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## **RIWAYAT KORESPONDENSI**



Aris Marjuni <aris.marjuni@mail.ugm.ac.id>

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## Notification to co-authors of submission to Journal of Big Data BIGD-D-19-00120

1 message

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**Journal of Big Data Editorial Office** <em@editorialmanager.com> Fri, Jul 19, 2019 at 9:45 AM  
Reply-To: Journal of Big Data Editorial Office <visalatchi.irudhayanathan@springernature.com>  
To: Aris Marjuni <aris.marjuni@mail.ugm.ac.id>

BIGD-D-19-00120

Unsupervised Software Defect Prediction using Median Absolute Deviation Threshold based Spectral Classifier with Signed Laplacian Matrix  
Aris Marjuni; Teguh Bharata Adji, Ph.D; Ridi Ferdiana

Dear author:

You are receiving this email because you have been listed as an author on a manuscript recently submitted to Journal of Big Data. The manuscript details are below.

Title: Unsupervised Software Defect Prediction using Median Absolute Deviation Threshold based Spectral Classifier with Signed Laplacian Matrix  
Authors: Aris Marjuni; Teguh Bharata Adji, Ph.D; Ridi Ferdiana  
Corresponding author: Mr. Teguh Bharata Adji

If you are not aware of the submission, or if you should not be listed as contributing author, please notify the Editorial Office. Contact details for the Editorial Office are available under "Contact Us" on the journal website.

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**Date:** 02 Aug 2019  
**To:** "Teguh Bharata Adji" adj@ugm.ac.id  
**From:** "Journal of Big Data Editorial Office" visalatchi.irudhayanathan@springernature.com  
**Subject:** Your submission to Journal of Big Data - BIGD-D-19-00120

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BIGD-D-19-00120  
Unsupervised Software Defect Prediction using Median Absolute Deviation Threshold based Spectral Classifier with Signed Laplacian Matrix  
Aris Marjuni; Teguh Bharata Adji, Ph.D; Ridi Ferdiana  
Journal of Big Data

Dear Mr. Adji,

Your manuscript "Unsupervised Software Defect Prediction using Median Absolute Deviation Threshold based Spectral Classifier with Signed Laplacian Matrix" (BIGD-D-19-00120) has been assessed by our reviewers. Although it is of interest, we are unable to consider it for publication in its current form. The reviewers have raised a number of points which we believe would improve the manuscript and may allow a revised version to be published in Journal of Big Data.

Their reports, together with any other comments, are below. Please also take a moment to check our website at <https://www.editorialmanager.com/bigd/> for any additional comments that were saved as attachments.

If you are able to fully address these points, we would encourage you to submit a revised manuscript to Journal of Big Data.

Once you have made the necessary corrections, please submit online.

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The due date for submitting the revised version of your article is 30 Aug 2019.

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Please be aware that we may investigate, or ask your institute to investigate, any unauthorised attempts to change authorship or discrepancies in authorship between the submitted and revised versions of your manuscript.

We look forward to receiving your revised manuscript soon.

Best wishes,

Borko Furht  
Journal of Big Data  
<https://journalofbigdata.springeropen.com/>

Reviewer reports:

Editors' comments ( if any ) :

Reviewer #1: Good Work.

Reviewer #2: Out of the Scope

This Manuscript is Focused on the importance of I declare that I have no competing interests  
this is not related to subject field of big data

Reviewer #3: author approach is good  
check formatting of paper  
some language corrections are need

Reviewer #4: 1- Needs some writing corrections:

in "Introduction" : {

Bishnu et al. [16] compared the quad tree based k-means algorithm with some algorithms, which are k-means, Naive Bayes, and Linear Discriminant Analysis to assess the clustering performance. The error rates of quad tree based k-means algorithm are fairly comparable to the k-means, Naive Bayes, and Linear Discriminant Analysis. The error rates of quad tree based k-means algorithm are fairly comparable with k-means, naive Bayes, and Linear Discriminant Analysis. Abaei et al.}

in "Introduction" :{

This paper is organized into several sections as follows. Introduction section presents the background, issues, and the motivation of this study}

You write about your previous publication in "introduction" and other sections such as?:

{The use of spectral clustering in unsupervised approach is also proposed by Marjuni et al. [13].}

2- There isn't great difference with your previous publication:

{13. Marjuni, A., Adji, T.B., Ferdiana, R.: Unsupervised software defect prediction using signed laplacian-based spectral classier. *Soft COmputing*, 1{12 (2019). doi:10.1007/s00500-019-03907-6}

Reviewer #5: The paper is recommended for publication.

Reviewer #6: Include concrete results in abstract.

Study area used in this study is not defined. Authors are requested to describe the study area and also why this 'area of interest' is selected for this type of study? Also, explain what other data this DBI model can work?

Authors have clearly explained the first algorithm in this manuscript. They are required to explain the second algorithm too.

What are the tools & systems, and programming languages used for this study?

How many of the validation points were ground-truthed for both the methods? What was the accuracy of the validation point interpretations? What are the other validation methods can be applied to this study?

How did your account for potential auto-correlation in the training data?

This part is explained in detail with the useful insights. What are the future prospects and limitation of the study conducted? Is it a pilot study or regular research study?

What are the future prospects and limitation of the study conducted? Is it a pilot study or regular research study?

Reviewer #7: . Research Idea initiative is good and slight extension of your previous work with limited novelty of work carried out

2. Academic Language and grammar should be improved .Avoid complex sentences

3. Abstract and Conclusion part states the major contribution towards the research objective and results

4. Error function analysis is needed for proving accuracy.

5. Recommended to enhance the features of dataset and statistical test is missing for proving accuracy.

There is additional documentation related to this decision letter. To access the file(s), please click the link below. You may also login to the system and click the 'View Attachments' link in the Action column.

\*\*\*\*\*

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Date: August 16, 2019

**Borko Furht**  
Editor-in-Chief  
Journal of Big Data

Subject: Response to Reviewers  
Re: Manuscript reference No. BIGD-D-19-00120

Dear Dr. Borko Furht,

Please find attached a revised version of our manuscript “Unsupervised Software Defect Prediction using Median Absolute Deviation Threshold based Spectral Classifier on Signed Laplacian Matrix”, which we would like to resubmit for publication as a research article in Journal of Big Data.

Your comments and those of reviewers were very insightful and enabled us to improve the quality of our manuscript significantly. We have carefully reviewed all the reviewer’s comments point-by-point. Our responses and explanations enclosed in the “Response to Reviewers” table. We also made some revisions in the old submission manuscript to follow the reviewer’s comments and suggestions, including the language and grammar improvement.

We hope the revised manuscript version is now suitable for publication. We shall look forward to hearing from you at your earliest convenience. Thank you.

Sincerely,

Teguh Bharata Adji, S.T., M.T., M.Eng., Ph.D.  
*Corresponding Author*

Address: Department of Electrical and Information Engineering  
Faculty of Engineering, Universitas Gadjah Mada  
Yogyakarta 55281, Indonesia  
Email: adji@ugm.ac.id

## Response to Reviewers

Title                    Unsupervised Software Defect Prediction using Median Absolute Deviation Threshold based Spectral Classifier on Signed Laplacian Matrix.  
Authors                Aris Marjuni, Teguh Bharata Adji, and Ridi Ferdiana.  
Submitted to        Journal of Big Data.

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<b>Reviewer #1</b>			
No.	Comment	Response and Explanation	Page: Line
1.	Good Work.	Thank you for your review.	-

<b>Reviewer #2</b>			
No.	Comment	Response and Explanation	Page: Line
1.	Out of the Scope. This Manuscript is focused on the importance of I declare that I have no competing interests this is not related to subject field of big data.	<p>The trend of current software inevitably leads to the big data era. There are much of large software developed from hundreds to thousands of modules. In software development projects, finding the defect proneness manually on each module in large software dataset is probably inefficient in resources. In this task, the use of a software defect prediction model becomes a popular solution with much more cost-effective rather than manual reviews.</p> <p>This study presents a specific machine learning algorithm, which is the spectral classifier, to develop a software defect prediction model using unsupervised learning approach.</p> <p>Thus, we expect that this study is relevant to the ‘machine learning algorithms for big data’ subject field and appropriates to the aim and scope of the Journal of Big Data.</p>	1: 13-19

<b>Reviewer #3</b>			
No.	Comment	Response and Explanation	Page: Line
1.	Author approach is good.	Thank you for your review.	-
2.	Check formatting of paper.	For paper submission, we used BioMedCentral’s article template, as preferred in the submission guidelines at the journal website.	-
3.	Some language corrections are need.	We have re-read our article and have tried to improve the language and grammar.	-



Reviewer #4			
No.	Comment	Response and Explanation	Page: Line
1.	Needs some writing corrections: in "Introduction" : { Bishnu et al. [16] compared the quadtree based k-means algorithm with some algorithms, which are k-means, Naive Bayes, and Linear Discriminant Analysis to assess the clustering performance. The error rates of quadtree based k-means algorithm are fairly comparable to the k-means, Naive Bayes, and Linear Discriminant Analysis. The error rates of quadtree based k-means algorithm are fairly comparable with k-means, naive Bayes, and Linear Discriminant Analysis. Abaei et al. }	These sentences have been revised and have changed using the correct sentence.  New sentences: Bishnu et al. [16] compared the quadtree based k-means algorithm with some algorithms, which are k-means, Naive Bayes, and Linear Discriminant Analysis to assess the clustering performance. The error rates of quadtree based k-means algorithm are reasonably comparable to the k-means, Naive Bayes, and Linear Discriminant Analysis.	2: 40-43
2.	Needs some writing corrections: in "Introduction" :  { This paper is organized into several sections as follows. Introduction section presents the background, issues, and the motivation of this study. }	This sentence has been revised and has changed using the correct sentence.  New sentences: This paper presents this study in several sections. Introduction section describes the background, issues, and motivation of this study.	3: 38-39
3.	You write about your previous publication in "Introduction" and other sections such as:  { The use of spectral clustering in unsupervised approach is also proposed by Marjuni et al. [13]. }	<ul style="list-style-type: none"> <li>- We cite our previous publication [13] to refer that this paper is the extension of our previous study with different issues.</li> <li>- Our previous study in [13] was performed to address the negative similarity issue in spectral classifier through the signed Laplacian based spectral classifier as the proposed method. This study uses our previous publication as the baseline of this study.</li> <li>- This study is performed to address both the dominantly partitioning and the eigenvector value's outlier issues of the zero value threshold in spectral classifier through the median absolute deviation threshold based spectral classifier.</li> </ul>	<p>3: 15-20 3: 29-37</p> <p>8: 23-46 9: 1-2 (Baseline method)</p> <p>7: 22-46 8: 1-21 (Proposed method)</p>
	There isn't great difference with your previous publication:  { 13. Marjuni, A., Adji, T.B., Ferdiana, R.: Unsupervised software defect prediction using signed laplacian-based spectral classier. Soft Computing, 1 {12 (2019). doi:10.1007/s00500-019-03907-6 }		

<b>Reviewer #5</b>			
<b>No.</b>	<b>Comment</b>	<b>Response and Explanation</b>	<b>Page: Line</b>
1.	The paper is recommended for publication.	Thank you for your review.	-

<b>Reviewer #6</b>			
<b>No.</b>	<b>Comment</b>	<b>Response and Explanation</b>	<b>Page: Line</b>
1.	Include concrete results in abstract.	The experimental results have been added in Abstract to complete the achievement.	1: 35-43
2.	Study area used in this study is not defined. Authors are requested to describe the study area and also why this 'area of interest' is selected for this type of study?	The area of interest has been added in Abstract to describe the study area.	1: 13-19
3.	Also, explain what other data this DBI model can work?	Davies Bouldin Index (DBI) is a universal internal validation measure for any clustering algorithm and represents the cluster compactness. So, this measure can work on any dataset and at any application areas where the clustering algorithm is applied.	9: 44-46
4.	Authors have clearly explained the first algorithm in this manuscript. They are required to explain the second algorithm too.	Additional explanations have been added in the Methods section to clarify the differences between both the proposed and baseline algorithms.	8: 23-46 9: 1-2 (Baseline method)  7: 22-46 8: 1-21 (Proposed method)
5.	What are the tools & systems, and programming languages used for this study?	We used Matlab programming to study both the proposed and baseline algorithms.	10: 6-7
6.	How many of the validation points were ground-truthed for both the methods?	We use ten software datasets from the NASA MDP software projects. Each dataset comes with the class labels (defective and clean classes) for all entities as the ground-truth points for classification model validation. Each dataset has different of the numbers of ground-truth points as each dataset has different sizes, as shown in Table 1.	9: 12-19 and Table 1

Reviewer #6			
No.	Comment	Response and Explanation	Page: Line
7.	What was the accuracy of the validation point interpretations?	The accuracy interprets the ability of the classifier to predict the entities class correctly. The accuracy value indicates the ratio of all correct predicted entities to all entities (Eq. 19), which are observed using the comparison between the predicted and ground-truth classes.	11: 7-10 14: 34-46 15: 1-5
8.	What are the other validation methods can be applied to this study?	<p>There are many other validation methods which can be applied to evaluate both clustering and classification.</p> <ul style="list-style-type: none"> <li>– For clustering, there are many methods for validation, such as BetaCV, C-Index, Normalized Cut, Modularity, Dunn Index, Davies Bouldin Index, Silhouette Coefficient, and Hubert Statistic. In this study, we use the Davies Bouldin Index to measure the cluster compactness.</li> <li>– For classification or labeling, there are many methods for classifier validation, such as confusion matrix, ROC, Cohen’s K score, and cross validation. In this study, so we use the confusion matrix and ROC to validate our proposed method.</li> <li>– To validate the performance significance between the proposed and the baseline methods, we use the two-tailed Wilcoxon signed-rank test at the 95% level of confidence.</li> </ul>	9: 42-46 10: 11-26  10: 1-5 10: 27-46 11: 1-24  11: 25-46 12: 1-6
9.	How did your account for potential auto-correlation in the training data?	<ul style="list-style-type: none"> <li>– This study is performed to address the zero threshold issues in the spectral based clustering process. We take out significant conclusions based on those issues. Thus, we don’t yet perform the feature analysis in this study.</li> <li>– However, we consider that feature analysis is essential in the classifier model. If it is possible, we will include the feature analyses, such as the auto-correlation or multi-collinearity between the software features/metrics/attributes through feature selection method as future work to improve the spectral classifier performance of this study.</li> </ul>	3: 29-37 16: 1-6

Reviewer #7			
No.	Comment	Response and Explanation	Page: Line
10.	This part is explained in detail with the useful insights. What are the future prospects and limitation of the study conducted? Is it a pilot study or regular research study?	<ul style="list-style-type: none"> <li>– The software defect prediction modeling is one of the alternative solutions for software engineers to predict the software defect prone during software testing. Due to the increase in software sizes and complexities, it usually needs more resources to exploit the large software dataset. In this case, there is a significant potential for the use of software defect prediction approach in future because this approach is much more cost-effective rather than manual reviews.</li> <li>– This study is a regular research study and not a pilot study, which as the extension of our previous study [13] in unsupervised software defect prediction with a different problem/issue. ----- [13]. Marjuni, A., Adj, T.B., Ferdiana, R.: Unsupervised software defect prediction using signed laplacian-based spectral classier. Soft Computing, 1-12 (2019). DOI:10.1007/s00500-019-03907-6.</li> </ul>	<p>1: 40-43 15: 39-46 16: 1-6</p> <p>8: 23-46 9: 1-2 (Baseline method)</p> <p>7: 22-46 8: 1-21 (Proposed method)</p>

Reviewer #7			
No.	Comment	Response and Explanation	Page: Line
1.	Research idea initiative is good and slight extension of your previous work with limited novelty of work carried out.	<p>This study is the extension of our previous study with different issues.</p> <ul style="list-style-type: none"> <li>– Our previous study in [13] was performed to address the negative similarity issue in spectral classifier through the signed Laplacian based spectral classifier as the proposed method.</li> <li>– This study is performed to address both the dominantly partitioning and the eigenvector value's outlier issues of the zero value threshold in spectral classifier through the median absolute deviation threshold based spectral classifier.</li> </ul>	<p>8: 23-46 9: 1-2 (Baseline method)</p> <p>7: 22-46 8: 1-21 (Proposed method)</p>





Aris Marjuni <aris.marjuni@mail.ugm.ac.id>

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## Fwd: Decision on your Submission to Journal of Big Data - BIGD-D-19-00120R1

1 message

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**Teguh Bharata Adji** <adji@ugm.ac.id>  
To: Aris Marjuni <aris.marjuni@mail.ugm.ac.id>

Mon, Sep 23, 2019 at 4:54 PM

----- Forwarded message -----

From: **Journal of Big Data Editorial Office** <em@editorialmanager.com>

Date: Sun, Sep 8, 2019 at 7:56 PM

Subject: Decision on your Submission to Journal of Big Data - BIGD-D-19-00120R1

To: Teguh Bharata Adji <adji@ugm.ac.id>

BIGD-D-19-00120R1

Unsupervised Software Defect Prediction using Median Absolute Deviation Threshold based Spectral Classifier on Signed Laplacian Matrix

Aris Marjuni; Teguh Bharata Adji, Ph.D; Ridi Ferdiana

Journal of Big Data

Dear Mr. Adji,

I am pleased to inform you that your manuscript "Unsupervised Software Defect Prediction using Median Absolute Deviation Threshold based Spectral Classifier on Signed Laplacian Matrix" (BIGD-D-19-00120R1) has been accepted for publication in Journal of Big Data.

Before publication, our production team will check the format of your manuscript to ensure that it conforms to the standards of the journal. They will be in touch shortly to request any necessary changes, or to confirm that none are needed.

Any final comments from our reviewers or editors can be found, below. Please quote your manuscript number, BIGD-D-19-00120R1, when inquiring about this submission.

We look forward to publishing your manuscript and I hope you will consider Journal of Big Data again in the future.

Best wishes,

Borko Furht

Journal of Big Data

<https://journalofbigdata.springeropen.com/>

Editors' comments ( if any ) :

Reviewer #3: Author addressed all reviewer comments

Reviewer #6: Authors have implemented the requested suggestions.  
But they need to include quantitative (value) results in the abstract also.

**FINAL ARTICLE**  
**PUBLISHED**

RESEARCH

Open Access



# Unsupervised software defect prediction using median absolute deviation threshold based spectral classifier on signed Laplacian matrix

Aris Marjuni<sup>1,2</sup>, Teguh B. Adji<sup>1\*</sup> and Ridi Ferdiana<sup>1</sup>

\*Correspondence:

adji@ugm.ac.id

<sup>1</sup> Department of Electrical and Information Engineering, Faculty of Engineering, Universitas Gadjah Mada, Jl. Grafika No. 2, Kampus UGM, Yogyakarta 55281, Indonesia  
Full list of author information is available at the end of the article

## Abstract

**Area of interest:** The trend of current software inevitably leads to the big data era. There are much of large software developed from hundreds to thousands of modules. In software development projects, finding the defect proneness manually on each module in large software dataset is probably inefficient in resources. In this task, the use of a software defect prediction model becomes a popular solution with much more cost-effective rather than manual reviews. This study presents a specific machine learning algorithm, which is the spectral classifier, to develop a software defect prediction model using unsupervised learning approach.

**Background and objective:** The spectral classifier has been successfully used in software defect prediction because of its reliability to consider the similarities between software entities. However, there are conditional issues when it uses the zero value as partitioning threshold. The classifier will produce the predominantly cluster when the eigenvector values are mostly positives. Besides, it will also generate low clusters compactness when the eigenvector contains outliers. The objective of this study is mainly to propose an alternative partitioning threshold in dealing with the zero threshold issues. Generally, the proposed method is expected to improve the spectral classifier based software defect prediction performances.

**Methods:** This study proposes the median absolute deviation threshold based spectral classifier to carry out the zero value threshold issues. The proposed method considers the eigenvector values dispersion measure as the new partitioning threshold, rather than using a central tendency measure (e.g., zero, mean, median). The baseline method of this study is the zero value threshold based spectral classifier. Both methods are performed on the signed Laplacian matrix to meet the non-negative Laplacian graph assumption. For classification, the heuristic row sum method is used to assign the entity class as the prediction label.

**Results and conclusion:** In terms of clustering, the proposed method can produce better cluster memberships that affect the cluster compactness and the classifier performances improvement. The cluster compactness average of both the proposed and baseline methods are 1.4 DBI and 1.8 DBI, respectively. In classification performance, the proposed method performs better accuracy with lower error rates than



the baseline method. The proposed method also has high precision but low in the recall, which means that the proposed method can detect the software defect more precisely, although in the small number in detection. The proposed method has the accuracy, precision, recall, and error rates with average values of 0.79, 0.84, 0.72, and 0.21, respectively. While the baseline method has the accuracy, precision, recall, and error rates with average values of 0.74, 0.74, 0.89, and 0.26, respectively. Based on those results, the proposed method able to provide a viable solution to address the zero threshold issues in the spectral classifier. Hence, this study concludes that the use of the median absolute deviation threshold can improve the spectral based unsupervised software defect prediction method.

**Keywords:** Unsupervised software defect prediction, Spectral classifier, Signed Laplacian matrix, Zero thresholding, Median absolute deviation thresholding

## Introduction

Software defect prediction model generally needs a prior software project repository dataset to train the model [1, 2]. Based on the dataset availability, there are two most common model development approaches. The first one is a supervised approach, where the software defect prediction model is developed from the training dataset and evaluated using the testing dataset. The second one is an unsupervised approach, where the software defect prediction model is developed using the current testing dataset without training dataset.

In the supervised approach, there are two types of defect prediction models, namely within project defect prediction and cross project defect prediction [3]. The within project defect prediction model uses a training dataset from the same prior software projects to develop a model. The homogeneity between the training and testing datasets in this model could produce the best performance compared to the other approaches. However, there is a difficulty to implement this model on a new software project. Because, a new software project does not have a prior training datasets [3, 4]. The cross project defect prediction model is then used to deal with this issue, where training dataset is taken from the other different software projects to develop a model [4–6]. However, taking similar training datasets from the other software projects is not an easy way because of the heterogeneity between the source and the target projects [3, 7, 8]. If there are no lacks in the training dataset availability, the supervised approach is the main alternative in the software prediction model development. Otherwise, the unsupervised approach offers an alternative solution to address the training dataset availability issue [9, 10].

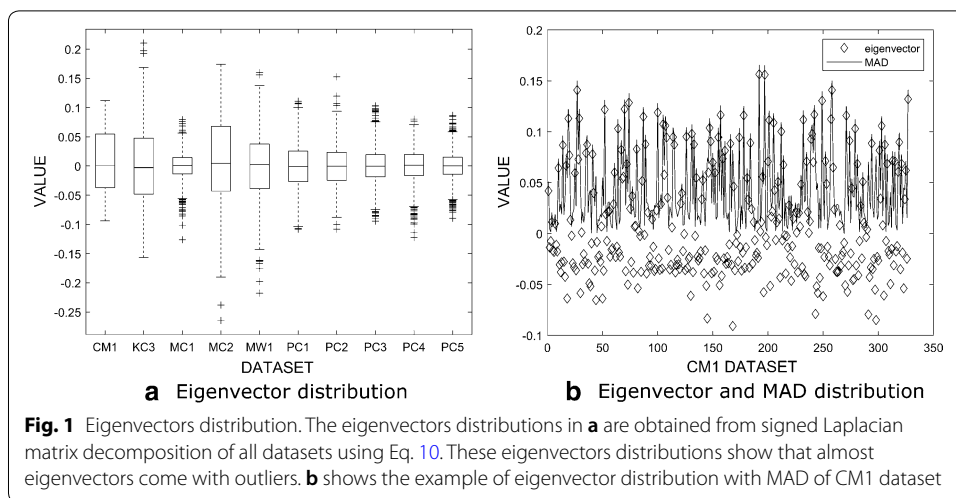
In the unsupervised approach, the predicted model can be developed directly using the current software dataset with no need training datasets [11]. This dataset contains a set of unlabeled software entities. The unsupervised approach relies on the clustering techniques to group the unlabeled dataset into the non-overlapped clusters, which are defective and non-defective [12]. All entities in each cluster are then labeled using a classifier [3, 13]. The use of the unsupervised software defect prediction has been studied using many methods. Zhong et al. [14] compared the  $k$ -means algorithm with the natural-gas algorithms to evaluate the clustering performance. Based on the mean square error values, the natural gas algorithm outperformed the  $k$ -means algorithm.

However, this method requires a software expert to decide whether the software belongs to either defective or clean class.

Catal et al. [15] applied  $x$ -means clustering algorithm to obtain defective and non-defective clusters using selected software metrics as the partitioning threshold. Those metrics are lines of code, cyclomatic complexity, operator, and operand. The software entity is predicted as defective if the metric values are higher than the threshold, and conversely. Bishnu et al. [16] proposed the quadtree based  $k$ -means algorithm and compare it to some clustering algorithms. The error rates of their proposed algorithm are reasonably comparable to the  $k$ -means, Naive Bayes, and Linear Discriminant Analysis. Abaei et al. [17] proposed the self-organizing map algorithm for software defect clustering. In the classes labeling, they used a similar threshold metrics that have been used in the  $x$ -means clustering algorithm, which is proposed by Catal et al. [15]. Their experimental results show that their algorithm outperformed both the Catal et al. [15] and Bishnu et al. [16] algorithms. Nam et al. [18] proposed the median-based partitioning threshold for clustering. All entities with higher values than the median are placed in one cluster, otherwise those entities are placed in the other cluster. The top half of the cluster members are then classified as defective, while the bottom half of the cluster members are classified as non-defective. Their proposed method has better performances compared to the Bayesian network, J48 decision tree, logistic model tree, logistic regression, Naive Bayes, random forest, and support vector machine algorithms.

Zhang et al. [3] applied the spectral graph to develop unsupervised based spectral classifier algorithm. The set of software entities and its connectivities represent the nodes and edges of the spectral graph, respectively. They used zero value threshold to obtain the predicted defective and clean clusters. For labeling all cluster members, they used the heuristic row sums criterion. Their experimental results show that the spectral classifier outperformed the random forest, logistic regression, Naive Bayes, and logistic model tree algorithms. Their proposed method is improved by Marjuni et al. [13] to ensure the use of spectral classifier requirements, especially in the non-negative Laplacian graph matrix. They applied the absolute of adjacency matrix to construct the signed Laplacian graph matrix. Their experiment shows that the use of signed Laplacian on spectral classifier can improve both the cluster compactness and classification.

Based on the preliminary experiment in this study, there are two conditional issues in the use of zero value threshold for spectral clustering. In case the eigenvector's values are mostly positives, then all of the entities will be placed predominantly into one cluster. The other case is if the eigenvector's values have balanced positives and negatives, then about half of all the entities will be predicted in the defective cluster. This kind of result might never occur in software project development. Second, the eigenvector's values sometimes come with outliers, as shown in Fig. 1. In the clustering process, this outliers issue may reduce the cluster compactness because of the wide distance between the eigenvector to the threshold. This study is motivated to address those issues by considering the characteristic of eigenvector values dispersion for partitioning. In this study, the median absolute deviation is chosen as the partitioning threshold to represent the eigenvector values dispersion. That means that the partitioning process will use the dispersion



measure belonging to the eigenvector position rather than using the central tendency measure (e.g., zero, mean, or median). Thus, the objective of this study is specifically to address the partitioning issues in spectral clustering such that could improve the spectral classifier based on the zero value threshold.

This paper is presented in several sections, as follows. "Introduction" section describes the background, issues, and motivation of this study. "Related works" section presents the current studies related to the use of spectral clustering and classifier in unsupervised based software defect prediction. "Basic theories" section presents a brief of median absolute deviation and signed Laplacian based spectral clustering as the basic theories of this study. "Methods" section presents the median absolute based spectral classifier as the proposed method and also the zero threshold based spectral classifier as the baseline of this study. "Experimental setup" section presents the dataset preparation, experimental design, performance evaluation, and validation method. The experimental results and discussion of this study are presented in "Results and discussion" section. Finally, "Conclusion" section summarizes the conclusions and future work of this study.

**Related works**

The unsupervised approach is usually used to avoid the limitation in the availability of the training datasets. A prediction model in this approach is commonly built using clustering methods. In this study, spectral clustering is chosen as the main study because it has the advantage to consider the similarities between entities. As a graph based clustering, spectral clustering also has better performance than distance based clustering methods [3, 4].

Zhang et al. [3] initially proposed the spectral classifier in unsupervised software defect prediction to address the heterogeneity issue of the dataset in cross project software defect prediction. The negative similarities of the adjacency matrix are transformed into zeros to ensure the non-negative Laplacian matrix assumption. The Laplacian matrix is then decomposed to obtain the eigenvalues and eigenvectors. The second smallest eigenvalue and its eigenvector are then selected to construct the clusters by comparing the eigenvector value with the zero value threshold [19]. All entities with the eigenvector value greater than zero are grouped in the predicted defective cluster. Otherwise, these entities are grouped in the

predicted non-defective cluster. The heuristic row sum criterion is then used to predict the label class of all the cluster members. The software entity is predicted as defective if its row sum greater than the row sum average of its cluster. Overall, their algorithm outperformed the  $k$ -means, partition around medoids, fuzzy  $C$ -means, and natural-gas algorithms.

However, there is a slight issue in the adjacency matrix construction. The adjacency matrix is a matrix that is used to represent the similarities between the pair of entities. This adjacency matrix can contain positive, zero, or negative values. If the adjacency matrix contains negative values, then this matrix might not be able to produce the non-negative Laplacian matrix, and the Laplacian matrix in spectral classifier becomes undefined. To address this issue, Marjuni et al. [13] proposed the signed Laplacian based spectral classifier. They performed an absolute transformation on the adjacency matrix to fulfill the non-negative Laplacian matrix assumption. Similar to the spectral classifier proposed by Zhang et al. [3], they also used the zero value thresholding for clustering and the heuristic row sum criterion for labeling. Their experimental results show that the use of signed Laplacian matrix can produce more cluster memberships, and affects to the classifier performances improvement.

There are some advantages of the use of signed Laplacian matrix in spectral clustering. First, the signed Laplacian matrix is positive semidefinite and fulfills the non-negative assumption of the Laplacian matrix. Second, the cluster memberships become increase because there are additional numbers of similarities obtained from negative similarities in the adjacency matrix. The increasing of the cluster memberships will affect the cluster compactness improvement [12, 13, 18, 20–23].

### Basic theories

The goal of this study is to improve the spectral classifier performance on signed Laplacian matrix by considering the eigenvector's values dispersion as the partitioning threshold. This dispersion is measured by the median absolute deviation, which is also known as a positional characteristic of data distribution.

#### Median absolute deviation

Median absolute deviation (MAD) is used for dispersion measurement corresponds to the absolute deviation from median [24–28]. This measure is robust against data outliers and suitable for both nonparametric estimator location and scale [29–32]. The MAD value of the sample dataset is obtained as follows.

Given  $x = (x_1, x_2, \dots, x_n)$  is an univariate vector of ordered increasingly quantitative dataset with size of  $n$ , and  $\tilde{x}$  is the median of  $x$ . The MAD value of  $x$  is computed in Eq. 1 as follows:

$$MAD = \text{median } |x_i - \tilde{x}|, \quad \text{for } x_i \in x. \quad (1)$$

The median  $\tilde{x}$  is calculated in Eq. 2 with odd and even cases in the number of data size of  $n$ :

$$\tilde{x} = \begin{cases} x_{\lceil \frac{n+1}{2} \rceil}, & \text{if } n \text{ is odd} \\ \frac{x_{\lceil \frac{n}{2} \rceil} + x_{\lfloor \frac{n}{2} \rfloor + 1}}{2}, & \text{if } n \text{ is even} \end{cases}, \quad (2)$$

where  $x_n$  is a data value of  $n$ th term, and satisfies the probabilities in Eq. 3, as follows:

$$P(x \leq \tilde{x}) \geq 1/2 \text{ and } P(x \geq \tilde{x}) \geq 1/2. \tag{3}$$

It means that the median  $\tilde{x}$  is the middle value of  $x$ , where a half of the values of  $x$  are greater than  $\tilde{x}$  and a half of the values of  $x$  are less than  $\tilde{x}$ . If  $|x_i - \tilde{x}|$  is a deviation between the data point and the median that are:  $|x_1 - \tilde{x}|, |x_2 - \tilde{x}|, \dots, |x_n - \tilde{x}|$ , then the median of  $|x_i - \tilde{x}|$  in Eq. 1 also satisfies the probabilities in Eq. 4, as follows:

$$P(|x_i - \tilde{x}| \leq \tilde{x}) \geq 1/2 \text{ and } P(|x_i - \tilde{x}| \geq \tilde{x}) \geq 1/2. \tag{4}$$

The median value represents the measure of the central tendency of the dataset that computes the rank of the data observations. Whilst the MAD value represents the measure of dataset dispersion through the location of item data to its medians. This characteristic makes the MAD more resistant to outliers than the median value itself [33].

There are two steps for determining the MAD value. First, computing the median of the dataset. Second, computing the median of the absolute distances between the instance values from the median. For example, given a set of a quantitative dataset of  $A = \{42, 25, 75, 40, 20\}$ , and  $B = \{42, 25, 75, 40, 20, 200\}$ . The ordered sets of A and B are  $A = \{20, 25, 40, 42, 75\}$ , and  $B = \{20, 25, 40, 42, 75, 200\}$ . The size of set A is  $n_A = 5$  (odd), while the size of set B is  $n_B = 6$  (even). The median of A is  $\tilde{x}_A = x_{\lceil \frac{5+1}{2} \rceil} = x_{[3]}$ , that is  $\tilde{x}_A = 40$ . Similarly, the median of B is  $\tilde{x}_B = \frac{x_{\lceil \frac{6}{2} \rceil} + x_{\lceil \frac{6}{2} + 1 \rceil}}{2} = \frac{x_{[3]} + x_{[4]}}{2} = \frac{40 + 42}{2}$ , that is  $\tilde{x}_B = 41$ . The absolute distances between the instances and its median of A and B, namely  $D_A$  and  $D_B$ , are  $D_A = \{20, 15, 0, 2, 35\}$  and  $D_B = \{21, 16, 1, 1, 34, 159\}$ , respectively. The ordered of  $D_A$  and  $D_B$  are  $D_A = \{0, 2, 15, 20, 35\}$  and  $D_B = \{1, 1, 16, 21, 34, 159\}$ . Hence, the MAD of A and B are  $MAD_A = 15$  and  $MAD_B = 18.5$ , respectively.

**Signed Laplacian based spectral clustering**

In software defect prediction, the terminology of a graph can be used to represent a software entities dataset. Software entities reflect the graph nodes, while the similarities between software entities reflect the graph edges. The software entities and their similarities are used to construct the adjacency matrix as graph representation. The adjacency matrix is then used to define the non-negative Laplacian graph matrix. In spectral classifier, the Laplacian graph matrix is decomposed to obtain the eigenvalues and eigenvector that are used for clustering process [29, 34]. This study uses the signed Laplacian graph matrix to apply the spectral classifier in unsupervised software defect prediction.

Given a software dataset  $D = \{x\}_{ij}^n$  that contains  $n$  entities with  $m$  metrics. An adjacency matrix  $W$  of  $D$  is defined as a matrix of similarities between entities as follows:

$$\begin{aligned}
 W &= \begin{bmatrix} w_{11} & w_{12} & \dots & w_{1n} \\ w_{21} & w_{22} & \dots & w_{2n} \\ \dots & \dots & \dots & \dots \\ w_{n1} & w_{n2} & \dots & w_{nn} \end{bmatrix}, \\
 &= x_i \cdot x_j \\
 &= \sum_{k=1}^m a_{ik} \cdot a_{kj}
 \end{aligned} \tag{5}$$

where  $x_i$  and  $x_j$  are the metric vectors of software entities  $i$  and  $j$ , respectively;  $a_{ik}$  is the  $k$ th metric value for the  $i$ th software entity; and  $m$  is the total number of metrics. The element  $w_{11} \in W$  can be positive, negative, or zero [20, 33]. This adjacency matrix is then used to construct the Laplacian matrix. In the signed graph, the Laplacian matrix must fulfill the positive semi-definite requirement [35]. In this study, the adjacency matrix  $W$  in Eq. 5 is transformed into its absolute to ensure the requirement, that is:

$$\overline{W} = |W|. \tag{6}$$

The signed Laplacian matrix is then defined as the following terms. For each entity  $i$ , let  $\overline{d}_i^{-1/2}$  denotes the signed degree of entity  $i$ th for  $\overline{w}_{ij} \in \overline{W}$ , which is defined as:

$$\overline{d}_i^{-1/2} = \sum_{j=1}^n |\overline{w}_{ij}|^{-1/2}, \tag{7}$$

and let  $\overline{D}^{-1/2}$  denotes the normalized diagonal matrix of signed degree  $\overline{d}_i^{-1/2}$ , which is defined as:

$$\overline{D}^{-1/2} = \text{diag}(\overline{d}_1^{-1/2}, \overline{d}_2^{-1/2}, \dots, \overline{d}_n^{-1/2}), \tag{8}$$

and also let  $I$  denotes the unity matrix with size of  $n$ . The signed Laplacian matrix is then defined as:

$$\overline{L}_{sym} = I - \overline{D}^{-1/2} \overline{W} \overline{D}^{-1/2}. \tag{9}$$

The signed Laplacian matrix  $\overline{L}_{sym}$  is then decomposed to get its spectrum that are both the eigenvalues and eigenvectors. A non-zero vector  $\overline{v}$  of size  $n$  is an eigenvector of a signed Laplacian matrix  $\overline{L}_{sym}$  of size  $n$  if there is an eigenvalues  $\overline{\lambda}$  such that satisfies the following linear equation:

$$\overline{L}_{sym} \overline{v} = \overline{\lambda} \overline{v}. \tag{10}$$

In spectral clustering, the second smallest eigenvalue (e.g.,  $\overline{\lambda}_1$ ) and the associated eigenvector (e.g.,  $\overline{v}_{1i}$ ) are then selected to generate clusters [19, 35].

## Methods

### Proposed method

In this study, the MAD value of the eigenvector dispersion is proposed as the partition threshold to generate the predicted defective and clean clusters. All software entities with eigenvector's value greater than the MAD value are grouped into the predicted defective cluster. Otherwise, these entities are grouped into the predicted clean cluster. This cluster membership is formulated as follows.

Suppose  $\overline{v}_{1i}$  is the eigenvector, which is associated to the second smallest eigenvalue  $\overline{\lambda}_1$  of the signed Laplacian matrix (i.e., obtained in Eq. 10), and  $MAD_i$  is the median absolute deviation of eigenvector  $\overline{v}_{1i}$  (i.e., computed using Eq. 1). The cluster membership of entity  $x_i$ , namely  $\hat{c}_i$ , is then defined as:

$$\hat{c}_i = \begin{cases} 1, & \text{for } \overline{v}_{1i} > MAD_i \\ 0, & \text{otherwise} \end{cases}. \tag{11}$$

All entities with  $\hat{c}_i = 1$  are placed as a member of the predicted defective cluster, namely  $C_d$ . Conversely, all entities with  $\hat{c}_i = 0$  are placed as a member of the predicted clean cluster, namely  $C_c$ .

Finally, each member in all of the predicted clusters will be labeled as defective or clean. The labeling method is adopted from the heuristic method using row sum criteria, which is proposed by Zhang et al. [3]. If the row sum of an entity greater than the average row sum of its cluster, the class of this entity will be labeled as defective, otherwise it will be labeled as clean. This criterion is formulated as follows. Suppose  $C_i$  is a predicted cluster of  $i$ th with  $i = 1, 2, \dots, k$ . In this study, the  $k$  value is 2 ( $k = 2$ ) because there are only two predicted clusters that are defective and clean clusters. Suppose  $r_{x_i} = \sum_{j=1}^m x_{ij}$  is the row sum of entity  $x_i \in C_i$  with  $m$  attributes. Suppose  $\bar{r}_{c_i} = \frac{1}{n} \sum_{i=1}^n r_{x_i}$  is the average row sum of cluster  $i$ th. The label of entity  $x_i$ , namely  $c_i$ , is defined as:

$$c_i = \begin{cases} 1, & \text{for } r_{x_i} > \bar{r}_{c_i} \\ 0, & \text{otherwise} \end{cases} \quad (12)$$

All entities with  $c_i = 1$  will be labeled as defective. Conversely, all entities with  $c_i = 0$  will be labeled as clean. The brief of this proposed method is summarized in Algorithm 1.

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**Algorithm 1** MAD threshold based spectral classifier.

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**Step 1.** Normalize dataset using  $z$ -score transformation.

**Step 2.** Construct the adjacency matrix  $W$ .

**Step 3.** Transform the adjacency matrix into its absolute (i.e.,  $\bar{W} = |W|$ ).

**Step 4.** Construct the signed Laplacian matrix  $\bar{L}_{sym}$ .

**Step 5.** Decompose  $\bar{L}_{sym}$  and select the second smallest eigenvalue with the associated eigenvectors.

**Step 6.** Compute the MAD threshold on the eigenvector.

**Step 7.** Perform clustering using MAD threshold.

**Step 8.** Labeling all entities in each cluster.

---

**Baseline method**

This study uses the zero threshold based spectral classifier [13] as the baseline method. All issues addressed in this study come from the baseline method. Thus, the proposed method in this study is generally designed to improve the performance of the baseline method. The baseline method algorithm is similar to the proposed method algorithm, with a slight difference in the partitioning threshold. In the baseline method, the cluster membership of entity  $x_i$ , namely  $\hat{c}_i$ , is defined as:

$$\hat{c}_i = \begin{cases} 1, & \text{for } \bar{v}_{1i} > 0 \\ 0, & \text{otherwise} \end{cases} \quad (13)$$

All entities with  $\hat{c}_i = 1$  are placed as a member of the predicted defective cluster, namely  $C_d$ . Conversely, all entities with  $\hat{c}_i = 0$  are placed as a member of the predicted clean cluster, namely  $C_c$ . Algorithm 2 summarizes the algorithm of the baseline method.

**Algorithm 2** Zero threshold based spectral classifier.

**Step 1.** Normalize dataset using  $z$ -score transformation.

**Step 2.** Construct the adjacency matrix  $W$ .

**Step 3.** Transform the adjacency matrix into its absolute (i.e.,  $\overline{W} = |W|$ ).

**Step 4.** Construct the signed Laplacian matrix  $\overline{L}_{sym}$ .

**Step 5.** Decompose  $\overline{L}_{sym}$  and select the second smallest eigenvalue with the associated eigenvectors.

**Step 6.** Perform clustering using zero threshold.

**Step 7.** Labeling all entities in each cluster.

**Experimental setup**

This experimental setup is prepared to conduct the experiments on both the proposed and baseline methods through Algorithm 1 and Algorithm 2, respectively. This experimental setup includes the dataset preparation, experimental design, performance evaluation, and validation method.

**Dataset preparation**

This experiment uses ten of the NASA Metrics Data Program (MDP) public dataset repositories that have been commonly used to study the software defect prediction. Table 1 summarizes those datasets. Each dataset consists of hundreds to thousands of program module entities [36, 37] with different numbers in the defect proportions. The original NASA MDP dataset repositories may need to be cleaned to remove the data redundancy and inconsistency to get more feasible for analyzing purposes [3, 38]. All

**Table 1** NASA MDP datasets

Dataset name	Dataset description	Programming language	Number of instance	Defect ratio (%)
CM1	Spacecraft instrument	C	327	12.8
KC3	Storage management for ground data	Java	194	18.6
MC1	Zero gravity experiment related to combustion	C and C++	1988	2.3
MC2	Video guidance system	C	125	35.2
MW1	Zero gravity experiment related to combustion	C	253	10.7
PC1	Flight software from an earth orbiting satellite	C	705	8.7
PC2	Dynamic simulator for attitude control systems	C	745	2.1
PC3	Flight software for earth orbiting satellite	C	1077	12.4
PC4	Flight software for earth orbiting satellite	C	1287	13.8
PC5	Flight software for earth orbiting satellite	C++	1711	27.5



public datasets in this study are online available, which are provided by Klainfo [39] and Shepperd et al. [40].

### Data preprocessing

As shown in Algorithm 1 and Algorithm 2, the preprocessing step using  $z$ -score transformation is added in both algorithms. The transformation aims to standardize the metric values so that all metrics have the same value scale on the model development. The  $z$ -score transformation is formulated in Eq. 14.

$$\hat{y}_i = \frac{y_i - \bar{y}_i}{s_i}, \quad (14)$$

where  $y_i$  is the metric vector of  $i$ ,  $\bar{y}_i$  is the mean of  $y_i$ ,  $s_i$  is the standard deviation of  $y_i$ , and  $\hat{y}_i$  is the normalized value of  $y_i$ .

### Experiment design

Both of the proposed and baseline methods in this study have two main processes that are clustering and labeling, as shown in Algorithm 1 and Algorithm 2. Before carrying out those main processes, the data preprocessing is performed to improve the dataset quality for analysis purpose. In this preprocessing, the dataset is standardized by the  $z$ -score transformation in Eq. 14 to improve the data normalization and features scaling. The main experiments are then sequentially taken as follows.

- Clustering: The clustering process is partitioning process to generate the defect and clean clusters. The clustering performance is evaluated using Davies Bouldin Index (DBI) that is computed using Eq. 15. The DBI is a universal measure that is usually used to validate any clustering performance as the internal validation.
- Labeling: The labeling process is classification process to obtain the predicted class of all entities in each cluster using the row sum criterion in Eq. 12. The labeling performance is evaluated using some measures, such as precision, recall, accuracy, AUC, and error rates that are computed using Eqs. 17–21, respectively.

The study is conducted using Matlab software to evaluate both Algorithm 1 and Algorithm 2 with the prepared datasets.

### Performance evaluations

For clustering performance, the clustering algorithm is evaluated using an internal validation that is measured by the DBI, as follows:

$$DBI = \frac{1}{k} \sum_{i=1}^k \max_{j \neq i} DB_{ij}, \quad (15)$$

where  $k$  is the number of all clusters, and  $DB_{ij}$  is calculated as follows:

$$DB_{ij} = \frac{\sigma_{\mu_i} + \sigma_{\mu_j}}{\delta(\mu_i, \mu_j)}, \quad (16)$$

**Table 2** Confusion matrix

Actual	Predicted	
	False (clean)	True (defective)
False (clean)	True negative (TN)	False positive (FP)
True (defective)	False negative (FN)	True positive (TP)

where  $\mu_i$  is the cluster mean of  $C_i$ ,  $\mu_j$  is the mean of cluster  $C_j$ ,  $\delta(\mu_i, \mu_j)$  is the distance between cluster mean of  $\mu_i$  and  $\mu_j$ , while  $\sigma_{\mu_i}$  and  $\sigma_{\mu_j}$  are the standard deviation of cluster  $C_i$  and  $C_j$ , respectively. The smaller DBI indicates the better clustering, which means the predicted clusters are well partitioned because of the distance between each cluster means is large [20, 33, 36].

For labeling performance, the predicted labels are compared to the actual labels using confusion matrix for labeling performance evaluation [41]. The confusion matrix presents the number of predicted labels in columns and the number of actual labels in rows [16, 33], as shown in Table 2, which is constructed by these following elements:

- True negative (TN): The number of clean entities that the classifier predicts as clean.
- False positive (FP): The number of clean entities that classifier predicts as defective.
- False negative (FN): The number of defective entities that classifier predicts as clean.
- True positive (TP): The number of defective entities that the classifier predicts as defective.

The prediction performance measurements [33, 41–43] are then computed using the confusion matrix elements, as follows:

- Precision (PRE), which is measured by the fraction of true positive with all predicted defective entities.

$$\text{PRE} = \frac{\text{TP}}{\text{TP} + \text{FP}} \quad (17)$$

- Recall (REC), which is measured by the fraction of true positive with all actual defective entities. This metric is also called as true positive rate (TPR).

$$\text{REC} = \frac{\text{TP}}{\text{TP} + \text{FN}} \quad (18)$$

- Accuracy (ACC), which is measured by the fraction of all correct predicted entities with all total entities.

$$\text{ACC} = \frac{\text{TP} + \text{TN}}{\text{TP} + \text{FP} + \text{FN} + \text{TN}} \quad (19)$$

- Area under curve (AUC), which is measured by the area under curve of ROC (the receiver operating characteristics). The ROC itself is constructed by the curves of true positive rate across false positive rate. The true positive rate (TPR) and false positive rate (FPR) are computed using Eqs. 18 and 20, respectively [20].

$$\text{FPR} = \frac{\text{FP}}{\text{FP} + \text{TN}} \quad (20)$$

- Error rates (ERR) or misclassified error, which is measured by the fraction all incorrect predicted entities with all total entities.

$$\text{ERR} = \frac{\text{FP} + \text{FN}}{\text{TP} + \text{FP} + \text{FN} + \text{TN}} = 1 - \text{ACC} \quad (21)$$

### Validation method

Validation method is performed by comparing the proposed method performances to the baseline method. The significance of performance differences between the proposed and baseline methods are validated using the Wilcoxon signed-rank test [44, 45] at the 95% level of confidence. The Wilcoxon signed-rank is a nonparametric statistical test. Thus it does not need any assumptions about the dataset distribution. The validation test uses the null hypothesis  $H_0$  based on the median difference between pairs of data samples. This hypothesis is then tested using the  $p$  value for a decision making whether the null hypothesis is accepted or rejected [36, 46]. The  $p$  value is a probability associated with the critical value that a Type I error (false positive) is allowed. These are the following hypotheses for this study.

- $H_{0(1)}$ : There is no difference in cluster compactness between the proposed and baseline methods.
- $H_{0(2)}$ : There is no difference in precision performance between the proposed and baseline methods.
- $H_{0(3)}$ : There is no difference in recall performance between the proposed and baseline methods.
- $H_{0(4)}$ : There is no difference in accuracy performance between the proposed and baseline methods.
- $H_{0(5)}$ : There is no difference in error rates between the proposed and baseline methods.
- $H_{0(6)}$ : There is no difference in AUC performance between the proposed and baseline methods.

All of the null hypotheses will be tested separately using the two-tailed Wilcoxon signed-rank test at the 95% level of confidence (i.e.,  $p < 0.05$ ). If the  $p$  value is less than 0.05, then the null hypothesis is rejected, and the performances of both methods are significantly different. Otherwise, the null hypothesis is accepted.

### Results and discussion

This experiment is conducted to evaluate the performances of the proposed method and comparing those performances to the baseline. The proposed and baseline methods are summarized in Algorithm 1 and Algorithm 2, respectively. As explained in "Experimental setup" section, both the proposed and baseline methods are performed on the signed Laplacian matrix, which is constructed by Eq. 6.

The data preprocessing, which follows step 1 in Algorithm 1 and Algorithm 2, is performed to standardize the dataset on the same scale. The clustering process of both the proposed and baseline algorithm follows step 2 to step 7 in Algorithm 1 and step 2 to step 6 in Algorithm 2. Step 2 and step 3 are constructing the adjacency matrix and the absolute of adjacency matrix using Eqs. 5 and 6, respectively. Step 4 is constructing the signed Laplacian matrix in Eq. 9 from the absolute of adjacency matrix in step 3. The signed Laplacian matrix is then decomposed to get the eigenvalues and eigenvectors using Eq. 10 correspond to the step 5. Figure 1a shows the eigenvectors distribution for all datasets. Almost all of the eigenvectors come with outliers in their distribution. If the zero value threshold is used to generate clusters on those eigenvectors, it may produce clusters with low compactness in cluster memberships.

In this study, the proposed method considers the eigenvector’s values dispersion measure using the MAD value to determine the new threshold. The MAD value is computed using Eq. 1. Figure 1b illustrates the comparison example between the eigenvector dispersion and its MAD on CM1 dataset. It shows that the MAD value overcomes the outliers in the eigenvector dispersion. The MAD value represents the eigenvector values position and does not reflect the changes of the eigenvector values. Table 3 presents the dispersion measures of eigenvectors for all datasets, such as range, interquartile range, standard deviation, and MAD.

The next step is the partitioning process to generate defective and clean clusters. The second smallest eigenvalue of the Laplacian matrix is chosen to define the selected eigenvector for the partitioning process. Besides, the partitioning process also needs a partitioning threshold. The proposed method uses the median absolute deviation of the selected eigenvector, while the baseline method used the zero value as the partition thresholds. The MAD value of the selected eigenvector is computed using Eq. 1, corresponds the step 6 in Algorithm 1. Both of these thresholds are then used to generate the clusters, corresponds the step 7 in Algorithm 1 and step 6 in Algorithm 2, respectively. The cluster memberships of both proposed and baseline methods follow Eqs. 11 and 13, respectively.

In the proposed method, the defective cluster is constructed to group all entities which its eigenvalue greater than the MAD threshold—otherwise, the clean cluster is

**Table 3 Eigenvector dispersion measures**

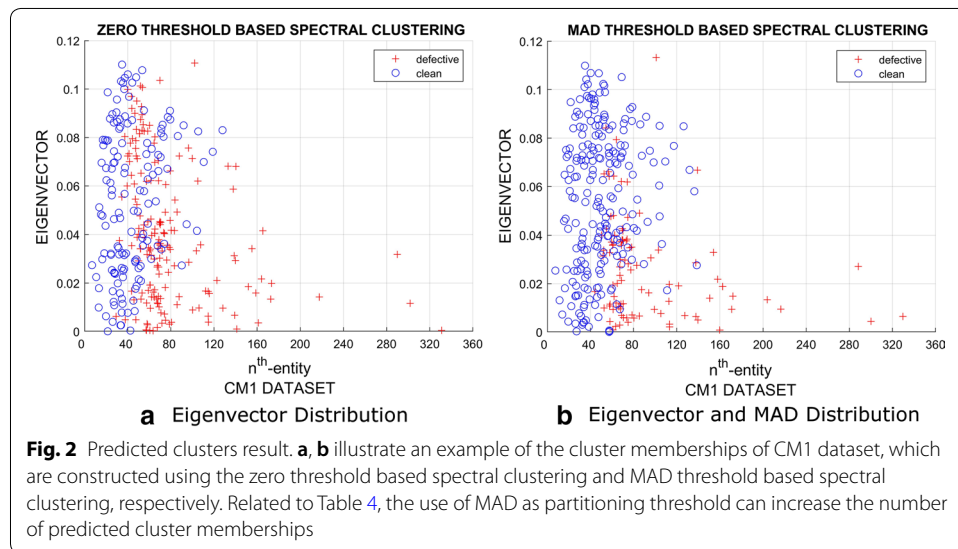
Dataset name	Range value	Interquartile range	Standard deviation	Median absolute deviation
CM1	0.247	0.085	0.054	0.028
KC3	0.278	0.119	0.070	0.054
MC1	0.313	0.020	0.022	0.013
MC2	0.369	0.127	0.088	0.060
MW1	0.311	0.091	0.062	0.035
PC1	0.197	0.063	0.037	0.030
PC2	0.252	0.052	0.036	0.027
PC3	0.255	0.071	0.041	0.035
PC4	0.142	0.090	0.027	0.018
PC5	0.104	0.042	0.024	0.021

**Table 4 Clusters membership and performance**

Dataset name	Zero threshold based spectral clustering			MAD threshold based spectral clustering		
	Cluster memberships (%)		Compactness (DBI)	Cluster memberships (%)		Compactness (DBI)
	$D_c$	$C_c$		$D_c$	$C_c$	
CM1	44.34	55.66	1.3	28.75	71.25	<i>1.1</i>
KC3	44.85	55.15	1.4	21.50	78.50	<i>1.2</i>
MC1	32.80	67.20	2.0	24.60	75.40	<i>1.8</i>
MC2	48.00	52.00	1.9	21.60	78.40	<i>1.1</i>
MW1	45.45	54.55	1.4	30.83	69.17	<i>1.0</i>
PC1	48.23	51.77	1.9	31.49	68.51	<i>1.3</i>
PC2	45.23	54.77	2.2	19.06	80.94	<i>1.5</i>
PC3	48.47	51.53	2.4	30.16	69.84	<i>1.4</i>
PC4	46.23	53.77	1.8	29.91	70.09	<i>1.2</i>
PC5	47.63	52.37	2.1	31.09	68.91	<i>1.9</i>
Average	45.12	54.88	1.8	26.90	73.10	<i>1.4</i>

$D_c$  is the predicted defective cluster;  $C_c$  is the predicted clean cluster

The italicized values indicate the better performance



constructed to group all entities which its eigenvector less or equals than the MAD threshold. In the baseline method, the defective cluster is constructed to group all entities which its eigenvalue greater than zero—otherwise, the clean cluster is constructed to group all entities which its eigenvector less or equals than zero. Table 4 shows the distribution of cluster memberships using both proposed and baseline methods.

The proposed method detects 26.90% of all entities as defective (i.e., in the defective cluster) and 73.10% of all entities as clean (i.e., in the defective cluster). Whereas, the baseline method detects 45.12% of all entities as defective and 54.88% of all entities as clean. Based on these results and compared to the number of defective in Table 1, the proposed method can produce clusters with better cluster memberships than the

**Table 5 Performance comparison between zero and MAD threshold based spectral classifiers**

Dataset name	Zero threshold based spectral classifier					MAD threshold based spectral classifier				
	PRE	REC	ACC	AUC	ERR	PRE	REC	ACC	AUC	ERR
CM1	0.72	<i>0.92</i>	0.71	0.68	0.29	<i>0.81</i>	0.71	<i>0.76</i>	<i>0.76</i>	<i>0.24</i>
KC3	<i>0.78</i>	<i>0.88</i>	<i>0.75</i>	0.75	<i>0.25</i>	0.72	0.61	0.71	<i>0.80</i>	0.29
MC1	0.76	<i>0.94</i>	0.78	0.67	0.22	<i>0.98</i>	<i>0.94</i>	<i>0.94</i>	<i>0.74</i>	<i>0.06</i>
MC2	<i>0.74</i>	<i>0.71</i>	<i>0.68</i>	0.75	<i>0.32</i>	0.60	0.57	0.67	<i>0.82</i>	0.33
MW1	0.69	<i>0.95</i>	0.69	0.70	0.31	<i>0.84</i>	0.58	<i>0.71</i>	<i>0.74</i>	<i>0.29</i>
PC1	0.73	<i>0.94</i>	0.75	<i>0.78</i>	0.25	<i>0.83</i>	0.64	<i>0.80</i>	0.77	<i>0.20</i>
PC2	0.72	<i>0.89</i>	0.77	<i>0.76</i>	0.23	<i>0.97</i>	0.63	<i>0.84</i>	0.74	<i>0.16</i>
PC3	0.72	<i>0.92</i>	0.79	<i>0.77</i>	0.21	<i>0.85</i>	0.86	<i>0.80</i>	0.77	<i>0.20</i>
PC4	0.79	<i>0.90</i>	0.72	0.67	0.28	<i>0.87</i>	0.80	<i>0.73</i>	0.77	<i>0.27</i>
PC5	0.78	0.81	0.78	<i>0.74</i>	0.22	<i>0.88</i>	<i>0.87</i>	<i>0.94</i>	<i>0.74</i>	<i>0.06</i>
Average	0.74	<i>0.89</i>	0.74	0.73	0.26	<i>0.84</i>	<i>0.72</i>	<i>0.79</i>	<i>0.77</i>	<i>0.21</i>

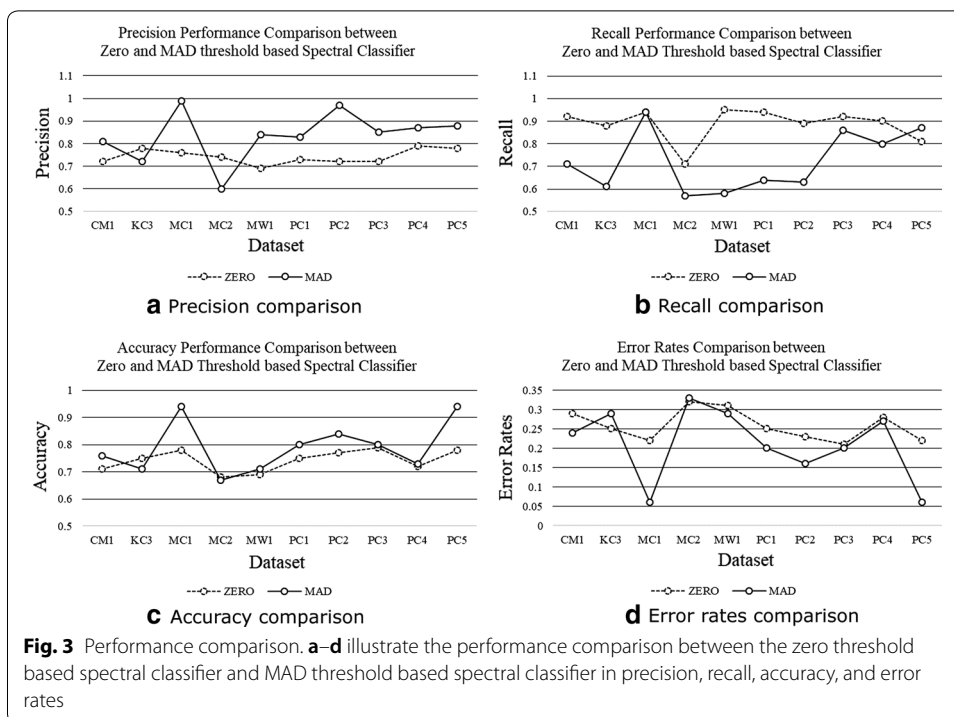
The italicized values indicate the better performance

baseline. The proposed method groups the entities based on their eigenvector positions that are measured by its distance to the median. Whereas, the zero threshold groups the entities based on their eigenvector difference values with the zero. Figure 2 shows the example of the cluster memberships on CM1 dataset that are produced by both the proposed and baseline methods.

The clustering performances of both methods are evaluated using the cluster compactness, which is measured by the DBI in Eq. 15. Table 4 shows the DBI values, where the proposed method outperforms the baseline for all datasets. The cluster compactness averages of both the proposed and baseline methods are 1.4 DBI and 1.8 DBI, respectively. This achievement corresponds with the increase in the number of cluster membership as referred on Table 4. The clustering performance difference between the proposed and baseline methods is validated using the two-tailed Wilcoxon signed-rank test at 95% level of confidence. The observed  $p$  value of this test is 0.001. Hence, the null hypothesis  $H_{0(1)}$  is rejected. This significance test concludes that the proposed method can produce better cluster compactness than the baseline method.

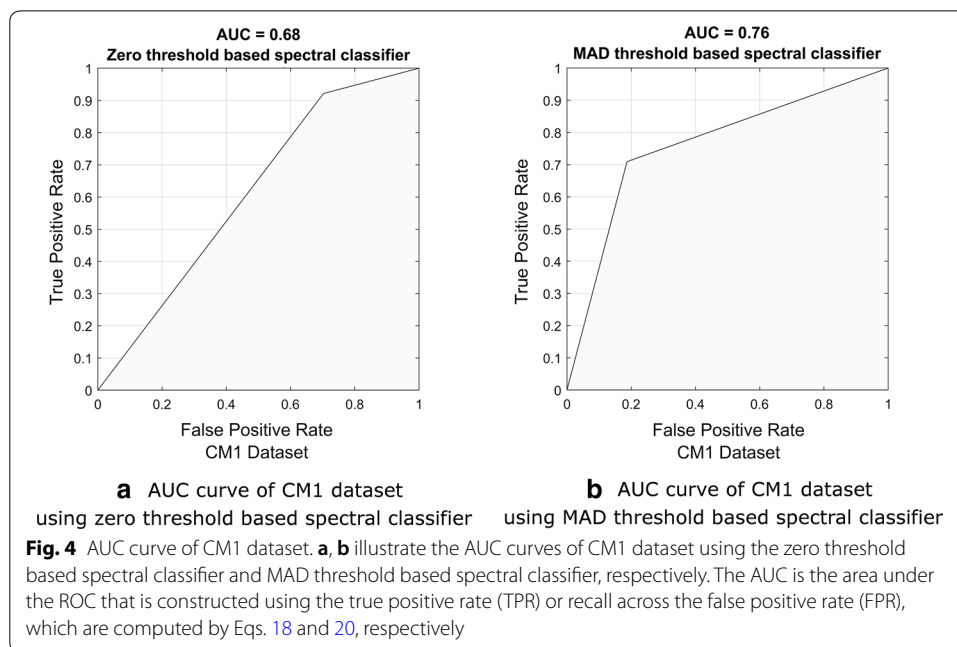
Following Algorithm 1 and Algorithm 2, the last step is labeling all cluster members as defective or clean classes. The label of each entity is obtained using the heuristic row sum criterion in Eq. 12 by comparing the row sum of the entity with the average row sum of its cluster. The classifier performances are then evaluated using precision, recall, accuracy, AUC, and also error rates, which are computed using Eq. 17–21, respectively. Table 5 shows all of these performances using both the proposed and baseline methods.

In precision performance, Table 5 shows that the proposed method can improve the precision of the baseline. The precisions averages of both methods are 0.84 and 0.74, respectively. While Fig. 3a illustrates the precision performance comparison between the proposed and baseline methods for all datasets. The precision performance difference has been validated using the two-tailed Wilcoxon signed-rank test at the 95% level of confidence. The observed  $p$  value is 0.005 and the null hypothesis  $H_{0(2)}$  is rejected. Hence, the precisions of both methods are significantly different. Table 5 also shows that the proposed method outperforms the baseline in the CM1, MC1, MW1, PC1,



PC2, PC3, PC4, and PC5 datasets. However, the proposed method underperformed in the KC3 and MC2 datasets. Related to the eigenvector’s values dispersion measure in Table 3, the KC3 and MC2 datasets have higher dispersion measures than the other datasets, especially in the interquartile range, standard deviation, and MAD values. Statistically, those eigenvector’s dispersion measures affect the entities distribution. For example, the interquartile range represents the range of the middle half of the eigenvector’s values distribution. The higher interquartile range indicates that the eigenvector’s values spread out from the central tendency point (e.g., median). Hence, the number of both defective and clean entities in the middle-half of the eigenvector’s values distribution becomes increasing. Consequently, the precision of the proposed method becomes decreasing in the KC3 and MC2 datasets and underperformed the baseline method.

In contrary to precision, the proposed method underperformed the baseline in the recall performance. As shown in Table 5, the recall values of the proposed method are smaller than the recall values of the baseline method. The recall averages of both methods are 0.72 and 0.89, respectively. Figure 3b illustrates the recall performance comparison between the proposed and baseline methods for all datasets. The recall performance difference is proven significant by the two-tailed Wilcoxon signed-rank test at the 95% level of confidence. The observed  $p$  value is 0.01 and the null hypothesis  $H_{0(3)}$  is rejected. Hence, the recall performance of the proposed method is significantly different from the baseline method. Table 5 also shows that recall of the proposed method outperforms the baseline only in the PC5 dataset. Related to the eigenvector’s values dispersion of this dataset in Table 3, the PC5 dataset has the lowest range of eigenvector values than the other datasets. It means that the difference value between the maximum and minimum of the eigenvectors in the PC5 dataset is too small. Hence, almost entities spread



around the central tendency point, including both the defective and clean entities. Consequently, the ratio of the correct predicted defective to all the actual defective entities (i.e., recall) in the PC5 dataset becomes increasing.

Based on the precision and recall, there is a trade-off performance between the proposed and baseline methods. The proposed method has high in precision but low in the recall, while the baseline method has low in precision but high in the recall. This trade-off indicates that the proposed method can predict more precisely the actual defective, although in the small number. Whereas the baseline method can predict more defective entities, but not all of the predicted defectives are actual defectives.

In terms of accuracy, Table 5 shows that the proposed method outperforms the baseline in almost all of datasets, but underperformed in the KC3 and MC2 datasets. These lower accuracies in those two datasets might be caused by the high dispersion measures in the interquartile range, standard deviation, and MAD, as shown in Table 3. Overall, the accuracy averages of both the proposed and baseline methods are 0.79 and 0.74, respectively. Hence, the error rate of both the proposed and baseline methods are 0.21 and 0.26, respectively. It means that 79% of all entities are classified correctly using the proposed method with the misclassified rates of 21%, while the baseline method can classify 74% of all entities correctly with the misclassified rates of 26%. Those accuracies and error rates indicate that the proposed method can predict the defective entities more accurate than the baseline. The performance comparison of accuracy and error rates between the proposed and baseline methods for all datasets are illustrated in Fig. 3c, d, respectively. The accuracy performance and error rates differences are proven significant by the Wilcoxon test at the 95% level of confidence. The observed  $p$  value of accuracy performance test is 0.03, and the null hypothesis  $H_{0(4)}$  is rejected. While the error rates difference test gives observed  $p$  value of this test is 0.04, and the null hypothesis  $H_{0(5)}$  is rejected.



In terms of AUC, Table 5 shows that the proposed method outperforms the baseline in almost all of datasets. The AUC averages of both methods are 0.77 and 0.73, respectively. It means that the abilities to distinguish the defective and clean classes of both methods are 77% and 73%, respectively. The AUC performance difference is proven significant by the two-tailed Wilcoxon signed-rank test at the 95% level of confidence. The observed  $p$  value of this test is 0.01, and the null hypothesis  $H_{0(6)}$  is rejected. Figure 4 illustrates the example of the AUC curve for the CM1 dataset. The AUC itself is the area under the ROC that is constructed using the true positive rate (TPR) across the false positive rate (FPR), which are computed by Eqs. 18 and 20, respectively.

## Conclusion

The median absolute deviation threshold has been presented in this paper to address the zero threshold issues in the spectral classifier based unsupervised software defect prediction. The median absolute deviation, as a dispersion measure, is proposed as the partitioning threshold to reduce the predominantly cluster issue when the eigenvector values of spectral graph matrix are mostly positives. The median absolute deviation threshold is also designed to address the eigenvector values outliers issue. Thus, the objective of this study is generally to improve the performances of zero value threshold based spectral classifier in unsupervised software defect prediction through the median absolute deviation threshold based spectral classifier.

Experimental results show that the proposed method can improve both clustering and classification. The proposed method produces better cluster memberships and increase cluster compactness. Which means there is an improvement in the numbers of entities that are separated correctly into the predicted clusters. This resulting clusters affect the classification performance improvement. The accuracy and misclassified rates of the median absolute deviation threshold based spectral classifier outperformed the zero threshold value based spectral classifier. In the AUC performance, the median absolute deviation threshold based spectral classifier also performs better in the ability to distinguish the defective and clean classes. In the precision and recall performances, the median absolute deviation threshold based spectral classifier high in precision although low in the recall. Based on those achievements, this study concludes that the median absolute deviation threshold based spectral classifier not only able to overcome the zero value threshold issues in the spectral classifier. But, it can also improve the zero value threshold based spectral classifier performances. The median absolute deviation threshold based spectral classifier method is an unsupervised approach with no need of training dataset. Thus, there is a significant potential to generalize the proposed method on the new software projects or an established software project with lacks in training dataset.

This study only deals with the zero threshold issues in the spectral classifier based unsupervised software defect prediction and does not perform any feature analyses, such as feature selection, auto-correlation analysis, or multi-collinearity analysis. As well known, feature analyses are essential to obtain the only relevant features in the classifier model development. Hence, for future work, the feature analyses would be considered to optimize the achievement of this study.

### Abbreviations

MAD: median absolute deviation; MDP: metrics data program; DBI: Davies Bouldin index; DB: Davies Bouldin; TP: true positive; TN: true negative; FP: false positive; FN: false negative; PRE: precision; REC: recall; TPR: true positive rates; FPR: false positive rates; ACC: accuracy; ERR: error rates; AUC: area under curve; ROC: receiver operating characteristics.

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### Authors' contributions

All authors contributed both the concepts and contents of this study. AM provided the manuscript under supervised by TBA and RF. All authors also performed discussion intensively for contents improvement. All authors read and approved the final manuscript.

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### Availability of datasets

The datasets used in this paper are publicly online available as described in "Dataset preparation" section.

### Competing interests

The authors declare that they have no competing interests.

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