A comparative analysis of data mining algorithms to mitigate spurious detections in Gaia

Author: Esther Pallares Guimera.

Abstract: Gaia is an ESA mission that observes about 50 million sources per day. A small part of these detections are considered spurious generated for example by cosmic rays. The main objective of this study is to perform a comparative analysis of several algorithms to automatically detect spurious detections. Successfully identifying these detections is important to prevent them from entering the cross-match stage where they create several problems and degrade resolution performance. We will use appropriate metrics to determine the execution and assess the algorithms. Finally, it will be discussed if any of these data mining algorithms could be a good solution to the spurious detection problem.

I. THE GAIA MISSION

Gaia is an European Space Agency (ESA) mission whose main goal is to scan our Galaxy and beyond to build a three dimensional map according to astrometric measurements of objects up to magnitude G=20.7. Moreover, photometric and spectroscopic measures are also taken with an unprecedented resolution only possible from space.

It is expected to observe more than a billion stars and solar system objects. Current estimations hint that processing all these observations will require more than $10^{21}$ flops [1].

Gaia has been operating since 2013 and a first version of the catalogue was published on September 2016. The final version is scheduled for 2022. The data is opened to the public and it is expected to make significant contributions to the knowledge of our Galaxy, the Milky Way, among other scientific targets.

A. The Gaia Instrument

Gaia has two optical telescopes which correspond to two fields of view (FoV). Both telescopes share the same focal plane depicted in FIG. 1. There are one hundred and six charged couple devices (CCDs) divided into seven rows and into seventeen strips comprising nearly a thousand million pixels. A deeper description of the focal plane is detailed in [2].

The Sky Mappers CCDs are in charge of the detection. Point-like sources passing by one telescope are detected. Afterwards, a window is placed around objects of interest. Only these windows are read out to reduce background noise. We refer to detections of the Sky Mappers CCDs as detections, observations or transits.

B. Spurious Detections

The processing power is limited and so the detection process has to be simple, and sometimes false positives happen, thus producing detections of non-real stars, what we refer to as spurious detections, which can reach up to 20% of all observations for some periods [3].

There are different types of spurious detections: spurious detections caused by cosmic rays, by background noise, due to unexpected light paths, as a result of internal reflections within the satellite and around bright sources of magnitude up to 16. We refer to bright sources causing spurious as parent detections. The spurious detections are located along the diffraction spikes, but for sources with magnitude brighter than 6, known as Very Bright Sources (VBS), spurious detections appear not
only along the diffraction spikes but also in more complicated patterns near the bright source.

This work is focused on the classification of spurious detections around bright sources because these are the most frequent spurious type representing around 90% of the total spurious detections. To simplify this work we restricted this initial study to spurious created by sources with magnitudes between G=7.5 and G=12.

A single transit of a VBS source can produce thousands of spurious detections. A transit of a source of magnitude G=8 produces hundreds of spurious detections. For sources of magnitude G=12 just a few spurious detections are produced. FIG.2. illustrates the detections of a single transit for a source of magnitude G=5.02 (top), G=8.18 (middle) and G=11.7 (bottom).

II. CLASSIFICATION ALGORITHMS

The two main types of machine learning algorithms are: supervised learning and unsupervised learning. Supervised learning algorithms learn from datasets with class known. On the other hand, unsupervised learning algorithms extract their own conclusions from the dataset, but in this case, class is not provided. Unsupervised learning algorithms have not been considered as we have several physical information of Gaia detections and we want to control the process.

The aim of a classification algorithm is to choose between a set of options. In this particular case, we intend to decide if an observation belongs to a source or to a spurious. This decision is made according to a model previously created for each algorithm after training it. We have chosen to work with Weka, which provides the implementation for several data mining algorithms [4].

The chosen algorithms for this spurious classification analysis are: REPTree, kstar and IBK, because they executed the best performance among others on a first test of magnitude G=8.

A training set is a data set where the classification class of each element is known. The algorithms are trained with this dataset. On the other hand, a test set is a data set where the classification class of each element is unknown. Models are evaluated with test sets. A model is the rule to distinguish an observation between a star and a spurious.

To build the training dataset it was necessary to manually classify observations as real or spurious. Thirty-six training sets were generated in total, with a minimum of seven training sets per magnitude between G=7.5 and G=12. In order to do that, Gaia detections were plotted and manually classified using TOPCAT[5].

Defining the dimensionality (number of attributes) of a training set is relevant as machine learning algorithms work better with low rather than high dimensional data sets. For this reason, each attribute has to be relevant. Moreover, if the set of attributes are not relevant the classification will not be performed correctly [6].

FIG. 2: Spurious detections around a source of magnitude G=5.02, G=8.18 and G=11.7 from top to bottom, for several scans. (Units: 10^{-8} rad)

A major part of this study has been to test and refine the necessary parameters for a correct classification. We concluded that absolute attributes (not referenced to any parent observation) such as alpha, delta, magnitude and scan number were not good enough for the classification. For this reason, we decided to reference the attributes to the parent observation.

The parent observation can be any source of magnitude
between \(G=7.5\) and \(G=12\). The dataset is created with detections in its surroundings selected with a fix window with different dimensions according to the parent magnitude.

The final set of attributes chosen per detection has been:

- Parent magnitude: it is a fundamental variable because this magnitude defines spurious distribution complexity around it. A brighter source has more spurious around it as it can be seen in FIG. 2.
- Parent field of view: it allows knowing which telescope observed the parent observation.
- Field of view: it informs about which telescope observed the detection. This allows algorithms to discard detections by FoV if they are not relevant.
- Magnitude difference between the observation and the parent observation: it was selected because spurious around bright sources are much fainter than the parent detection.
- AL and AC relative position difference in pixels between an observation and the parent observation: spurious appear in certain patterns, as a result, spurious are more likely to be in a certain region. Therefore, we use these variables to have relative location to the parent observation.

### A. REPTree

REPTree is a regression tree algorithm. A tree algorithm is based on decision points, named nodes, which are produced by splitting data into smaller subsets. Each data division adds new branches to the tree. Each path from the root to the leaf constitutes a region of data. Each region is fitted with a constant value \(k\).

In regression trees there are three main characteristics: splitting rule, termination criterion and assigning a constant value to a terminal node. In this case, REPTree uses information gain (it calculates the expected decrease of entropy) as splitting rule and mean square error as termination rule also known as pruning [7].

### B. Kstar

Kstar is a cluster analysis algorithm as well as an instance based learner. An instance, \(x\), is assigned to the cluster with the nearest mean measured by entropic distance which has some benefits including handling of real valued attributes and missing values [8].

### C. IBK

IBK is an incremental instance-based learning algorithm whose leit moti is: similar instances have similar classification. This type divides training instances depending on their category attribute’s value in order to save a set of representative instances, which will allow similarity function and classification function to predict the class of a new instance [9].

### III. PERFORMANCE METRICS

In order to compute precision, sensitivity and accuracy we need four numbers: true positives (TP) which means the number of spurious correctly classified as spurious, true negatives (TN) counting the number of sources correctly classified as sources, false positive (FP) meaning the number of sources incorrectly classified as spurious and finally, false negative (FN) meaning the number of spurious incorrectly classified as sources.

We have used the following set of metrics to analyse the performance of the three selected algorithms:

- Execution time: it is the result of building the model from training sets and the classification of a test set.
- Precision: percentage of correctly classified spurious detections out of total, false or not, of predicted as spurious.

\[
\text{Precision} = \frac{TP}{TP + FP} \tag{1}
\]

- Sensitivity: percentage of correctly classified spurious detections out of total number of real spurious.

\[
\text{Sensitivity} = \frac{TP}{TP + FN} \tag{2}
\]

- Accuracy: percentage of correct classification.

\[
\text{Accuracy} = \frac{TP + TN}{TP + TN + FP + FN} \tag{3}
\]

Neither precision nor sensitivity depend on the size of the test set, however, accuracy does depend on the number of instances of the test set, but this metric is the most basic measure of the performance of a classifier [10].

In order to validate the classification, two methods have been used, ten fold cross-validation and hold out cross-validation.

Ten fold cross-validation splits the training set in two parts in order to use one of them to compute the model and the other to evaluate it. In our case, data set is divided into ten subsets (nine for training and one for testing).

Hold out cross-validation trains the model with a training set, and tests it with a different dataset [11].
Overfitting is an issue that can happen with these algorithms. It occurs when a model reflects excessively a singular training set. The model works perfectly with that training set but it is not trained to be able to correctly perform the classification of a different data set.

The metrics of ten fold cross-validation method were successful with the wrong set of attributes (absolute attributes) because the test and training sets used by this method belonged to a unique original set, according to its definition explained before. Thereby, both sets of instances, training and test were pretty similar. However, hold out cross-validation showed very bad results due to the model only reflected some specific scenarios.

IV. RESULTS

In order to analyse each algorithm, we started performing a test to determine the minimum number of training sets to obtain good results. We executed the same test several times but increasing the number of training sets. We started to build the model with one training set and each time more independent training sets were concatenated to the initial set, always trying to maintain a proportionality between magnitudes, until the number of spurious stabilized around thirty-eight percent (orange line in FIG. 3). The percentage of spurious detections is the discriminatory parameter, as it is the real number of spurious of our test set. The sum of the numerous independent training sets builds the final training set.

FIG. 3: Graphic evolution for IBK (yellow), REPTree (blue) and kstar (green). The percentage of spurious detections of the input test set of magnitude G=7.77 according to the number of training sets used to build the model.

FIG. 3 shows the percentage of spurious corresponding to a test set of magnitude G=7.7. Therefore, if the model behaves correctly in this magnitude, it is expected to behave even better in the rest of magnitudes, due to brighter parents produce more complex spurious detections patterns.

REPTree and kstar show a constant performance from thirty training sets. However, IBK never arrives to converge with REPTree and kstar. Hence, the minimum number of training sets has to be bigger than thirty. Despite this, in theory as more training instances are used to build a model, a better performance is achieved, always avoiding overfitting. Provided that, thirty-six datasets were used to compare the data mining algorithms.

Once we knew the number of training sets, we proceeded to evaluate their performance with ten fold cross-validation and hold out cross-validation, see FIG. 4. The results of these metrics can be seen in TABLE I and II.

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>Precision</th>
<th>Sensitivity</th>
<th>Accuracy</th>
<th>Time (s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>REPTree</td>
<td>0.950</td>
<td>0.972</td>
<td>0.979</td>
<td>1.38</td>
</tr>
<tr>
<td>kstar</td>
<td>0.765</td>
<td>0.738</td>
<td>0.875</td>
<td>1109</td>
</tr>
<tr>
<td>IBK</td>
<td>0.936</td>
<td>0.971</td>
<td>0.976</td>
<td>1.45</td>
</tr>
</tbody>
</table>

TABLE I: Measures of precision, sensitivity, accuracy and execution time with ten fold cross-validation per analysed algorithm.

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>Precision</th>
<th>Sensitivity</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>REPTree</td>
<td>1</td>
<td>0.828</td>
<td>0.932</td>
</tr>
<tr>
<td>kstar</td>
<td>0.922</td>
<td>0.888</td>
<td>0.983</td>
</tr>
<tr>
<td>IBK</td>
<td>0.400</td>
<td>1</td>
<td>0.588</td>
</tr>
</tbody>
</table>

TABLE II: Measures of precision, sensitivity and accuracy with hold out cross-validation per analysed algorithm.

In this particular case, our training set has more than fourteen hundred instances and the test set has three hundred eighty-nine instances. The necessary time to build the model of an algorithm and perform the classification of a test set is the same for both cross-validation methods.
In table II we can see that some metrics achieve its maximum value, one. In the case of IBK, the sensitivity is maximum because it has over classified data as spurious. On the other hand, the precision of REPTree is also maximum because there are not stars classified as spurious.

In table I and II, we can see the difference on IBK precision between ten fold and hold out cross-validation, we assure the degradation in the performance must be due to IBK will require a larger training set. In FIG. 3, IBK do not converge to the expected value, thirty-eight percent. Our assumption is that with IBK the training set still needs more data for a correct training.

Comparing table I and II, REPTree and kstar show some difference between ten fold and hold out cross-validation. In the case of kstar, its metrics are higher in hold out cross-validation, contrary to REPTree. Having into account hold out cross-validation is obtained testing an independent test, we will give more weight to this method.

Table I shows that kstar is much slower than REPTree and IBK. The time difference rose to more than three hours between kstar and REPTree, and it neither had stopped growing nor stabilized with the number of independent training sets.

V. CONCLUSIONS

It can be said that data mining algorithms are successfully classifying spurious detections from the Gaia data. The research of the right set of attributes has taken half of this work as they are the base for a correct classification. From the first attempts to the final set of attributes has been an evolution, modifying the dimensionality of the set and adjusting each parameter to arrive to the final set. In this process, we found out that Gaia data had to be considered by scan instead of multi scan, for example. Moreover, we concluded that increasing the training set size do not improve the performance if the eight attributes are not appropriate. However, the right set of attributes classified much better with a much reduced size training set.

We conclude that REPTree is the best algorithm because its hold out cross-validation metrics are excellent and also its speed. We discard kstar because it is very time consuming, and IBK because executed the worst performance.

As future work, in order to use data mining classification technique for all Gaia data, the model should be created using training datasets covering the missing magnitude ranges G=5 to G=8, and G=12 to G=16. Moreover, additional training sets should be manually created and tested for all magnitudes in order to improve the current results. Furthermore, a chance should be given to machine learning algorithms not considered because they might obtain better results with the total range of Very Bright Sources.

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