A Probabilistic Analysis Method for Functional Qualification under Mutation Analysis

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Abstract—Mutation Analysis (MA) is a fault-based simulation technique that is used to measure the quality of testbenches in error (mutant) detection. Although MA effectively reports the living mutants to designers, it suffers from the high simulation cost. This paper presents a probabilistic MA preprocessing technique, Error Propagation Analysis (EPA), to speed up the MA process. EPA can statically estimate the probability of the error propagation with respect to each mutant for guiding the observation-point insertion. The inserted observation-points will reveal a mutant’s status earlier during the simulation such that some useless testcases can be discarded later. We use the mutant model from an industrial EDA tool, Certitude, to conduct our experiments on the OpenCores’ RT-level designs. The experimental results show that the EPA approach can save about 14% CPU time while obtaining the same mutant status report as the traditional MA approach.

I. INTRODUCTION

Functional verification is a process to ensure that the implementation is in accord with the design intent [24][26]. When designs are increasingly complex, functional verification are getting more difficult for designers. In general, simulation-based verification methods are the mainstream in the current design flow, especially for large designs. Due to the fact that exhaustive simulation is infeasible, coverage metrics are proposed to measure the quality of verification and thus reduce the simulation cost. However, the completeness problem is still an issue. This is because even all coverage metric reports show that there is no bug in the design under verification, it cannot ensure that additional verification tasks would never discover a new bug.

Coverage metrics can be generally classified into two categories, and they are structural coverage (or code coverage) [18][23] and functional coverage [9][12][23]. Most structural coverage metrics only evaluate the “reachability” of the stimuli to the design content, but do not consider stimuli’s abilities to propagate bugs to observation locations. Therefore, the higher scores in structural coverage metrics do not indicate the better qualities of the test vectors. On the other hand, the functional coverage metric considers the semantic interpretation of the functionality to exercise the design. Since verification engineers need to manually set up various verification plans and a set of coverage points based on the specifications, it is a challenge to develop an automatic method to deal with it.

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Fault-based verification technique [5–7][10][11] is an effective approach that evaluates the quality of the verification environment. By injecting artificial faults into the original implementation, designers can check whether the verification environment can differentiate the faulty and original designs. If all the artificial faults can be detected, it implies that the verification environment is effective to reveal potential bugs; otherwise, the verification environment exists some weakness. Thus, fault-based verification does consider the error propagation issue that the structural coverage does not. The fault-based verification is similar to gate-level fault simulation where the stuck-at fault model is used for calibration of manufacturing test vectors.

Mutation Analysis (MA) is such a fault-based verification technique, originating from the software engineering field in the 1970s [8][13]. MA targets at only a subset of all potential faults and assumes that these faults are sufficient to represent all faults. This is because MA approach is based on two hypotheses: the Competent Programmer Hypothesis [8] and Coupling Effect Hypothesis [8][20][22]. Competent Programmer Hypothesis states that programmers develop their programs very close to the correct version. Coupling Effect Hypothesis assumes that complex faults can be coupled by simple faults in such a way that a test set detecting all simple faults in a program will detect a very high percentage of the complex faults [22]. As a result, MA approach only considers some simple syntactical changes instead of generating all potential faults. For example, CertitudeTM [3][14] is a commercial EDA tool implementing MA approach in RT-level designs for testbench qualification.

In the process of MA, each simple syntactical change is called a mutant. For example, given a program segment $a \leq b \mid c$, after changing the operator, the new program segment becomes $a \leq b \& c$. The replacement of “$\mid$” with “$\&$” is a mutant for the original program. In the traditional process of MA, a set of mutants will be generated first, and then all test data are simulated. If we observe any difference caused by a mutant at the primary outputs, we call the mutant is killed, otherwise this mutant is a living mutant. The existence of living mutants indicates some weaknesses in the verification environment.

The MA makes the fault-based verification technique become a systematic verification methodology. However, it has suffered from some problems. The surveys of MA in literature [16][21] addressed that the MA induces extremely high computational cost in the state-of-the-art. For each injected
mutant, we need to perform the whole design simulation for all the testcases once. Therefore, the computational cost will rapidly increase due to a great amount of injected mutants and testcase.

In this work, we propose a preprocessing technique for accelerating MA process named Error Propagation Analysis (EPA), which analyzes the error propagation ability of each mutant before performing the simulation in RT-level designs. Then, we determine a few observation-points [17] to observe the effect of error propagation. The analyzed results can be embedded into the MA cost reduction technique, hence, we can reduce the simulation cost of MA.

II. COST REDUCTION TECHNIQUES IN MA

In the traditional MA process, a set of mutated programs are generated first, then all the testcases are applied to each mutated program to determine if each mutant is killed or not. The simulation cost of this MA approach is much more than that in the original design simulation.

In the latest survey of MA [16], cost reduction techniques are divided into two types: one is mutant reduction, and the other is simulation reduction. The mutant reduction is to reduce the number of generated mutant without having significant loss of the testbench quality. The simulation reduction is to minimize the simulation cost of a mutated program. In this work, we focus on the issue of simulation reduction. Thus, in the next paragraph, we will introduce the background of a widely used simulation reduction technique, “Strong, Weak, and Firm Mutation (SWFM)”, that is the basis of this work.

SWFM proposed three different degrees of mutation with respect to the observation-points, and they are Strong Mutation, Weak Mutation, and Firm Mutation. In the traditional MA process, when a mutant is “killed” after the simulation, it means that the error effect caused by this mutant is propagated to the primary outputs. This type of mutation is called Strong Mutation [8]. Strong Mutation can reveal the potential weaknesses thoroughly, but it suffers from the high simulation cost due to the whole design simulation. To reduce the simulation cost, Weak Mutation [15] was proposed. Weak Mutation assumes that each statement in a design is an observation-point. Thus, if a mutant causes any statement’s output changed, the mutant is said killed. As a result, Weak Mutation reduces the simulation cost but sacrificing the quality of testbench. When the observation-points lie between the mutated statements (Weak Mutation) and the primary outputs (Strong Mutation), this mutation is called Firm Mutation [25], which is a tradeoff between Strong and Weak Mutations. A Verilog example about these three types of mutations is shown in Fig. 1 and Table I.

In Fig. 1, the original operation $b \& c$ is replaced with $b \wedge c$ in line 8, and it is an injected mutant. We apply an input vector $\{a, b, c\} = \{0, 0011, 0001\}$ to both the mutated and the original programs. In this example, we need to simulate this design for 5 clock cycles to get the value of $out$. The results are shown in Table I. Since these three types of mutations have different observation-points, they may result in different mutant statuses. The Strong Mutation performed the complete simulation, and determined the mutant status after simulating 5 cycles. This mutant status is living because the primary output $out$ of mutated program is the same as that of the original program. In the Weak Mutation, the observation-point is the site of the mutant. Thus, this mutant status is killed after one cycle simulation because of the different results between the statements of $x = b \& c$ and $x = b \wedge c$. For the Firm Mutation, however, the location of observation-point has many choices. Assume we set $y$ as the observation-point in this example, then the mutant is still living after one cycle simulation by comparing the value $y$ in the mutated and the original programs. From this example, we found that the advantage of Firm Mutation is that it can determine the living mutant after fewer simulation cycles. This is because a living mutant under the Firm Mutation is still a living mutant under the Strong Mutation.

Due to the incomplete simulation, killed mutants under the Weak Mutation are very likely to be living mutants under the Strong Mutation. To avoid sacrificing the quality of results, we need to check whether all killed mutants under the Weak Mutation also can be killed under the Strong Mutation. Although the Weak Mutation Analysis cannot decide the mutants’ actual statuses, it can be used to eliminate some useless testcases for the further Strong Mutation. For example, when applying 100 testcases to test a mutant A, we need to perform the whole design simulation 100 times under the Strong Mutation. Suppose we applied the Weak Mutation first, and found that 80 out of 100 testcases cannot kill the mutant A. Then these 80 testcases cannot kill the mutant A under the Strong Mutation, either. Therefore, only the remaining 20 testcases need to be applied for the mutant A under the Strong Mutation. Thus, the total simulation cost can be reduced by first applying the Weak Mutation.

Although the Weak Mutation effectively reduces the simulation cost, the remaining simulation cost for the Strong Mutation is still large. A killed mutant under the Weak Mutation but living under the Strong Mutation means that this mutant causes the error in the mutated statement but cannot be propagated to the primary outputs. We called such a mutant Non-Propagation Mutant (NPM). The Firm Mutation is proposed to mainly target at this propagation issue. It can also be designed to insert the internal observation-points to find the potential NPMs earlier. However, the determination of the locations and the number of the observation-points is a

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**Figure 1.** An example for demonstrating Strong, Weak, and Firm Mutations.

**Table I**

<table>
<thead>
<tr>
<th>Mutation type</th>
<th>Observation-Point</th>
<th>Expected Result</th>
<th>Mutated Result</th>
<th>Mutant Status</th>
<th>Determined at # cycle</th>
</tr>
</thead>
<tbody>
<tr>
<td>Strong</td>
<td>out</td>
<td>out=5</td>
<td>out=5</td>
<td>Living</td>
<td>5</td>
</tr>
<tr>
<td>Weak</td>
<td>x</td>
<td>$x=0001$</td>
<td>$x=0011$</td>
<td>Killed</td>
<td>1</td>
</tr>
<tr>
<td>Firm</td>
<td>y</td>
<td>$y=0000$</td>
<td>$y=0000$</td>
<td>Living</td>
<td>1</td>
</tr>
</tbody>
</table>

Input testcases $\{a, b, c\} = \{00011, 00001\}$ and the mutant $b \& c$. The results can be reduced to a significant extent.
critical issue. This is because too many observation-points will increase the simulation overhead, and improper observation-points will decrease the probability of finding potential NPMs. To solve this problem, we propose a probabilistic analysis to automatically determine the observation-points based on the Firm Mutation mechanism, which will be detailed in Section III.

III. PROPOSED APPROACH

A. Overview

The EPA intends to statically analyze the structure of HDL design and evaluate the error propagation ability for a given mutant. We propose Mutant Controllability (MC) to represent the changing probability of each signal. We also propose the MC estimation formulae for various operations that are used to determine the MCs of the signals in the propagation path with respect to each mutant. Using the estimated MC, we derive the Decreasing Rate (DR) of MC, and select the locations with the largest DR for inserting observation-points.

In the Firm Mutation approach, the inserted observation-points will affect a mutant’s status during the simulation. As a result, the testcases that cannot kill the mutant are excluded from the original testcase set for speeding up the succeeding Strong Mutation simulations. Finally, we collect the living mutants under the Firm Mutation, and analyze their related MCs to point out the possible causes for their liveness. Note that our approach in this work targets at the word-level MC estimation in its propagation path with respect to each mutant. Using the estimated MC, we derive the DR for inserting observation-points.

B. Terminology

1) Mutant Controllability (MC): When MA is performed, the signals in the propagation path from the site of mutant to the primary outputs will be possibly changed. If one of the primary output changes, the mutant is killed as desired. However, if the mutant effect is blocked such that all primary outputs are intact, the mutant is living. Here, we define the Mutant Controllability (MC) of each signal as its changing probability in its propagation path with respect to a given mutant. Specifically, \( MC(m, s) \) represents the changing probability of a signal \( s \) with respect to a mutant \( m \). \( MC(m, s) = 1 \) if the signal \( s \) is on the mutated statement. \( MC(m, s) = 0 \) if the mutant \( m \) does not change the value of the signal \( s \). The signal with a higher MC means that the mutant is able to change the signal with a higher probability. For the example in Fig. 3, the mutant \( m \) is on \( tmp1 \), hence, we set \( MC(m, tmp1) = 1 \). Since the error effect will be propagated and the signals \( x, y \) are on the propagation path from \( m \), we can derive the values of \( MC(m, x) \) and \( MC(m, y) \) after the MC calculation. On the other hand, \( tmp2 \) is not on the propagation path, hence, we can directly set \( MC(m, tmp2) = 0 \).

2) Control Data Flow Graph (CDFG): In this work, the input is a Verilog design represented as a CDFG [4][19]. A CDFG representation is composed of four types of nodes: operational nodes, control nodes, call nodes, and storage nodes. Operational nodes are responsible for arithmetic, logical, or relational operations. Control nodes are responsible for branch conditions such as if-else or case statements. Call nodes and storage nodes represent calls to subprogram and assignment operations associated with variables and signals, respectively. Since most mutant effects are propagated through the operational nodes and control nodes, our MC estimation formulae only focus on these two types of nodes. Although storage nodes do not affect the propagation of mutant effects, they are the MC recomputation points for the sequential part of designs. Fig. 4 shows a mapping example from a Verilog design to its CDFG.

C. Mutant Controllability Estimation Formulae

MC estimation formulae are used to estimate how difficult a mutant effect can be propagated out. Operational nodes can be divided into unary and binary operational nodes and are discussed as follows:

1) Unary operational nodes: Consider a unary operation "\( out = op \ a \)", where \( a \) is an operand, \( op \) is an unary operator as shown in Fig. 5(a). The MC estimation formula for a unary operation is shown as EQ(1).

\[
MC(m, out) = MC(m, a) \times [1 - P_{mask}[v'_a, op]]
\]  

(1)

Figure 2. Functional qualification flow with the Error Propagation Analysis.

Figure 3. A propagation path from the mutant \( m \).

Figure 4. An example about the mapping from a Verilog design to its CDFG.
In EQ(1), we get the MC of \( \text{out} \) from the MC of \( a \). Here, suppose that for the operand \( a \), \( v_a \) is its correct value and \( v_a' \) is the new value caused by the injected mutant. \( P_{\text{mask}}(v_a', op) \) represents the probability of the "same" result in the operations of \( op \) \( v_a \) and \( op \) \( v_a' \). This probability is called masking probability. Assume \( v_{\text{out}} \) and \( v'_{\text{out}} \) are the results of the unary operation with respect to the input \( v_a \) and \( v_a' \), then \( P_{\text{mask}}(v_a', op) = P(v_{\text{out}} = v'_{\text{out}}) \). To derive this masking probability, we need to consider the operator \( op \) and the width \( w \) of the operand \( a \).

To simplify the computation, we assume that \( v_a \) and \( v_a' \) are evenly distributed from 0 to \( 2^w - 1 \), and \( v_a \neq v_a' \) (the mutant effect has been activated). For example, suppose the \( op \) is a unary AND "\&", which will output 1 iff all bits of the operand are 1; 0 otherwise. The masking probability is the probability that \( v_{\text{out}} \) and \( v'_{\text{out}} \) are the same. Thus, \( P_{\text{mask}}(v_a', \&) \) is derived as \( P_1((v_{\text{out}} = 1) \land (v'_{\text{out}} = 1)) + P_2((v_{\text{out}} = 0) \land (v'_{\text{out}} = 0)) \). Because only one vector \{111...1\} results in the output 1, it is not possible that \&\(v_a' = v_{\text{out}} = \&v_a = v_{\text{out}} = 1 \). As a result, \( P_1 \) is 0. On the other hand, the probability of \&\(v_a' = v_{\text{out}} = 0 \) is \( 1 - \frac{1}{2^w} \), and since \( v_a' \neq v_a \), the probability of \&\(v_{\text{out}} = 0 \) is \( 1 - \frac{1}{2^w} \). Also, since these two events are independent, \( P_2 \) is the product of them. Hence, we can get \( P_{\text{mask}}(v_a', \&) = P_1 + P_2 = 0 + (1 - \frac{1}{2^w})(1 - \frac{1}{2^w}) \). Finally, we get \( MC(m, out) = MC(m, a) \times [1 - (1 - \frac{1}{2^w})(1 - \frac{1}{2^w})] \). This result shows that the \( MC(m, out) \) will be smaller when \( w \) is larger, which implies that the error effect will be difficultly propagated. The masking probabilities of the common unary operations are summarized in the bottom of Table II.

2) Binary operational nodes: Consider a binary operation "\( out = a \; op \; b \)" where \( a \) and \( b \) are the operands. The estimation formula is shown as EQ(2).

\[
MC(m, out) = MC(m, a) \times MC(m, b) \times [1 - P_{\text{mask}}(v_a', v_b, op)] + \frac{MC(m, a) \times MC(m, b) \times [1 - P_{\text{mask}}(v_a, v_b, op)]}{MC(m, a) \times MC(m, b) \times [1 - P_{\text{mask}}(v_a', v_b', op)]}
\]

In EQ(2), \( MC(m, a) \) and \( MC(m, b) \) represent the probability of the correct values on the input signals and can be derived from \( 1 - MC(m, a) \) and \( 1 - MC(m, b) \), respectively. In a binary operation as Fig. 5(b) shows, there are three conditions that the error effects will be propagated out with respect to the incorrect values on \( a \), \( b \), or both. Because the masking probabilities in these three conditions are different, we separately consider these conditions and multiply them with their corresponding \( MC \) or \( \overline{MC} \) values. The first term in EQ(2) shows that \( v_a' \) is an incorrect value and \( v_b \) is a correct value, hence, the equation is involved with the factors \( MC(m, a) \) and \( \overline{MC}(m, b) \). Similarly, the second term is involved with \( \overline{MC}(m, a) \) and \( MC(m, b) \) with respect to \( v_a \) and \( v_b' \). The last term represents the condition of the simultaneous incorrect values \( v_a' \) and \( v_b' \).

For example, suppose the operator \( op \) is "\( + \)" and the width of the operands is \( w+ \). To derive the masking probability for each term in EQ(2), we assume that the error effect will cause a change with \( \Delta_a \) and \( \Delta_b \) such that \( v_a' = v_a + \Delta_a \) and \( v_b' = v_b + \Delta_b \), where \( \Delta_a \neq 0 \) and \( \Delta_b \neq 0 \). Thus, both \( P_{\text{mask}}(v_a', v_b', +) \) and \( P_{\text{mask}}(v_a, v_b', +) \) will be 0 because of \( v_a + v_b \neq v_a' + v_b' = (v_a + \Delta_a) + v_b \) and \( v_a + v_b \neq v_a' + v_b' = v_a + (v_b + \Delta_b) \).

For the last term, \( v_a + v_b = v_a' + v_b' = v_a + v_b + \Delta_a + \Delta_b \) is true iff \( \Delta_a = -\Delta_b \) is true. Therefore, the masking probability \( P_{\text{mask}}(v_a', v_b', +) \) in the last term is the same as the probability of \( \Delta_a = -\Delta_b \), which is \( \frac{1}{2^w} \). Finally, we get \( MC(m, out) = MC(m, a) \times MC(m, b) \times (1 - \frac{1}{2^w}) \). The masking probabilities of the common binary operations are also summarized in Table II.

3) Control nodes: Consider a control node which contains \( n \) inputs \{\( d_0, d_1, ..., d_{n-1} \)\}, a data output \( out \), and a selector \( s \) with the range from 0 to \( n - 1 \). There are two conditions that the error effect will be propagated to the data output \( out \): one is the error effect on the selector \( s \) and the incorrectly selected input has a different value with that of the expected input; the other is the error effect is on \( d_i \) and \( s = i \). To simplify our computation, we assume \( s \) is an integer evenly distributed from 0 to \( n - 1 \), which means that each input signal will be selected with the same probability. Additionally, the width of each \( d_i \) is \( w \) and its value is also evenly distributed from 0 to \( 2^w - 1 \). The estimation formula for the control nodes is shown as EQ(3).

\[
MC(m, out) = \left( \frac{MC(m, s) \times 2^w - 1}{2^w} \right) + \left( \frac{MC(m, s) \times \sum_{d=0}^{n-1} MC(m, d_i)}{n} \right)
\]

We divide EQ(3) into two terms with respect to whether the selector is erroneous or not. The first term is for the case that the error effect is on the selector \( s \). Suppose \( i, j \) are the correct and incorrect values of the selector \( s \), we only need to consider whether the expected input \( d_i \) and the incorrectly selected input \( d_j \) have the same value. Since \( d_j, d_j \) are evenly distributed from 0 to \( 2^w - 1 \), the probability of \( d_j \neq d_j \) is \( \frac{2^w - 1}{2^w} \). Therefore, the first term is \( MC(m, s) \times \frac{2^w - 1}{2^w} \) where \( MC(m, s) \) is the probability of erroneous selector. The second term is for the case that the value of the selector \( s \) is correct. The error effect will be propagated when the selected input has an incorrect value. Because we assume each input will be selected with the same probability, we use the averaged MC value \( \frac{\sum_{d=0}^{n-1} MC(m, d_i)}{n} \) to represent the probability of the
error effect on the selected input. Thus, the second term is the product of $MC(m, s)$ and the averaged MC value among all inputs.

**D. MC Estimation Considering the Sequential Circuits**

When considering sequential circuits, we can see that there exists feedback loops in a CDFG. Because a mutant is likely to affect some signals after a few cycles, we need to recompute the MC values of signals in the feedback loop for several times. Consider the feedback case as shown in Fig. 6, after deriving the $MC(m, E)$ with a given mutant $m$, the value of $MC(m, A)$ will be updated in the next clock cycle. Because assignment operations do not affect the propagation of error effects, we get updated $MC(m, A) = M(m, E)$. Therefore, we need to recompute $MC(m, D)$ for the operation $D = A op B$. To avoid too much computation cost for the feedback, we terminate the recomputation when the difference between the current and the next MC values is under a given threshold.

**E. The Observation-Point Selection**

For any node in the CDFG, if the error effect can be observed on its inputs but not on its output, this node is an error-masking node. If there exists one error-masking node on each propagation path of a mutant, this mutant is an NPM. The observation-point is used to observe the mutants’ statuses under the Firm Mutation. Since we desire to reduce the unnecessary simulation effort, a proper observation-point should be lied on the output of an error-masking node.

To determine these proper observation-points, we must find the error-masking nodes first. Because of the enormous number of nodes in the CDFG, finding all the error-masking nodes for an NPM is time-consuming. Thus, we use the results of the MC estimation to find the potential error-masking nodes.

Next, we detail the observation-point selection method. In the first step, we extract all the propagation paths of a given mutant without considering feedback signals. For a given mutant $m$, assume that there are $n$ propagation paths from $m$ to the primary outputs. $PP_{m,i}$ denotes the $i^{th}$ propagation path of the mutant $m$ where $1 \leq i \leq n$. $PP_{m,i}$ consists of some ordered signals on this propagation path. $|PP_{m,i}|$ denotes the number of signals on this path. For example, considering a design segment in Fig. 7, if there exists an error effect on $A$ from a mutant $m$, we can identify two propagation paths $PP_{m,1} = \{A, tmp, X, Y, O_1\}$, $PP_{m,2} = \{A, tmp, X, Y, O_2\}$, and $|PP_{m,1}| = |PP_{m,2}| = 5$.

We propose the Decreasing Rate (DR) as shown in EQ(4) to represent the degree of the error-masking effect between adjacent signals on a given propagation path $PP_{m,i}$.

$$DR_{s_k}(PP_{m,i}) = \frac{MC(m, s_{k-1}) - MC(m, s_k)}{MC(m, s_{k-1})}, 2 \leq k \leq |PP_{m,i}|$$

(4)

In EQ(4), $DR_{s_k}(PP_{m,i})$ represents the DR of the $k^{th}$ signal, $s_k$, on the $PP_{m,i}$. For example, given a $PP_{m,i}$ with three signals $\{a, b, c\}$, $DR_{s_3}(PP_{m,i})$ is the DR of the third signal $c$ on the $PP_{m,i}$ and it can be derived by $\frac{MC(m,b) - MC(m,c)}{MC(m,b)}$. The larger $DR_{s_k}(PP_{m,i})$ implies the node between the signals $s_{k-1}$ and $s_k$ is a potential error-masking node. Therefore, we select the signals with the largest DR in each propagation path to be the observation-point candidates. Finally, we choose the most frequently selected signals among all candidates to be the final observation-points.

**IV. EXPERIMENTAL RESULTS**

We implemented the EPA analyzer in C++ and the testbench in Verilog, respectively. The experiments were conducted on an Intel Core™2 Quad 2.5 GHz Linux platform (Ubuntu 10.10).

To integrate the MA and our EPA approach into the verification environment, we establish a framework to automate this technique as shown in Fig 8. The EPA analyzer analyzes the mutated design and determines its observation-points. It also automatically generates the monitors to observe the simulation data on the observation-points. Here, we use a random generator to generate the testcases for the designs. The comparator compares the simulation data between the original and the mutated designs, and reports the results. Because the generated monitors can be easily configured to observe different signals for performing the Strong, Weak, or Firm Mutation, this framework makes the testbench reusable for different mutated designs. The threshold value for terminating the recomputation in feedback loops is set 0.1.

The commercial EDA tool, Certitude [14], has proposed various types of mutants. In our experiments, we adopt three types of mutants, and they are changing operator, dead assignment, and condition stuck-at-true (false). Our simulation engine is Icarus Verilog [1], an open-source Verilog simulator. The benchmark comes from the OpenCore website [2]. We only conduct the experiments for three designs in our preliminary implementation. The results of EPA are shown in Table III. Columns 2 and 3 show the number of lines in the benchmarks and the number of injected mutants, respectively. Column 4 is the spent CPU time of the EPA approach.

**Table III**

**The EPA results.**

| Design | REL Times | | CPU (s) |
|--------|-----------|-------------|
| double_fpu | 1971 | 100 | 13 |
| core_uhs | 1232 | 130 | 10 |
| reed_solomon_decoder | 3802 | 300 | 31 |
Table IV: The Weak and Firm Mutation Analysis Results.

<table>
<thead>
<tr>
<th>Design</th>
<th>[TC]</th>
<th>[JM]</th>
<th>[TSR]</th>
<th>Preprocessing Method</th>
<th>Strong Mutation Analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Pre. Types</td>
</tr>
<tr>
<td>double_fpu</td>
<td>1000</td>
<td>100</td>
<td>1.0 × 10^5</td>
<td>Weak</td>
<td>5.72 × 10^4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Firm</td>
<td>4.07 × 10^4</td>
</tr>
<tr>
<td>ecu64</td>
<td>1000</td>
<td>150</td>
<td>1.5 × 10^5</td>
<td>Weak</td>
<td>4.37 × 10^4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Firm</td>
<td>4.92 × 10^4</td>
</tr>
<tr>
<td>reed_solomon_decoder</td>
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<td>300</td>
<td>3.0 × 10^4</td>
<td>Weak</td>
<td>1.36 × 10^4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Firm</td>
<td>1.82 × 10^4</td>
</tr>
</tbody>
</table>

Averaged CPU time reduction of the Strong Mutation Analysis with the Firm Mutation preprocessing: 0.86

To compare the performance of our preprocessing method, we apply the Weak and the Firm Mutation to the benchmark, respectively, followed by the Strong Mutation. The Firm Mutation is embedded in the proposed EPA approach. The simulation results are shown in Table IV. Columns 2 and 3 list the number of testcases (ITC) and the number of injected mutants (IMI), respectively. Column 4 is the number of the total simulation rounds (ITSR). Here, we use a Simulation Round (SR) to represent a unit cost in the mutation simulation, which is the time cost of applying a testcase to test a mutant. Thus, the total number of SRs in performing the MA without any preprocessing method is the product of the number of testcases and the number of injected mutants. Columns 5 to 9 show the results of the two MA preprocessing methods, Weak and Firm Mutation. Because both preprocessing methods can find the living mutants and the corresponding useless testcases, the numbers of SRs and the number of injected mutants applying for the further Strong Mutation are reduced. Columns 6 and 7 show the number of the reduced SRs (iTSR) and the percentage of that to the ITSR (RSP%). Column 8 is the number of living mutants reported by the preprocessing methods. Column 9 is the required CPU time measured in second. Note that the Firm Mutation is embedded into the EPA approach, the time for the EPA as shown in Table III has been accounted in Column 9. Columns 10 to 12 show the results of overall flow including the preprocessing method. Column 10 is the number of living mutants (Nsp). Column 11 is the total CPU time including the preprocessing time. Column 12 shows the ratio of total CPU time of the Strong Mutation embedding the Firm Mutation against the Weak Mutation for the preprocessing (Tsp%).

According to Table IV, we found that the Weak Mutation cost less CPU time as compared to the Firm Mutation for the preprocessing. It is because for the Weak Mutation, it can determine a mutant is killed when the mutated statement was activated, without considering the propagation issue. On the other hand, the Firm Mutation embedded in our EPA method identified more living mutants and the corresponding useless testcases for the succeeding Strong Mutation. Thus, although the Firm Mutation cost more CPU time, its larger number of RSRs makes computation cost in the further Strong Mutation less as shown in Column 11. The averaged CPU time reduction in the last row shows that the Firm Mutation embedded in our EPA method can save 14% CPU time on average.

V. Conclusion

In this paper, we propose an Error Propagation Analysis approach that deals with the error propagation issues in Mutation Analysis. This approach can probabilistically analyze the HDL designs and add the proper observation-points. Based on the Firm Mutation approach, we can determine the mutant status by monitoring these selected observation-points. Therefore, the Firm Mutation can discover more living mutants and discard the useless testcases of each mutant to reduce the simulation cost of the further Strong Mutation. The experimental results show that our approach is more efficient than the Weak Mutation, which does not consider the error propagation issue.

REFERENCES