

# Explanation vs Performance in Data Mining: A Case Study with Predicting Runaway Projects

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**Abstract**—Often, the *explanatory power* of a learned model must be traded off against model *performance*. In the case of predicting runaway software projects, we show that the twin goals of high performance and good explanatory power are achievable after applying a variety of data mining techniques (discretization, feature subset selection, rule covering algorithms). This result is a new high water mark in predicting runaway projects. Measured in terms of precision, this new model is as good as can be expected for our data. Other methods might out-perform our result (e.g. by generating a smaller, more explainable model) but no other method could out-perform the precision of our learned model.

## 1 INTRODUCTION

*Seek simplicity, and distrust it.*  
— Alfred North Whitehead

EVERY teacher knows that generating succinct explanations means skipping over tedious details. Such explanations can be quickly communicated, but can miss the details needed to apply that knowledge in a real world setting.

An analogous situation occurs with data miners. All data miners are *performance systems*; i.e. they can reach conclusions about a test case. However, only some data miners are *explanation systems* that offer a high-level description of how the learned model functions.

The ability to *explain how* a conclusion was reached is a very powerful tool for helping users to understand and accept the conclusions of a data miner. Despite this, sometimes explanatory power must be *decreased* in order to *increase* the efficacy of the predictor. For example, previously Abe, Muzono, Takagi, et.al. used a Naïve Bayes classifier to generate a predictor for runaway software projects [1], [2], [3]. That model *performs* well but, as shown below, cannot easily *explain* how it reaches its conclusions.

This paper repairs the *explainability* of that prior result. Using an iterative exploration of data mining techniques (cross-validation, different rule learners, discretization, feature subset selection), we found a particular combination of methods that yielded succinct explanations of how to predict for runaway software projects while out-performing the Naïve Bayes classifier. In hold-out experiments, this new model exhibited perfect precision; i.e. precision = 1.0. Other methods *might* be able to out-perform this new result (e.g. by finding a more succinct and explainable model) but no other method could be more precise (since  $0 \leq \text{precision} \leq 1$ ).

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The rest of this paper is structured as follows. First, the software runaway problem is defined and the explanation problems of prior results are discussed. Next, the general problem of explaining a learned model is explored using a range of data miners, and examples from the software engineering literatures (in summary, the best *performing* models may be very poor at *explaining* how those models make their conclusions). A class of data miners called *rule learners* will then be introduced and applied to our data via various *treatments* (some combination of discretizer, feature selector, and learner). The subsequent discussion will review (a) related work; (b) the external validity of these results; as well as (c) general principles of building explainable models via data mining.

## 2 RUNAWAY SOFTWARE

GLASS defines a “runaway software project” as “a project that goes out of control primarily because of the difficulty of building the software needed by the system” [4]. For Glass “out of control” means “schedule, cost, or functionality that was twice as bad as the original estimates”.

Many software projects suffer from runaways:

- In 2001, the Standish group reported that 53% of U.S. software projects ran over 189% of the original estimate [5]. This 189% is not the 200% required by Glass’ definition, but it is close enough and large enough to be alarming.
- Figure 1 shows data from 31 real-world projects, 10 of which (32%) are classified as “runaway”.

Figure 1 was collected by [1], [2], [3] as follows:

- Questions covering the various aspects of software development (see Figure 2) areas were delivered to development companies and collected one month later. These projects are actual industrial software development projects of embedded systems in the period 1996 to 1998.
- The questions were distributed to the project managers or project leaders of various target projects. The detail and purpose of the questionnaire was explained. Answers were coded Strongly agree, Agree, Neither agree nor disagree, and Disagree as 3, 2, 1, and 0, respectively.
- All of these projects had completed their development. As a result, some of the projects could be classified as “runaways”. Takagi et.al. took care to ensure that all developers held a consensus view that some prior project had been a runaway. Also, to be classified as a runaway, the researchers used other objective measures such as cost and duration.

Using manual methods, Takagi et.al. [1] found four features from Figure 1 (e3, e5, p3, p5) that seemed promising predictors for runaways. The coefficients of those terms (found via logistic regression)

Requirements					Estimation					Planning						Team Organizations			Management			class
R1	R2	R3	R4	R5	E1	E2	E3	E4	E5	P1	P2	P3	P4	P5	P6	O1	O2	O3	M1	M2	M3	class
0	0	0	0	0	2	3	3	2	0	2	0	0	0	0	0	2	1	0	0	0	0	ok
0	0	0	0	0	0	0	0	0	0	0	0	0	2	0	0	0	0	0	0	0	0	ok
0	0	0	0	3	0	0	2	3	0	0	0	0	2	0	0	0	0	0	0	0	0	ok
3	3	2	2	3	0	0	2	2	0	2	2	0	0	0	1	2	0	0	0	0	0	ok
0	0	0	0	2	0	0	0	0	0	0	2	0	2	2	0	0	0	0	0	2	0	ok
0	3	2	0	0	2	2	2	0	2	0	2	0	0	0	0	0	0	0	0	0	2	ok
0	0	2	3	2	0	0	0	0	0	0	2	0	3	0	0	0	0	0	0	0	0	ok
0	2	3	3	0	1	0	2	0	0	2	2	0	0	2	2	0	0	1	3	0	0	ok
0	2	0	2	3	0	0	0	0	0	2	2	0	2	2	0	0	0	0	0	0	2	ok
0	0	0	0	2	0	2	2	0	0	0	2	0	0	2	0	0	0	0	2	0	0	ok
0	3	3	2	0	0	0	3	3	0	0	0	0	0	0	0	0	0	2	0	0	0	ok
0	2	2	2	0	0	2	0	0	0	0	2	0	2	0	0	0	0	0	0	0	2	ok
0	2	0	2	0	0	0	0	0	0	2	3	3	0	2	2	2	2	0	2	2	1	ok
0	0	0	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	ok
0	2	2	2	2	0	2	2	0	0	0	0	0	0	0	0	3	2	0	3	0	0	ok
0	0	0	0	2	0	2	0	2	3	3	2	0	2	3	2	3	2	0	2	2	2	ok
0	0	0	0	0	0	2	0	0	0	2	2	2	3	2	2	0	0	0	2	2	0	ok
0	0	0	0	1	0	0	0	0	0	0	2	2	0	0	0	0	0	0	0	0	0	ok
0	0	0	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	ok
0	2	3	2	3	0	0	0	0	0	3	0	0	0	3	0	2	0	0	0	3	3	ok
0	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	ok
3	2	3	3	2	2	1	3	2	1	0	2	2	2	0	1	3	1	2	2	2	0	ok
2	2	0	2	3	0	0	2	3	0	2	0	2	2	3	2	0	0	0	0	2	3	runaway
2	2	3	3	3	2	2	3	2	3	3	3	3	2	3	2	3	3	0	2	2	2	runaway
3	2	0	0	3	0	0	0	0	0	3	0	0	3	3	0	0	0	0	0	0	0	runaway
0	2	3	2	2	3	0	2	2	1	0	2	0	0	2	2	0	2	2	2	0	2	runaway
0	2	2	2	2	0	3	2	3	3	0	2	2	0	0	2	2	2	0	0	0	0	runaway
2	3	3	2	2	0	0	3	3	2	3	0	3	0	2	3	2	0	2	0	2	2	runaway
3	2	3	2	0	3	2	2	2	0	0	2	2	2	3	0	2	0	2	0	3	3	runaway
2	2	3	3	2	0	0	2	0	2	2	2	2	2	2	0	3	0	2	0	2	0	runaway
0	0	0	0	0	0	0	0	0	0	3	3	3	3	3	3	3	3	0	0	3	3	runaway
2	3	3	3	2	2	3	3	3	3	3	3	3	2	3	3	3	3	2	3	3	0	runaway

Fig. 1. Data used in this study, collected using the methods. For an explanation of the columns features, see Figure 2. From [1].

were combined as follows:

$$X(e3, e5, p3, p5) = -8,834 \cdot e3 + 1,577 \cdot e5 + 0,964 \cdot p3 + 1,228 \cdot p3 + 2,222 \cdot p5 \quad (1)$$

$$P(\text{runaway}|X) = \frac{e^x}{1+e^x}$$

Unlike prior results [4], [6], [7], this model is *operational*; it is possible to precisely characterize the strengths and weaknesses of its performance:

- For *high* and *low* values of  $P(\text{runaway}|X)$ , Equation 1 is a perfect predictor for runaways in Figure 1. No project with  $P \leq 0.03$  is a “runaway” and no project with  $P \geq 0.81$  is “ok”. This is the majority ( $\frac{22}{33} = 67\%$ ) of the data in Figure 1.
- In the minority case ( $\frac{11}{33}$ ),  $P$  is mid-range ( $0.03 < P(\text{runaway}|X) < 0.81$ ) and Equation 1 yields incorrect predictions in  $\frac{4}{11}$  rows.

While an important result, Equation 1 has several drawbacks:

- *Not automatic*: Equation 1 was created after a manual inspection of the data by a team of skilled mathematicians. Such a manual analysis is hard to reproduce or apply to a new data set. Subsequent work by Abe, Takagi, et.al [2] automated the method with a Naïve Bayes classifier, but this compromised the *explainability* of the predictive model (see below).
- *Only explores one subset*: Takagi et.al. did not compare the feature subset  $\{e3, e5, p3, p5\}$  with other feature subsets. Hence, while they showed that this subset was useful, they did not demonstrate that it was the *most* useful subset.
- *Ambiguous*: At low and high  $P$  values, Equation 1 sends a clear signal about what is, and is not, a potentially runaway project. However, at middle-range  $P$  values, Equation 1’s conclusions are ambiguous and, hence, hard to explain.

### 3 THE EXPLANATION PROBLEM

LEARNING explainable models is harder than it may appear. This section offers examples where learned models *perform* well, but *explain* themselves poorly.

Features of Figure 1.
<p><i>Requirements</i> features relate to the understanding and commitment of the requirements among the project members:</p> <p>R1: Ambiguous requirements;                      R2: Insufficient explanation of the requirements;                      R3: Misunderstanding of the requirements;                      R4: Lack of commitment regarding requirements between the customer and the project members;                      R5: Frequent requirement changes;</p>
<p><i>Estimation</i> features relate to the technical methods for carrying out the estimation, and the commitment between project members and customers:</p> <p>E1: Insufficient awareness of the importance of the estimation;                      E2: Insufficient skills or knowledge of the estimation method;                      E3: Insufficient estimation of the implicit requirements;                      E4: Insufficient estimation of the technical issues;                      E5: Lack of stake holder’ commitment of estimation.</p>
<p><i>Planning</i> features relate to the planning or scheduling activity and the commitment to the project plan among project members:</p> <p>P1: Lack of management review for the project plan;                      P2: Lack of assignment of responsibility;                      P3: Lack of breakdown of the work products;                      P4: Unspecified project review milestones;                      P5: Insufficient planning of project monitoring and controlling;                      P6: Lack of project members’ commitment for the project plan.</p>
<p><i>Team organization</i> features relate to the state of the projects; e.g. the fundamental skills or experience and morale of project members:</p> <p>O1: Lack of skills and experience;                      O2 ]: Insufficient allocation of resources;                      O3 ]: Low morale.</p>
<p><i>Project management</i> factors about management activities:</p> <p>M1: Project manager lack of resource management throughout a project;                      M2: Inadequate project monitoring and controlling;                      M3: Lack of data needed to keep objective track of a project.</p>

Fig. 2. Explanation of the features seen in Figure 1.

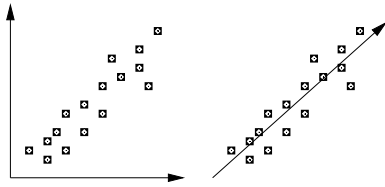


Fig. 3. The two features in the left plot can be transferred to the right plot via one latent feature.

### 3.1 Learning Latent Features

Numerous data mining methods check if the available features can be combined in useful ways. In this way, *latent* features within a data set can be discovered.

For example, principal components analysis (PCA) [8] has been widely applied to resolve problems with structural code measurements; e.g. [9]. PCA identifies the distinct orthogonal sources of variation in a data sets, while mapping the raw features onto a set of uncorrelated features that represent essentially the same information contained in the original data. For example, the data shown in two dimensions of Figure 3 (left-hand-side) could be approximated in a single latent feature (right-hand-side).

Since PCA combines many features into fewer latent features, the structure of PCA-based models may be very simple. For example, previously [10], we have used PCA and a decision tree learner to find the following predictor for defective software modules:

```

if  $domain_1 \leq 0.180$ 
then NoDefects
else if  $domain_1 > 0.180$ 
then if  $domain_1 \leq 0.371$  then NoDefects
     else if  $domain_1 > 0.371$  then Defects

```

Here, “ $domain_1$ ” is one of the latent features found by PCA. This tree seems very simple, yet is very hard to explain to business clients users since “ $domain_1$ ” is calculated using a very complex weighted sum (in this sum,  $v(g)$ ,  $ev(g)$ ,  $iv(g)$  are McCabe or Halstead static code metrics [11], [12] or variants on line counts):

$$\begin{aligned}
domain_1 = & 0.241 * loc + 0.236 * v(g) \\
& + 0.222 * ev(g) + 0.236 * iv(g) + 0.241 * n \\
& + 0.238 * v - 0.086 * l + 0.199 * d \\
& + 0.216 * i + 0.225 * e + 0.236 * b + 0.221 * t \\
& + 0.241 * LOCcode + 0.179 * LOComment \\
& + 0.221 * LOBlank + 0.158 * LOCcodeAndComment \\
& + 0.163 * uniqOp + 0.234 * uniqOpnd \\
& + 0.241 * totalOp + 0.241 * totalOpnd \\
& + 0.236 * branchCount
\end{aligned} \quad (2)$$

As we shall see below, other learners can yield effective models that are simpler to explain without using complex latent features.

### 3.2 Ensemble Learning

Data mining for SE means summarizing the complex behavior of a group of developers struggling to build intricate artifacts. Data mining over such complex multi-dimensional data often requires *fusing* together the results from multiple learners [13]. Such ensembles may *perform* well but, as we shall see, are hard to *explain*.

In *basic ensemble method* (BEM),  $l$  learners are run on various subsets of the available data. These learners use some features  $r$  to predict for classes  $s$  to find a function  $x_j(r, s)$  that returns the probability of the target classes  $s$ . BEM returns the mean probability:

$$\hat{x}_{BEM} = \frac{1}{l} \sum_{j=1}^l x_j(r, s) \quad (3)$$

The *linear generalized ensemble method* (GEM) returns a weighted sum of the conclusions of each learner  $x$  in the ensemble.

$$\hat{x}(\alpha)_{GEM} = \sum_{j=1}^l \alpha_j x_j(r, s) \quad (4)$$

where  $\alpha_j$  is the normalized performance score of  $x_j$  on the training data (so learners that performed the worst, contribute the least).

For some data sets, the combination rule is non-linear and complex. For example, Toh et.al. [13]’s variant of Equation 4 uses a Jacobian matrix for  $\hat{x}$  with different coefficients for each feature  $r_i \in r$  and target class  $s_m \in s$ . These coefficients are learned via multivariate polynomial regression. Toh et.al. report that their resulting ensemble *performs* better than simpler schemes. However, it may be harder to *explain* the ensemble since that explanation must cover:

- The learning methods used to generate  $x_j$ ;
- The combination rule that computes  $\hat{x}$ ; and
- The regression method used to tune the coefficients used in the combination method.

Such an explanation is not required if the users are willing to accept the conclusions of the learner, without explanation. However, for data sets as small Figure 1, it seems reasonable to expect that a simple explanation of runaway projects should be possible. Also, if managers are to use the results of the learner as part of their deliberations, they need some succinct structures that they can reflect over.

### 3.3 Naïve Bayes Classifiers

It is hardly surprising that complex latent features (e.g. Equation 2) or intricate combinations of multiple learners (e.g. Equation 4) are hard to explain. What is surprising is how hard it is to explain the results of even a single, supposedly simple, learner. For example, this section offers a complete description of how a Naïve Bayes classifiers makes its conclusions. The reader is asked to consider how many users would understand this description (in our experience, we have yet to meet a single one).

A Naïve Bayes classifier [14] is based on Bayes’ Theorem. Informally, the theorem says  $next = old * new$  i.e. what we’ll believe  $next$  comes from how  $new$  evidence effects  $old$  beliefs. More formally:

$$P(H|E) = \frac{P(H)}{P(E)} \prod_i P(E_i|H) \quad (5)$$

i.e. given fragments of evidence  $E_i$  and a prior probability for a class  $P(H)$ , the theorem lets us calculate a posterior probability  $P(H|E)$ .

When building predictors for runaways, the posterior probability of each hypothesis class ( $H \in \{\text{“ok” or “runaway”}\}$ ) is calculated, given the features extracted from a project such “ambiguous requirements” or “low morale” or any other of the features shown in Figure 2. The classification is the hypothesis  $H$  with the highest posterior  $P(H|E)$ .

Naïve Bayes classifiers are called “naïve” since they assume independence of each feature. While this assumption simplifies the implementation (frequency counts are required only for each feature), it is possible that correlated events are missed by this “naïve” approach. Domingos and Pazzani show theoretically that the independence assumption is a problem in a vanishingly small percent of cases [15]. This explains the repeated empirical result that, on average, Naïve Bayes classifiers perform as well as other seemingly more sophisticated schemes (e.g. see Table 1 in [15]).

Equation 5 offers a simple method for handling missing values. Generating a posterior probability means of tuning a prior probability to new evidence. If that evidence is missing, then no tuning is needed. In this case Equation 5 sets  $P(E_i|H) = 1$  which, in effect, makes no change to  $P(H)$ .

When estimating the prior probability of hypothesis  $H$ , it is common practice [16] to use an  $M$ -estimate as follows. Given that the total number of hypothesis is  $C$ , the total number of training instances is  $I$ , and  $N(H)$  is the frequency the hypothesis  $H$  within  $I$ , then

$$P(H) = \frac{N(H) + m}{I + m \cdot C} \quad (6)$$

Here  $m$  is a small non-zero constant (often,  $m = 1$ ). Three special cases of Equation 6 are:

- For high frequency hypothesis in large training sets,  $N(H)$  and  $I$  are much larger than  $m$  and  $m \cdot C$ , so Equation 6 simplifies to  $P(H) = \frac{N(H)}{I}$ , as one might expect.
- For low frequency classes in large training sets,  $N(H)$  is small,  $I$  is large, and the prior probability for a rare class is never less than  $\frac{1}{I}$ ; i.e. the inverse of the number of instances. If this were not true, rare classes would never appear in predictions.
- For very small data sets,  $I$  is small and  $N(H)$  is even smaller. In this case, Equation 6 approaches the inverse of the number of classes; i.e.  $\frac{1}{C}$ . This is a useful approximation when learning from very small data sets when all the data relating to a certain class has not yet been seen.

The prior probability calculated in Equation 6 is a useful lower bound for  $P(E_i|H)$ . If some value  $v$  is seen  $N(f = v|H)$  times in feature  $f$ 's observations for hypothesis  $H$ , then

$$P(E_i|H) = \frac{N(f = v|H) + l \cdot P(H)}{N(H) + l} \quad (7)$$

Here,  $l$  is the  $L$ -estimate and is set to a small constant (Yang & Webb [16] recommend  $l = 2$ ). Two special cases of are:

- A common situation is when there are many examples of an hypothesis and numerous observations have been made for a particular value. In that situation,  $N(H)$  and  $N(f = v|H)$  are large and Equation 7 approaches  $\frac{N(f=v|H)}{N(H)}$ , as one might expect.
- In the case of very little evidence for a rare hypothesis,  $N(f = v|H)$  and  $N(H)$  are small and Equation 7 approaches  $\frac{l \cdot P(H)}{l}$ ; i.e. the default frequency of an observation in a hypothesis is a fraction of the probability of that hypothesis. This is a useful approximation when very little data is available.

For numeric features it is common practice for N ave Bayes classifiers to use the Gaussian probability density function [17]:

$$g(x) = \frac{1}{\sqrt{2\pi}\sigma} e^{-\frac{(x-\mu)^2}{2\sigma^2}} \quad (8)$$

where  $\{\mu, \sigma\}$  are the feature's {mean, standard deviation}, respectively. To be precise, the probability of a continuous feature having exactly the value  $x$  is zero, but the probability that it lies within a small region, say  $x \pm \epsilon/2$ , is  $\epsilon \times g(x)$ . Since  $\epsilon$  is a constant that weighs across all possibilities, it cancels out and needs not be computed.

N ave Bayes classifiers are frustrating tools in the data mining arsenal. They exhibit excellent performance, but offer few clues about the structure of their models. The means and standard deviations for Figure 1 are shown in Figure 4. Note that this figure is an incomplete characterization of Figure 1. For example, row 1 of Figure 4 suggests that  $r1$  ("ambiguous requirements") for "ok" is a Gaussian distribution with a mean of 0.27 and a standard deviation of 0.86. A visual inspection of column one values for "ok" projects in Figure 1 shows that this is not true:  $r1$  is usually zero except in two cases where it takes the value of three.

One method of handling non-Gaussians like  $P(r1 = X|ok)$  is Johns and Langley's *kernel estimation* technique [18]. This technique approximates a continuous distribution sampled by  $n$  observations  $\{ob_1, ob_2, \dots, ob_n\}$  as the sum of multiple Gaussians with means

feature	P(ok) = 0.68		P(runaway) = 0.32	
	mean	sd	mean	sd
r1	0.2727	0.8624	1.35	1.0500
r2	0.9545	1.0650	1.65	0.8078
r3	0.9545	1.1571	1.95	1.3500
r4	0.8864	1.0759	1.65	1.0500
r5	1.4091	1.2670	1.90	1.0440
e1	0.3182	0.6998	1.00	1.2649
e2	0.7273	1.0082	1.00	1.2649
e3	0.8182	1.0824	1.65	1.0500
e4	0.5455	0.9642	1.65	1.2460
e5	0.2727	0.7497	1.40	1.2806
p1	0.6818	0.9833	1.80	1.3077
p2	0.9545	0.8516	1.50	1.1619
p3	0.3409	0.7744	1.80	1.1225
p4	0.6818	0.9833	1.35	1.0500
p5	0.7500	0.9857	2.25	1.0062
p6	0.4545	0.7820	1.70	1.1874
o1	0.6818	1.0824	1.65	1.2460
o2	0.3636	0.7100	1.30	1.3454
o3	0.2273	0.5979	1.00	1.0000
m1	0.6136	0.9762	0.60	0.9950
m2	0.4773	0.8323	1.50	1.1619
m3	0.5455	0.9404	1.50	1.2845

Fig. 4. Means and standard deviations from Figure 1.

$\{ob_1, ob_2, \dots, ob_n\}$  and standard deviation  $\sigma = \frac{1}{\sqrt{n}}$ . In this approach, to create a highly skew distribution like  $P(r1 = X|ok)$ , multiple Gaussians would be added together at  $r1 = 0$ . Conclusions are made by asking all the Gaussians which class they believe is most likely.

### 3.4 N ave Bayes and Software Engineering

N ave Bayes classifiers are widely used in the SE literature for several reasons. N ave Bayes classifiers summarize the training data in one frequency table per class. Hence, they consume very little memory and can quickly modify their knowledge by incrementing the frequency count of feature ranges seen in new training examples. Also, many studies (e.g. [15], [19], [20]) report that N ave Bayes exhibit excellent performance compared to other learners.

For example, recently Menzies, Greenwald & Frank [21] have built predictors for software detectors using a N ave Bayes classifier and two explanation systems- the OneR rule learner and the J4.8 decision tree learner. In that study, the learner with the *worst* explanation power (N ave Bayes) had the *best* performance, by far. For the data sets explored by Menzies, Greenwald & Frank, the median advantage of N ave Bayes, the C4.5 decision tree learner [22], and the OneR rule learner [23] over the other learners was 52.4%, 0%, -16.7%, respectively (see Figure 5). On analysis, Menzies, Greenwald & Frank concluded that N ave Bayes worked so well because of the the product calculation of Equation 5. They reasoned as follows. Many static code features have similar information content. Hence, minor changes in how the training data was sampled yielded different "best" features for predicting defects. The best predictions come from mathematical methods like N ave Bayes that accumulate the signal from many code features (using Equation 5's product rule). Decision tree learners like C4.5 and rule learners like OneR, on the other hand, do not perform well in this domain since they assume hard and fast boundaries between what is defective and what is not.

In summary, when mining software engineering data, there are many reasons to start with a N ave Bayes classifier. Abe, Muzono, Takagi, et.al. [2] used such classifiers to extend their prior work on runaway software projects [1], [3]. However, this classifier was only a *performance system*. not an *explanation system*, so it could not offer insights into, say, how to best change a software project in order to avoid runaways. As shown above, N ave Bayes classifiers do not generate such succinct generalizations. This is a problem since what developers really want to know is what should be done to avoid runaway status.

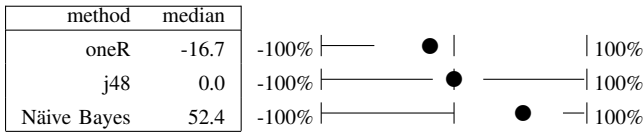


Fig. 5. Quartile charts from Menzies, Greenwald & Frank [21]. The charts show the *differences* when learners were applied to the same the training and test data. Performance was measured using *recall*; i.e. the percent of the defective modules found by the learners. The the upper and lower quartiles are marked with black lines. The median is marked with a black dot. Vertical bars are added to mark (i) the zero point and (ii) the minimum possible value and (iii) the maximum possible value. The median performance of Näive Bayes was much higher than the other methods.

### 3.5 Discussion of the Explanation Problem

As the mathematics gets more elaborate, it becomes harder to explain a Näive Bayes classifier to a typical business user:

- Many users are not trained mathematicians. Hence, they may be confused by Equation 5, Equation 6, Equation 7 and Equation 8.
- Presenting the internal statistics (e.g. Figure 4) is uninformative, at least for the business users we have worked with.
- The problem is compounded if the data is non-Gaussian (like Figure 1) since this requires explaining kernel estimation.
- Worse, a standard Näive Bayes classifier (with our without kernel estimation) can not answer business-level questions such as “what minimal changes should be make to most decrease the odds of runaway projects?”.

To be fair, Näive Bayes’ explanation problems are seen in other kinds of data miners:

- The problems with PCA and ensemble-based learners were discussed above.
- Tree learners such as C4.5 [22] or CART [24] execute in local top-down search, with no memory between different branches. Hence, the same concept can be needlessly repeated many times within the tree. Such trees can be cumbersome, needlessly large, and difficult to understand.
- Clustering algorithms [25] and nearest neighbor methods [26], [27] do not condense their working memory into succinct descriptions. Rather, inferences on new information are made by a query over all the old information.
- Simulated annealers [28] learn constraints to an input space that results in higher values in the output space. However, there is no generalization or summarization in a simulated annealer such as which subset of the input space is most important to control.
- Neural networks store their knowledge as weights distributed across a network. Concepts have no centralized location so it is impossible to inspect, say, all the information about one idea at one location in a network [29].

The problem of explaining the performance of these learners to end-users has been explored extensively in the literature (see the review in [30]). Often, some *post-processor* is used to convert an opaque model into a more understandable form:

- Towell and Shavlik generate refined rules from the internal data structures of a neural network [29].
- Quinlan implemented a post-processor to C4.5 called C45rules that generates succinct rules from cumbersome decision tree branches via (a) a greedy pruning algorithm followed by (b) duplicate removal then (c) exploring subsets of the rules relating to the same class [22].

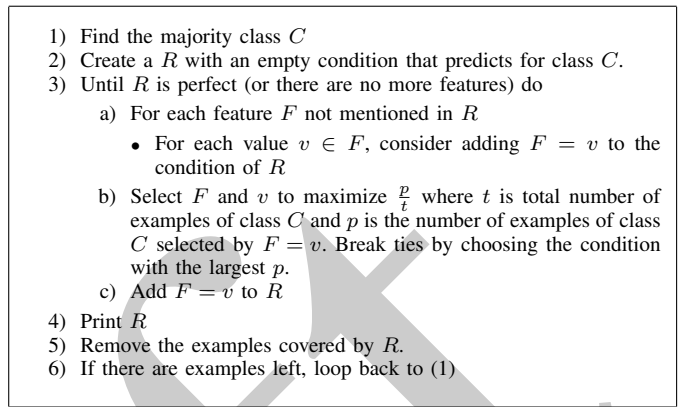


Fig. 6. PRISM pseudo-code.

- TARZAN was another post-processor to C4.5 that searched for the smallest number of decisions in decision tree branches that (a) pruned the most branches to undesired outcomes while (b) retaining branches leading to desired outcomes [31].

## 4 LEARNING METHODS

### 4.1 Rule Learners

RATHER than patch an opaque learner with a post-processor, it may be better to build learners that directly generate succinct high-level descriptions of a domain. For example, RIPPER [32] is one of the fastest *rule learners* known in the literature. The generated rules are of the form *condition*  $\rightarrow$  *conclusion*:

$$\underbrace{Feature_1 = Value_1 \wedge Feature_2 = Value_2 \wedge \dots}_{condition} \rightarrow \underbrace{Class}_{conclusion}$$

The rules generated by RIPPER perform as well as C45rules, yet are much smaller and easier to read [32].

Rule learners like RIPPER and PRISM [33] generate small, easier to understand, symbolic representations of the patterns in a data set. PRISM is a less sophisticated learner than RIPPER and is not widely used. It was initially added to this study to generate a lower bound on the possible performance. However, as we shall see, it proved surprisingly effective.

Like RIPPER, PRISM is a *covering* algorithm that runs over the data in multiple passes. As shown in the pseudo-code of Figure 6, PRISM learns one rule at each pass for the *majority class* (e.g. in Figure 1, at pass 1, the majority class is *ok*). All the examples that satisfy the condition are marked as *covered* and removed from the data set. PRISM then recurses on the remaining data.

The output of PRISM is an ordered *decision list* of rules where *rule<sub>j</sub>* is only tested if all conditions in *rule<sub>i<j</sub>* fail. PRISM returns the conclusion of the first rule with a satisfied condition.

One way to visualize a covering algorithm is to imagine the data as a table on a piece of paper. If there exists a clear pattern between the features and the class, define that pattern as a rule and cross out all the rows covered by that rule. As covering recursively explores the remaining data, it keeps splitting the data into:

- what is easiest to explain, and
- any remaining ambiguity that requires a more detailed analysis.

PRISM is a naïve covering algorithm and has problems with *residuals* and *over-fitting*. If there are rows with similar patterns and similar frequencies occur in different classes, then:

- these *residual* rows are the *last* to be removed for each class;
- so the *same* rule can be generated for *different* classes.

In *over-fitting*, a learner fixates on spurious signals that do not predict for the target class. PRISM's over-fitting arises from part 3.a of Figure 6 where the algorithm loops through all features. If some feature is poorly measured, it might be noisy (contains spurious signals). Ideally, a rule learner knows how to skip over noisy features.

RIPPER addresses residuals and over-fitting problem three techniques: *pruning*, *description length* and *rule-set optimization* For a full description of these techniques, see [34]. In summary:

- *Pruning*: After building a *rule*, RIPPER performs a back-select to see what parts of a *condition* can be deleted, without degrading the performance of the rule. Similarly, after building a *set of rules*, RIPPER performs a back-select to see what *rules* can be deleted, without degrading the performance of the rule set. These back-selects remove features/rules that add little to the overall performance. For example, back pruning could remove the residual rules.
- *Description length*: The learned rules are built while minimizing their *description length*. This is an information theoretic measure computed from the size of the learned rules, as well as the rule errors. If a rule set is over-fitted, the error rate increases, the description length grows, and RIPPER applies a rule set pruning operator.
- *Rule set optimization* tries replacing rules straw-man alternatives (i.e. rules grown very quickly by some naïve method).

### 4.2 Performance Measures

Our results are presented in terms of the following performance measures. Suppose we have some historical log, like Figure 1 that can comment on the correct classification of each row. By comparing the historical log with the output of the learner, we can define several measures of success. Let  $\{A, B, C, D\}$  denote the true negatives, false negatives, false positives, and true positives (respectively) found by a binary detector (binary detectors work on data sets with two classes, like Figure 1).  $A, B, C, D$  can be combined in many ways. For example, accuracy (or *acc*) is the percentage of true positives ( $D$ ) and negatives ( $A$ ) found by the detector.

$$acc = accuracy = (A + D)/(A + B + C + D) \quad (9)$$

Also, recall (or *pd*) comments on how much of the target was found.

$$pd = recall = D/(B + D) \quad (10)$$

Precision (or *prec*) comments on how many of the instances that triggered the detector actually containing the target concept.

$$prec = precision = D/(D + C) \quad (11)$$

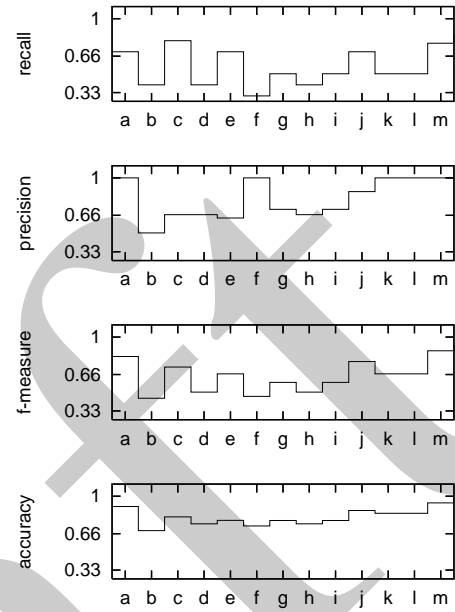
The *f*-measure is the harmonic mean of precision and recall. It has the property that if *either* precision or recall is low, then the *f*-measure is decreased. The *f* measure is useful for dual assessments that include *both* precision and recall.

$$f\text{-measure} = \frac{2 \cdot prec \cdot pd}{prec + pd} \quad (12)$$

All these measures fall in the range  $0 \leq \{pd, prec, f, acc\} \leq 1$ . Also, the *larger* these values, the better the model.

### 4.3 Experiments With the Learning Methods

VARIOUS combinations of the learning method described above were applied to Figure 1. The results are shown in Figure 7. In all 13 *treatments* where applied to Figure 1. Each *treatment* is some combination of a data filter, a learner, and a assessment method. This



	bins	learner	#features	#tests
a	n/a	ripper	22	1
b	n/a	ripper	22	10
c	n/a	nb	22	10
d	3	ripper	22	10
e	3	nb	22	10
f	3	prism	22	10
g	3	ripper	1 (r1)	10
h	3	ripper	2 (r1 + p5)	10
i	3	bayes	1 (r1)	10
j	3	bayes	2 (r1 + p5)	10
k	3	prism	1 (r1)	10
l	3	prism	2 (r1 + p6)	10
m	3	prism	3 (r1 + p6 + o3)	10

Fig. 7. Results from this study. The four plots, shown at top, come from the 13 treatments shown at bottom.

section discusses how each treatment was designed using results from the proceeding treatments.

Before moving on, we call attention to the *accuracy* results of Figure 7. Observe how *accuracy* can be an remarkably insensitive performance measure; i.e. it remained roughly constant, despite large changes in recall and precision. This result has been seen in many other data sets [21], [35]. Hence, accuracy is deprecated by this paper.

#### 4.3.1 Cross-Validation

Treatment *a* is a simple application of RIPPER to Figure 1. The learned theory was applied back on the training data used to generate it; i.e. all of Figure 1. As shown in Figure 7, this produced one of the largest *f*-measures seen in this study.

Treatment *a* assessed a learned model using the data that generated it. Such a *self-test* can lead to an over-estimate of the value of that model. Cross-validation, on the other hand, assesses a learned model using data *not* used to generate it. The data is divided into, say, 10 buckets. Each bucket is set aside as a test set and a model is learned from the remaining data. This learned model is then assessed using the test set. Such cross-validation studies are the preferred evaluation method when the goal is to produce predictors intended to predict future events [17].

In treatment *b*, a cross-validation experiment was applied to the data. The treatment *b* results shows how badly treatment *a* over-estimated the performance: changing the training data by as little as

feature	range	frequency	
o3	0	24	████████████████████
	1	1	█
	2,3	7	██████
p6	0	19	██████████████████
	1,2	10	██████████
	3	3	███
r1	0,1	23	██████████████████
	2	5	█████
	3	4	████

Fig. 8. Some 3bin results from Figure 1.

10% nearly halved the precision and recall. Clearly, the conclusions from the self-test from this data set are *brittle*; i.e. unduly altered by minor changes in the training data.

Treatment *c* illustrates the explanation vs performance trade-off discussed in the introduction. As mentioned above, the output from rule learners can be far easier to explain than the output of treatment *c*; i.e. a Naïve Bayes classifier (with kernel estimation) running on data sets with non-Gaussian distributions like Figure 1. So, if optimizing for *explainability*, an analyst might favor rule learners over Bayes classifiers. On the other hand, Figure 7 shows treatment *c* out-performing treatment *b*, especially in terms of recall. So, if optimizing for *performance* an analyst might favor a Bayes classifier.

Note that treatment *c* uses the method favored by the previous high water mark in this research [2]. In the sequel, we show how this study found data mining methods that significantly out-perform that prior work.

### 4.3.2 Discretization

Treatments *d, e* and *f* explore *discretization*. Discretization clumps together observations taken over a continuous range into a small number of regions. Humans often discretize real world data. For example, parents often share tips for “toddlers”; i.e. humans found between the breaks of *age* = 1 and *age* = 3. Many researchers report that discretization improves the performance of a learner since it gives a learner a smaller space to reason about, with more examples in each part of the space [16], [20], [36], [37].

After Gama and Pinto [38], we say that discretization is the process of converting a continuous range into a histogram with *k* break points  $b_1 \dots b_k$  where  $(\forall i < j : b_i \leq b_j)$ . The histogram divides a continuous range into bins (one for each break) and many observations from the range may fall between two break points  $b_i$  and  $b_{i+1}$  at frequency counts  $c_i$ .

Simple discretizers are *unsupervised* methods that build their histograms without exploiting information about the target class; e.g.

- *equal width*:  $(\forall i, j : (b_i - b_{i-1}) = (b_j - b_{j-1}))$ ;
- *equal frequency*:  $(\forall i, j : (c_i = c_j))$ . For Naïve Bayes classifiers working on *n* instances, Yang & Webb [16] advocate equal frequency with  $c_i = c_j = \sqrt{n}$ .

For example, Figure 1 holds 32 instances so a  $b = 3$  equal frequency discretion hopes to place  $\frac{32}{3} \approx 10$  values into each part of the histogram. However, Figure 1 does not have ten instances for each feature value so, as shown in Figure 8, a skewed histogram is generated.

More sophisticated discretizers are *supervised* methods that build their histograms using knowledge of the target class. Specifically, the continuous range is explored looking for a break that is a *cliff*; i.e. a point where the class frequencies are most different above and below the cliff. Once a top-level *cliff* is found, this method usually recurses into each region above and below the *cliff* to find the next best sub-cliff, sub-sub-cliff, and so on.

For example, the Fayyad & Irani [37] supervised discretizer assumes that the best cliff is the one that most divides target classes.

In terms of information theory, this can be measured using *entropy*; i.e. the number of bits required to encode the class distribution. If the classes in a sample of *n* instances occur at frequencies counts  $c_1, c_2, \dots$ , then the entropy of that sample is

$$Ent(c_1, c_2, \dots) = -\frac{c_1}{n} \cdot \log_2 \left( \frac{c_1}{n} \right) - \frac{c_2}{n} \cdot \log_2 \left( \frac{c_2}{n} \right) - \dots$$

If a break divides *n* numbers into two regions of size  $n_1, n_2$ , then the best cliff is the one that minimizes the sum of the entropy below and above the cliff; i.e.  $\frac{n_1}{n} \cdot Ent_1 + \frac{n_2}{n} \cdot Ent_2$ .

Various discretizers were explored, with disappointing results:

- Yang & Webb’s rule ( $c_i = \sqrt{n} = \sqrt{33} \approx 6$ ) was not useful here since our data has less than 6 distinct values per feature.
- Fayyad&Irani’s method reduced most features to a single bin; i.e. it found no information gain in any parts of our ranges.
- Best results were seen with a simple 3bin equal frequency scheme (i.e.  $|b| = 3$ ) in Treatment *f* where PRISM achieved precisions as high as the RIPPER self-test (treatment *a*). However, the same experiment saw the worst recall.
- The same 3bin scheme offered little help to RIPPER or Naïve Bayes (see treatments *d, e*).

Since the precision results were the most promising seen to date, 3bin was retained for the rest of our experiments. Other methods were then employed to achieve the benefits of 3bin (high precision) without its associated costs (low recall).

### 4.3.3 Feature Subset Selection

The remaining treatments (*g, h, i, j, k, l, m*) explore how different *feature subsets* change the performance of the learning. A repeated result in the data mining community is that simpler models with equivalent or higher performance can be built via *feature subset selection* algorithms that intelligently prune useless features [19]. Features may be pruned for several reasons:

- They may be noisy; i.e. contain spurious signals unrelated to the target class;
- They may be uninformative; e.g. contain mostly one value, or no repeating values;
- They may be correlated to other variables- in which case, they can be pruned since their signal is also present in other variables.

The reduced feature set has many advantages:

- Miller has shown that models generally containing fewer variables have less variance in their outputs [39].
- The smaller the model, the fewer are the demands on interfaces (sensors and actuators) to the external environment. Hence, systems designed around small models are easier to use (less to do) and cheaper to build.
- In terms of this article, the most important aspect of learning from a reduced features set is that it produces smaller models. Such smaller models are easier to explain (or audit).

One such feature subset selector is Kohavi & Johns’ WRAPPER algorithm [40]. Starting with the empty set, WRAPPER adds some combinations of features and asks some target learner to build a model using just those features. WRAPPER then grows the set of selected features and checks if a better model comes from learning over the larger set of features.

If we applied WRAPPER to our three learners (RIPPER, PRISM, Naïve Bayes), then WRAPPER’s search through the 22 features of Figure 1 could require  $3 \cdot 2^{22} = 12,582,912$  calls to a learner. In practice, a heuristic search drastically reduce this search space. WRAPPER stops when there are no more features to select, or there has been no significant improvement in the learned model for the last five additions (in which case, those last five additions are deleted).

feature	Learner			average	
	PRISM	Näive Bayes	RIPPER		
group #1 usually selected	r1 o3 p5	10 7	10 8	6 4	8.7 7 6
group #2: sometimes selected	p6 m3 r2 p2 e1 o2	8	1 3	2 2 1 1 1	4.5 3 2 1.5 1.3 1.3
group #3: rarely selected	e2 e3 m2 o1 p1 p3 p4 r3	1 1	1	1 1 1 1 1 1 1	1 1 1 1 1 1 1
group #4: never selected	r4 r5 e4 e5 m1				

Fig. 9. Number of times WRAPPER selected features in ten experiments on 90% samples of the data.

```

1: If o3 = 1 then ok
2: If r1 = 0,1 and p6 = 1 then ok
3: If r1 = 3 and p6 = 1,2 then ok
4: If r1 = 0,1 and p6 = 1,2 and o3 = 0 then ok
5: If r1 = 1,2 then runaway
6: If p6 = 3 then runaway
7: If r1 = 3 and p6 = 0 then runaway
8: If r1 = 0,1 and p6 = 1,2 and o3 = 2,3 then runaway
9: If r1 = 0,1 and p6 = 1,2 and o3 = 0 then runaway
    
```

Fig. 10. Rules generated by treatment *m*.

Technically speaking, this is a hill-climbing forward select search with a “stale” parameter set to 5. For data sets as small as Figure 1, WRAPPER terminates in a under a minute (but for large data sets, other feature selectors would be required- see [19] for a survey).

Figure 9 shows the results of running 10 WRAPPER experiments on Figure 1 (discretized via *3bin*) for our three learners. In each experiment, 10% of Figure 1 (selected at random) was ignored:

- Group #1 shows the features that, on average, were selected in the majority of ten runs (on average, 6 times or more).
- Group #2 shows the features that were selected 2 to 5 times.
- Group #3 shows the features that were selected only once.
- Group #4 shows the features that were never selected.

There are only three features in Group #1 suggesting that many of the Figure 1 features could be ignored. This has implications for the cost of data collection and the explaining runaway projects:

- Data collection could be constrained to just Group #1, and perhaps *p6* (which PRISM selected eight times). Such a constrained data collection program would be cheaper to conduct, especially over a large organization.
- Figure 10 shows a rule predicting runaway projects found by PRISM using just the features recommend by WRAPPER (*r1, p6, o3*) on *3bin* discretized data. The figure shows that just using the top-ranked features of Figure 9 yields a very succinct, easy to explain model.

Treatments *g, h, ...m* show the results of applying the top-ranked features to the discretized data. For each learner, if WRAPPER usually selected *N* features, then that learner was tested in a 10-way cross-validation using the top ranked feature, the second-top ranked features, and so on up to using *N* features.

#### 4.3.4 Best Results

The best results were obtained in treatment *m*. That treatment applied PRISM using the three features usually selected by WRAPPER+PRISM: *r1, o3, p6*. This resulted in Figure 10.

Figure 8 showed  $r1 \in \{0, 1\}$ ,  $p6 \in \{1, 2\}$ ,  $o3 = 0$  is a frequent pattern in our data. Hence, after a covering algorithm removes all other more interesting structures, the residual rows can contain this frequent pattern. This, in turn, means that identical rules could be generated for different classes; e.g. rules 4&9 of Figure 10 (this is the *residual rule* problem discussed above).

It is important to read these rules top to bottom since a rule fires *only* if *all* the rules above it *fail*. In practice, this means that the residual rule 9 is never used (it is blocked by rule 4).

A 10-way cross-validation study showed that this rule generation method yields an average precision, recall, and *f*-measure across the 10-way of 1,0.85, and 0.92 (respectively). This result is actually much better than it appears. To achieve average precisions and recalls of 1 and 0.85 in such a 10-way is something of an accomplishment. In a 10-way cross-validation on the 33 records of Figure 1, the test set is of size three or four. In such a small test set, a single outlier project can have a large and detrimental result on the collected statistics.

#### 4.3.5 User Studies

To test the *explainability* of Figure 10, we ran a session with eight software engineers managing large software verification projects.

Pseudocode for Näive Bayes (with kernel estimation) and PRISM (Figure 6) was introduced. PRISM was summarized this way: “each rule handles some examples, which are then removed, and the algorithm repeats on the remaining data”.

Within an hour, the engineers were hand-simulating PRISM. Using a pen and ruler, all the rows of Figure 1 that matched rule #1 (in Figure 10) were identified and crossed off. The rows that matched rule #2 were identified, then crossed off. The engineers stopped after simulating PRISM’s activities on two or three rules, making comments like “I see what is going on- the learner is finding and handling the most obvious next thing.” Significantly, none of the engineers tried to apply Näive Bayes; i.e. *m*-estimates, *l*-estimates, the  $\epsilon$  approximation, and the Gaussians of kernel estimation.

In summary, the simplicity of PRISM the rules of Figure 10 allowed them to be explained to one focus group, all within a one hour session.

## 5 DISCUSSION

### 5.1 Related Work

THIS research aims at producing a precise, explainable, operational definition of a runaway project. Other work in this area is less precise and not operational.

For example, in 1997, Glass [4] had informally sampled several high-profile software disasters and found the following features to be predictive for runaways:

- Project objectives not fully specified (in 51% of the sample);
- Bad planning and estimating (48%);
- Technology new to the organization (45%);
- Inadequate/no project management methodology (42%);
- Insufficient senior staff on the team (42%);
- Poor performance by suppliers of hardware/software (42%)
- Other- performance (efficiency) problems (42%)

Glass did not offer an clear operational method for combining their features into an effective predictor. Other work carefully documented the software risk problem, but did not offer automatic tool support:

- Jiang et.al. [6] studied 40 features collected from questionnaires posted to personnel with recent experience with an IS project.



Their study is an exemplary example of software engineering research: after clearly defined six hypotheses about software risk, they identify those hypotheses not supported by their data.

- Ropponen & Lyytinen [7, p110] studied self-reported data from 83 project managers and 1,110 projects to find 26 software risk components: six scheduling and timing risks; four system functionality risks; three subcontracting risks; four requirements management risks; four resource usage and performance risks; and five personnel management risks.

Both reports have the same limitations: their conclusions contain a somewhat ill-defined and manual procedure for managers to explore the above risks. For example, both reports list risks and their weighted contribution to total risk. However, no combination rule is offered on how to best combine evidence of *multiple* risks.

Another aspect that sets this work apart from other studies is *reproducibility*. Neither the Jiang et.al. nor Ropponen & Lyytinen [7, p110] studies are reproducible since they did not made their data available to other researchers. Reproducibility is an important methodological principle in other disciplines since it allows a community to confirm, refute, or even improve prior results. In our view, in the field of software engineering, there are all too few examples of reproduced, and extended, results<sup>1</sup>. This current report began when the second and third authors published their data [1] and defined a research challenge: how to better explain the results of their learning to developers [2]. We would strongly encourage software engineering researchers to share data, define challenges, and to take the time to rework the results of others.

## 5.2 External Validity

This study has produced:

- 1) A recommended *feature subset* for predicting runaways ( $r1, p6, o3$ );
- 2) A recommended *model* that combines those features (Figure 10); and
- 3) A recommended *method* for generating that subset and that model:
  - 3bin discretization;
  - a WRAPPER around PRISM;
  - 10-way cross-validation using PRISM on the subsets found by WRAPPER.

It is good practice to question the external validity of these recommendations.

WRAPPER selected different features than the manual method that produced Equation 1. That is, the recommended feature subset learned by our recommended method is different to that found by our earlier work. This raises a concern about external validity: why do our conclusions keep changing?

We endorse the conclusions of this study over our prior work [1] for two reasons. Firstly, this study explored far more feature subsets than before:

- Equation 1 was generated after a manual analysis of a few features.
- Figure 10 was generated after an automatic search through thousands of subsets.

Secondly, the results of this study perform better than our prior results:

- Equation 1 offers ambiguous conclusions in the range ( $0.03 < P(\text{runaway}|X) < 0.81$ ).

1. Exception: see the reports of the PROMISE workshop <http://promisedata.org/repository/papers.html>

- Figure 10 offers categorical conclusions about the runaway status of a project. Further, it does so with perfect precision.

A more serious validity threat comes from the data used in this study. Any inductive process suffers from a *sampling bias*; i.e. the conclusions of the study are a function of the data used in that study. In that regard, we have evidence that our results are stable across small to medium-sized changes to our project sample. In a 10-way cross-validation experiment, 10% of the data (in our case, 3 to 4 records) is set aside and the model is learned from the remaining information. Our learned model had an average precision of 1.0 in a 10-way; i.e. the precision of our model remained perfect, despite a 10% change in the training data.

Also, Figure 1 does not show all the data available to this study. Some of the data available to this research group is proprietary and cannot be generally released. In order to check the external validity of our methods, these ten extra records were not analyzed until *after* we reached the above conclusions regarding the recommended data mining method for this data. When our recommended method was applied to Figure 1, plus the extra ten records, WRAPPER still found the features shown in Figure 9. Further, the performance of the rule set learned from the extended data had the same properties as Figure 10; i.e.

- It out-performed NaïveBayes;
- It exhibited perfect precision (precision=1.0) over the 10-way cross-validation.

In summary, despite the data set size changing by a small to medium amount (-10% to +33%), there is:

- no instability in the recommended features;
- no instability in the performance of of the recommended model;
- no instability in the recommended method.

## 5.3 Method Selection for Quirky Data

Several times we found that certain widely regarded methods (RIPPER; discretization using Fayyad&Irani; discretization with Yang & Webb's  $\sqrt{n}$  rule) did not yield the best results for this data set. The reason for this is simple: software engineering data sets are often small:

- Figure 1 is one table with only 22\*33 cells;
- Elsewhere we have published results on even smaller data sets [41], [42].

It is hard to know *a priori* what are the quirks of small software engineering data sets. Hence, we recommend trying many methods, even supposedly out-dated ones. For example, in this study, a very simple rule-learner (PRISM) produced the best performance while being most understandable to our users.

More generally, Fayyad [43] argues persuasively that data mining should be viewed as a *small part* of the knowledge and data discovery (KDD) cycle shown in Figure 11. For example, in this report we used discretization and feature subset-selection for *pre-processing* and *selection* steps shown in Figure 11. Also, we looped through the KDD cycle 13 times: each time, the results from the previous round informed our work for the next round.

## 5.4 Data Mining Methods

Based on this work, and certain standard texts in the data mining field [17], [43], we offer the following advice to other researchers data mining on SE data.

It is important to understand the *goals* of the data mining task. If the learned model only needs to *perform*, and not *explain* then any data mining method might do ranging from

- Naïve Bayes classifiers

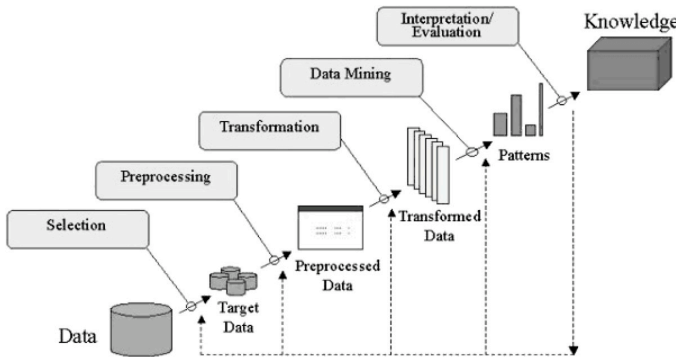


Fig. 11. The KDD (Knowledge Discovery in Databases) cycle, adapted from [43].

- to clustering algorithms, decision tree learners, neural nets, etc
- or, as explored in Equation 4, ensembles of the above.

The simplest of the above is N aive Bayes. Such classifiers scale to very large data sets and, in many domains, have performed very well [15], [19], [20]. Also, in at least one SE domain [21], they far out-performed other methods.

However, if the goal is to generate an *explainable* theory, then:

- Many business users do not have the background required to understand mathematical-based learners. For such users, the rule learners (e.g. RIPPER) may be most useful since they produce succinct summaries of the data.
- It is useful to reduce the range of number variables with discretization. Once reduced, the learned model can be simpler since it only needs to comment on a few discrete ranges rather than the entire number line.
- It is also useful to reduce the number of features with feature subset selection. A repeated result in the literature [19], [39], [40] is that the majority of the features can be pruned away and the resulting model is either simpler, performs better or both. For example, in this case study, the best performance and the most succinct/explainable model were found using just  $\frac{3}{22}$  of the available data.

As to the choice of feature subset selector:

- Hall and Holmes [19] compare WRAPPER to several other variable pruning methods including the principal component analysis (PCA) method used by Ropponen & Lyytinen and Munson [9] (amongst others). Feature selection methods can be grouped according to (a) whether or not they make special use of the target variable in the data set such as “runaway”; (b) whether or not pruning uses the target learner. PCA does not make special use of the target variable. Also, unlike other pruning methods, WRAPPER *does* use the target learner as part of its analysis. Hall and Holmes found that PCA was one of the worst performing methods (perhaps because it ignored the target variable) while WRAPPER was the best (since it can exploit its special knowledge of the target learner).
- For large data sets, WRAPPER can be too slow. When WRAPPER is not possible, see the conclusion of the Hall & Holmes study [19] for recommendations on two other feature subset selection methods.
- If the data set is small enough (e.g. Figure 1), use WRAPPER around a rule learner. WRAPPER is the slowest feature subset selector but it is the only one that can tune itself to the target learner.

Regarding performance measures, we have two recommendations:

- Comparing the *f*-measures in treatment *a* and *b* of Figure 7, it is clear that self-tests can over-estimate the value of a learned model. Hold-out sets are the recommended way to assess a learned model.
- Accuracy is a widely used measure for assessing a learned theory. Figure 7 shows that it can be remarkably uninformative. In that figure, large changes in precision and recall make very little impact on the accuracy. Hence, we strongly recommend against the use of accuracy.

The above issues are widely discussed in the data mining literature (e.g. [17], [43], [44], [45]). Nevertheless, our reading of the literature is that multiple traversals of the KDD cyclic application using a range of techniques (e.g. different learners, discretizers, and feature subset selectors) is quite rare. Often researchers take one learner, apply it once, then report the conclusion. Also, despite many positive empirical studies, feature selection is rarely seen in software engineering (exceptions: [21], [46]). Further, it is still standard practice for software engineers to present their data mining results in terms of accuracy of non-hold-out experiments (e.g. [47, p6]). We hope our results encourage a change in that standard practice.

## 6 CONCLUSION

INTUITIVELY, it seems reasonable that optimizing for performance can compromise explainability. Software engineering data can be complex, noisy, or confusing. Such complex data may require complex and arcane learning strategies; e.g. the defect data sets studied by Menzies, Greenwald, and Frank. Complex and arcane learning strategies will be hard to explain. That is, good performance in a learned model may imply poor explanatory power, especially for real world software engineering data.

This paper is a counter-argument to such pessimism. We show that at least for predicting runaway software projects, certain standard data mining methods resulted in models with both:

- High performance: i.e. precision=1.0; and
- good explainability: i.e. small rule sets, understandable by our users;

This result is a new high water mark in predicting runaway projects. This new predictor out-performs prior results in several ways:

- Our results are fully reproducible: the data for our analysis comes from Figure 1; the software used is freely available<sup>2</sup>.
- Prior work by other researchers [4], [6], [7] has carefully documented the influence of features on software risk, but did not offer an operational model (by “operational”, we mean that the model can generate performance statistics like Figure 7).
- As to our own prior results, the logistic regression method [1] required some manual intervention on the part of the analyst. In contrast to that, the techniques described here are automatic. Also, due to ambiguities in the middle *P* ranges of Equation 1, or the inner complexities of our N aive Bayes classifier [2], our prior mathematical results were much harder to explain than the new rules of Figure 10.
- Comparing treatment *c* and treatment *m* in Figure 7, we see that our new data mining method (treatment *m*: 3bin, WRAPPER, PRISM) has similar recall but much higher precision than our old data mining method (treatment *c*: N aiveBayes, from [2]).
- Measured in terms of precision, this new model is as good as can ever be expected for our data. Other combination data mining methods could out-perform our result (e.g. by generating a smaller, more explainable model with higher recall) but no

2. <http://www.cs.waikato.ac.nz/~ml/weka/>

other method could be more precise (since precision's maximum value is 1.0).

- Prior results conducted a manual exploration of a few subsets of the features [1]. Here, we employed a feature subset selector that explored thousands of feature subsets. Hence, we have far more confidence that the following factors are most useful in recognizing runaways: ambiguous requirements; low morale; lack of project members' commitment to the project plan.

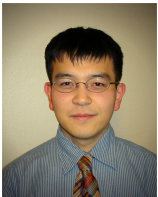
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