Ensemble missing data techniques for software effort prediction

Bhekisipho Twala\textsuperscript{a,*} and Michelle Cartwright\textsuperscript{b}
\textsuperscript{a}Department of Electrical and Electronic Engineering Science, University of Johannesburg, P.O. Box 524, Auckland Park, Johannesburg 2006, South Africa
\textsuperscript{b}Brunel Software Engineering Research Centre, School of Information Systems, Computing and Mathematics, Brunel University, Uxbridge, UK

Abstract. Constructing an accurate effort prediction model is a challenge in software engineering. The development and validation of models that are used for prediction tasks require good quality data. Unfortunately, software engineering datasets tend to suffer from the incompleteness which could result to inaccurate decision making and project management and implementation. Recently, the use of machine learning algorithms has proven to be of great practical value in solving a variety of software engineering problems including software prediction, including the use of ensemble (combining) classifiers. Research indicates that ensemble individual classifiers lead to a significant improvement in classification performance by having them vote for the most popular class. This paper proposes a method for improving software effort prediction accuracy produced by a decision tree learning algorithm and by generating the ensemble using two imputation methods as elements. Benchmarking results on ten industrial datasets show that the proposed ensemble strategy has the potential to improve prediction accuracy compared to an individual imputation method, especially if multiple imputation is a component of the ensemble.

Keywords: Machine learning, supervised learning, decision tree, software prediction, incomplete data, imputation, missing data techniques, ensemble

Abbreviations

- CART: Classification and Regression Trees
- CCCS: Command, Control and Communications System
- CMI: Class Mean Imputation
- DA: Data Augmentation
- DSI: Defence Security Institute
- DT: Decision Tree
- DTSI: Decision Tree Single Imputation
- EM: Expectation-Maximization
- EMSI: Expectation-maximization Single Imputation
- FC: Fractioning of Cases
- FIML: Full Information Maximum Likelihood
- HDSI: Hot-Deck Single Imputation
- ISBSG: International Software Benchmarking Standards Group

\textsuperscript{*}Corresponding author: Bhekisipho Twala, Department of Electrical and Electronic Engineering Science, University of Johannesburg, P.O. Box 524, Auckland Park, Johannesburg 2006, South Africa. Tel.: +27 11 559 4404; Fax: +27 11 559 2357; E-mail: btwala@uj.ac.za.
1. Introduction

Software engineering (SE) researchers and practitioners remain concerned with prediction accuracy when building prediction systems. Lack of adequate tools to evaluate and estimate the cost for a software project is one of the main challenges in SE. We use datasets of past projects to build and validate estimation or prediction systems of software development effort, for example, which allows us to make management decisions, such as resource allocation. Or we may use datasets of measurements describing software systems to validate metrics predicting quality attributes.

Various ML techniques have been used in SE to predict software cost [5], software (project) development effort [6,26,61], software quality [17], and software defects [18,45]. Reviews of the use of ML in SE [37,38] report that ML in SE is a mature technique based on widely-available tools using well understood algorithms. The decision tree (DT) classifier is an example of a ML algorithm that can be used for predicting continuous attributes (regression) or categorical attributes (classification). Thus, software prediction can be cast as a supervised learning (SL) problem, i.e., the process of learning to separate samples from different classes by finding common features between samples of known classes. An important advantage of ML over statistical analysis as a modelling technique lies in the fact that the interpretation of, say, decision rules is more straightforward and intelligible to human beings than, say, principal component analysis (a statistical tool for finding patterns in data of high dimension). In recent years, there has been an explosion of papers in the ML and statistics communities discussing how to combine models or model predictions.

Most techniques for predicting attributes of a software system or project data require past data from which models will be constructed and validated. An important and common issue faced by researchers who use industrial and research datasets is incompleteness of data. Even if part of a well thought out measurement programme, industrial datasets can be incomplete, for a number of reasons. These include inaccurate or non reporting of information (without a direct benefit, a project manager or developer might see data collection as an overhead they can ill afford, for example), or, where data from a number
of different types of projects or from a number of companies are combined, certain fields may be blank because they are not collectable for all projects. Often data is collected either with no specific purpose in mind (i.e. it is collected because it might be useful in future) or the analysis being carried out has a different goal than that for which the data was originally collected. In research datasets, e.g. experiments on human subjects to assess the effectiveness of a new SE technique, say, dropout or failure to follow instructions may lead to missing values. The relevance of this issue is strictly proportional to the dimensionality of the collected data. SE researchers have become increasingly aware of the problems and biases which can be caused by missing or incomplete data. Moreover, many SE datasets tend to be small with many different attributes – software project datasets grow slowly, for example – and the numbers of available human subjects limit the size of many experimental datasets. Thus, we can ill afford to reduce our sample size further.

Many works in both the ML and statistical pattern recognition communities have shown that combining (ensemble) individual classifiers is an effective technique for improving classification accuracy. An ensemble is generated by training multiple learners for the same task and then combining their predictions. There are different ways in which ensembles can be generated, and the resulting output combined to classify new instances. The popular approaches to creating ensembles include changing the instances used for training through techniques such as bagging [1,4], boosting [19], changing the features used in training [21], and introducing randomness in the classifier itself [13]. The interpretability of classifiers can produce useful information for experts responsible for making reliable classification decisions, making DTs an attractive scheme.

There are various reasons why DTs and not other SL algorithms were utilized to investigate the problem considered in this paper. Despite being one of the well known algorithms from the ML and statistical pattern recognition communities, DTs are non-parametric and are relatively fast to construct; classification is very fast too. They work for almost all classification problems and can achieve good performance on many tasks. However, one property that sets DTs apart from all other ML methods is their invariance to monotone transformations of the predictor variables. For example, replacing any subset of the predictor variables \( \{x_j\} \) by (possible different) arbitrary strictly monotone functions of them \( \{x_j \leftarrow m_j(x_j)\} \), gives rise to the same tree model. Thus, there is no issue of having to experiment with different possible transformations \( m_j(x_j) \) for each individual predictor \( x_j \) to try to find the best ones. This invariance provides immunity to the presence of extreme values (“outliers”) in the predictor variable space. In addition, DTs incorporate a pruning scheme that partially addresses the outlier (noise) removal problem. DTs also make good candidates for combining because they are structurally unstable classifiers and produce diversity in classifier decision boundaries. DTs have also been one of the tools of choice for building classification models in the SE field [29,30,46–48,58,64–66,70–72]. We assume that the readers are familiar with DT learning [4,49]. However, a brief introduction on DTs is given in Section 3.

To the best of our knowledge, our research group was the first to present the effect of ensemble missing data techniques (MDTs) on predictive accuracy in the SE field [70,72]. In this paper, we expand on the preliminary work by first assessing the robustness of eight MDTs on the predictive accuracy of resulting DTs. We then propose three ensemble methods with each ensemble having two MDTs as elements. The proposed method utilizes probability patterns of classification results.

The following section briefly discusses missing data patterns and mechanisms that lead to the introduction of missing values in datasets. Section 3 gives a brief introduction on DTs, followed by a presentation of techniques that have been used to handle incomplete data when using DTs. The proposed ensemble procedure is also described. Section 5 reviews some related work to the problem of incomplete data in
the SE area. Section 6 empirically explores the robustness and accuracy of eight MDTs to ten industrial datasets. This section also presents empirical results from the application of the proposed ensemble procedure with current MDTs. We close with conclusions and directions for future research.

2. Patterns and mechanisms of missing data

The two other crucial tasks when dealing with incomplete data are to investigate the pattern of missing data and the mechanism underlying the missingness. This is to get an idea of the process that could have generated the missing data, and to produce sound estimates of the parameters of interest, despite the fact that the data are incomplete.

The pattern simply defines which values in the data set are observed and which are missing. The three most common patterns of non-response in data are univariate, monotonic and arbitrary. When missing values are confined to a single variable we have a univariate pattern; monotonic pattern occurs if a subject, say $Y_j$, is missing then the other variables, say $Y_{j+1}, \ldots Y_p$, are missing as well or when the data matrix can be divided into observed and missing parts with a “staircase” line dividing them; arbitrary patterns occur when any set of variables may be missing for any unit.

The law generating the missing values seems to be the most important task since it facilitates how the missing values could be estimated more efficiently. If data are missing completely at random (MCAR) or missing at random (MAR), we say that missingness is ignorable. In other words, the analyst can ignore the missing data mechanism provided the values are MCAR or MAR. For example, suppose that you are modelling software defects as a function of development time. If missingness is not related to the missing values of defect rate itself and also not related on the values of development time, such data is considered to be MCAR. For example, there may be no particular reason why some project managers told you their defect rates and others did not.

Secondly, software defects may not be identified or detected due to a given specific development time. Such data are considered to be MAR. MAR essentially says that the cause of missing data (software defects) may be dependent on the observed data (development time) but must be independent of the missing value that would have been observed. It is a less restrictive model than MCAR, which says that the missing data cannot be dependent on either the observed or the missing data. MAR is also a more realistic assumption for data to meet, but not always tenable. The more relevant and related attributes one can include in statistical models, the more likely it is that the MAR assumption will be met.

Finally, for data that is informatively missing (IM) or not missing at random (NMAR) then the mechanism is not only non-random and not predictable from the other variables in the dataset but cannot be ignored, i.e., we have non ignorable missingness [36,55]. In contrast to the MAR condition outlined above, IM arise when the probability that defect rate is missing depends on the unobserved value of defect rate itself. For example, software project managers may be less likely to reveal projects with high defect rates. Since the pattern of IM data is not random, it is not amenable to common MDTs and there are no statistical means to alleviate the problem.

MCAR is the most restrictive of the three conditions and in practice it is usually difficult to meet the MCAR assumption. Generally you can test whether MCAR conditions can be met by comparing the distribution of the observed data between the respondents and non-respondents. In other words, data can provide evidence against MCAR. However, data cannot generally distinguish between MAR and IM without distributional assumptions, unless the mechanisms is well understood. For example, right censoring (or suspensions) is IM but is in some sense known. An item or unit, which is removed from a reliability test prior to failure or a unit which is in the field and is still operating at the time the reliability of these units is to be determined is called a suspended item or right censored instance.
Table 1

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Possible values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adjusted function points (ADJFPs)</td>
<td>continuous</td>
</tr>
<tr>
<td>Duration in months (DUR)</td>
<td>continuous</td>
</tr>
<tr>
<td>Experience of project manager in years (EXPPM)</td>
<td>continuous</td>
</tr>
<tr>
<td>Number of transactions (TRANS)</td>
<td>continuous</td>
</tr>
<tr>
<td>Skill level of development team (SKILL)</td>
<td>beginner, intermediate, advanced</td>
</tr>
</tbody>
</table>

3. Decision trees

DT induction is one of the simplest and yet most successful forms of SL algorithm. It has been extensively pursued and studied in many areas such as statistics [4] and machine learning [49–51] for the purposes of classification and prediction.

DT classifiers have four major objectives. According to Safavian and Landgrebe [53], these are: 1) to classify correctly as much of the training sample as possible; 2) generalise beyond the training sample so that unseen samples could be classified with as high accuracy as possible; 3) be easy to update as more training samples become available (i.e., be incremental; 4) and have as simple a structure as possible. Objective 1) is actually highly debatable and to some extent conflicts with objective 2). Also, not all tree classifiers are concerned with objective 3).

DTs are non-parametric (no assumptions about the data are made) and a useful means of representing the logic embodied in software routines. A DT takes as input a case or example described by a set of attribute values, and outputs a Boolean or multi-valued “decision”. For the purpose of this paper, we shall stick to the Boolean case (thus, binary DTs) for several reasons given below.

The power of the Boolean approach comes from its ability to split the instance space into subspaces where each subspace is fitted with different models. Other distinctive advantages of Boolean expressions are their time complexity and space complexity, i.e., as the input size as Boolean functions grow linearly, computer time and storage requirements of algorithms to represent and manipulate DTs do not necessarily grow exponentially (as is the case for traditional representation techniques). Thus, both the memory space to store DT and the time of the learning and using phase are decreased [41]. Furthermore, information-based attribute selection measures used in the induction of DTs are all biased in favour of attributes that have many possible values [74].

The use of Boolean attributes is one strategy for handling multi-valued attributes and as a result overcomes this problem, i.e., attributes with fewer levels and those with many levels are all selected on a competitive basis.

Consider the following fictitious SE example, where the categorical attribute specifies whether a project is likely to require low, medium or high software effort (EFFORT) based on attributes related to software development. The attributes are summarized in Table 1.

The training data is used to build the model or classifier (a DT) and a corresponding hierarchical partitioning shown in Figs 1 (a) and 1 (b), respectively. The DT has classes low, medium and high. This is a classifier in the form of a tree, a structure that is either: a leaf (rectangular box), indicating a class, or a decision node (ovals) that specifies some test to be carried out on a single attribute value, with one branch and sub-tree for each possible outcome of the test. Notice that Test 1, Test 2 and Test 3 are tests on the real-valued attributes. Only test 4 uses the nominal attribute. Further note that the attribute DUR has been not utilised in the DT even though we have it as one of the continuous attributes in the training data. It is also easy to see that as the depth of the tree increases, the resulting partitioning becomes more and more complex.
The prediction problem is handled by partitioning classifiers. These classifiers split the space of possible observations into partitions. For example, when a project manager needs to make a decision whether a project is likely to require low, medium or high effort based on several software development factors, a classification tree can help identify which factors to consider and how each factor has historically been associated with different outcomes of the decision.

4. Decision trees and missing data

Several methods have been proposed in the literature to treat missing attribute values (incomplete data) when using DTs. Missing values can cause problems at two points when using DTs: 1) when deciding on a splitting point (when growing the tree), and 2) when deciding into which daughter node each instance goes (when classifying an unknown instance). Methods for taking advantage of unlabelled classes can
Specific DT techniques for handling incomplete data are now going to be discussed. These missing data techniques (MDTs) are some of the widely recognized and they are divided into three categories: ignoring and discarding data, imputation and SL or ML. The MDTs are briefly described in the following sections and summarized in Table 2.

4.1. Ignoring and discarding data (Listwise deletion)

Incomplete data are often dealt with by using several general approaches, like, deleting the instances with missing data which aim to modify up the data so that they can be analysed by methods designed for complete data. Basically, listwise deletion (LD) means that any individual with missing data on any variable is deleted from the analysis under consideration. This approach can drastically reduce the sample size since it can sacrifice a large amount of data leading to a severe lack of statistical power [36]. It can even lead to complete case loss if many variables are involved. However, due to its simplicity and ease of use, LD is the default in most statistical packages.

4.2. Imputation techniques

Imputation techniques fill in (impute) missing items with plausible values in the dataset, thus making it possible to analyze the data as if it were complete. Two kinds of imputation strategies are discussed in this paper: single imputation (SI) and multiple imputation (MI). SI refers to filling in a missing value with a single replacement value while multiple values are simulated for MI. A brief description of each imputation method is given below.

4.2.1. Single imputation

4.2.1.1. Mean or mode imputation

Some researchers have used arbitrary methods like mean imputation for addressing the missing value problem, i.e., replacing the missing values of a variable by the mean of its observed values. Mean substitution also assumes a MCAR mechanism. The strength of mean imputation is that it preserves the data and it is easy to use. However, mean imputation can be misleading because it produces biased
and inconsistent estimates of both coefficients and standard errors). Little and Rubin [36] point out that variance parameter estimates under mean imputation are generally negatively biased. Also, substitution of the simple (grand) mean will reduce the variance of the variable and its correlation with other variables. Somewhat better is substitution of the group or global mean (or mode in the case of nominal data) for a grouping variable known to correlate as highly as possible with the variable which has missing values. We shall now call this technique mean or mode single imputation (MMSI).

4.2.1.2. k-nearest neighbour imputation

We use the k-nearest neighbour (k-NN) approach to determine the imputed data, where nearest is usually defined in terms of a distance function based on the auxiliary variable(s). In this method a pool of complete instances is found for each incomplete instance and the imputed values for each missing cell in each recipient is calculated from the values of the respective field in complete instances [10]. The strength of k-NN lies in its ability to predict both discrete attributes (the most frequent value among the k nearest neighbours) and continuous attributes (the mean among the k nearest neighbours). Also, k-NN can be easily adapted to work with any attribute as class, by just modifying which attributes will be considered in the distance metric. This single imputation approach, which also assumes a MCAR mechanisms, shall now be called k−NNSI.

4.2.1.3. Decision tree imputation

The unordered attribute DTs approach, which can also be considered a machine learning technique as it uses a DT algorithm to impute missing values, is another strategy that has been used for handling missing values in tree learning. This technique was suggested by Shapiro [60]. The method builds DTs to determine the missing values of each attribute, and then fills the missing values of each attribute by using its corresponding tree. Separate trees are built using a reduced training set for each attribute, i.e., restricting your analysis to only those instances that have known values. The original class is treated as another attribute, while the value of the attribute becomes the “class” to be determined. The attributes used to grow the respective trees are unordered. These trees are then used to determine the unknown values of that particular attribute.

For this strategy, it is not clear what the probability generating the missingness is for both strategies. However due to the fact that the dependencies between attributes is not considered, we shall assume that the data is MCAR.

4.2.2. Missingness incorporated in attributes

This approach proposed by Twala [68] and followed up by Twala et al. [69] is closely related to the technique of treating “missing” as a category in its own right, generalizing it for use with continuous as well as categorical variables as described below:

Let \( X_n \) be an ordered or numeric attribute variable with unknown attribute values, the proposed approach searches essentially over all possible values of \( x_n \) for binary splits of the following form:

1. Split A: \((X_n \leq x_n \text{ or } X_n = \text{missing}) \) versus \((X_n > x_n)\)
2. Split B: \((X_n \leq x_n) \) versus \((X_n > x_n \text{ or } X_n = \text{missing})\)
3. Split C: \((X_n = \text{missing}) \) versus \((X_n = \text{not missing})\)

The idea is to find the best split from the candidate set of splits given above, with the goodness of split measured by how much it decreases the impurity of the sub-samples.

If \( X_n \) is a nominal attribute variable (i.e., a variable that takes values in an unordered set), the search is over all splits of the form:
1. Split A: \((X_n \in Y_n \text{ or } X_n = \text{missing})\) versus \((X_n \notin Y_n)\)
2. Split B: \((X_n \in Y_n)\) versus \((X_n \notin Y_n \text{ or } X_n = \text{missing})\)
3. Split C: \((X_n = \text{missing})\) versus \((X_n = \text{not missing})\)

where \(Y_n\) is the splitting subset at node \(n\).

So, if there were \(\partial\) options for splitting a branch without missingness, there are \(2\partial + 1\) options to be explored with missingness present.

This missingness incorporated in attributes (MIA) algorithm is very simple, natural and applicable to any method of constructing DTs, regardless of that method’s detailed splitting/stopping/pruning rules. It has a very close antecedent: the approach of handling unknown attribute values by treating all attributes as categorical and adding missingness as a further category. The two approaches are superficially the same but differ a little in their treatment of continuous attributes: rather than categorizing continuous variables, missingness is directly incorporated in splits of continuous variables. The approach can be expected to be particularly useful when missingness is not random but informative. Classifying a new individual whose value of a branching attribute is missing is immediately provided, especially if the attribute in the training set that was used to construct the DT had missing values.

4.2.3. Multiple imputation

MI is one of the most attractive methods for general purpose handling of missing data in multivariate analysis. Rubin [52] described MI as a three-step process. First, sets of plausible values for missing instances are created using an appropriate model that reflects the uncertainty due to the missing data. Each of these sets of plausible values can be used to “fill-in” the missing values and create a “completed” dataset. Second, each of these datasets can be analyzed using complete-data methods. Finally, the results are combined. For example, replacing each missing value with a set of five plausible values or imputations would result to building five DTs, and the predictions of the five trees would be averaged into a single tree, i.e., the average tree is obtained by MI.

MI retains most of the advantages of single imputation and rectifies its major disadvantages as already discussed. There are various ways to generate imputations. Schafer [55] has written a set of general purpose programs for MI of continuous multivariate data (NORM), multivariate categorical data (CAT), mixed categorical and continuous (MIX), and multivariate panel or clustered data (PNA). These programs were initially created as functions operating within the statistical languages S and SPLUS [73]. NORM includes and EM algorithm for maximum likelihood estimation of means, variance and covariances. NORM also adds regression-prediction variability by a procedure known as data augmentation [55].

Although not absolutely necessary, it is almost always a good idea to run the expectation maximization (EM) algorithm [12] before attempting to generate MIs. The parameter estimates from EM provide convenient starting values for data augmentation (DA). Moreover, the convergence behaviour of EM provides useful information on the likely convergence behaviour of DA. In brief, EM is an iterative procedure where a complete dataset is created by filling-in (imputing) one or more plausible values for the missing data by repeating the following steps: 1.) In the E-step, one reads in the data, one instance at a time. As each case is read in, one adds to the calculation of the sufficient statistics (sums, sums of squares, sums of cross products). If missing values are available for the instance, they contribute to these sums directly. If a variable is missing for the instance, then the best guess is used in place of the missing value. 2.) In the M-step, once all the sums have been collected, the covariance matrix can simply be calculated. This two step process continues until the change in covariance matrix from one iteration to the next becomes trivially small. This is the approach we follow in this paper.

Details of the EM algorithm for covariance matrices are given in [12,36,55]. EM requires that data are MAR.
4.2.4. Supervised learning imputation

SL techniques are those that deal with missing values using ML algorithms. ML algorithms learn a function from training data. The training data consist of pairs of input objects (vectors) and desired output. The output of the function can be a continuous value (called regression) or can predict a class label of the input object (called classification). ML techniques are generally more complex than statistical techniques. This section will describe missing data imputation using two supervised machine learning systems: surrogate variable splitting and fractioning of cases.

4.2.4.1. Surrogate variable splitting

One sophisticated but refined method worthy of note and study is the surrogate variable splitting (SVS), which has been used for the classification and regression trees (CART) system [4]. CART handles missing values in the database by substituting "surrogate splitters". Surrogate splitters are predictor variables that are not as good at splitting a group as the primary splitter but which yield similar splitting results; they mimic the splits produced by the primary splitter; the second does second best, and so on. The surrogate splitter contains information that is typically similar to that which would be found in the primary splitter. The surrogates are used for tree nodes when there are values missing. The surrogate splitter contains information that is typically similar to what would be found in the primary splitter. Both values for the dependent variable (response) and at least one of the independent variables (attributes) take part in the modelling. The surrogate variable used is the one that has the highest correlation with the original attribute (observed variable most similar to the missing variable or a variable other than the optimal one that best predicts the optimal split). The surrogates are ranked. Any observation missing on the split variable is then classified using the first surrogate variable, or if missing that, the second is used, and so on. The CART system only handles missing values in the testing case but RPART handles them on both the training and testing cases.

SVS makes no assumption about the law generating the missingness. However, due to the fact that it uses the dependencies between attributes, we shall assume that the data is MAR.

4.2.4.2. Fractioning of cases

Quinlan [49] borrows the probabilistic complex approach by Cestnik et al. [8] by “fractioning” cases or instances based on a priori probability of each value determined from the cases at that node that have specified values. Quinlan starts by penalising the information gain measure by the proportion of unknown cases and then splits these cases to both subnodes of the tree.

The learning phase requires that the relative frequencies from the training set be observed. Each case of, say, class C with an unknown attribute value A is substituted. The next step is to distribute the unknown examples according to the proportion of occurrences in the known instances, treating an incomplete observation as if it falls down all subsequent nodes. The evaluation measure is weighted with the fraction of known values to take into account that the information gained from that attribute will not always be available (but only in those cases where the attribute value is known). During training, instance counts used to calculate the evaluation heuristic include the fractional counts of instances with missing values. Instances with multiple missing values can be fractioned multiple times into numerous smaller and smaller “portions”.

For classification, Quinlan’s technique is to explore all branches below the node in question and then take into account that some branches are more probable than others. Quinlan, once again, borrows Cestnik et al.’s [8] strategy of summing the weights of the instance fragments classified in different ways at the leaf nodes of the tree and then choosing the class with the highest probability or the most probable
1. Partition original dataset into \( n \) incomplete training datasets, \( ITR_1, ITR_2, \ldots, ITR_n \).

2. Construct \( n \) individual DT models (\( DT_1, DT_2, \ldots, DT_n \)) with the different incomplete training datasets \( ITR_1, ITR_2, \ldots, ITR_n \) to obtain \( n \) individual DT classifiers (ensemble members) generated by different imputation algorithms, thus different.

3. Select \( m \) de-correlated DT classifiers from \( n \) DTs using de-correlation maximization algorithm.

4. Using Eq. 3, obtain \( m \) DTs’ output values (misclassification error rates) of unknown instance.

5. Transform the output value to reliability degrees of positive class and negative class, given the imbalance of some datasets.

6. Fuse the multiple DTs into aggregate output in terms of majority voting. When there is a tie in the predicted probabilities, choose the class with the highest probability or else use a random choice when the probabilities between the two methods are equal.

Fig. 2. The ensemble imputation methods algorithm.
Fig. 3. The general formation process of missing data ensemble learning model.
training data selection strategies generate multiple classifiers by applying a single learning algorithm, to different versions of a given dataset. Two different methods for manipulating the dataset are normally used: random sampling with replacement (also called bootstrap sampling) in bagging and re-weighting of the misclassified training instances in boosting. In our study, bagging is selected.

4.3.3. Decision tree learning

A brief description of DTs has already been given in Section 3 of the paper. Many variants of DT algorithms have concentrated on DTs in which each node checks the value of a single attribute. This class of DTs may be called axis – parallel because the tests at each node are equivalent to hyper-planes that are parallel to axes in the attribute space, i.e., they correspond to partitioning the parameter space with a set of hyper-planes that are parallel to all the features axes except for the one being tested and are orthogonal to that one.

In axis – parallel decision methods, a tree is constructed in which at each node a single parameter is compared to some constant. If the feature value is greater than the threshold, the right branch of the tree is taken; if the value is smaller, the left branch is followed.

There are DTs that test a linear combination of the attributes at each internal node. They allow the hyper-planes at each node of the tree to have any orientation in parameter space [42,43]. Mathematically, this means that at each node a linear combination of some or all the parameters is computed (using a set of feature weights specific to that node) and the sum is compared with a constant. The subsequent branching until a leaf node is reached is just that used for axis parallel trees.

Since these tests are equivalent to an oblique orientation to the axes, we call this class of trees oblique DTs. Note that oblique DTs produce polygonal (polyhedral) partitioning of the attribute space. Oblique DTs are also considerably more difficult to construct than axis parallel trees because there are so many more possible planes to consider at each node. As a result, the training process is slower. This is one the major reasons why axis-parallel DTs are considered in our study.

4.3.4. Selecting appropriate ensemble members

After training, each individual DT grown by using different MDTs has generated its own result. However, if there is a great number of individual members (i.e., MDTs, hence, in part, DTs), we need to select a subset of representatives in order to improve ensemble efficiency. Furthermore, it does have to follow the rule ‘the more the better’ rule as mentioned by some researchers. In this study, a de-correlation maximization method [24,34] is used to select the appropriate number of DTs ensemble members. The idea is that the correlations between the selected MDTs, thus, DT classifiers, should be as small as possible. The de-correlation matrix can be summarized in the following steps:

1. Compute the variance-covariance matrix and the correlation matrix
2. For the $i^{th}$ DT classifier ($i = 1, 2, \ldots, p$), calculate the plural-correlation coefficient $i$
3. For a pre-specified threshold $\theta$, if $i < \theta$, then the $i^{th}$ classifier should be deleted from the $p$ classifiers.
   Conversely, if $i > \theta$, then the $i^{th}$ classifier should be retained.
4. For the retained classifiers, perform Eqs (1–3) procedures iteratively until satisfactory results are obtained.

4.3.5. Performance measure evaluation

In the previous phase the DT classifier outputs are used as performance evaluation measures (in terms of misclassification error rates). It has often been argued that selecting and evaluating a classification model based solely on its error rates is inappropriate. The argument is based on the issue of using
both the false positive (rejecting a null hypothesis when it is actually true) and false negative (failing to reject a null hypothesis when it is in fact false) errors as performance measures whenever classification models are used and compared. Furthermore, in the business world, decisions (of the classification type) involve costs and expected profits. The classifier is then expected to help making the decisions that will maximise profits. For example, predicting development effort of software systems involves two types of errors: 1) predicting software effort as likely to be high when in fact it is low, and 2) predicting software effort development as likely to be low when in fact it is high. Now, mere misclassification rate is simply not good enough to predict software effort. To overcome this problem and further make allowances for the inequality of mislabelled classes, variable misclassification costs are incorporated in our attribute selection criterion via prior specification for all our experiments. This also solves the imbalanced data problem. Details about how misclassification costs are used for both splitting and pruning rules are presented in [4].

4.3.6. Integrating multiple classifiers into ensemble output

Depending upon the work in the previous stages, a set of appropriate number of ensemble members can be identified. The subsequent task is to combine these selected members into an aggregated classifier in an appropriate ensemble strategy. Common strategies to combine these single DT results and then produce the final output are simple averaging; weighted averaging, ranking and majority voting. For more information on these strategies, the reader is referred to [14], which are otherwise briefly discussed below.

Simple averaging is one of the most frequently used combination methods. After training the members of the ensemble, the final output can be obtained by averaging the sum of each output of the ensemble members. Some experiments have shown that simple averaging is an effective approach [3].

Weighted averaging is where the final ensemble result is calculated based on individual ensemble members’ performances and a weight attached to each individual member’s output. The gross weight is 1 and each member of an ensemble is entitled to a portion of this gross weight according to their performance or diversity.

Ranking is where members of the ensemble are called low level classifiers and they produce not only a single result but a list of choices ranked according to their likelihood. Then the high level classifier chooses from these set of classes using additional information that is not usually available to or well represented in a single low level classifier.

Majority voting is the most popular combination method for classification problems because of its easy implementation. Members of trees voting decide the value of each output dimension. It takes over half the ensemble to agree a result for it to be accepted as the final output of the ensemble (regardless of the diversity and accuracy of each tree generalization). Majority voting ignores the fact that some trees that lie in a minority sometimes do produce correct results. However, this is the combination strategy approach we follow in our study.

The following example demonstrates the mechanics of the proposed procedure as presented in Table 3.

Suppose that there are two methods used to handle incomplete data when growing decision trees, resulting in two trees. Also, there are only two classes in the data: 1 and 2. Using the predicted probabilities, there are four possible classification patterns from the training data such as (1, 1), (1, 2), (2, 1) (2, 2). The first element in each pair denotes the class predictions by DT\textsubscript{1} and the second by DT\textsubscript{2}. If the instance has predictions (1, 1) for both methods, it is simply assigned class 1 anyway. However, if the instance has predictions (1, 2) it is assigned to the class with the higher overall probability which in this case happens to be 2, and so on.
Table 3  An example pattern table

<table>
<thead>
<tr>
<th>Predicted probabilities (for each method)</th>
<th>Predicted class (for each method)</th>
<th>Predicted class (combining methods)</th>
</tr>
</thead>
<tbody>
<tr>
<td>{(0.6, 0.4); (0.7, 0.3)}</td>
<td>(1, 1)</td>
<td>1</td>
</tr>
<tr>
<td>{(0.6, 0.4); (0.3, 0.7)}</td>
<td>(1, 2)</td>
<td>2</td>
</tr>
<tr>
<td>{(0.4, 0.6); (0.7, 0.3)}</td>
<td>(2, 1)</td>
<td>1</td>
</tr>
<tr>
<td>{(0.4, 0.6); (0.3, 0.7)}</td>
<td>(2, 2)</td>
<td>2</td>
</tr>
</tbody>
</table>

5. Related work

Several researchers have examined various techniques to solve the problem of incomplete data in SE. One popular approach includes discarding instances with missing values and restricting the attention to the completely observed instances, which is also known as listwise deletion (LD). This is the default in commonly used statistical packages such as SPLUS [73]. Another common way uses imputation (estimation) approaches that fill in a missing value with an efficient single replacement value, such as the mean, mode, hot deck, and approaches that take advantage of multivariate regression and k-NN models. Another technique for treating incomplete data is to model the distribution of incomplete data and estimate the missing values based on certain parameters. Specific results are discussed below.

Lakshminarayan et al. [35] performed a simulation study comparing two ML methods for missing data imputation using an industrial process maintenance dataset of 82 variables and 4383 instances. Their results show that for the single imputation task, the SL algorithm C4.5, which utilizes the fractioning of cases (FC) strategy, performed better than Autoclass [9], a strategy based on unsupervised Bayesian probability. For the MI task, both methods performed comparably.

MI was used by El-Emam and Birk [16] to handle missing values in their empirical study that evaluated the predictive validity of the capability measures of the ISO/IEC 15504 software development processes (i.e., develop software design, implement software design, and integrate and test). The study was conducted on 56 projects. For large organizations their results provided evidence of predictive validity which was found to be strong while for small organizations the evidence was rather weak.

The pioneering work by Strike et al. [62] performed a comprehensive simulation study to evaluate three MDTs in the context of software cost modelling. These techniques are LD, mean or mode single imputation (MMSI) and eight different types of hot deck single imputation (HDSI). A dataset composed of 206 software projects and 26 different companies was used. Their results show LD as not only having a severe impact on regression estimates but yielding a small bias as well. However, the precision of LD worsens with increases in missing data proportions. Their results further show that better performance would be obtained from applying imputation techniques.

Another comparative study of LD, mean or mode single imputation (MMSI), similar response pattern imputation (SRPI) and full information maximum likelihood (FIML) in the context of software cost estimation was carried out by Myvreit et al. [44]. The simulation study was carried out using 176 projects. Their results show FIML performing well for data that is not MCAR. LD, MMSI and SRPI are shown to yield biased results for data other than MCAR.

The performance of $k$-NNSI and sample mean imputation (SMI) was analyzed by Cartwright et al. [7] using two industrial datasets; one dataset from a bank (21 completed projects) and the other from a multinational company (17 projects). Their results show both methods yielding good results with $k$NNSI providing a more robust and sensitive method for missing value estimation than SMI.

Song and Shepperd [63] evaluated $k$NNSI and class mean imputation (CMI) for different patterns and mechanisms of missing data. The dataset used for the study is the ISBSG database with 363 complete
instances. Their results show $k$NNSI slightly outperforming CMI with the missing data mechanisms having no impact on either of the two imputation methods.

The $k$–NNSI method was evaluated using a likert dataset with 56 cases in the SE context by Jönsson and Wohlin [25]. Their results not only showed that imputing missing likert data using the $k$-nearest neighbour method was feasible but also showed that the outcome of the imputation depends on the number of complete cases more than the proportion of missing data. Their results further showed the importance of choosing an appropriate $k$ value when using such a technique.

The use of multinomial logistic regression imputation (MLRI) for handling missing categorical values on a dataset on 166 projects of the ISBSG multi-organizational software database was proposed by Sentas et al. [59]. Their proposed procedure was compared with LD, MMSI, Expectation-Maximization single imputation (EMSI) and regression-based single imputation (RBSI). Their results showed LD and MMSI as efficient when the percentage of missing values is small while RBSI and MLRI were shown to outperform LD and MMSI as the amount of missing values increased. Overall, MLRI gave the best results, especially for MCAR and IM data. For MAR data, MLRI compared favourably with RBSI.

Twala et al. [71] evaluates the impact of seven MDTs {LD, EMSI, 5-NNSI, MMSI, MI, FC and surrogate variable splitting (SVS)} on eight industrial datasets by artificially simulating three different proportions, two patterns and three mechanisms of missing data. Their results show MI achieving the highest accuracy rates with other notably good performances by methods such as FC, EMSI and $k$NNSI. The worst performance was by LD. Their results further show MCAR data as easier to deal with compared with IM data. Twala [68] further found missing values as more damaging when they are in the test sample than in the training sample. MIA was shown to be the most effective method when dealing with incomplete data using DTs especially for IM data [69]. The superior performance of MI and the severe impact of IM data on predictive accuracy is also observed in [67].

Despite the scarcity and small sizes of software data and the fact that the LD procedure involves an efficiency cost due to the elimination of a large amount of valuable data, most SE researchers have continued to use it due to its simplicity and ease of use. However, by sacrificing a large amount of data, the sample size is severely reduced. There are other problems caused by using the LD technique. For example, elimination of instances with missing information decreases the error degrees of freedom in statistical tests such as the student t distribution. This decrease leads to reduced statistical power (i.e. the ability of a statistical test to discover a relationship in a dataset) and larger standard errors. In extreme cases this may mean that there is insufficient data to draw any useful conclusions from the study. Other researchers have shown that randomly deleting 10% of the data from each attribute in a matrix of five attributes can easily result in eliminating 59% of instances from analysis [27,33].

As for the SI techniques, the results are not so clear, especially for small amounts of missing data. However, the performance of each technique differs with increases in the amount of missing data. Several shortcomings of single imputation have been documented by Little and Rubin [36], Schafer and Graham [57] and others. The obvious limitation is that it cannot reflect sampling variability under one model for nonresponse or uncertainty about the correct model for nonresponse, i.e. it does not account for the uncertainty about the prediction of the imputed values. The lack of sampling variability could distort estimates, standard errors, and hypothesis tests, leading to statistically invalid inferences [36].

MI, which overcomes limitations of single imputation methods by creating an inference that validly reflects uncertainty of prediction of the missing data, seem not to have been widely adopted by researchers. This is despite the fact that MI has been shown to be flexible, and software for creating MIs is available and some downloadable free of charge.¹

¹Methodology Centre, Penn State University, USA [40].
6. Experimental set-up

6.1. Introduction

Incomplete data has been shown to have a negative impact in reducing ML performance in terms of predictive accuracy while the use of an ensemble of classifiers strategy has been shown to improve predictive accuracy by aggregating the predictions of many classifiers. In this regard, two sets of experiments based on ten industrial datasets (see Table 4) are carried out. Of these datasets, a couple were obtained from the Promise SE repository [54] and others, like ISBSG [23], from researchers at various software companies. Nearly all the datasets used for the experiments have missing values. The objective is to have control over of how different missing data patterns and mechanisms are simulated on a complete dataset. Thus, all instances with missing values were initially removed using LD (which assumes missing values are MCAR) before starting the experiment.

The response or class attribute (software effort) is continuous for all the datasets. However, in this paper we are dealing with a classification-type of problem that predicts values of a categorical dependent attribute from one or more continuous or categorical attribute values. Therefore, software effort was made discrete into a set of three disjoint categories for all datasets. The rule induction technique [15, 49] was utilized for this task. The cut-points were determined in a way that was not blinded to the class attribute, hence, achieving unbiased effect estimates. Monte Carlo simulations which take into account both multiplicities and uncertainty in the choice of cut-points were also utilized for this task [22]. For example, for the Company X dataset, the three categories used were: (low effort, when EFFORT $< 5000$; medium effort, when $5000 < EFFORT <= 10000$; high effort, when EFFORT $> 10000$).

5-fold cross validation was used for tree induction for the experiments. For each fold, four of the parts of the instances in each category were placed in the training set, and the remaining one was placed in the corresponding test as shown in Table 5.

Since the distribution of missing values among attributes (pattern) and the missing data mechanism were two of the most important dimensions of this study, three suites of data were created corresponding to MCAR, MAR and IM. In order to simulate missing values on attributes, the original datasets are run using a random generator (for MCAR) and a quantile attribute-pair approach (for both MAR and IM, respectively). Both of these procedures have the same percentage of missing values as their parameters. These two approaches were also run to get datasets with four levels of proportion of missingness $p$, i.e.,
Table 5
Partitioning of dataset to training and test sets

<table>
<thead>
<tr>
<th>Fold</th>
<th>Training set</th>
<th>Test set</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Part II + Part III + Part IV + Part V</td>
<td>Part I</td>
</tr>
<tr>
<td>2</td>
<td>Part I + Part III + Part IV + Part V</td>
<td>Part II</td>
</tr>
<tr>
<td>3</td>
<td>Part I + Part II + Part IV + Part V</td>
<td>Part III</td>
</tr>
<tr>
<td>4</td>
<td>Part I + Part II + Part III + Part V</td>
<td>Part IV</td>
</tr>
<tr>
<td>5</td>
<td>Part I + Part II + Part III + Part IV</td>
<td>Part V</td>
</tr>
</tbody>
</table>

0%, 15%, 30% and 50% missing values. The experiment consists of having p% of data missing from both the training and test sets.

6.2. Modelling of missing data mechanisms

The missing data mechanisms were constructed by generating a missing value template (1 = present, 0 = missing) for each attribute and multiplying that attribute by a missing value template vector. Our assumption is that the instances are independent selections.

For each dataset, two suites were created. First, missing values were simulated on only half of the attributes. Second, missing values were introduced on all the attribute variables. For both suites, the missingness was evenly distributed across the attributes. This was the case for the three missing data mechanisms, which from now on shall be called MCAR_{half}, MAR_{half}, IM_{half} (for the first suite) and MCAR_{unifo}, MAR_{unifo}, IM_{unifo} (for the second suite). These procedures are described below.

For MCAR, each vector in the template (values of 1’s for non-missing and 0’s for missing) was generated using a random number generator utilising the Bernoulli distribution. The missing value template is then multiplied with the attribute of interest, thereby causing missing values to appear as zeros in the modified data.

Simulating MAR values was more challenging. The idea is to condition the generation of missing values based upon the distribution of the observed values. Attributes of a dataset are separated into pairs, say, (A_x, A_Y), where A_Y is the attribute into which missing values are introduced and A_X is the attribute on the distribution of which the missing values of A_Y is conditioned, i.e., P(A_Y = miss|A_x = observed). The pairing of attributes was based on how highly correlated to one another they were. So, highly correlated attributes was paired against each other. Since we want to keep the percentage of missing values at the same level overall, we had to alter the percentage of missing values of the individual attributes. Thus, in the case of k% of missing values over the whole dataset, 2k% of missing values were simulated on A_Y. For example, having 10% of missing values on two attributes is equivalent to having 5% of missing values on each attribute. Thus, for each of the A_x attributes its 2k quantile was estimated. Then all the instances were examined and whenever the A_X attribute has a value lower than the 2k quantile a missing value on A_Y is imputed with probability 0, and 1 otherwise. More formally, P(A_Y = miss| A_x < 2k) = 0 or P(A_Y = miss| A_x > 2k) = 1. This technique generates a missing value template which is then multiplied with A_Y. Once again, the attribute chosen to have missing values was the one highly correlated with the class variable. Here, the same levels of missing values are kept. For multi-attributes, different pairs of attributes were used to generate the missingness. Each attribute is paired with the one it is highly correlated to. For example, to generate missingness in half of the attributes for a dataset with, say, 12 attributes (i.e. A_1, . . . , A_{12}), the pairs (A_1, A_2), (A_3 ,A_4) and (A_5, A_6) could be utilised. We assume that A_1 will be highly correlated with A_2; A_3 highly correlated with A_4, and so on. For the (A_1, A_2) pairing, A_1 is used to generate a missing value template of zeros and
ones utilizing the quantile approach. The template is then used to “knock off” values (i.e., generating missingness) in $A_2$, and vice versa.

In contrast to the MAR situation outlined above where data missingness is explainable by other measured variables in a study, IM data arise due to the data missingness mechanism being explainable, and only explainable by the very variable(s) on which the data are missing. For conditions with data IM, a procedure identical to MAR was implemented. However, for the former, the missing values template was created using the same attribute variable for which values are deleted in different proportions.

For consistency, missing values were generated on the same attributes for each of the three missing data mechanisms. This was done for each dataset. For split selection, the impurity approach was used. For pruning, we use 5-fold cross validation cost complexity pruning and 1 Standard Error (1-SE) rule in CART to determine the optimal value for the complexity parameter [4]. The same splitting and pruning rules when growing the tree were carried out for each of the ten industrial datasets.

6.3. Performance evaluation

A classifier was built on the training data and the predicted accuracy is measured by the smoothed classification error rate of the tree, and was estimated on the test data. Instead of summing terms that are either zero or one as in the error-count estimator, the smoothed misclassification error rate uses a continuum of values between zero and one in the terms that are summed. The resulting estimator has a smaller variance than the error-count estimate. Also, the smoothed error rate can be very helpful when there is a tie between two competing classes. This is the main reason why it was considered for our experiments.

It was reasoned that the condition with no missing data should be used as a baseline and what should be analysed is not the error rate itself but the increase or excess error induced by the combination of conditions under consideration. Therefore, for each combination of method for handling incomplete data, the number of attributes with missing values, proportion of missing values, and the error rate for all data present was subtracted from each of the three different proportions of missingness. This would be the justification for the use of differences in error rates analysed in some of the experimental results.

All statistical tests were conducted using the MINITAB statistical software program [39]. Analyses of variance, using the generalized linear model procedure [32] were used to examine the main effects and their respective interactions. This was done using a 4-way repeated measures design (where each effect was tested against its interaction with datasets). The fixed effect factors were the: missing data techniques; number of attributes with missing values (missing data patterns); missing data proportions; and missing data mechanisms. A 1% level of significance was used because of the many number of effects. Results were averaged across five folds of the cross-validation process before carrying out the statistical analysis. The averaging was done as a reduction in error variance benefit.

Numerous measures are used for performance evaluation in ML. In predictive knowledge discovery, the most frequently used measure is predictive accuracy. To measure the performance of MDTs and the ensemble MDTs, the training set/test set methodology is employed. Unfortunately, an operational definition of accurate prediction is hard to come by. However, predictive accuracy is mostly operationally defined as the prediction with the minimum costs (the proportion of misclassified instances). The need for minimizing costs, rather than the proportion of misclassified instances, arises when some predictions that fail are more catastrophic than others, or when some predictions that fail occur more frequently than others. Minimizing costs, however, does correspond to minimizing the proportion of misclassified instances when priors are taken to be proportional to the class sizes and when misclassification costs are taken to be equal for every class. This is the approach we follow in the paper.
Khoshgoftaar and Seliya [28] and Khoshgoftaar et al. [31] argue that in the context of the software development, selecting and evaluating a classification model (like a DT) based solely on its error rates is inappropriate. Their argument is based on the issue of using both the false positive (rejecting a null hypothesis when it is actually true) and false negative (failing to reject a null hypothesis when it is in fact false) errors as performance measures whenever classification models are used and compared. Furthermore, in the business world, decisions (of the classification type) involve costs and expected profits. The classifier is then expected to help making the decisions that will maximise profits. CART is able to handle this problem in terms of taking into account misclassification costs when constructing the DT.

6.4. Programs and codes for methods

No software or code was used for LD. Instead, all instances with missing values on that particular attribute were manually excluded or dropped, and the analysis was applied only to the complete instances.

S-PLUS code was also developed for the MMSI approach. The code was developed in such a way that it replaced the missing data for a given attribute by the mean (for a numerical or quantitative attribute) or mode (for a nominal or qualitative attribute) of all known values of that attribute.

An index structure M-tree [11] was used as a representative of the $k$-NN approach for handling missing attribute values in both the training and test samples. This technique can organize and search datasets based on a generic metric space. In addition, it can drastically reduce the number of distance computations in similarity queries. The missing values were estimated using 1, 3, 5, 11, 15, 21 nearest neighbours. However, only results with 5-nearest neighbour will be showed in this work.

From the MIA method, an unknown (missing) value is considered an additional attribute value. Hence, the number of values is increased by one of each attribute that depicts an unknown value in the training or test set. S-PLUS code for this method was developed.

There are many implementations of MI. Schafer’s [55] set of algorithms (headed by the NORM program) that use iterative Bayesian simulation to generate imputations was an excellent option. NORM was used for datasets with only continuous attributes. A program called MIX written was used for mixed categorical and continuous data. For strictly categorical data, CAT was used. All three programs are available as S-PLUS routines.

For the SVS method, the RPART routine, which implements within S-PLUS many of the ideas found in the CART book and programs of Breiman et al. [4] was used for both training and testing DTs.

The DTSI method uses a DT for estimating the missing values of an attribute and then uses the data with filled values to construct a DT for estimating or filling in the missing values of other attributes. An S-PLUS code to estimate missing attribute values using a DT for both incomplete training and test data was developed.

The DT learner C4.5 was used as a representative of the FC or probabilistic technique for handling missing attribute values. This technique is probabilistic in the sense that it constructs a model of the missing values, which depends only on the prior distribution of the attribute values for each attribute tested in a node of the tree. The main idea behind the technique is to assign probability distributions at each node of the tree. These probabilities are estimated based on the observed frequencies of the attribute values among the training instances at that particular node.

S-PLUS codes for all the ensembles were developed.
6.5. Experiments

6.5.1. Experiment I

In order to empirically evaluate the performance of the eight MDTS on predictive accuracy, an experiment on ten industrial datasets is given in Table 2 is used. This experiment is carried out in order to rank individual MDTs and also assess the impact of missing data (at various levels) on software predictive accuracy.

6.5.1.1. Experimental results I

Experimental results on the effects of current methods for handling both incomplete training and test data on predictive accuracy using DTs are described. The behaviour of these methods is explored for different patterns, levels of missing values, and for the mechanism of missing data. The error rates of each method of dealing with the introduced missing values are averaged over the ten datasets. Also, all the error rates increases over the complete data case are formed by taking differences. From these experiments the following results are observed.

Main effects

All the main effects were found to be significant at the 1% level of significance ($F = 61.95$, df = 7 for MDTs; $F = 19.36$, df = 1 for number of attributes with missing values; $F = 201.54$, df = 2 for missing data proportions, $F = 112.80$, df = 2 for missing data mechanism; $p < 0.01$ for each).

From Fig. 4, MIA is the overall best technique for handling incomplete data with an excess error rate of 8.2%, closely followed by MI, FC and $k$-NNSI, with excess error rates of 9.6%, 9.9% and 10.8, respectively. The worst technique is LD, which exhibits an error rate of 14.7%.

Tukey’s multiple comparison tests showed no significant differences between DTSI and MMSI (on the one hand) and MIA and MI (on the other hand). However, significant differences are observed between MI and the other single imputation strategies like DTSI, $k$-NNSI and MMSI. The two SL strategies (FC and SVS) were found to be significantly different from each other. 5-NNSI and FC were found to be not significantly different from each other. The significance level for all the comparison tests is 0.01. All interaction effects we found to be insignificant at the 1% level of significance. Hence, they are not discussed in the paper.
6.5.2. Experiment II

The main objective of this experiment is to compare the performance of the new ensemble method with current approaches to deal with the problem of incomplete data, especially the top three MDTs that exhibited higher accuracy rates in the previous experiment. These include MI, MIA and FC. Also, we thought it would be interesting to test the effectiveness of the proposed approach with a statistical imputation technique (MI), a new approach (MIA) and a SL technique (FC). From the combination of three MDTs, three different ensembles methods are also proposed as given below:

i. MIAMI, a component of MIA and MI
ii. MIAFC, a component of MIA and FC; and
iii. MIFC, a component of MI and FC

The above three ensemble methods are also compared with each of the three MDTs, individually.

6.5.2.1. Experimental results II

Main effects

All the main effects were found to be significant at the 1% level of significance (F = 84.5, df = 5 for existing and ensemble missing data methods; F = 26.3, df = 1 for number of attributes with missing values; F = 73.8 df = 2 for missing data proportions, F = 36.4, df = 2 for missing data mechanism; p < 0.01 for each).

Figure 5 shows the average results of 180 experiments (3 MDTs plus 3 ensemble methods x 2 missing data patterns x 3 missing data proportions x 3 missing data mechanisms) which summarizes imputation accuracy of each method. The accuracies of each method are averaged over the ten datasets. Figure 5 further shows that the MIAMI ensemble has on average the best accuracy throughout the entire spectrum missing data proportions, patterns, and mechanisms. Tukey’s multiple comparison tests showed significant differences between MIAMI and the other individual MDTs at the 1% level.

Interaction effects

Only two two-way interactions effects were found to be statistically significant at the 1% level. These are: the interaction between current and new MDTs, ensemble methods and missing data pattern (F = 8.45, df = 12; p < 0.01), and the interaction between current and new MDTs and ensemble methods and missing data mechanisms (F = 13.8, df = 18; p < 0.01).
The interaction effect between the ensemble and MDTs and the missing data patterns is shown in Fig. 6. This figure shows all the methods performing better when the missing values are in all the attributes compared with when they are only in half of the attributes.

In addition, the impact of missing data patterns appears to differ by the type of missing data method. For example, MI exhibits one of the biggest increases in error rates (as the number of attributes with missing values increases) compared with only a very small increase achieved by MIA. Figure 7 shows all techniques achieving bigger error rates when dealing with IM data compared with either MCAR or MAR data. From the three ensemble methods, the most affected is MIAFC, especially for IM data. In addition, some techniques appear to be severely impacted by the different missing mechanisms than others.

6.5.3. Results for Individual Datasets: Current and New MDTs Vs. Ensemble Methods

The results that illustrate specific deviations from the overall results of the effectiveness of the new ensemble methods against the current and new MDTs for constructing and classifying incomplete vectors on different database characteristics, especially on datasets where the new method yielded superior performance compared with MIA, MI and FC, are given below.
6.5.3.1. Results on a dataset with mainly nominal attributes and small sample size: test equipment

For the test equipment problem, the effects of missing values on classification accuracy for MCAR\textsubscript{half} data are summarised in Fig. 8A. MIAFC performs slightly better than MIAMI at the 30% level of missing values. However, there are no clear differences in error rates between all the individual MDTs and ensemble methods, especially at higher levels of missing values.

Once again, MIAFC outperforms MIAMI at the 30% level of missing values (Fig. 8B). However, the difference in performances by all the methods is now noticeable.

Performances by all the different methods for handling MAR\textsubscript{half} data follow a similar pattern to that of MCAR\textsubscript{half} data (Fig. 8C).

For MAR\textsubscript{all} data, both MIAFC and MIAMI are the most effective methods as shown in Fig. 8D. The poor performance of MIFC (especially at lower levels of missing values) is also observed.

The impact of IM\textsubscript{half} data on classification accuracy is shown in Fig. 8E. One again there appears to be no significant differences between methods although the impact of missing data appears to be more severe for this type of mechanism.

In this suite of IM\textsubscript{all} experiments, the behaviour of the methods shown in Fig. 8F is not different from the one observed in the MAR\textsubscript{all} case. However, MIAMI slightly outperforms MIAFC at the 15% level but its performance deteriorates as the proportion of missing data increases.

For this kind of dataset it appears that MIAFC handles small datasets with mainly numerical attributes well, especially for MCAR or IM data. This seems to be the case when missing values are distributed among all attributes. When missing values are only in half of the attributes, we observe good performances by MIAMI even though the difference in performances by all the methods appears not to be significant.

6.5.3.2. Results on a dataset with purely numerical attributes: CCCS

As can be seen from Fig. 9A, the overall best performance for MCAR\textsubscript{all} data is by MIAFC, with MIAMI as a serious competitor.

The results for the MCAR\textsubscript{all} suite suggest prominent increases in error rates in some cases compared with MCAR\textsubscript{half} (Fig. 9B). MIAMI and MIAFC yield the best performances with no clear ‘winner’ between the three ensemble methods at the 50% level.

In the MAR\textsubscript{half} suite, the ensemble methods show slightly superior performances compared with individual MDTs (Fig. 9C).

Figure 9D shows the performance of MIAMI as improving from being the second best method (in the MCAR\textsubscript{half} case) to being the best method at the 50% level of missing values for handling MAR\textsubscript{all} data.

For the IM\textsubscript{half} suite, the results illustrated in Fig. 9E show a relatively superior performance by MIAMI over MIAFC and MIAFC, especially at lower levels of missing values. At the 50% level, there are no significant differences in performance between the ensembles and MIA.

In the IM\textsubscript{all} case (Fig. 9F), the behaviour by all the methods is similar to the one observed in IM\textsubscript{half}. MIAMI performs as good as MIAFC at all levels of missing data with MIAFC struggling at the 50% level.

MIAMI exhibits a very good performance for this dataset which contains purely numerical attributes. This is the case when missing values are in all the attributes and for the MCAR and MAR situations. Also, the performance of MIAMI seems to be better on average when missing values are distributed among half of the attributes. However, the poor performance of MIAFC for handling MCAR\textsubscript{all} and MAR\textsubscript{all} data compared to its superior performance for handling IM\textsubscript{half} is rather surprising while the good performance of MIAMI is expected due to the slightly bigger size of the data.
Fig. 8. Comparative results of current, new MDTs and ensemble methods for the test equipment data. A) MCAR_{half}, B) MCAR_{all}, C) MAR_{half}, D) MAR_{all}, E) IM_{half}, F) IM_{all}.
Ensemble Imputation Methods

Fig. 9. Comparative results of current, new MDTs and ensemble methods for the CCCS data. A) MCAR_{half}, B) MCAR_{all}, C) MAR_{half}, D) MAR_{all}, E) IM_{half}, F) IM_{all}.
6.5.3.3. Results on a dataset with mainly nominal attributes and bigger sample size: Company X

From Fig. 10A it appears that when both training and test data has missing values due to the MCAR half mechanism, MIAMI achieves slightly better accuracy compared to the other methods. This is the case at the 15% level of missing values. At higher levels of missing values, the difference in performances of methods is not significant.

For MCAR all data, the results show MIAMI achieving the best results at the 50% level while the performance of MIAFC deteriorates with increases in missing data (Fig. 10B).

Error rates of methods for the Company X data problem for MAR half data are presented in Fig. 10C. At lower levels of missing values, the best overall performance is by MIAMI, closely followed by MIAFC. At the 50% level, MIAFC rises from last to third position.

Results by methods for MAR all data are displayed in Fig. 10D. Once again, MIAMI achieves higher accuracy rates at all levels of missing values. The poor performance of MIAFC compared to individual MDTs is also observed.

In this suite of IM half experiments, the results show each method performing best at each respective level of missing values. MIAMI performs best at the 15 and 50% levels while MIAFC achieves the best result with MIAMI at the 30% level. At the 50% level, MIAFC produces a much better performance compared to its poor performance at lower levels of missing values (Fig. 10E).

In the IM all case, as shown in Fig. 10F, MIAMI exhibits the highest accuracy rates at all levels of missing values, closely followed by MIAFC. The poor performance of MIAFC is also noticeable.

For this kind of dataset, it seems that MIAMI handles any form of missing data pattern and mechanism better but with serious competition from MIAFC. MIFC seems not only to be ineffective as a method for handling much bigger datasets but also ineffective for handling missing values when they are distributed among half of the attributes (on the one hand) and when they are distributed in all the attributes (on the other hand).

7. Discussion and conclusions

The major contribution of the paper is the development of an ensemble of techniques for imputation of missing data algorithm in domain specific datasets for the prediction of software development effort. The referred techniques are well known, but using the ensemble of them is an original contribution. Furthermore, imputation methods are not of widespread use in SE, so showing the possibility of using the techniques on software process data is another contribution of this paper for SE.

The empirical study is based on ten SE datasets, and the results suggest that the proposed technique can be successfully applied. Based on preliminary evidence, it has been found that MIAMI (with MIA and MI as its components) improves the prediction accuracy of the baseline imputation methods (MI, MIA and FC). This improvement is achieved mainly in SE datasets with mostly nominal attributes and whose values are either MAR or IM missing. Individually, MIA is effective for datasets with mostly nominal attributes and whose values are IM while MI achieves higher accuracy rates for MAR data. This is the case for both big and small datasets. The performance of MIAMI was also good for those SE datasets with mostly categorical attributes, especially for the big datasets whose values were IM. MIAFC is more effective for small datasets with mainly numerical attributes, especially when missing values are distributed in all the attributes while MIFC struggles with any type of dataset and its respective attributes.

An important question is why does MIAMI outperforms other ensemble methods by such significant margins, and in particular what differentiates it from, say MIAFC or MIFC? One reason could be the level of “inertia” displayed by each system. All the ensemble algorithms display adaptive behaviour and look
Fig. 10. Comparative results of current, new MDTs and ensemble methods for the CCCS data. A) MCAR_{half}, B) MCAR_{all}, C) MAR_{half}, D) MAR_{all}, E) IM_{half}, F) IM_{all}.
to perturb the data at each iteration of the imputation algorithm. With fractioning of cases based on MI methods or missing incorporated in attributes, once the weight associated with an observation becomes trivial it may as well be excluded from further iterations of the algorithm because the weight is unlikely to recover to a significant level. With missingness incorporated into attributes based on MI methods this is not the case. An observation can flip between inclusion and exclusion at each iteration. Another reason may be the performance of the individual imputation methods comprising each ensemble methods. MI and MIA consistently dominated the other methods. One strength of MIA that is worth emphasis is the ease with which it handles IM data while MI requires the data be MAR. However, even when the MAR assumption is violated, MI performs better than several imputation methods. As Schafer [55] points out, the richness of the observed data is related to the plausibility of the MAR assumption. In a multivariate dataset, if the associations among the observed data are reasonably strong, then the observed data may provide good information for constructing a model for imputing the missing data.

There are two other advantages of MI, and both involve correcting for the uncertainty of the imputations. One advantage is in the parameter estimates, by essentially taking the mean of the parameter estimate from each of the \( m \) imputations (where \( m \) is the number of imputations). The next advantage of MI is the correction of standard errors for within and between imputation variance, which allows for better estimates of confidence intervals and \( p \)-values. The strengths of MIA and MI (individually) could be the reason why the combination (ensemble) of the two methods consistently dominates the other methods.

The poor performance of any ensemble with FC as a component could be attributed to the fact that not only does the strategy fail to exploit or take into account the interrelationships among the attributes, the mechanism generating the missing attribute values is unknown. Also, The FC approach works well when most of the attributes are independent, because it depends only on their prior distribution of the attribute values for each attribute being tested in a node of the tree. In other words, the different imputation treatments differ in how they take advantage of statistical dependencies between attributes.

Our results also show the impact on the performance of methods is caused by the pattern and mechanism of missing values, especially at lower levels of missingness. However, as the proportion of missing values increases, the major determining factor on the performance of methods is how the missing values are distributed among attributes. All methods yield lower accuracy rates when missing values are distributed among all the attributes compared with when only one attribute has missing values. The performance of MI and FC degrade faster as the number of attributes with missing values increases. This was the case for all the datasets. Also, given that the classification performance of each method varies by mechanism of missing data, it appears that the treatment of missing values not only heavily depends on the missing data proportions but on the nature of the missing data pattern as well.

The worst performance achieved by methods is for IM data, followed by MCAR and MAR data, respectively. This was a surprising result, which is in not in accordance with statistical theory which considers MCAR as easier to deal with and IM data as very difficult to handle [36].

It was also observed that the impact of missing values depends not just upon a missing data technique, missing data pattern or missing data mechanism but upon a combination of the three. Therefore, neither can be considered in isolation.

From both experiments, there exists threats to the validity of the results. Potential threats include the initial exclusion from the datasets of all the instances with missing values, which could have involuntarily introduced biases, especially if those missing values contained improtant information and they were not MCAR as we assumed. However, for two of the biggest datasets (Company X and Finnish) the experimental results were carefully validated. For example, the experiments were conducted under the supervision of a domain expert who had a deep understanding for each respective underlying dataset.
This was a time consuming exercise on ourselves and the experts. The experts were mainly involved in the cleaning up of the datasets in terms of attribute and class noise and outliers. They were also able to give us reasons why some attributes values were missing. For example, some project managers were reluctant to show projects that had, say, high defect rates. This kind of nonresponse results in IM data which otherwise we could not have known.

The averaging of results of the 10 domains/datasets (which is more like mixing datasets from different companies into one dataset) is another potential issue. Some empirical studies in SE assume that domain-and-process factors can be best accounted for within homogenous data, i.e., data produced by the same company, with an almost stable process, in the same environment. Hence, in the case of our study, it would be interesting to see if the results are consistent thorough the ten datasets although results for individual datasets with certain attribute characteristics was looked at but not into great detail.

The issue of determining whether or not to apply the ensemble strategy to a given dataset must be considered. For the work described here, the data were artificially corrupted, i.e. missing values were artificially simulated on the attributes. Unfortunately, this type of information is rarely known for most “real-world” applications. In some situations, it may be possible to use domain knowledge to determine the mechanism generating the missing values. For situations where this knowledge is not available, the conservative nature of the consensus ensemble dictates that the data will be missing randomly. In addition, fundamentally there are three different classes of missingness mechanism which cover whatever situation one might encounter in practice. Our approach is general since one can apply our results as appropriately. For example, a relevant part of our paper could be taken if an empirical SE researcher thinks he/she has MCAR data. In addition, it is possible to formally test for the MCAR assumption. However, as much as MAR and IM are important, it is hard to test their assumptions, especially for IM data, which requires one to test for missingness that was deliberately created when data was collected or the data could be missing due to an unknown censoring mechanism.

This paper has looked at the performance of ensemble imputation methods in terms of smoothed misclassification error rate. A natural extension would be to consider the impact of such ensemble methods on other measures of performance, and in particular measures of group separation such as GINI or the magnitude of relative error that is also commonly used to assess classifier performances in the SE industry. MIAMI also deserves further investigation on a number of fronts. First, in terms of the training parameters and the combination rules that can be employed. Second, empirical studies of the application of MIAMI to datasets from other areas of data mining should be undertaken to assess its performance across a more general field. Third, comparisons should be made between MIAMI and other forms of imputation methods ensembles not investigated in this paper using non-tree-based methods. For example, artificial neural networks and naïve Bayes classifier.

We leave the above issues to be investigated in the future.

In sum, this paper provides the beginnings of a better understanding of the relative strengths and weaknesses of ensemble imputation methods and using decision trees as their component classifier. It is hoped that it will motivate future theoretical and empirical investigations into incomplete data and software prediction, and perhaps reassure those who are uneasy regarding the use of imputed data in software prediction.

Acknowledgements

The authors would like to thank the UK Engineering and Physical Sciences Research Council (under grant GR/S55347) for funding this research and to the University of Johannesburg, South Africa for
additional financial support to finish the research. Further, the authors would like to thank Professor David Hand and the anonymous reviewers for their invaluable comments on the paper and also for the contribution from a third software organization who provided one of the biggest datasets for the simulation study but has requested that its identity remains anonymous.

References


