Impact of Mutation Operators on Mutant Equivalence

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Abstract: The presence of equivalent mutants is a recurrent source of aggravation in mutation-based studies of software testing, as it distorts our analysis and precludes assertive claims. But the determination of whether a mutant is equivalent to a base program is undecidable, and practical approaches are tedious, error-prone, and tend to produce insufficient or unnecessary conditions of equivalence. We argue that an attractive alternative to painstakingly identifying equivalent mutants is to estimate their number. This is an attractive alternative for two reasons: First, in most practical applications, it is not necessary to identify equivalent mutants individually; rather it suffices to know their number. Second, even when we need to identify equivalent mutants, knowing their number enables us to single them out with little to moderate effort.

1 EQUIVALENT MUTANTS

1.1 A Survey of Equivalent Mutants

The issue of equivalent mutants has mobilized the attention of researchers for a long time; mutation is used in software testing to analyze the effectiveness of test data or to simulate faults in programs, and is meaningful only to the extent that the mutants are semantically distinct from the base program (Jia and Harman, 2011; Just et al., 2014; Andrews et al., 2005; Namin and Kakarla, 2011). But in practice mutants may often be semantically undistinguishable from the base program while being syntactically distinct from it (Yao et al., 2014; Schuler and Zeller, 2012; Gruen et al., 2009; Just et al., 2013; Just et al., 2014; Wang et al., 2017; Papadakis et al., 2014).

Given a base program P and a mutant M, the problem of determining whether M is equivalent to P is known to be undecidable (Budd and Angluin, 1982). In the absence of a systematic/algorithmic procedure to determine equivalence, researchers have resorted to heuristic approaches. In (Offutt and Pan, 1997) Offutt and Pan argue that the problem of detecting equivalent mutants is a special case of a more general problem, called the feasible path problem; also they use a constraint-based technique to automatically detect equivalent mutants and infeasible paths. Experimentation with their tool shows that they can detect nearly half of the equivalent mutants on a small sample of base programs. Program slicing techniques are proposed in (Voas and McGraw, 1997) and subsequently used in (Harman et al., 2000; Hierons et al., 1999) as a means to assist in identifying equivalent mutants. In (Ellims et al., 2007), Ellims et al. propose to help identify potentially equivalent mutants by analyzing the execution profiles of the mutant and the base program. Howden (Howden, 1982) proposes to detect equivalent mutants by checking that a mutation preserves local states, and Schuler et al. (Schuler et al., 2009) propose to detect equivalent mutants by testing automatically generated invariant assertions produced by Daikon (Ernst et al., 2001); both the Howden approach and the Daikon approach rely on local conditions to determine equivalence, hence they are prone to generate sufficient but unnecessary conditions of equivalence; a program P and its mutant M may well have different local states but still produce the same overall behavior; the only way to generate necessary and sufficient conditions of equivalence between a base program and a mutant is to analyze the programs in full (vs analyze them locally).

1.2 Counting Equivalent Mutants

It is fair to argue that despite several years of
research, the problem of automatically and efficiently detecting equivalent mutants remains an open challenge. In this paper we are exploring a way to address this challenge, not by a painstaking analysis of individual mutants, but rather by estimating the number of equivalent mutants; more precisely, we are interested to estimate the ratio of equivalent mutants (abbr: REM) that a program is prone to generate, for a given mutant generation policy. This is an attractive alternative to current research, for two reasons:

- First because for most applications it is not necessary to identify equivalent mutants individually, but rather to estimate their number. If, for example, we generate 100 mutants of program P and we estimate that the ratio of equivalent mutants of P is 0.2 then we know that approximately 80 of these mutants are semantically distinct from P. Then we can assess the thoroughness of a test data set T by the ratio of mutants it kills over 80, not over 100.

- Second, even when we need to identify equivalent mutants, having an estimate of their number enables us to identify them to an arbitrary level of confidence with relatively little effort. If we have 100 mutants of program P and we estimate that 20 of them are equivalent to P, then we can use testing to kill as many of the 100 mutants as we can; with each killed mutant, the probability that the surviving mutants are equivalent to P increases.

### 1.3 Mutant Generation Policy

In order to estimate the number of equivalent mutants that a program P is prone to generate under a given mutant generation policy, we must analyze program P and the mutant generation policy.

- The impact that a program P has on the number of equivalent mutants generated for a given mutant generation policy is currently under investigation; we have already published evidence to the effect that the amount of redundancy in a program is an important factor that strongly affects the ratio of equivalent mutants generated from this (Marsit et al., 2017). To model the impact of a program on the ratio of equivalent mutants that it is prone to generate, we run an empirical experiment where we analyze relevant redundancy metrics of a sample set of programs, then apply a fixed mutant generator to each of these programs and observe the number of equivalent mutants that are generated for each. Using analytical and statistics-based empirical arguments, we show that the ratio of equivalent mutants has a significant correlation with the selected metrics; also, using the selected metrics as independent variables, we derive a regression model that estimates the ratio of equivalent mutants.

The regression model discussed above is valid for the mutation generation policy that we have used in the empirical experiment, but is not necessarily meaningful if a different mutation generation policy is used. Hence in order for our results to be of general use we need to understand and integrate the impact of the mutation generation policy on the ratio of equivalent mutants. The brute force approach to this problem is to select a set of common mutation generation policies and build a specific regression model for each. For the sake of generality and breadth of application, we propose an alternative approach whereby we analyze the impact of individual mutation operators on the ratio of equivalent mutants; then we investigate how the ratio of equivalent mutants produced by a combination of operators can be derived from those of the individual operators. The purpose of this paper is to explore what relation links the ratio of equivalent mutants obtained by a combination of operators to the ratios obtained by the individual operators. For the sake of simplicity, we first consider this problem in the context of two then three operators. The investigation of combinations of four operators or more is under way, at the time of writing.

In section 2 we discuss how to estimate the ratio of equivalent mutants of a base program P by quantifying several dimensions of redundancy of P under a fixed mutant generation policy. In section 3, we derive a regression model that enables us to estimate the REM of a program (dependent variable) from the redundancy metrics of the program (independent variables), which are derived by static analysis of the program’s source code; this regression model is derived empirically using a uniform mutant generation policy. In section 4 we discuss how to estimate the REM of a program under an arbitrary mutant generation policy, and design an experiment which may enable us to do so automatically; in section 5 we present the results of our experiment and analyze them, and in section 6 we summarize our findings and sketch our plans for future research.
2 A FIXED MUTANT GENERATION POLICY

The agenda of this paper is not to identify and isolate equivalent mutants, but instead to estimate their number. To estimate the number of equivalent mutants, we consider question RQ3 raised by Yao et al. in (Yao et al., 2014): What are the causes of mutant equivalence? Two main attributes may cause a mutant M to be equivalent to a base program P: the mutation operator(s) that are applied to P to obtain M, and P itself. In this section, we consider a fixed mutation generation policy, specifically that of the default operators of PiTest (http://pitest.org/), and we reformulate the question as: For a selected mutation generation policy, what attributes of a program P determine the REM of the program? Or, equivalently, what attributes of P make it prone to generate equivalent mutants?

To answer this question, consider that the attribute that makes a program prone to generate equivalent mutants is the exact same attribute that makes a program fault tolerant: indeed, a fault tolerant program is a program that continues to deliver correct behavior (by, e.g. maintaining equivalent behavior) despite the presence and sensitization of faults (introduced by, e.g. application of mutation operators). We know what feature causes a program to be fault tolerant: redundancy. Hence if only we could find a way to quantify the redundancy of a program, we could conceivably relate it to the ratio of equivalent mutants generated from that program. Because mutants that are found to be distinct from the base program are usually said to be killed, we may refer to the ratio of equivalent mutants as the survival rate of the program’s mutants, or simply as the program’s survival rate.

Because our measures of redundancy use Shannon’s entropy function (Shannon, 1948), we briefly introduce some definitions and notations related to this function, referring the interested reader to more detailed sources (Csiszar and Koerner, 2011). Given a random variable X that takes its values in a finite set which, for convenience we also designate by X, the entropy of X is the function denoted by H(X) and defined by:

\[
H(X) = -\sum_{x_i \in X} p(x_i) \log(p(x_i)),
\]

where \( p(x_i) \) is the probability of the event \( X = x_i \). Intuitively, this function measures (in bits) the uncertainty pertaining to the outcome of \( X \), and takes its maximum value \( H(X) = \log(N) \) when the probability distribution is uniform, where \( N \) is the cardinality of \( X \).

We let \( X \) and \( Y \) be the two random variables; the conditional entropy of \( X \) given \( Y \) is denoted by \( H(X|Y) \) and defined by:

\[
H(X|Y) = H(X,Y) - H(Y),
\]

where \( H(X,Y) \) is the joint entropy of the aggregate random variable \( (X,Y) \). The conditional entropy of \( X \) given \( Y \) reflects the uncertainty we have about the outcome of \( X \) if we know the outcome of \( Y \). All entropies (absolute and conditional) take non-negative values. Also, regardless of whether \( Y \) depends on \( X \) or not, the conditional entropy of \( X \) given \( Y \) is less than or equal to the entropy of \( X \) (the uncertainty on \( X \) can only decrease if we know \( Y \)). Hence for all \( X \) and \( Y \), we have the inequality:

\[
0 \leq \frac{H(X|Y)}{H(X)} \leq 1.0.
\]

3 A REGRESSION MODEL

The purpose of this section is to build a regression model that enables us to estimate the REM of a program using its redundancy metrics. To this effect, we review a number of redundancy metrics of a program, and for each metric, we discuss, in turn:

- How we define this metric.
- Why we feel that this metric has an impact on the REM.
- How we compute this metric in practice (by hand for now).

Because our ultimate goal is to derive a formula for the REM of the program as a function of its redundancy metrics, and because the REM is a fraction that ranges between 0 and 1, we resolve to let all our redundancy metrics be defined in such a way that they range between 0 and 1.

3.1 State Redundancy

**What is State Redundancy?** State redundancy is the gap between the declared state of the program and its actual state. Indeed, it is very common for programmers to declare much more space to store
their data than they actually need, not by any fault of theirs, but due to the limited vocabulary of programming languages. State redundancy arises whenever we declare a variable that has a broader range than the set of values we want to represent, and whenever we declare several variables that maintain functional dependencies between them.

**Definition: State Redundancy.** Let P be a program, let S be the random variable that takes values in its declared state space and σ be the random variable that takes values in its actual state space. The state redundancy of Program P is defined as:

$$H(S) - H(\sigma)$$

Typically, the declared state space of a program remains unchanged through the execution of the program, but the actual state space grows smaller and smaller as execution proceeds, because the program creates more and more dependencies between its variables with each assignment. Hence we define two versions of state redundancy: one pertaining to the initial state, and one pertaining to the final state.

$$SR_I = \frac{H(S) - H(\sigma_I)}{H(S)}$$

$$SR_F = \frac{H(S) - H(\sigma_F)}{H(S)}$$

where σ_I and σ_F are (respectively) the initial state and the final state of the program, and S is its declared state.

**Why is state redundancy correlated to the REM?**
State redundancy measures the volume of data bits that are accessible to the program (and its mutants) but are not part of the actual state space. Any assignment to/ modification of these extra bits of information does not alter the state of the program.

**How do we compute state redundancy?** We must compute the entropies of the declared state space ($H(S)$), the entropy of the actual initial state ($H(\sigma_I)$) and the entropy of the actual final state ($H(\sigma_F)$). For the entropy of the declared state, we simply add the entropies of the individual variable declarations, according to the following table (for Java):

<table>
<thead>
<tr>
<th>Data Type</th>
<th>Entropy (bits)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Boolean</td>
<td>1</td>
</tr>
<tr>
<td>Byte</td>
<td>8</td>
</tr>
<tr>
<td>Char, short</td>
<td>16</td>
</tr>
<tr>
<td>Int, float</td>
<td>32</td>
</tr>
<tr>
<td>Long, double</td>
<td>64</td>
</tr>
</tbody>
</table>

For the entropy of the initial state, we consider the state of the program variables once all the relevant data has been received (through read statements, or through parameter passing, etc.) and we look for any information we may have on the incoming data (range of some variables, relations between variables, assert statements specifying the precondition, etc.); the default option being the absence of any condition. For the entropy of the final state, we take into account all the dependencies that the program may create through its execution. As an illustration, we consider the following simple example:

```java
public void example(int x, int y)
{assert (1<=x && x<=128 && y>=0);
 long z = reader.nextInt(); // initial state
 Z = x+y;} // final state
```

We find:

- $H(S) = 32 + 32 + 64 = 128$ bits
- $H(\sigma_I) = 10 + 31 + 64 = 105$ bits
- $H(\sigma_F) = 10 + 31 = 41$ bits

Hence

$$SR_I = \frac{128 - 105}{128} = 0.18,$$

$$SR_F = \frac{128 - 41}{128} = 0.68.$$
arguments into the same image. A sorting routine applied to an array of size N, for example, maps N! different input arrays (corresponding to N! permutations of N distinct elements) onto a single output array (the sorted permutation of the elements). A natural way to define non-injectivity is to let it be the conditional entropy of the initial state given the final state: if we know the final state, how much uncertainty do we have about the initial state? Since we want all our metrics to be fractions between 0 and 1, we normalize this conditional entropy to the entropy of the initial state. Hence we write:

\[
 NI = \frac{H(\sigma_I | \sigma_F)}{H(\sigma_I)}.
\]

Since the final state is a function of the initial state, the numerator can be simplified as \(H(\sigma_I) - H(\sigma_F)\). Hence:

**Definition: Non Injectivity.** Let P be a program, and let \(\sigma_I\) and \(\sigma_F\) be the random variables that represent, respectively its initial state and final state. Then the non-injectivity of program P is denoted by NI and defined by:

\[
 NI = \frac{H(\sigma_I | \sigma_F)}{H(\sigma_I)},
\]

**Why is non-injectivity correlated to the REM?** Of course, non-injectivity is a great contributor to generating equivalent mutants, since it increases the odds that the state produced by the mutation be mapped to the same final state as the state produced by the base program.

**How do we compute non-injectivity?** We have already discussed how to compute the entropies of the initial state and final state of the program; these can be used readily to compute non-injectivity.

## 3.3 Functional Redundancy

**What is Functional Redundancy?** A program can be modeled as a function from initial states to final states, as we have done in sections 0 and 3.2 above, but can also be modeled as a function from an input space to an output space. To this effect we let X be the random variable that represents the aggregate of input data that the program receives, and Y the aggregate of output data that the program returns.

**Definition: Functional Redundancy.** Let P be a program, and let X be the random variable that ranges over the aggregate of input data received by P and Y the random variable that ranges over the aggregate of output data delivered by P. Then the functional redundancy of program P is denoted by FR and defined by:

\[
 FR = \frac{H(Y)}{H(X)}.
\]

**Why is Functional Redundancy Related to the REM?** Functional redundancy is actually an extension of non-injectivity, in the sense that it reflects not only how initial states are mapped to final states, but also how initial states are affected by input data and how final states are projected onto output data.

**How do we compute Functional Redundancy?** To compute the entropy of X, we analyze all the sources of input data into the program, including data that is passed in through parameter passing, global variables, read statements, etc. Unlike the calculation of the entropy of the initial state, the calculation of the entropy of X does not include internal variables, and does not capture initializations. To compute the entropy of Y, we analyze all the channels by which the program delivers output data, including data that is returned through parameters, written to output channels, or delivered through return statements.

## 3.4 Non Determinacy

**What is Non Determinacy?** In all the mutation research that we have surveyed, mutation equivalence is equated with equivalent behavior between a base program and a mutant; but we have not found a precise definition of what is meant by behavior, nor what is meant by equivalent behavior. We argue that the concept of equivalent behavior is not precisely defined: we consider the following three programs,

- **P1:** \{int x,y,z; z=x; x=y; y=z;\}
- **P2:** \{int x,y,z; z=y; y=x; x=z;\}
- **P3:** \{int x,y,z; x=x+y; y=x-y; x=x-y;\}

We ask the question: are these programs equivalent? The answer to this question depends on how we interpret the role of variables x, y, and z in these...
programs. If we interpret these as programs on the space defined by all three variables, then we find that they are distinct, since they assign different values to variable \( z \) (x for P1, y for P2, and z for P3). But if we consider that these are actually programs on the space defined by variables x and y, and that z is a mere auxiliary variable, then the three programs may be considered equivalent, since they all perform the same function (swap x and y) on their common space (formed by x, y).

Rather than making this a discussion about the space of the programs, we wish to turn it into a discussion about the test oracle that we are using to check equivalence between the programs (or in our case, between a base program and its mutants). In the example above, if we let \( x_P, y_P, z_P \) be the final values of x, y, z by the base program and \( x_M, y_M, z_M \) the final values of x, y, z by the mutant, then oracles we can check include:

\[ O_1:\{ \text{return } x_P==x_M \text{ && } y_P==y_M \text{ && } z_P==z_M; \} \]
\[ O_2:\{ \text{return } x_P==x_M \text{ && } y_P==y_M; \}

Oracle \( O_1 \) will find that P1, P2 and P3 are not equivalent, whereas oracle \( O_2 \) will find them equivalent. The difference between \( O_1 \) and \( O_2 \) is their degree of non-determinacy; this is the attribute we wish to quantify. To this effect, we let \( \sigma^P \) be the final state produced by the base program for a given input, and we let \( \sigma^M \) be the final state produced by a mutant for the same input. We view the oracle that tests for equivalence between the base program and the mutant as a binary relation between \( \sigma^P \) and \( \sigma^M \). We can quantify the non-determinacy of this relation by the conditional entropy \( H(\sigma^M|\sigma^P) \): Intuitively, this represents the amount of uncertainty (or the amount of latitude) we have about (or: we allow for) \( \sigma^M \) if we know \( \sigma^P \). Since we want our metric to be a fraction between 0 and 1, we divide it by the entropy of \( \sigma^M \). Hence the following definition.

**Definition: Non Determinacy.** Let O be the oracle that we use to test the equivalence between a base program P and a mutant M, and let \( \sigma^P \) and \( \sigma^M \) be, respectively, the random variables that represent the final states generated by P and M for a given initial state. The **non-determinacy** of oracle O is denoted by ND and defined by:

\[
ND = \frac{H(\sigma^M|\sigma^P)}{H(\sigma^M)}.
\]

**Why is Non Determinacy correlated with the REM?** Of course, the weaker the oracle of equivalence, the more mutants pass the equivalence test, the higher the survival rate.

**How do we compute non determinacy?** All equivalence oracles define equivalence relations on the space of the program, and \( H(\sigma^M|\sigma^P) \) represents the entropy of the resulting equivalence classes. As for \( H(\sigma^M) \), it represents the entropy of the whole space of the program.

### 3.5 Empirical Study: Experimental Conditions

In order to validate our conjecture, to the effect that the REM of a program P depends on the redundancy metrics of the program and the non-determinacy of the oracle that is used to determine equivalence, we consider a number of sample programs, compute their redundancy metrics then record the REM that they produce under controlled experimental conditions. Our hope is to reveal significant statistical relationships between the metrics (as independent variables) and the ratio of equivalent mutants (as a dependent variable).

We consider functions taken from the Apache Common Mathematics Library (http://apache.org/); each function comes with a test data file. The test data file includes not only the test data proper, but also a test oracle in the form of assert statements, one for each input datum. Our sample includes 19 programs. We use PITEST (http://pitest.org/), in conjunction with maven (http://maven.apache.org/) to generate mutants of each program and test them for possible equivalence with the base program. The mutation operators that we have chosen include the following:

- Increments_mutator.
- Void_method_call_mutator,
- Return_vals_mutator,
- Math_mutator,
- Negate_conditionals_mutator,
- Invert_negs_mutator,
- Conditionals_boundary_mutator.

When we run a mutant M on a test data set T and we find that its behavior is identical to that of the base program P, we may not conclude that M is
equivalent to P unless we have some assurance that T is sufficiently thorough. In practice, it is impossible to ascertain the thoroughness of T short of letting T be all the input space of the program, which is clearly impractical. As an alternative, we mandate that in all our experiments, line coverage of P and M through their execution on test data T equals or exceeds 90%. In order to analyze the impact of the non-determinacy of the equivalence oracle on the ratio of equivalent mutants, we revisit the source code of PITEST to control the oracle that it uses. As we discuss above, the test file that comes in the Apache Common Mathematics Library includes an oracle that takes the form of assert statements in Java (one for each test datum). These statements have the form:

\[ \text{Assert.assertEqual}(y_P, M(x)) \]

where x is the current test datum, y_P is the output delivered by the base program P for input x, and M(x) is the output delivered by mutant M for input x. For this oracle, we record the non-determinacy (ND) as being zero. To test the mutant for other oracles, we replace the clause \text{assertEqual}(y_P, M(x)) with \text{assertEquivalent}(y_P, M(x)) for various instances of equivalence relations.

### 3.6 Statistical Analysis: Regression

Since the dependent variable, the REM, is a proportion (number of equivalent mutants over the total number of generated mutants), we use a logistic linear model for the survival rate so that the response will be constrained to be between 0 and 1. More specifically, the logarithm of the odds of equivalence (\[ \frac{REM}{1-REM} \]) is a linear function of the predictors:

\[ \log \left( \frac{\text{REM}}{1-\text{REM}} \right) = \alpha + X\beta. \]

For any model M consisting of a set of the covariates X, we can obtain a residual deviance D(M) that provides an indication of the degree to which the response is unexplained by the model covariates. Hence, each model can be compared with the null model of no covariates to see if they are statistically different. Furthermore, any pair of nested models can be compared (using a chi-squared test). We fit the full model with all five covariates, which was found to be statistically significant, and then successively dropped a covariate, each time testing the smaller model (one covariate less) with the previous model. We continued until the smaller model was significantly different, i.e. worse than the previous model.

Using the procedure described above, we find that the final model contains the metrics FR, NI and ND, with coefficient estimates and standard errors given in the table below:

<table>
<thead>
<tr>
<th>Metric</th>
<th>Estimate</th>
<th>Standard Error</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>-2.765</td>
<td>0.246</td>
<td>&lt;&lt; 0.001</td>
</tr>
<tr>
<td>FR</td>
<td>0.459</td>
<td>0.268</td>
<td>0.086</td>
</tr>
<tr>
<td>NI</td>
<td>2.035</td>
<td>0.350</td>
<td>&lt;&lt; 0.001</td>
</tr>
<tr>
<td>ND</td>
<td>0.346</td>
<td>0.152</td>
<td>0.023</td>
</tr>
</tbody>
</table>

Hence, the model is

\[ \log \left( \frac{\text{REM}}{1-\text{REM}} \right) = -2.765 + 0.459FR + 2.035NI + 0.235ND. \]

The plot below shows the relative errors of the model estimates with respect to the actuals; virtually all the relative errors are within less than 0.1 of the actuals.

4 ARBITRARY MUTATION POLICY

4.1 Analyzing the Impact of Individual Operators

For all its interest, the regression model we present above applies only to the mutant generation policy.
that we used to build the model. This raises the question: how can we estimate the REM of a base program P under a different mutant generation policy? Because there are dozens of mutation operators in use by different researchers and practitioners, it is impossible to consider building a different model for each combination of operators. We could select a few sets of operators, that may have been the subject of focused research (Andrews et al., 2005; Just et al., 2014; Namin and Kakarla, 2011; Laurent et al., 2018) and select a specific model for each. While this may be interesting from a practical standpoint, it presents limited interest as a research matter, as it does not advance our understanding of how mutation operators interact with each other. What we are interested to understand is: if we know the REM’s of a program P under individual mutation operators \( o_{P_1}, o_{P_2}, \ldots, o_{P_n} \), can we estimate the REM of P if all of these operators are applied jointly?

Answering this question will enable us to produce a generic solution to the automated estimation of the REM of a program under an arbitrary mutant generation policy:

- We select a list of mutation operators of interest (e.g. the list suggested by Laurent et al (Laurent et al., 2018) or by Just et al. (Just et al., 2014), or their union).
- Develop a regression model (similar to the model we derived in section 3) based on each individual operator.
- Given a program P and a mutant generation policy defined by a set of operators, say \( o_{P_1}, o_{P_2}, \ldots, o_{P_n} \), we apply the regression models of the individual operators to compute the corresponding ratios of equivalent mutants, say \( \text{REM}_{P_1}, \text{REM}_{P_2}, \ldots, \text{REM}_{P_n} \).
- Combine the REM’s generated for the individual operators to generate the REM that stems from their simultaneous application.

### 4.2 Combining Operators

For the sake of simplicity, we first consider the problem above in the context of two operators, say \( o_{P_1}, o_{P_2} \). Let \( \text{REM}_{P_1}, \text{REM}_{P_2} \) be the REM’s obtained for program P under operators \( o_{P_1}, o_{P_2} \). We ponder the question: can we estimate the REM obtained for P when the two operators are applied jointly? To answer this question, we interpret the REM as the probability that a random mutant generated from P is equivalent to P. At first sight, it may be tempting to think of REM as the product of \( \text{REM}_1 \) and \( \text{REM}_2 \) on the grounds that in order for mutant \( M_{12} \) (obtained from P by applying operators \( o_{P_1}, o_{P_2} \)) to be equivalent with P, it suffices for \( M_1 \) to be equivalent to P (probability: \( \text{REM}_1 \)), and for \( M_{12} \) to be equivalent to \( M_1 \) (probability: \( \text{REM}_2 \)). This hypothesis yields the following formula of REM:

\[
\text{REM} = \text{REM}_1 \times \text{REM}_2.
\]

But we have strong doubts about this formula, for the following reasons:

- This formula assumes that the equivalence of P to \( M_1 \) and the equivalence of \( M_1 \) to \( M_{12} \) are independent events; but of course they are not. In fact we have shown in section 3 that the probability of equivalence is influenced to a considerable extent by the amount of redundancy in P.
- This formula ignores the possibility that mutation operators may interfere with each other; in particular, the effect of one operator may cancel (all of or some of) the effect of another.
- This formula assumes that the ratio of equivalent mutants of a program P decreases with the number of mutation operators; for example, if we have five operators that yield a REM of 0.1 each, then this formula yields a joint REM of \( 10^{-5} \).

For all these reasons, we expect \( \text{REM}_1 \times \text{REM}_2 \) to be a loose (remote) lower bound for REM.

Elaborating on the third item cited above, we argue that in fact, whenever we deploy a new mutation operator, we are likely to make the mutant more distinct from the original program, hence it is the probability of being distinct that we ought to compose, not the probability of being equivalent. This is captured in the following formula:

\[
(1 - \text{REM}) = (1 - \text{REM}_1)(1 - \text{REM}_2),
\]

which yields:

\[
\text{REM} = \text{REM}_1 + \text{REM}_2 - \text{REM}_1 \times \text{REM}_2.
\]

In the following section we run an experiment to test which formula of REM is borne out in practice.
4.3 Empirical Analysis

In order to evaluate the validity of our proposed models, we run the following experiment:

- We consider the sample of seventeen Java programs that we used to derive our model of section 3.
- We consider the sample of seven mutation operators that are listed in section 3.5.
- For each operator Op, for each program P, we run the mutant generator Op on program P, and test all the mutants for equivalence to P. By dividing the number of equivalent mutants by the total number of generated mutants, we obtain the REM of program P for mutation operator Op.
- For each mutation operator Op, we obtain a table that records the programs of our sample, and for each program we record the number of mutants and the number of equivalent mutants, whence the corresponding REM.
- For each pair of operators, say (Op1, Op2), we perform the same experiment as above, only activating two mutation operators rather than one. This yields a table where we record the programs, the number of mutants generated for each, and the number of equivalent mutants among these, from which we compute the corresponding REM.
- For each pair of operators, we build a table that shows, for each program P, the REM of P under each operator, the REM of P under the joint combination of the two operators, and the residuals that we get for the two tentative formulas:

  F1: \[ REM = REM_1 REM_2, \]
  F2: \[ REM = REM_1 + REM_2 - REM_1 REM_2. \]

  At the bottom of each such table, we compute the average and standard deviation of the residuals for formulas F1 and F2.
- We summarize all our results into a single table, which shows the average of residuals and the standard deviation of residuals for formulas F1 and F2 for each (of 21) combination of two operators. The following section presents the results of our experiments, and the preliminary conclusions that we may draw from them.

5 EMPIRICAL OBSERVATIONS

The final result of this analysis is given in Table 1. The first observation we can make from this table is that, as we expected, the expression \[ F1: REM = REM_1 REM_2 \] is indeed a lower bound for \[ REM \], since virtually all the average residuals (for all pairs of operators) are positive, with the exception of the pair (Op1, Op2), where the average residual is virtually zero. The second observation is that, as we expected, the expression \[ F2: REM = REM_1 + REM_2 - REM_1 REM_2 \] gives a much better approximation of the actual REM than the F1 expression; also, interestingly, the F2 expression hovers around the actual REM, with half of the estimates (11 rows) below the actuals and half above (10 rows). With the exception of one outlier (Op4, Op5), all residuals are less than 0.2 in absolute value, and two thirds (14 out of 21) are less than 0.1 in absolute value. The average (over all pairs of operators) of the absolute value of the average residual (over all programs) for formula F2 is 0.080.

6 CONCLUSION AND PROSPECTS

6.1 Summary

The presence of equivalent mutants is a constant source of aggravation in mutation testing, because equivalent mutants distort our analysis and introduce biases that prevent us from making assertive claims. This has given rise to much research aiming to identify equivalent mutants by analyzing their source code or their run-time behavior. Analyzing their source code usually provides sufficient but unnecessary conditions of equivalence (as it deals with proving locally equivalent behavior); and analyzing run-time behavior usually provides necessary but insufficient conditions of equivalence (just because two programs have comparable run-time behavior does not mean they are functionally equivalent). Also, static analysis of mutants is generally time-consuming and error-prone, and wholly impractical for large and complex programs and mutants.
Table 3: Residuals for Candidate Formulas.

<table>
<thead>
<tr>
<th>Operator Pairs</th>
<th>Residuals, F1</th>
<th>Residuals, F2</th>
<th>Abs(Residuals)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>average Std dev</td>
<td>average Std dev</td>
<td>F1 F2</td>
</tr>
<tr>
<td>Op1, op2</td>
<td>0.1242467 0.1884347</td>
<td>-0.0163621 0.0459150</td>
<td>0.1242467 0.0163621</td>
</tr>
<tr>
<td>Op1, op3</td>
<td>-0.0008928 0.0936731</td>
<td>0.0241071 0.0740874</td>
<td>0.0008928 0.0241071</td>
</tr>
<tr>
<td>Op1, op4</td>
<td>0.3616666 0.4536426</td>
<td>0.1797486 0.5413659</td>
<td>0.3616666 0.1797486</td>
</tr>
<tr>
<td>Op1, op5</td>
<td>0.1041666 0.2554951</td>
<td>0.0260416 0.3113869</td>
<td>0.1041666 0.0260416</td>
</tr>
<tr>
<td>Op1, op6</td>
<td>0.0777777 0.2587106</td>
<td>0.0777777 0.2587106</td>
<td>0.0777777 0.0777777</td>
</tr>
<tr>
<td>Op1, op7</td>
<td>0.0044642 0.0178571</td>
<td>-0.0625 0.25</td>
<td>0.0044642 0.0625</td>
</tr>
<tr>
<td>Op2, op3</td>
<td>0.1194726 0.122395</td>
<td>0.0658514 0.1397070</td>
<td>0.1194726 0.0658514</td>
</tr>
<tr>
<td>Op2, op4</td>
<td>0.1583097 0.1416790</td>
<td>-0.1246387 0.2763162</td>
<td>0.1583097 0.1246387</td>
</tr>
<tr>
<td>Op2, op5</td>
<td>0.1630756 0.1588262</td>
<td>-0.0535737 0.1469494</td>
<td>0.1630756 0.0535737</td>
</tr>
<tr>
<td>Op2, op6</td>
<td>0.2479740 0.4629131</td>
<td>0.0979913 0.332460</td>
<td>0.2479740 0.0979913</td>
</tr>
<tr>
<td>OP2,op7</td>
<td>0.1390082 0.1907661</td>
<td>-0.0535258 0.2445812</td>
<td>0.1390082 0.0535258</td>
</tr>
<tr>
<td>Op3, op4</td>
<td>0.1601363 0.1411115</td>
<td>0.1436880 0.3675601</td>
<td>0.1601363 0.1436880</td>
</tr>
<tr>
<td>Op3, op5</td>
<td>0.0583333 0.0898558</td>
<td>-0.0447916 0.1019656</td>
<td>0.0583333 0.0447916</td>
</tr>
<tr>
<td>Op3, op6</td>
<td>0.0166666 0.1409077</td>
<td>-0.0083333 0.0845893</td>
<td>0.0166666 0.0083333</td>
</tr>
<tr>
<td>OP3,op7</td>
<td>0.0152173 0.0504547</td>
<td>-0.0642468 0.2496315</td>
<td>0.0152173 0.0642468</td>
</tr>
<tr>
<td>Op4, op5</td>
<td>0.5216666 0.4221049</td>
<td>0.2786375 0.4987435</td>
<td>0.5216666 0.2786375</td>
</tr>
<tr>
<td>Op4, op6</td>
<td>0.3166666 0.2855654</td>
<td>0.1347486 0.4101417</td>
<td>0.3166666 0.1347486</td>
</tr>
<tr>
<td>OP4,op7</td>
<td>0.3472951 0.3530456</td>
<td>0.125903 0.3530376</td>
<td>0.3472951 0.125903</td>
</tr>
<tr>
<td>Op5, op6</td>
<td>0.075 0.1194121</td>
<td>-0.003125 0.132247</td>
<td>0.075 0.003125</td>
</tr>
<tr>
<td>Op5, op7</td>
<td>0.078125 0.1760385</td>
<td>-0.0669642 0.2494466</td>
<td>0.078125 0.0669642</td>
</tr>
<tr>
<td>Op6, op7</td>
<td>0.0349264 0.0904917</td>
<td>-0.0320378 0.2735720</td>
<td>0.0349264 0.0320378</td>
</tr>
<tr>
<td>Averages</td>
<td>0.1487287 0.0297332</td>
<td>0.1488137 0.0802188</td>
<td></td>
</tr>
</tbody>
</table>

In this paper, we submit the following premises:

- First, for most practical purposes, determining which mutants are equivalent to a base program (and which are not) is not important, provided we can estimate their number.
- Second, even when it is important to single out mutants that are equivalent to the base program, knowing (or estimating) their number may be of great help in practice. With every mutant that is killed (i.e. found to be distinct from the base), the probability that the remaining mutants are equivalent increases as their number approaches the estimated number of equivalent mutants.
- Third, what makes a program prone to produce equivalent mutants is the same attribute that makes it fault tolerant, since fault tolerance is by definition the property of maintaining correct behavior (e.g. by being equivalent) in the presence of faults (e.g. artificial faults, such as mutations). The attribute that makes programs fault tolerant is well-known: redundancy. Hence if we can quantify the redundancy of a program, we ought to be able to correlate it statistically to the ratio of equivalent mutants.
- Fourth, what determines the ratio of equivalent mutants of a program includes not only the
program itself, of course, but also the mutation operators that are applied to it. We are exploring the possibility that the ratio of equivalent mutants of a program $P$ upon application of a set of mutation operators can be inferred from the REM of the program $P$ under each operator applied individually. If we can produce and validate a generic formula to this effect, we can conceivably estimate the REM of a program for an arbitrary set of mutation operators.

### 6.2 Prospects

One of the most inhibiting obstacles to the large scale applicability of our approach is that the calculation of the redundancy metrics presented in section 3 is done by hand. Hence the most urgent task of our research agenda is to automate the calculation of the redundancy metrics on a program of arbitrary size and complexity. This is currently under way, using routine compiler generation technology; we are adding semantic rules to a skeletal Java compiler, which keeps track of variable declarations (to compute state space entropies), and assignment statements (to keep track of functional dependencies between program variables). Once we have a tool that can automatically compute our redundancy metrics, we can revisit the experimental process that we have followed in section 3.6 to derive a detailed/ accurate statistical model for estimating the REM of a program for a specific mutant generation policy.

In order to make the model adaptable to different mutant generation policies, we envision to continue exploring the relationship between the REM of a program for individual mutation operators and the REM of the same program for a combination of operators. In particular, we are investigating the validity of the conjecture that the REM of a program for a set of $N$ mutation operators can be obtained by solving the equation:

$$(1 - REM) = \prod_{i=1}^{N} (1 - REM_i).$$

We are currently running an experiment to check this formula for $N = 3$, which yields:

$$REM = REM_1 + REM_2 + REM_3 - REM_1REM_2 - REM_2REM_3 - REM_1REM_3 + REM_1REM_2REM_3.$$  

If the proposed formula of the REM of a program $P$ for an arbitrary number of mutation operators is validated, then we can estimate the REM

Let us consider $n$ mutation operators, say $op_1, op_2, ..., op_n$. Let us assume that we have a regression model for each individual operator, which estimates the REM of a program $P$ using its redundancy metrics as independent variables. If the formula

$$(1 - REM) = \prod_{i=1}^{n} (1 - REM_i).$$

Is validated, then we can use it to compute the REM of a program $P$ for any subset of the $n$ operators for which we have a regression model. In other words, we can compute the REM of a program for $2^n$ different sets of mutation operators, using merely $n$ regression models.

### 6.3 Assessment and Threats to Validity

The task of identifying and weeding out equivalent mutants by painstaking analysis of individual mutants is inefficient, error prone, and usually yields unnecessary or insufficient conditions of equivalence. We feel confident that our approach to this problem, which relies on estimating the number of equivalent mutants rather than singling them out individually, is a cost-effective alternative. Indeed, most often it either obviates the need to identify equivalent mutants individually, or makes the task significantly simpler.

We do not consider the work reported on in this paper as complete:

- The regression model that we present in section 3.6 is not an end in itself; rather it is a means to show that the REM of a program is statistically related to its redundancy metrics, and that it is possible to estimate the former using the latter. Our end goal is to automate the calculation of redundancy metrics, then use these to derive a large scale regression model that is based on a large sample of large and complex software artifacts.

- The empirical investigation of section 4.3 is not an end in itself; rather it is a means to prove that it may be possible to estimate the REM of a program for a composite mutation generation strategy from the REM’s obtained for its member mutation operators. We have some tentative results to this effect, and we are pursuing tantalizing venues.

Hence we view this paper as putting forth research questions rather than providing definitive answers; our research agenda is to investigate these questions.
REFERENCES


Just, R., Ernst, M. D. and Fraser, G. (2014). Efficient Mutation Analysis by Propagating and Partitioning Infected Execution States. In *Proceedings, ISSTA '14, San Jose, CA USA*.


