calling the ExitProcess function.

The last sample in the cluster was submitted only to tune the threshold and for this reason there were no important differences with the second sample. The control
The behavior described so far suggest malicious intents. This is also confirmed by other cluster metadata. For instance, while the first sample in the cluster was unknown to VirusTotal, the last one was clearly identified as a common Trojan application. This suggests that the original sample, without the timing check, has never been used in the wild. Once more, the fact that all three samples have been submitted days before the trojan was first observed in the wild strongly supports the fact that the person who submitted them was indeed the malware author.

Example II: Testing a Trojan Dropper

The second cluster we want to describe is composed of five samples. Our report indicates that the first four are written in Delphi and the last one is written in Visual Basic. This is already a strange fact, since the two programming languages are quite different and it is unlikely that they could generate similar binaries.

In this case the cluster timeline does not provide useful information as all the Delphi samples share exactly the same compilation time: 20th of June, 1992. Only the Visual Basic sample had a compilation time consistent with the submission. On the contrary, the submission times provide an interesting perspective. All the samples have been submitted in few hours and this might indicate a possible development. In addition, there are two IP addresses involved: one for the four Delphi samples and one for the final Visual Basic version. The static features of the first four samples show very little differences, suggesting that these are likely just small variations of the same program. In average, they share 169 out of 172 functions and 7 out of 8 PE sections. By inspecting the changes, we notice that the attacker was adding some thread synchronization code to a function responsible for injecting code into a different process. The control flow graph similarity reported by our tool was over 98%, confirming the small differences we observed between each versions. Once the author was happy with the result, she submitted one more sample, this time completely different from the previous ones. Despite the obvious differences in most of the static analysis features, the fuzzyhash similarity with sample 4 was 100%. A rapid analysis showed that this perfect match was due to the fact that the Visual Basic application literally embedded the entire binary of the fourth Delphi program. In addition, the behavior report confirmed that, once executed, the Visual Basic Trojan dropped the embedded executable that was later injected inside a target process. None of the Antivirus software used by VirusTotal recognized the first four samples as malicious. However, the last one was flagged by 37 out of 50 AVs as a trojan dropper malware.

It is important to stress that a clear advantage of our system is that it was able to automatically reconstruct the entire picture despite the fact that not all samples were submitted from the same IP address (even though all located in the same geographical area). Moreover, we were able to propagate certain metadata extracted by our system (for example the username of the author extracted from the binary compiled with Visual Studio) from one sample to the others in which that information was missing. This ability to retrieve and propagate metadata between different samples can be very useful during an investigation.

Another very interesting aspect of this malware devel-
opment is the fact that after the process injection, the program used a well-known dynamic DNS service (no-1p) to resolve a domain name. The IP address returned by the DNS query pointed exactly to the same machine that was used by the author to submit the sample. This suggests that the attacker was indeed testing his attack before releasing it, and this information could be used to locate the attacker machine.

We identified a similar connect-back behavior in other 1817 clusters. We also noticed how most of these clusters contain samples generated by known trojan builders, like Bifrost [8] or PoisonIvy [9]. While this may seem to prove that these are mostly unsophisticated attacks, Fire-Eye [22] recently observed how the Xtremerat builder [7] (which appeared in 28 of our clusters) was used to prepare samples used in several targeted attacks.

**Example III: Probe Development**

In this last example we show an attacker fingerprinting the analysis environment and how, at the end, she manages to create her own successful anti-sandbox check. The cluster consists of two samples, both submitted from France in a time span of 23 hours by the same IP address. The two samples have the same size, the same number of functions (164), and of sections (4). There is only one function (_start) and two sections (.text and .rdata) presenting some differences. The two programs perform the same actions, they create an empty text file and then they retrieve the file attributes through the API GetFileAttributes. The only differences are on the API version they use (GetFileAttributesA or GetFileAttributesW) and on the file name to open.

At a first look, this cluster did not seem very interesting. However, the inter-cluster connections pointed to other six loosely correlated samples submitted by the same author in the same week. As explained in Section 4, these files have not been included in the core cluster because the binary similarity was below our threshold. In this case, these samples were all designed either to collect information or to test anti-virtualization/emulation tricks. For instance, one binary implemented all the known techniques based on idt, gdt and ldt to detect a virtual machine monitor [48, 47, 42]. Another one simply retrieved the computer name, and another one was designed to detect the presence of inline hooking. Putting all the pieces together, it is clear that the author was preparing a number of probes to assess various aspects of the sandbox environment.

This example shows how valuable the inter-clusters edges can be to better understand and link together different submissions that, while different between each other at a binary level, are likely part of the same organized “campaign”.

## 6.3 Malware Samples in the Wild

As we already mentioned at the beginning of the section, out of 3038 clusters reported as malware development candidates by our machine learning classifier, 1474 (48%) contained binaries that were detected by the antivirus signatures as malicious (according to VirusTotal).

A total of 228 of the files contained in these clusters were later detected in the wild by the Symantec’s antivirus engine. The average time between the submission to our sandbox and the time the malware was observed in the wild was 135 days – i.e., it took between four and five months for the antivirus company to develop a signature and for the file to appear on the end-users machines. Interestingly, some of these binaries were later detected on more than 1000 different computers in 13 different countries all around the world (obviously a lower bound, based on the alerts triggered on a subset of the Symantec’s customers). This proves that, while these may not be very sophisticated malware, they certainly have a negative impact on thousands of normal users.

## 7 Limitations

We are aware of the fact that once this research is published, malware authors can react and take countermeasures to sidestep this type of analysis systems. For instance, they may decide to use “private” malware checkers, and avoid interacting with public sandboxes altogether. First of all, this is a problem that applies to many analysis techniques ranging from botnet detection, to intrusion prevention, to malware analysis. Despite that, we believe that it is important to describe our findings so that other researchers can work in this area and propose more robust methodologies in the future.

Moreover, as we mentioned in the introduction, after we completed our study someone noticed that some known malware development groups were testing their creation on VirusTotal [52, 27]. This confirms that what we have found is not an isolated case but a widespread phenomenon that also affects other online analysis systems. Second, now that the interaction between malware developers and public sandboxes is not a secret anymore, there is no reason that prevents us from publishing our findings as well.

We are aware of the fact that our methodology is not perfect, that it can be evaded, and that cannot catch all development cases. However, we believe the key message of the paper is that malware authors are abusing public sandboxes to test their code, and at the moment we do not need a very sophisticated analysis to find them. Since this is the first paper that tries to identify these cases, we found that our approach was already sufficient to detect
thousands of them. Certainly more research is needed in this area to develop more precise monitoring and early warning system to analyze the large amounts of data automatically collected by public services on a daily basis.

8 Related Work

While there has been an extensive amount of research on malware analysis and detection, very few works in the literature have studied the datasets collected by public malware dynamic analysis sandboxes. The most comprehensive study in this direction was conducted by Bayer et al. [24]. The authors looked at two years of Anubis [10] reports and they provided several statistics about malware evolution and about the prevalent types of malicious behaviors observed in their dataset.

Lindorfer et al. [43] conducted the first study in the area of malware development by studying the evolution over time of eleven known malware families. In particular, the authors documented the malware updating process and the changes in the code for a number of different versions of each family. In our study we look at the malware development process from a different angle. Instead of studying different versions of the same well known malware, we try to detect, on a large scale, the authors of the malware at the moment in which they interact with the sandbox itself. In a different paper, Lindorfer et al. [44] proposed a technique to detect environment sensitive malware. The idea is to execute each malware sample multiple times on several sandboxes equipped with different monitoring implementations and then compare the normalized reports to detect behavior discrepancies.

A similar research area studies the phylogeny [30] of malware by using approaches taken from the biology field. Even if partially related to our work, in our study we were not interested in understanding the relationship between different species of malware, but only to detect suspicious submissions that may be part of a malware development activity.

In a paper closer to our work, Jang et al. [34] studied how to infer the software evolution looking at program binaries. In particular, the authors used both static and dynamic analysis features to recover the software lineage. While Jang’s paper focused mostly on benign programs, some experiments were also conducted on 114 malicious software with known lineage extracted from the Cyber Genome Project [12]. Compared to our work, the authors used a smaller set of static and dynamic features especially designed to infer the software lineage (e.g., the fact that a linear development is characterized by a monotonically increasing file size). Instead, we use a richer set of features to be able to distinguish malware developments from variations of the same samples collected on the wild and not submitted by the author.

While our approaches share some similarities, the goals are clearly different.

Other approaches have been proposed in the literature to detect similarities among binaries. Flake [29] proposed a technique to analyze binaries as graphs of graphs, and we have been inspired by his work for the control flow analysis described in Section 4. Kruegel et al. [39] proposed a similar technique in which they analyzed the control flow graphs of a number of worms and they used a graph coloring technique to cope with the graph-isomorphism problem.

Finally, one step of our technique required to cluster together similar malware samples. There are several papers in the area of malware clustering [31, 33, 51, 32]. However, their goal is to cluster together samples belonging to the same malware family as fast as possible and with the highest accuracy. This is a crucial task for all the Antivirus companies. However, our goal is different as we are interested in clustering samples based only on binary similarity and we do not have any interest in clustering together members of the same family based on their behavior.

9 Conclusion

Public dynamic analysis sandboxes collect thousands of new malware samples every day. Most of these submissions belong to well known malware families, or are benign files that do not pose any relevant security threat. However, hidden in this large amount of collected data, few samples have something special that distinguishes them from the rest. In this paper, we discussed the importance of looking at these samples from an intelligence and threat prevention point of view.

We show that several binaries used in the most famous targeted attack campaigns had been submitted to our sandbox months before the attack was first reported. Moreover, we propose a first attempt to mine the database of a popular sandbox, looking for signs of malware development. Our experiments show promising results. We were able to automatically identify thousands of developments, and to show how the authors modify their programs to test their functionalities or to evade detections from known sandboxes. Around 1,500 of them were real malware developments – some of which have been later observed on thousands of infected machines around the world.

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References


### A: File Features
- **Filename**: The original name of the file submitted by the user
- **File size**: The size of the file
- **MD5**: Simple hash used for lookup in other data sources
- **Fuzzy Hashes**: Using SSDeep algorithm

### B: Timestamps
- **Submission time**: Time in which the sample was submitted to Anubis Sandbox
- **Compile time**: Time in which the binary was compiled
- **Symantec first**: Time the sample was first observed in the wild by Symantec
- **VirusTotal first**: Time in which the binary was first submitted to VirusTotal

### C: AV Features
- **AV-Detection**: Number of AV that flag the samples as malicious (according to VirusTotal)
- **AV-Labels**: List of AV labels associated to the sample (according to VirusTotal)

### D: User-based Features
- **User Agent**: User agent of the browser used to submit the sample
- **Languages**: Languages accepted by the user browser (according to the `accept-language` HTTP header)
- **IP**: IP address of the user who submitted the file
- **IP Geolocation**: Geolocation of the user IP address
- **Email address**: Optional email address specified when the sample was submitted
- **Proxy**: Boolean value used to identify submission through popular anonymization proxies

### E: Binary Features
- **N.Sections**: Number of sections in the PE file
- **N.Functions**: Number of functions identified by the disassembly
- **Code Coverage**: Fraction of `.text` segment covered by the identified functions
- **Programming Language**: Programming language used to develop the binary
- **Metadata**: Filenames and username extracted from the PE file

### F: Behavioral Features
- **Duration**: Duration in seconds of the analysis
- **Errors**: Error raised during the analysis
- **Evasion**: Known anti-sandbox techniques detected by the sandbox itself
- **Behavior Bitstring**: Sequence of 24 boolean flags that characterize the behavior of the sample. (has_popups, has_udp_traffic, has_http, has_tcp_address_scan, modified_registry_keys, ...)

Table 5: List of Individual Features associated to each sample