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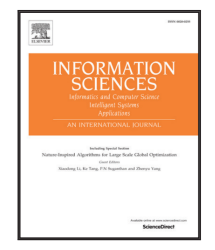
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A Genetic Algorithm Approach to Optimising Random Forests Applied to Class Engineered Data

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Abstract

In numerous applications and especially in the life science domain, examples are labelled at a higher level of granularity. For example, binary classification is dominant in many of these datasets, with the positive class denoting the existence of a particular disease in medical diagnosis applications. Such labelling does not depict the reality of having different categories of the same disease; a fact evidenced in the continuous research in root causes and variations of symptoms in a number of diseases. In a quest to enhance such diagnosis, datasests were decomposed using clustering of each class to reveal hidden categories. We then apply the widely adopted ensemble classification technique Random Forests. Such class decomposition has two advantages: (1) diversification of the input that enhances the ensemble classification; and (2) improving class separability, easing the follow-up classification process. However, to be able to apply Random Forests on such class decomposed data, three main parameters need to be set: number of trees forming the ensemble, number of features to split on at each node, and a vector representing the number of clusters in each class. The large search space for tuning these parameters has motivated the use of Genetic Algorithm to optimise the solution. A thorough experimental study on 22 real datasets was conducted, predominantly in a variety of life science applications. To prove the applicability of the method to other areas of application, the proposed method was tested on a number of datasets from other

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domains. Three variations of Random Forests including the proposed method as well as a boosting ensemble classifier were used in the experimental study. The results prove the superiority of the proposed method in boosting up the accuracy.

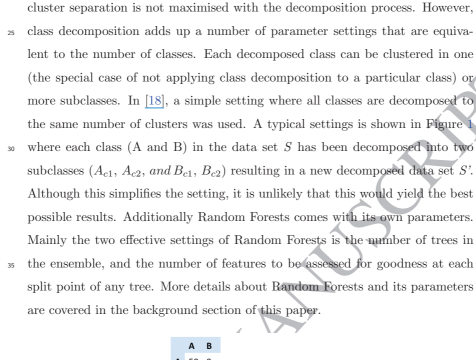
Keywords: Random Forests, Genetic Algorithm, Class Decomposition, Life Science

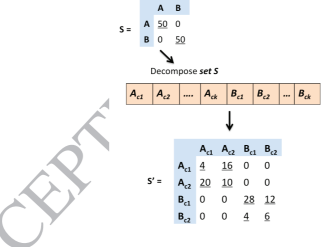
1. Introduction

Class decomposition is the process breaking down labelled datasets to a larger number of subclasses by means of applying clustering to the instances that belong to one class at a time. As such, the decomposition can be applied to one or more classe(s) in the data set. A typical scenario is illustrated in Figure 1 where a binary dataset S has been decomposed into multiple class problem (S'). Class decomposition can be traced back to 2003 when suggested to mitigate the issue of low variance classification methods [37]. However, it has been proposed in the context of biomedical data mining, as a data preprocessing phase for supervised learning. The motive is that genuine subclasses 10 can be detected, and as such the accuracy of the classification process can be enhanced. Taking two stages of development in this area of application, the work reported in [31] represents the first stage, it has bee applied to the positive class only of a number of biomedical datasets. In [18], the second stage is represented by generalising the class decomposition to all the classes in medical diagnosis 15 data sets.

In [18], Random Forests over class decomposed medical diagnosis data sets has been adopted as recent experimental studies showed its favourable results over other state-of-the-art methods [20]. In addition to the motive of finding genuine subclasses, the diversification of the data set originated from the process of class decomposition can further enhance the performance of ensemble classification methods, which in this case are represented by Random Forests. As such, it is desirable to apply class decomposition to all classes, even if the

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Figure 1: Class decomposition

Realising that setting the parameters for Random Forests over class decomposed datasets with its settings of number of clusters is an optimisation problem with a large search space, *Genetic Algorithm* is adopted to set all the parameters. Genetic Algorithm is superior to other optimisation methods when there are a relatively large number of local optima, which is the case in this problem. The search space is exponential in the number of classes available in the data set. If the range of setting the number of subclasses is r, where $r \in \mathbb{N}$, the

- ⁴⁵ number of classes in the data set is *nClasses*, the |mtry| is the range for the number of features to use to split on at each node, and |ntree| is the range of the number of trees in the Random Forests ensemble, the search space is in $O(r^{nClasses}|mtry||ntree|)$. For example, for a modest classification problem, if r = 10 (the number of subclasses attempted for each class ranges from 1 to 10),
- c = 5 (the number of classes is 5), |mtry| = 10 (the range of the number of features used to split on at each node), |ntree| = 100 (the range of the number of trees that form the ensemble), the search space is $10^5 \times 10 \times 100$, resulting in a large search space of 10^8 solutions. The contributions of this paper can be summarised as follows.
- Optimisation of Random Forests parameters applied to class decomposed datasets using Genetic Algorithm. These are the number of trees and the number of features;
 - optimisation of the class decomposition parameters by varying the setting of number of classes; and
 - experimental validation of the proposed technique when applied to 22 datasets, mostly in the area of life sciences with emphasis on biomedical datasets, with exception of a number of datasets from other domains to prove the general applicability of the method.

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The paper is organised as follows. Section 2 gives the necessary background of the computational intelligence and machine learning methods adopted in this research, namely, Random Forests and Genetic Algorithm. Section 3 reviews related work and contextualise the research accordingly. The proposed methods used in conducting the work have been detailed in Section 4. Section 5 provides a presentation of the experimental work and the results. Finally, the paper is concluded in Section 6 with a summary and possible directions for future work.

2. Background

2.1. Random Forests

Ensemble classification methods have passed the test of time, and proved to be highly accurate prediction and classification techniques. According to the ⁷⁵ winning solutions in *Kaggle*¹, the state-of-the-art ensemble methods are Random Forests [11, 13] and Gradient Boosting trees [22]. Random Forests has proved superiority experimentally when compared with all widely adopted classifiers including Gradient Boosting trees [20]. As an ensemble method, Random Forests adopts two methods for model diversification: (1) bootstrap sampling

- that applies sampling with replacement generating what is known as data replicas; and (2) each tree in the random forests chooses its node splits from a subset of the total number of features. The bootstrap sampling in the context of ensemble classification is referred to as *Bagging* [10]. Typically Random Forests would need two parameters to set, namely, the number of trees and number
- of features assessed for goodness of split at each node in the tree. As a rule of thumb, the number of trees is set between 100 and 500, and the number of features is set \sqrt{n} or $\log_2(n)$ where n is the total number of features in a data set.

A number of extensions have been proposed to further enhance the performance of Random Forests [19]. In [15], the authors addressed a number of Big Data problems adopting Random Forests arguing for its robustness as a classifier. Problems addressed are oversampling, undersampling, cost sensitivity resulting, in class imbalancing. MapReduce was used varying a number of settings, None of the adopted methods has shown superiority out of the extensive experimental study conduced in this work. In [39], the authors reported an improvement in the accuracy of going-concern prediction by using a hybrid Random Forests and rough set theory approach. Random Forests is used for feature

¹Kaggle is a platform that hosts and runs machine learning competitions (https://www.kaggle.com/)

selection, before the rough set method generates meaningful rules. However, none of these methods prevailed to the point to replace the original technique,

despite the slightly enhanced results reported in various papers. In a recent publication it has been reported that Random Forest outperformed most of the state-of-the-art machine learning techniques in security related-application. The authors in [33] used Random Forest with weighted voting scheme along with Principle Component Analysis (PCA) to detect database access anomalies, results showed that the proposed method improved false positive and false negative rates and the overall accuracy of the classifier.

More recently, Random Forest gained more popularity in machine vision and visual classification tasks classification tasks [28], [32] and [25]. For example in [25] the authors proposed to decompose the multi-class classification problem into a binary classification problem in order to be solved by standard binary classifiers. Evaluation on visual classification related tasks showed improvement in the accuracy.

2.2. Genetic Algorithm

Genetic Algorithm (GA) is the most widely used meta-heuristic approach
for hard optimisation problems [9, <u>38</u>, <u>16</u>]. As the name suggests, GA tries to emulate the genetic evolutionary process. It starts by an initial population with each individual solution is represented by a chromosome. The chromosome is decoded in most cases as a fixed-length binary string. Each chromosome is evaluated using a fitness function designed to measure the goodness of the solution. The value of this fitness function identifies the survivors of the current population, that represent the parents for the following population. Two basic operations are usually adopted as the solution strategy to generate the new population, namely, crossover and mutation.

Crossover is applied on two chromosomes at a time from the parents. Mainly a point in the binary string is identified randomly cutting the two chromosomes, each into two pieces, and a swap between the chromosomes is applied with crossing (i.e., the first part of one chromosome is used as the second part in the other one). After crossover is applied, mutation is used to generate randomness in the solution space. It is used on one chromosome at a time flipping one of its bits. As such, the (healthy) parents generate a new population. The process is then repeated for a pre-set number of populations. Numerous variants of this process have been proposed in the literature [14].

3. Related Work

Class decomposition was first proposed as a way to reduce bias in classifiers with high bias and low variance [37]. Noting that such classifiers cannot 135 draw boundaries among complex class structures. Clustering is applied to ease such complexity. The technique is applied only on single classifier systems, namely, Naive Bayes, and Support Vector Machine (SVM). Clustering was applied with a process that allows possible merging of the generated clusters, such that boundaries can be easily drawn among the new classes. This allows high 140 bias classifiers to perform better. As such the quality of the clustering process in terms of cluster separation is not the ultimate goal for this research. However, more recently, in [31], the notion of class decomposition applied to biomedical datasets was proposed. The motivation in this case was that the intuition that genuine subclasses can be found in positive classes in binary biomedical datasets 145 (medical diagnosis). As such, the work in [31] used class separation as the main criteria for determining the value of the class decomposed clusters. Also it was assumed that genuine subclasses can only be found in positive classes. Those two arguments were debated in [18]. Applying class decomposition to only the positive class does not address the problem of false alarms, when a negative 150 example is classified as positive. Although finding clusters with a good separation can enhance the classification process, it is not always desirable as the performance relies on the adopted classifier. As it is shown in [37], high bias classifiers can perform better if the decomposed clusters are re-merged. In [18], it has also been argued that class decomposition coupled with Random Forests 155 can give the advantage of enhancing the diversification of the data set. Thus, cluster separation was deemed insignificant. Favourable results were reported for applying class decomposition on medical diagnosis datasets in [18, 31]. However, the issue of optimal setting of class decomposition and Random Forests parameters remains an open research question. This issue is addressed in this

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paper by adopting Genetic Algorithm.

Class decomposition was also reported in [23] by a means of a very fast neural network based method. Applying this technique, neurons adjust themselves with each incoming observation, allowing classification of non-linearly separable classes through incremental adjustment and addition of neurons. Although the method needs only one pass over the data set, it can be easily affected by noise and may lead to overfitting. The model itself can decompose classes to its components, but as noise is modelled, the number of subclasses is not optimised. Furthermore, decoupling of decomposition and classification processes as applied in this paper allows only genuine clusters to be detected.

Genetic Algorithm has been used in different ways for optimising Random Forests. For example, in [8], each chromosome was a Random Forests solution with a variety of trees. Applying the solution strategy described in the background section of this paper, different solutions are generated and assessed.

- ¹⁷⁵ However, the optimisation of the number of features was not addressed. Also the number of trees was not optimised directly, but a variable length chromosome was used allowing navigation in this solution space. Favourable results were reported. Another example is reported in [6], where Genetic Algorithm was used as a feature selection phase to find the optimal set of features before
 ¹⁸⁰ applying Random Forests on the reduced feature space. The proposed method was applied to a lymph disease data set. The method has proved its applicability with a clear boost in accuracy over a number of other feature selection methods.
- Applying machine learning in medical diagnosis has been widely reported in the literature. Recently, Azar et al [5] have thoroughly experimented a number of support vector machines (SVM) classifiers applied to breast cancer mammography data. It was concluded from this study that linear programming support

vector machines (LPSVM) is superior in diagnosis aid. Results of adopting decision trees and ensemble of trees on the same data set have been reported in [4].

The study concluded that Random Forests is the most accurate method when compared with single tree classifiers and ensemble of boosted trees. In this paper, it is argued that we can further boost the accuracy of Random Forests when an optimised setting of class decomposition is applied. As aforementioned, class decomposition can detect subclasses, resulting in better organisation of class separability.

4. Methods

The two most critical parameters that define the performances of Random Forests are the number of trees (*ntrees*) used in each forest, and the number of features used at each split (*mtry*). In this paper, it is aimed to optimise (ntrees, mtry) along with the parameter k which defines the number of clusters per class in the data set. The hypothesis deriving the work reported in the paper is that by decomposing the observations within each class of a particular data set, the structure of non-linearly separable data is eased, and hence the predictive accuracy of Random Forests is boosted up.

205 4.1. Class Decomposition

Decomposing the classes of a particular data set into subclasses will be achieved by means of k-means clustering algorithm. Here, clustering will be used to decompose a particular class into a set of k-subclasses. By decomposing the class into a set of subclasses (clusters), the aim is to find the within-class similarities between different instances/observations of a data set and group them accordingly. With this approach, diversity is enhanced in the data set, and thereby the classification accuracy is improved.

To illustrate the idea of diversifying a search space by decomposing class labels within a particular data set, consider the classical hand-written digit recognition set. In such a data set, a digit 8 could be written in so many different ways, which may or may not share common characteristics, hence decomposing the set of instances that are labelled as 8 into a set of clusters that share certain characteristics may certainly improve diversity and consequently improve classification accuracy. Similarly, in a medical data set with hundreds

or thousands of observations, assume that each of these observations is labelled to indicate whether a disease is present or not (i.e. 1 or 0 respectively). Further class decomposition could be applied and may lead to better representation of the data (i.e. a disease is present and mild, present and severe, etc).

Formally, consider the scenario depicted in Equation 1, where X represents a set of observations, each is defined by a set of n features, and Y is the class labels set

$$X = \begin{bmatrix} x_{11} & x_{12} & \dots, & x_{1n} \\ \dots & x_{22} & \dots, & \dots \\ \dots & \dots & \dots & \dots \\ x_{m1} & \dots & \dots, & x_{mn} \end{bmatrix}, \quad Y = \begin{bmatrix} y_1 \\ \dots \\ \dots \\ y_m \end{bmatrix}, \quad (1)$$

Now, for simplicity, lets assume that this is a binary classification that represents a medical data set and that $Y \in \{1, 0\}$, which respectively represents the presence or absence of a certain type of cancerous disease. Clearly, decomposing the set Y into Y' will result in a larger set of classes that captures more 230 variations within classes, i.e. |Y'| > |Y| (where |Y'|, |Y| represent the number of unique class labels in the sets Y', Y respectively) and hence more diverse search space. Some techniques reported in the literature have already shown some improvement in classification accuracy when applying class-decomposition to datasets, such as [18] and [31], however, one of the main unanswered questions in this respect, is which class to decompose, and the number of subclasses in the decomposed class. For example in [31] only positive classes were considered, while in [18] all classes were decomposed using a fixed number of clusters that was experimentally set, as previously discussed in the related work section. To answer the aforementioned questions, in this paper, the well-known stochastic 240 meta-heuristic algorithm, namely Genetic Algorithm (GA) is utilised.

Consider Equation 1, for any machine learning algorithm, the objective is to find a function h(x), that maps each instance in X to its label in the set Y correctly. The ultimate aim is to maximise the accuracy of h(x) by optimising a set of parameters. Among these parameters are the k values that will define the new class set (Y'),

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$$\boldsymbol{Y}^{'} = (y_{1}^{kvalue},...,y_{i}^{kvalue},...y_{m}^{kvalue},...,y_{L}^{kvalue})$$

Where L represents the number of discrete classes in the data set and y_i^{kvalue} implies that the i^{th} class in the set Y will be decomposed into kvalue subclasses and kvalue is defined as in Equation 3

$$1 \le kvalue \le max, \quad kvalue \in \mathbb{N} \tag{3}$$

(2)

Notice that *kvalue* here could take any value that ranges from 1 which means apply no decomposition (i.e. clustering) to this class, all the way up to a maxK as will be defined in the following sections. It is worth pointing out that with such an arrangement, for any classifier h(x) where x belongs to class y_i , $h(x) = y_{ij}$ is considered as a correct classification $\forall j \in y_i$ subclasses. For further illustration, lets consider a binary classification problem (i.e. X in Equation 1) and suppose that X contains 100 observation with a label set $Y \in \{a, b\}$, and suppose that we decomposed its first class label into two subclasses, and the 2^{nd} class label into 3 subclasses. In addition, let's assume that a machine learning algorithm ϕ is applied which resulted in a classifier h_c with a 100% accuracy

represented in the form of a confusion matrix as can be seen in Equation 4.

$$h_{c} = \begin{bmatrix} a_{1} & a_{2} & b_{1} & b_{2} & b_{3} \\ a_{1} & \mathbf{10} & \mathbf{5} & 0 & 0 & 0 \\ a_{2} & \mathbf{4} & \mathbf{31} & 0 & 0 & 0 \\ b_{1} & 0 & 0 & \mathbf{8} & \mathbf{6} & \mathbf{10} \\ b_{2} & 0 & 0 & \mathbf{9} & \mathbf{1} & \mathbf{16} \\ b_{3} & 0 & 0 & \mathbf{2} & \mathbf{17} & \mathbf{7} \end{bmatrix}$$
(4)

Notice that, the confusion matrix shown in Equation 4 is often used to compute the accuracy of a classifier by summing all elements at the diagonal and dividing it by the total number of observations (i.e. $\frac{\sum_{i=1}^{n} h_{ii}}{M}$). However, the accuracy of h_c denoted by $Accuracy(h_c)$ is computed in a slightly different way to account for the decomposition of the classes in the data set (Equation 5).

$$Accuracy(h) = \frac{\sum_{i=0}^{nClasses} \sum_{j=0}^{k_i} h_c(i, j + [k_{i-1} * i])}{m}$$
(5)

Where m is the number of observations (i.e. 100), and nClasses represents the number of discrete classes in the data set, while k_i represents the number of clusters applied to each class as will be discussed in the next section. In short, Equation 5 will result in summing all the **bold** elements of the confusion matrix in Equation 4.

4.2. Optimised Random Forests

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As discussed earlier the two most critical parameters that define the performance of Random Forests are the number of trees (*ntrees*) used in each forest, and the number features used at each split (*mtry*). Recall that the aim is to optimise these two parameters along with the set of clusters to be applied at each class label in a particular data set (i.e. the *kvalue*/s as formulated in Equation 2). In doing so, Genetic Algorithm is adopted to optimise these parameters.

4.2.1. Chromosome representation

For any particular data set X with a set of observations and a set of classes Y that defines these observations (see Equation 1), and assuming that Y has m unique classes, then a real-valued feature vector V that represents the set of parameters to be optimised in order to maximise the accuracy of Random Forests is as shown in Equation 6.

$$V = \begin{bmatrix} y_1^{k1} & y_2^{k2} & \dots & y_{m-1}^{kL} & y_m^{kL} & mtry & ntrees \end{bmatrix}$$
(6)

where $y_i \in Y$ and k_i represents the k value that will be set to cluster the $i^t h$ class into k subsets. It is clear that by this arrangement, we are not only deciding the optimal k value for each class, but also which class will be decomposed. For example, if k was set to be equal to 1, then this simply means that no class decomposition will be applied to this particular class.

4.2.2. Solution Population

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- Equation 6 represents the solution representation that will be used to populate the GA population. In other words, an initial random set of solutions will be generated to represent different settings in order to optimise Random Forests, this population of solutions will then be evolved over a set of generations to improve and optimise the parameters and reach *near-optimal* settings.
- Let's consider the Parkinson data set [27], which contains a set of observations about people, each of them is defined by a set of attributes (23) and labelled as either healthy or having a Parkinson Disease (PD). A typical initial populations of solution may look like the one in Table 2.

Table 1	Typical	Solutions	in a	ĠΑ	population

	No	$Healthy_{\mathbf{kvalue}}$	$PD_{\mathbf{kvalue}}$	nTrees	MTRY
	1	1	2	390	5
3 3 2 350	2	5	1	450	12
	3	3	2	350	7
		· · ·			
size	size	· ·			

As can be seen in Table 3 $Healthy_{kvalue}$ column represents the kvalues that the first class in the data set may take. Similarly, PD_{kvalue} denotes the kvaluesthat may be applied to the second class in the data set (PD). It is also clear that the solution space will depend on the total number of classes that represents the data set. It is important to stress out here that a set of constraints are applied for the values that can appear within the solution representations. These include, the range of values that kvalue can take, which was constrained as follows:

$$1 \le kvalue \le 10 \tag{7}$$

According to Equation 7 The max k value has been set to equal 10, because conducted experiments in this work in addition to previous work (i.e [18] and [31]) have shown that increasing the k value beyond 10 will not have significant impact on improving the results.

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The number of trees has been set to range between 100 and 1000 (i.e. $100 \leq ntrees \leq 1000$). It was proven experimentally that the accuracy of the Random Forests do not significantly improve when increasing the number of trees beyond 500 to 1000 trees [11] therefore we set the maximum number of trees to be 1000. At the same time we set the minimum number of trees to 100.

Finally, in Equation 8, the set of values that can be assigned to *mtry* is set as follows:

$$\lceil 0.2 \times n \rceil \le mtry \le \lceil 0.8 \times n \rceil \tag{8}$$

where n is the total number of attributes in the data set. Notice that this range will include the default settings for the Random Forests (i.e. \sqrt{n} , or $\log_2(n)$) all the way up to 80% of the total number of attributes.

4.2.3. Fitness Function

GA evolves the solutions iteratively and often starts with a randomly generated set of solutions (population), as aforementioned. Then, this population is evolved over a set of iterations, where at each of them the fitness (quality) of each solution is evaluated. The design of the fitness function is critical to the success and convergence of the GA to a good solution. In this paper, this function (Algorithm 1) is designed to compute the classification accuracy of the Random Forests.

Algorithm 1 Compute the fitness of a solution Data: Dataset, Chromosome

Result: Accuracy of the Random Forests

begin

 $\begin{array}{l} A \longleftarrow Dataset; \\ & /* \text{ Decode the chromosome solution} \\ kvalues, ntrees, mtry \longleftarrow decode(Chromosome); \\ A_c \longleftarrow decomposeSet(A, kvalues); \\ model \longleftarrow fitRF(A_c, ntrees, mtry); \\ Accuracy \longleftarrow evaluate(model) \\ return(accuracy); \end{array}$

end

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The fitness function outlined in Algorithm 1 simply decodes the chromosome solution (decode(Chromosome)) to extract the set of kvalues along with the *ntrees* and *mtry* values. Following the decoding and as outlined in Algorithm 1 class decomposition (decomposeSet(..., kvalues)) is applied to the data set according to the solution's genes kvalues, then a Random Forests model will be fit (fitRF(...)) on the new clustered data set and subject to the optimised number of trees of the forest and the number of features used at each split (ntrees and mtry).

4.3. Algorithm

To wrap up this section, and before discussing our experimental setup and results, we briefly outline the GA workflow given the above arrangements as can be seen in Algorithm 2.

$\frac{Algorithm \ 2 \ Genetic \ Algorithm}{GA \ (iterations, n, GA \ Parameters)}$

 $\begin{array}{c} \textbf{begin} \\ c \leftarrow 0 \ ; \\ i \leftarrow 0 \ ; \\ Generation_c \leftarrow \text{generate random } \textbf{n} \text{ solutions } ; \\ fitness \leftarrow computeFitness(s) \forall s \in Generation_c \ ; \\ \textbf{while } fitness \ not \ reached \ and \ i \leq iterations \ \textbf{do} \\ | \ Generation_{c+1} \leftarrow evolve(Generation_c) \ ; \\ fitness \leftarrow computeFitness(s) \forall s \in Generation_c; \\ i \leftarrow i + 1; \\ \textbf{end} \\ \textbf{return } (\text{fittest solution}) \\ \textbf{end} \end{array}$

The *evolve*(*population*) outlined in Algorithm 2 refers to the application of the GA operators on the individuals (solutions) of a particular generation. This means the selection mechanism of solutions in the current generation, and the application of crossover and mutation. The parameter settings of the GA will be discussed in the following section. Notice that given Algorithm 2, the aim is to obtain the solution that yields the most accurate classification results (fittest solution).

340 5. Experiments

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This section provides details about the different experiments that have been carried out to evaluate the proposed method. In the following sections, RF will be used to refer to the classical Random Forests model while the proposed method will be referred to by RFGA. Secondly, RFTuned will be used to refer to the method of tuning the RF parameters using Genetic Algorithm without class decomposition. Finally, Adaboost will be used to refer to the AdaBoost Ensemble classifier which was compared against the proposed method (RFGA).

The extensive experimental study reported in this section aims at establishing the following:

• Class decomposition leads to a more accurate Random Forests classifier.

- Optimising class decomposition and Random Forests parameters is a key factor to a successful class-decomposed Random Forests.
- Affirming that class decomposition coupled with Genetic Algorithm as an optimiser is the best performing classifier among possible variations of solutions (i.e., variations of enabling or disabling decomposition, and
- enabling or disabling parameter optimisation using Genetic Algorithm).
- The proposed method is superior when compared with state-of-the-art classifiers.

In order to establish the validity and stability of the proposed method, all experiments discussed below have been replicated 10 times. Details of the average classification accuracy along with standard deviation are detailed in the following sections.

5.1. Datasets

In total, 22 datasets from the UCI repository have been used in this paper [7]. As can be seen in Table 2, these sets vary in terms of number of observations (from 150 to 7200 instances), number of attributes (from 3 to 34 attribute) and number of class labels (from 2 to 7).

The sets shown in Table 2 have been selected from different domains including 14 set from the life science domain. These are mostly medical and include the
followings: Breast Cancer Wisconsin (Diagnostic) [30], Contraceptive Method Choice [7], Dermatology [7], Diabetic Retinopathy Debrecen [3], Haberman's Survival set [7], Statlog (Heart) Data Set [7], Indian Liver Patient set (ILPD) [7], Mammographic Mass [17], Parkinsons [27], Pima Indians Diabetes Data Set (PID) [7], Thoracic Surgery [41], Thyrodid [7], Seeds [12] and Iris set [7]. The
remaining sets have been selected from other categories including: Computer

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Dataset	Size	Attributes No	Classes No
Balance	625	4	3
Bank Notes	1372	5	2
Blood Transfusion	748	5	2
Breast Cancer	569	32	2
Climate Model	540	18	2
Contraceptive	1473	9	3
Dermatology	366	33	6
Diabetic Retinopathy	1151	20	2
Haberman's	306	3	2
Heart	270	13	2
Liver	583	10	2
Ionosphere	351	34	2
Iris	150	4	3
Mammographic	961	6	2
Page Blocks	5473	10	5
Parkinsons	197	23	2
Pima Indians Diabetes (P	ID) 768	8	2
Seeds	210	7	3
Statlog	2310	19	7
Thoracic	470	17	2
Thyrodid	7200	21	3
User Knowledge	403	5	4

Table 2: Details of the datasets used in the experiments

Science and Engineering (User Knowledge Modelling [24], Statlog, Page Blocks Classification and Bank Notes Authentication sets [7]), Physical and Social Science (Ionosphere Data Set [7], Climate Model Simulation Crashes [29], Balance Scale Data Set [7]) and one set from business category (Blood Transfusion Service Center [40]).

5.2. Pre-processing & Experiments Setup

The main objective of this experiment is to establish the importance of decomposing class labels in improving the performance of Random Forests. Every set used in this experiment was subject to pre-processing where appropriate, in particular handling missing values in some sets using [35] and normalisation where feature's values are standardised in the range of 0 to 1 as can be seen in Equation 9

$$z_i = \frac{x_i - \min(x)}{\max(x) - \min(x)}$$

Where x_i represents the i^{th} value of feature/attribute x in the set, and max(x), min(x) represent the maximum and minimum values in feature x, respectively. This step was necessary to suppress the sensitivity of k-means algorithms to outliers [18]. Once sets were pre-processed, each set has been split into two subsets, training and testing sets. The size of the training set is set to equal 80% of the original set and was divided into further two subsets (training and validation, with the validation set size set to be 20% of the original training set).

Figure 2 depicts the workflow of the proposed method *RFGA*. Notice that the training set has been used during the optimisation process (i.e. applying GA to optimise RF) while the validation set has been used to test the optimised RF during the training process. The testing set in turn has only been used to ⁴⁰⁰ asses the resulting model (i.e. RFGA). In other words the testing set was only used upon the conclusions of the training and optimisation processes, mainly to test the resulting optimised RF model.

Genetic Algorithm (GA) was implemented using [34]. GA settings used in this experiment are outlined in Table 3. No other settings have been used in this paper as the optimisation of GA settings is beyond the scope of this work.

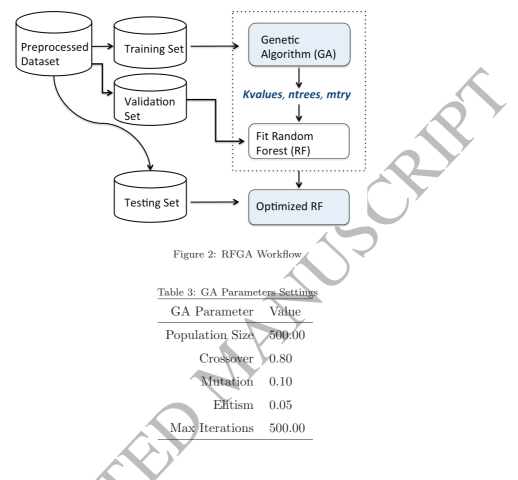
In order to asses the benefits of decomposing class labels on Random Forest performance, three different sets of experiments have been carried out on each set. Each of these experiments apply different methods and were replicated 10 times:

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• First, RF with the default settings was applied on each set,

• in the second experiment, *RFGA* was applied on the same sets,



• and finally, *RFTuned* was applied which includes disabling class decomposition (i.e. setting the *kvalue* to 1) and optimising RF parameters (*mtrees*, *mtry*) using GA.

These experimental settings are depicted in Figure 3 which shows the results of the replicated experiments across the three different methods. Notice that for RF, the default parameters were held constant and no decomposition was applied. It is also worth noting that the ten runs in case of the RF is represented by seven red dots in Figure 3 instead of ten, this is because some runs have produced the same results. In *RFGA* however, the proposed method was applied, and it can be noticed from the solution chromosomes (shown in the y - axis of the plot) that class decomposition have been applied to both classes in this case (Breast Cancer set). In the third experiment RFT uned, the optimisation was only applied to the *mtry* and *ntrees* while *kvalue* was set to equal 1 (no decomposition).

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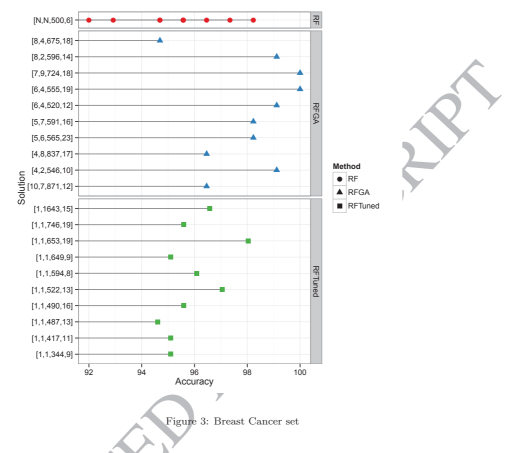
The following two sections discuss and compare the results of RFGA (the proposed method) against RF and RFT uned, where results are reported by means of average and standard deviation of the 10 replications on each set. RF_{Avg} , $RFGA_{Avg}$ and $RFTuned_{avg}$ denote the average runs of RF, RFGAand RFT uned respectively, while X_{SD} denotes the respective method standard deviation.

The experiments will be finally concluded by comparing the performance of the RFGA against a different and rival ensemble classifier. In particular, Adaboost was used for this purposed because it proves to be one of the stateof-the-art methods in achieving high predictive accuracy [21]. 435

5.3. RFGA Versus RF

Comparing the predictive accuracy of both the proposed method (RFGA)and the traditional Random Forests (RF), the results are presented in Table 4. The table reports the optimal setting of the parameters that achieved the best 440 predictive accuracy for the proposed method using Genetic Algorithm. It also reports the average and standard deviation in predictive accuracy of all the 10 runs for the traditional Random Forests and the proposed method. For a fair comparison, the average predictive accuracy is used in the discussion.

It can be shown that consistent boost in the accuracy has been achieved by the proposed method. In 18 out of the 22 datasets used in the experiment, the proposed method outperformed the traditional Random Forests. It can also be shown that class decomposition has been applied to the majority of classes in all datasets. In fact, all datasets have had at least one class decomposed to its subclasses. We have also measured the statistical significance of the results using



the paired t-test technique. With 95% confidence, the p-value for paired t-test is 0.003331, showing clear statistical significance. Accounting for the possibility of the results not following the normal distribution, we also computed the Wilcoxon Signed-Rank test adopting the 95% confidence. The p-value for this test is 0.001455. This also confirmed the statistical significance of the achieved results. As the proposed method is composed of a number components including Genetic Algorithm and class decomposition over Random Forests, it is important to establish whether only Genetic Algorithm has the main effect, or in fact, coupling class decomposition with Genetic Algorithm is the optimal solution.
460 This is tested in the following subsection of this experimental study.

Dataset	K_{values}	ntrees	mtry	RF_{Avg}	$RFGA_{Avg}$	RF_{SD}	RFGA _{SD}
Balance	[2,3,1,1]	520	1	85.25	86.01	1.71	1.49
BankNotes	[6, 6]	608	2	99.27	100.00	0.33	0.00
Blood Transfusion	[2,6]	549	1	74.16	77.69	1.87	1.51
Breast Cancer	[6,4]	555	19	95.31	98.14	1.88	1.74
Contrace ptive	[2,4,5]	492	3	53.21	52.05	1.44	1.95
Dermatology	$[2,\!2,\!2,\!2,\!2,\!1]$	768	11	97.66	98.36	0.74	1.13
Diabetic Retinopathy	[3,2]	619	9	67.39	68.26	1.31	1.80
Haberman	[3,5]	645	1	72.78	72.63	3.29	3.02
Heart	[7,3]	768	4	80.08	82.83	4.18	4.74
Ionosphere	[1,8]	841	14	92.86	93.73	1.35	1.42
IRIS	[4,7,2]	792	1	95.19	97.04	1.56	1.56
Liver	[3,2]	679	6	72.76	74.66	2.76	3.71
Mammographic	[3,5]	817	2	83.07	83.39	2.04	2.36
Page Blocks	$[5,\!1,\!5,\!2,\!2]$	696	7	97.34	97.34	0.27	0.30
PARKINSONS	[3,5]	700	12	91.32	96.32	4.12	3.55
PID	[7,5]	556	3	75.75	74.05	2.29	1.82
Seeds	[2,4,3]	616	3	91.07	94.13	3.88	2.28
Simulated Climate	[1,7]	776	7	91.71	92.40	0.00	0.97
Statlog	[9, 6, 3, 6, 6, 7, 7]	881	9	97.88	97.93	0.48	0.29
Thoratic	[[5,4]	664	6	84.61	84.67	0.56	1.49
Thyroid	[3,3,3]	651	9	99.55	99.56	0.12	0.15
User knowledge	[3,2,1,1]	746	3	92.44	94.00	2.21	2.68

Table 4: Experiments Results

5.4. RFGA Versus RFTuned

In this set of experiments, the class decomposition is disabled allowing Genetic Algorithm to tune only RF's two main parameters, namely, the number of features to split on at each node and the number of trees. This method is compared with the proposed method of applying class decomposition with Genetic Algorithm used to optimise all the parameters (i.e., RF parameters, and number of subclasses in each class). Table 5 reports the average of 10 runs for both methods. In 12 out of the 22 datasets, the proposed method outperformed

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the alternative one. Using the paired t-test with 95% confidence, the *p*-value is 0.9328, and for the Wilcoxon signed rank test (also 95% confidence), the p-470 value is 0.8736, the results are not statistically significant. However, the results suggest that it is recommended to run the optimised RF without class decomposition as the first step before decomposing the classes in the data set. Then the results can be compared. This can then lead to the best possible predictive accuracy. This suggested procedure aims at distilling the cases when optimising the Random Forests parameters can yield the best performance. Collectively both methods were the best performer among all the variations. As such, the practice of running both and select the best outcome has the potential of producing the strongest classifier in this family of methods. As the results show, over 3% accuracy boost can be achieved when applying class decomposition (e.g., the Parkinsons set). In life science related applications, this can be an important achievement, especially those related to medical diagnosis as reported in the Parkinsons set when the optimal setting suggested a class decomposition of both the positive and the negative classes of 3 and 5 respectively.

The results reported so far assert the positive impact of class decomposition on predictive accuracy of Random Forests. To establish the superiority of the proposed method over state-of-the-art ensemble methods, represented by AdaBoost, the following subsection discusses this comparative experimental study.

5.5. RFGA Versus AdaBoost

AdaBoost is an ensemble learning method that uses boosting of classifiers, having each classifier modelled to focus on examples misclassified by previously constructed classifiers in the sequence [21]. It is among the state-of-the-art methods in achieving a high predictive accuracy. To validate the proposed method in this paper, a comparison between the two methods is conducted.
⁴⁹⁵ Using the average of 10 runs for both methods the results are reported in Table
6. The results clearly suggest the superiority of the proposed method over AdaBoost. In 17 out of the 22 datasets, the proposed method outperformed AdaBoost. With 95% confidence, the *p*-value for both the paired t-test and the

Dataset	$RFTuned_{avg}$	$RFGA_{avg}$	$RFTuned_{SD}$	$RFGA_{SD}$
Balance	89.24	86.01	0.87	1.49
Bank Notes	99.75	100.00	0.46	0.00
Blood Transfusion	76.80	77.69	1.61	1.51
Breast Cancer	95.88	98.14	1.06	1.74
Contraceptive	54.34	52.05	1.48	1.95
Dermatology	97.89	98.36	2.13	1.13
Diabetic Retinopathy	69.35	68.26	2.22	1.80
Haberman	72.82	72.63	2.76	3.02
Heart	83.96	82.83	3.58	4.74
Ionosphere	93.33	93.73	3.02	1.42
IRIS	95.56	97.04	1.56	1.56
Liver	72.11	74.66	2.89	3.71
Mammographic	82.19	83.39	3.43	2.36
Page Blocks	97.31	97.34	0.00	0.30
PARKINSONS	92.89	96.32	3.93	3.55
PID	76.96	74.05	1.59	1.82
Seeds	93.33	94.13	3.13	2.28
Climate Model	93.83	92.40	1.60	0.97
Statlog	97.65	97.93	0.00	0.29
Thoratic	85.21	84.67	1.62	1.49
Thyroid	99.64	99.56	0.13	0.15
User Knowledge	94.49	94.00	2.16	2.68

Table 5: Tuned RF versus RFGA Performance

Wilcoxon signed rank test are 0.07397 and 0.03289, respectively. This shows
satisfactory statistical significance of the results. The AdaBoost classifier used in this experiment was set to iterate 100 times generating the same number of trees, avoiding overfitting when a large number of trees are generated.

5.6. Results Discussion

Concluding this experimental study, after analysing all the results of the three comparisons between the proposed method, and its three identified competitors, it is found that only the proposed method is able to consistently out-

Dataset	$AdaBoost_{avg}$	$RFGA_{avg}$	$AdaBoost_{SD}$	$RFGA_{SD}$
Balance	84.40	86.01	2.45	1.49
Bank Notes	99.68	100.00	0.33	0.00
Blood Transfusion	74.02	77.69	2.50	1.51
Breast Cancer	96.38	98.14	0.86	1.74
Contraceptive	56.00	52.05	1.90	1.95
Dermatology	96.41	98.36	1.54	1.13
Diabetic Retinopathy	67.27	68.26	1.48	1.80
Haberman	66.54	72.63	3.21	3.02
Heart	80.71	82.83	3.34	4.74
Ionosphere	93.65	93.73	1.99	1.42
IRIS	94.98	97.04	2.15	1.56
Liver	70.81	74.66	3.09	3.71
Mammographic	79.69	83.39	2.04	2.36
Page Blocks	97.29	97.34	0.25	0.30
PARKINSONS	92.63	96.32	2.41	3.55
PID	73.66	74.05	1.85	1.82
Seeds	93.99	94.13	3.08	2.28
Climate Model	94.42	92.40	1.28	0.97
Statlog	98.11	97.93	0.43	0.29
Thoratic	81.90	84.67	1.26	1.49
Thyroid	99.66	99.56	0.07	0.15
User Knowledge	99.66	94.00	1.94	2.68

Table 6: Adaboost versus RFGA Performance

perform the other methods. As shown in Table 7, in 11 datasets, the proposed method achieved the highest predictive accuracy. It is worth noting that the next best method found to be the optimised Random Forests (*RFTuned*) with a superior performance in only 4 datasets. All the results for the three variations of Random Forests and AdaBoost experimented in this project are summarised in Figures 4 and 5 by categorising the datasets to life science datasets, and non-life science datasets. It is clear that the proposed method is superior than its competitors in both categories. Our method has shown particular high predictive accuracy over medical diagnosis datasets like Parkinsons and Liver.

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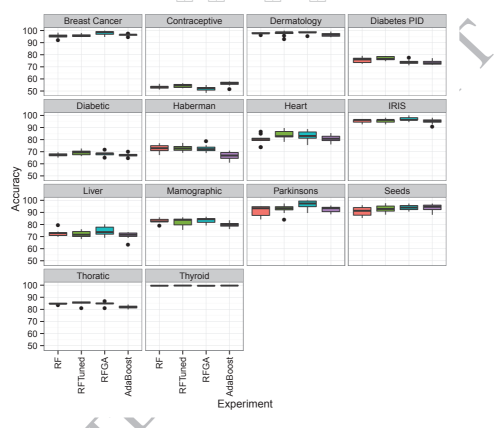
This can be attributed to the complexity of the problem, and that indeed these datasets can be naturally decomposed to its subclasses, that in turn facilitates classification using Random Forests.

D. I. I				
Dataset	$RFGA_{avg}$	RF_{avg}	$RFTuned_{avg}$	$AdaBoost_{avg}$
Bank Notes	100.00	99.27	99.75	99.68
Blood Transfusion	77.69	74.16	76.80	74.02
Breast Cancer	98.14	95.31	95.88	96.38
Dermatology	98.36	97.66	97.89	96.41
Ionosphere	93.73	92.86	93.33	93.65
IRIS	97.04	95.19	95.56	94.98
Liver	74.66	72.76	72.11	70.81
Mammographic	83.39	83.07	82.19	79.69
Page Blocks	97.34	97.34	97.31	97.29
PARKINSONS	96.32	91.32	92.89	92.63
Seeds	94.13	91.07	93.33	93.99

5.7. Implementation

A framework was implemented using **R** where several packages have been utilised. These include amongst other libraries: *randomForest* package [26] which implements Brieman and Cutler Random Forests for Classification and Regression, and the GA package [34] which allows parallel implementation of the Genetic Algorithm. Table 3 shows the parameters settings that have been used for this experiment. AdaBoost package [1] which has been used to build the AdaBoost ensemble. For handling missing values [35] and [36] were used to impute missing values.

The framework was designed to make use of the multicore facilities by utilising R packages that enable parallel execution of the code (i.e. [2]). It is worth noting that the proposed method is scalable, as individual chromosomes (Random Forests solutions) in each generation is constructed in parallel. Furthermore, each tree in any single Random Forests grows in parallel with all



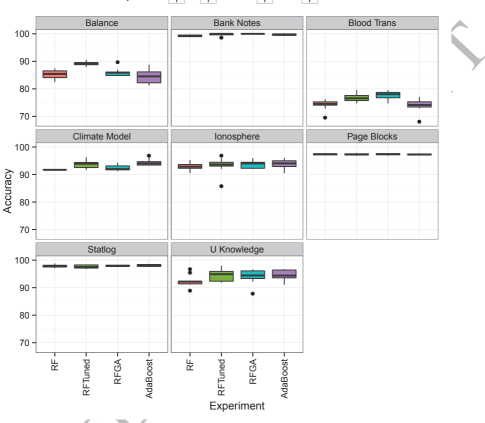
Experiment 🛑 RF 🛑 RFTuned 🚔 RFGA 🚔 AdaBoost

Figure 4: Life science Datasets Results

other trees. Consequently, only the number of iterations of the Genetic Algorithm is the main factor in the time needed to find the final solution. This is the case with all evolutionary optimisation methods, that are built in a sequence of generations.

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Dell Poweredge R730 running 56 Intel Xeon Processors E5-2695 v3 @ 2.30Ghz with 255Gb RAM connected to a tier 3 4Tb datastore (SATA Disk) was used to run the experiments reported in this paper. TThese experiments were carried



Experiment 🚔 RF 🚔 RFTuned 🚔 RFGA 🚔 AdaBoost

Figure 5: Non-Life science Datasets Results

540 out on 24 core VMWare Virtual Server with 48GB of RAM.

6. Conclusion and Future Work

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The paper proposed a three-component system for enhancing the classification accuracy in Random Forests. The first component is the class decomposition where clustering is applied to examples that belong to each class, resulting in a number of clustering sessions which is equal to the number of classes in the

data set. Setting the number of clusters for each class has its own effect on the predictive accuracy. Random Forests which is a highly accurate classification method is the second component of the proposed system. It requires two main parameters to be set: (1) number of trees in the ensemble, and (2) the number of features sampled randomly at each node split of each tree. Collectively the 550 number of parameters to set is equal to number of classes in the data set, in addition to the two Random Forests parameters. Realising the large search space generated from setting all these parameters which is exponential in the number of classes in a a data set, there is a clear need for an effective optimisation method. Thus, Genetic Algorithm is used as our third component. The system 555 was applied to 22 datasets predominantly in the area of life sciences, and the results proved the effectiveness of the proposed hybrid machine learning technique in enhancing the predictive accuracy. We can identify a number of future directions for this research as follows.

⁵⁶⁰ Experimenting the hybrid method to other application domains in life sciences such as gene expression datasets is one direction. The optimisation of GA parameters is another direction which may lead to further improvements of the RF performance. Also the adoption of other population-based meta-heuristic methods can be used to compare the effectiveness of a number of optimisation techniques . Finally, the use of other high performing machine learning algorithms like Gradient Boosting trees, or Support Vector Machines (SVM) can be explored instead of Random Forests.

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