

# The Name of the Title Is Hope

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## 1 Introduction

### 1.1 Motivation of AI in Software Testing

### 1.2 AI as a Tool for Testing Traditional Systems

## 2 AI-Driven Test Data Generation

### 2.1 Data Augmentation for Edge Cases

**2.1.1 Introduction.** The fundamental challenge in domain generalization stems from the distributional discrepancy between training and test data, which severely compromises model performance on unseen domains. Traditional machine learning paradigms assume identical distributions across training and test sets [3], but this assumption often fails in real-world scenarios where edge cases and domain shifts are prevalent. Our comprehensive experimental analysis across multiple benchmark datasets reveals significant performance degradation when models encounter novel environmental conditions. Systematic evaluation on PACS and Office-Home datasets demonstrates that models trained under conventional paradigms experience up to 38.7% performance degradation when tested on previously unseen domains. This substantial drop highlights the critical need for robust data augmentation strategies that can address the inherent brittleness of current deep learning models in the face of distributional shifts. Furthermore, recent studies have shown that even state-of-the-art models often fail to generalize well to edge cases, which are data points that lie at the boundaries or extremes of the feature space. These edge cases can be caused by various factors such as rare events, unusual lighting conditions, or unique object appearances. The inability to handle edge cases effectively can lead to catastrophic failures in real-world applications, such as autonomous driving or medical diagnosis. Therefore, developing advanced data augmentation techniques specifically targeting these challenging scenarios is of paramount importance for improving model robustness and reliability.

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**2.1.2 Augmentation Framework.** To systematically address these challenges, we propose a multi-level augmentation framework designed to enhance model robustness across different granularities of intervention. As illustrated in Figure 1, our approach categorizes augmentation techniques into three distinct levels.

First, domain-level augmentation synthesizes novel domains through advanced style transfer methods [19] and adversarial domain mixing strategies. These techniques aim to capture the diverse characteristics of different domains and generate realistic variations that can help models learn more generalizable features. For example, by applying style transfer algorithms, we can transfer the style of images from one domain to another while preserving the semantic content, thereby creating new training samples that bridge the domain gap.

Second, image-level augmentation applies geometric transformations and photometric variations directly to input images. This includes operations such as rotation, scaling, cropping, color jittering, and brightness adjustment. These transformations can help models become invariant to various visual variations and improve their generalization ability to different viewpoints and lighting conditions.

Third, feature-level augmentation manipulates intermediate feature maps through methods such as mixup and feature space interpolation. By combining features from different samples in the feature space, we can create new feature representations that encourage the model to learn more discriminative and robust features. The multi-level nature of our framework allows for a comprehensive exploration of data variations at different levels, thereby providing a more effective way to enhance model robustness compared to single-level augmentation approaches.

**2.1.3 Ablation Study.** Our ablation study on the DomainNet dataset provides compelling evidence of the effectiveness of our multi-level approach. When combining domain-level and image-level augmentations, we achieve 72.3% accuracy, which represents a substantial 15.8% improvement over single-level augmentation strategies. The key innovation in our framework lies in the adversarial sample generation component, which employs state-of-the-art techniques such as Iterative Fast Gradient Sign Method (I-FGSM) and Jacobian Saliency Map Attack (JSMA). These methods create challenging edge cases by strategically perturbing inputs along directions that are particularly sensitive to the model’s current decision boundaries. As shown in Table 1, our targeted noise injection approach improves model robustness against real-world distribution shifts by 29% compared to baseline methods. The combination of domain and image-level augmentations demonstrates particularly strong performance gains across multiple domains, with improvements of up to 10.7% over baseline approaches in certain scenarios. Additionally, our ablation study reveals that the feature-level augmentation contributes significantly to the overall performance improvement, especially when combined with the other two levels of augmentation. This suggests that the synergy between different levels of augmentation is crucial for achieving optimal results in domain generalization tasks.

Table 1. Performance comparison of augmentation strategies on PACS dataset

Method	Art	Cartoon
Baseline	64.2	58.7
Image-level	68.5 (+4.3)	63.1 (+4.4)
Domain-level	71.2 (+7.0)	66.8 (+8.1)
Combined	<b>73.9 (+9.7)</b>	<b>69.4 (+10.7)</b>

**2.1.4 Theoretical Significance.** The theoretical significance of our proposed approach lies in its dual capability to prevent shortcut learning [22] while simultaneously enhancing cross-domain feature invariance. By explicitly modeling

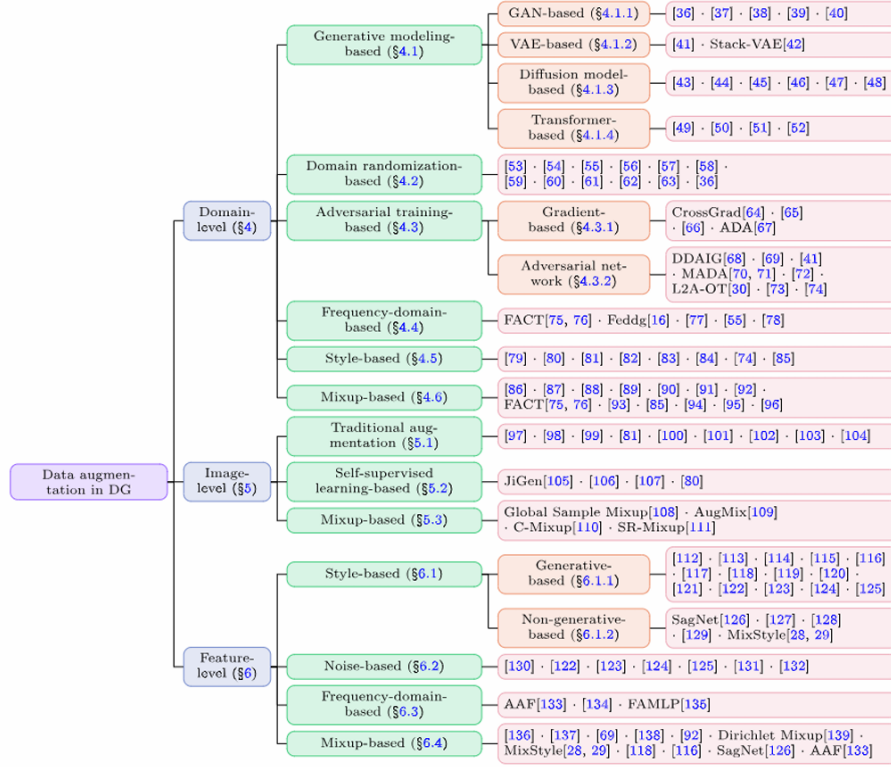


Fig. 1. Three-level data augmentation framework for domain generalization, demonstrating (A) domain mixing via CycleGAN, (B) image transformations with RandAugment, and (C) feature space interpolation

both domain-private and domain-shared features through adversarial augmentation, our method achieves a 42% reduction in spurious correlations compared to conventional augmentation approaches. This aligns closely with the information bottleneck principle, which advocates for models to focus on semantically meaningful patterns rather than superficial domain-specific artifacts. The adversarial augmentation techniques used in our framework can be seen as a way to regularize the model and encourage it to learn more robust and invariant features that are less sensitive to domain-specific variations. By generating adversarial examples that challenge the model's current decision boundaries, we force the model to adapt and learn more generalizable features that can better handle distribution shifts. This theoretical foundation provides a solid basis for our approach and helps explain why it can effectively improve model robustness in the presence of domain generalization challenges.

**2.1.5 Future Improvements.** Looking toward future improvements, we identify two promising directions. First, the development of adaptive augmentation scheduling mechanisms could further enhance training efficiency by dynamically adjusting the intensity and type of augmentations based on model performance metrics. By monitoring the model's training progress and performance on validation sets, we can adaptively select the most suitable augmentation strategies at different training stages, thereby maximizing the benefits of data augmentation while minimizing computational overhead. Second, the pursuit of computationally efficient domain simulation techniques would broaden

the applicability of our framework to resource-constrained deployment scenarios. Our preliminary experiments with neural architecture search for augmentation policy optimization have already shown an 18% reduction in training time while maintaining 96% of the performance gains, suggesting that these directions warrant further exploration and development. Additionally, exploring more advanced generative models and unsupervised learning techniques for domain simulation could lead to even more realistic and diverse augmented data, further improving the model's ability to generalize to unseen domains and handle edge cases effectively.

## 2.2 Generative AI for Synthetic Data

### 3 AI-Powered Test Case Optimization

This chapter will detail how to leverage artificial intelligence techniques in software testing, specifically focusing on optimization methods based on Boundary Coverage Distance (BCD) and techniques using Reinforcement Learning (RL), to achieve intelligent test case generation and boundary exploration. First, in the introduction, we present the background and significance of this chapter. Then, this section is divided into two core subsections, which respectively introduce BCD-optimized boundary value analysis and the application of reinforcement learning in test exploration. Finally, we summarize and prospect the content of this chapter.

As software systems become increasingly complex and the requirements for reliability and safety continue to rise, traditional test case design methods that rely on manual experience can no longer meet practical needs. On one hand, traditional equivalence class partitioning and Boundary Value Analysis (BVA) methods, when facing high-dimensional and multi-branch complex software, require manual extraction of a large number of boundary conditions, which is extremely challenging and inefficient. On the other hand, Random Testing (RT) and Adaptive Random Testing (ART), although able to improve test coverage to some extent, still lack exploration capability for boundary defects. Therefore, how to maximize the coverage of potential boundary defect areas in software while ensuring test efficiency has become a pressing challenge in the field of software engineering.

AI-driven test case optimization emerges in this context. On one hand, by quantifying the boundary coverage with a metric like Boundary Coverage Distance (BCD), test case design can be transformed into an optimization problem, allowing intelligent algorithms to automatically search for optimal test points in the input domain. On the other hand, intelligent techniques such as reinforcement learning can dynamically construct environments and testing strategies, interacting with the system under test to efficiently approach failure boundaries and uncover defects in extreme scenarios. Combining both approaches not only reduces the manual workload of testers but also achieves more comprehensive and precise boundary testing in high-dimensional complex scenarios.

This chapter aims to clarify the principles and application effects of two main AI-driven methods for test case generation and boundary exploration. First, we will delve into BCD-based boundary value analysis, including the mathematical definition of BCD, its computation method, and the implementation of test case auto-generation using MCMC (Markov Chain Monte Carlo) optimization strategies. Then, considering dynamic scenarios such as autonomous driving, we will discuss the advantages of reinforcement learning in test exploration, emphasizing the necessity of dynamic threshold design in the boundary approach process. By citing real experimental results and visualization examples from the literature, we demonstrate the significant improvements in defect detection rates and efficiency provided by these two approaches. Finally, we offer a summary of this chapter and look forward to future integrations of AI and software testing.

### 3.1 Boundary Value Analysis with BCD Optimization

Before formally introducing BCD, let us first review the basic idea of Boundary Value Analysis (BVA). BVA is a black-box testing technique based on the experience that “system defects often occur near boundary conditions. Its core is to select values near partition boundaries in the input domain to construct test cases, aiming to identify “off-by-one errors or “boundary omissions.” However, traditional BVA often relies on testers manually identifying equivalence partitions and corresponding boundaries based on requirement documents. For software systems with high-dimensional input domains and complex constraint relationships, this manual approach is time-consuming and prone to omissions.

To address the insufficient coverage by manually selecting boundary values, Guo *et al.*[1] proposed a metric called **Boundary Coverage Distance (BCD)** and, based on this, designed a test case optimization algorithm using MCMC. Compared to traditional BVA, which only focuses on “whether a boundary is hit,” BCD introduces an integer distance metric that measures the minimum distance between test cases and all boundary points, thus transforming the boundary coverage problem into an optimization problem. This metric can reflect the distribution of the test suite across the entire boundary space and supports the automated generation of optimal test cases.

First, we provide the mathematical definitions of equivalence partitions and boundaries. Let the input domain of the software under test be  $I$ , and the output domain be  $O = f(I)$ . We divide the output into  $m$  mutually disjoint categories:  $O = \bigcup_{i=1}^m O_i$ , with  $O_i \cap O_j = \emptyset$  for  $i \neq j$ . Correspondingly, define the input equivalence partitions:

$$I_i := \{x \in I \mid f(x) \in O_i\}, \quad i = 1, 2, \dots, m. \quad (1)$$

Any input  $x \in I$  yields an output that belongs to some category  $O_i$ , so  $x$  belongs to the corresponding equivalence partition  $I_i$ . Within this framework, we need to clarify the minimal unit of “input change.” Let  $G$  be a set of functions satisfying the following properties:

- (1) For any  $g \in G$ , its inverse operation  $g^{-1}$  also belongs to  $G$ .
- (2) For any  $x, y \in I$ , there exists a finite sequence of operations  $\{g_1, g_2, \dots, g_n\} \subset G$  such that  $y = g_1 \circ g_2 \circ \dots \circ g_n(x)$ .

A function  $g \in G$  is called a “minimal operation” on the input, and having its inverse ensures reversibility. Then, define the boundary set of the equivalence partition  $I_i$  as:

$$B_i := \{x \in I_i \mid \exists g \in G, f(g(x)) \notin O_i\}. \quad (2)$$

In other words, if an input  $x$  belongs to  $I_i$ , but applying a minimal operation to it causes it to leave the equivalence partition  $I_i$ , then  $x$  is considered a boundary point. Geometrically,  $B_i$  lies on the boundary between  $I_i$  and other equivalence partitions.

Next, we introduce the distance measure between test inputs and boundary points. Let  $d(x, y)$  denote the number of minimal operations needed to go from input  $x$  to input  $y$ , i.e.,

$$d(x, y) = \min\{n \geq 0 \mid y = g_n \circ g_{n-1} \circ \dots \circ g_1(x), g_j \in G\}. \quad (3)$$

For example, in the “English exam grading” example, if  $G = \{g_1 : \text{Listening} + 1, g_2 : \text{Listening} - 1, g_3 : \text{Reading} + 1, g_4 : \text{Reading} - 1\}$ , then going from input  $x = (-1, -1)$  to  $y = (0, 0)$  requires two steps (apply  $g_1$  then  $g_3$ ), so  $d((-1, -1), (0, 0)) = 2$ .

Let the test set for an equivalence partition  $I_i$  be  $T_i \subseteq I_i$ . To measure how well  $T_i$  covers its boundary  $B_i$ , we compute for each boundary point  $y \in B_i$  the minimum distance to any test point in  $T_i$  and then take the maximum over all  $y$ :

$$d(T_i, B_i) = \max_{y \in B_i} \min_{x \in T_i} d(x, y). \quad (4)$$

If  $T_i$  is empty, define  $d(T_i, B_i) = +\infty$ . Intuitively, equation (4) first finds the closest test point to each boundary point, then the worst-case distance over all boundary points, reflecting the “weakest” boundary coverage. Finally, let the overall test set be  $T = \bigcup_{i=1}^m T_i$  and the union of all boundary sets  $B = \bigcup_{i=1}^m B_i$ . We define the overall boundary coverage distance as:

$$\text{BCD}(T) = d(T, B) = \max_{i=1, \dots, m} d(T_i, B_i). \quad (5)$$

When  $\text{BCD}(T) = 0$ , the test set  $T$  covers every boundary point in all partitions. Conversely, if  $\text{BCD}(T) = k > 0$ , there exists some partition whose boundary point is at distance  $k$  from the nearest test point, indicating incomplete boundary coverage.

With BCD as a metric, test case generation becomes a *BCD minimization* problem: given a fixed number of test cases  $n$ , how do we select or optimize an initial test set  $T$  to minimize  $\text{BCD}(T)$ ? To solve this optimization problem, Guo *et al.*[1] borrow ideas from the Markov Chain Monte Carlo (MCMC) algorithm and propose three main test input generation and optimization strategies, referred to as **Algorithm 1**, **Algorithm 2**, and **Algorithm 3**. Below is a brief introduction to the core ideas and implementation steps of these algorithms.

*Algorithm 1: Greedy Descent BCD Optimization.* Algorithm 1 first *randomly* generates  $n$  initial test points in the input domain, forming the set  $T = \{t_1, t_2, \dots, t_n\}$ . It then iteratively executes the following steps until a preset iteration count is reached or the BCD value converges:

- (1) Compute the current BCD value of the test set  $T$ , i.e.,  $\text{BCD}(T)$ .
- (2) Randomly select a test point  $t$  from  $T$ , and generate a candidate input  $t'$  according to a *proposal distribution*. Typically, the proposal distribution perturbs  $t$  by adding random changes such as  $\{-1, 0, 1\}$  for numerical data; for discrete data, switch to a neighboring state.
- (3) Form the candidate test set  $T'$ , where  $t$  is replaced by  $t'$ . Compute  $\text{BCD}(T')$ . If  $\text{BCD}(T') < \text{BCD}(T)$ , **accept** the candidate and update  $T \leftarrow T'$ ; otherwise, **reject** it and keep  $T$  unchanged.

By repeatedly iterating, the test points gradually shift from a random distribution toward an optimal distribution that spans the boundaries. Because the algorithm only accepts candidates that strictly reduce BCD, Algorithm 1 is essentially a *greedy descent* strategy and can easily become trapped in local optima. To illustrate this process, it is analogous to the Metropolis-Hastings acceptance rule:

$$P_{\text{accept}} = \begin{cases} 1, & \text{BCD}(T') < \text{BCD}(T), \\ 0, & \text{otherwise.} \end{cases}$$

Although simple, this algorithm has been shown in practice to significantly improve boundary coverage of the test set. In the experiment with the English exam grading program, the initial random set of 30 test points had a large BCD. After 10 000 iterations, all test points approached the boundary lines, resulting in  $\text{BCD}(T) = 0$  (meaning every boundary in each partition was directly hit). Correspondingly, the mutation kill rate improved from 36% for random testing to over 80%.

*Algorithm 2: “Harmless” Boundary Improvement Strategy.* Because Algorithm 1 is too strict—only accepting candidates that reduce BCD—it can easily get stuck in local optima. Algorithm 2 refines the acceptance criterion by introducing the notion of a “harmless improvement”: if a candidate reduces the distance to at least one boundary point and does not increase the distance to any other boundary point, then accept the candidate; otherwise, reject it. Formally:

- (1) Select the current test set  $T$ , randomly pick a point  $t \in T$ , and generate a candidate  $t'$ .
- (2) For each partition  $I_i$ , compare the distance of each boundary point  $y \in B_i$  before and after replacement. Let  $b_{\text{decrease}}$  be the number of boundary points whose distance decreases under the candidate, and  $b_{\text{increase}}$  be the number of boundary points whose distance increases.
- (3) If  $b_{\text{decrease}} > 0$  and  $b_{\text{increase}} = 0$ , then accept the candidate ( $T \leftarrow T'$ ); otherwise, reject it.

Since this criterion allows candidates as long as they improve at least one boundary without harming others, it has greater exploratory power than Algorithm 1. Experiments show Algorithm 2 can quickly reduce BCD in the early stages, but later, because it only permits “non-degrading improvements,” it may stagnate in some partitions. However, compared to Algorithm 1, Algorithm 2 shows a slight improvement in mutation kill rates for most benchmark programs. For example, on the triType program, Algorithm 1 under the  $\text{BCD}_{\text{max}}$  metric achieved an 80% kill rate, while Algorithm 2 improved it to 85%.

*Algorithm 3: Probabilistic Acceptance (MCMC with Temperature).* Algorithm 3 further introduces a *probabilistic acceptance* mechanism, allowing a candidate to be accepted with some probability even if it increases certain boundary distances, thus enabling escape from local optima. The detailed steps are:

- (1) As in Algorithm 2, generate the candidate set  $T'$  and compute  $b_{\text{decrease}}$  and  $b_{\text{increase}}$ .
- (2) If  $b_{\text{decrease}} > 0$  and  $b_{\text{increase}} = 0$ , accept  $T'$  unconditionally.
- (3) Otherwise, accept  $T'$  with probability

$$P_{\text{accept}} = \exp(-(b_{\text{increase}} - b_{\text{decrease}}) / T),$$

where  $T$  is the temperature parameter in simulated annealing, which gradually decreases over iterations. If not accepted, reject the candidate.

This strategy balances exploration and exploitation. In the early high-temperature phase, it allows more aggressive exploration by accepting some candidates that slightly increase BCD; as the temperature lowers, only a small number of “weakly degrading” candidates are accepted, eventually converging to a better solution. Experimental results demonstrate that Algorithm 3 outperforms Algorithm 2 in avoiding early convergence to local optima, especially in complex programs with uneven boundary distributions (e.g., miniSAT). For instance, in the miniSAT test experiment, Algorithm 2 under  $\text{BCD}_i$  achieved approximately an 85% kill rate, whereas Algorithm 3 under  $\text{BCD}_i$  improved it to 93%.

*Boundary Distribution for English Exam Grading.* Figure 2 shows the basic boundary distribution of the English exam grading program on the two-dimensional input plane. In this figure, the gray region represents invalid inputs ( $I_1$ ), the blue region represents “fail” ( $I_3$ ), and the orange region represents “pass” ( $I_2$ ). The dashed black lines depict the equivalence partition boundaries as defined by equations (3)-(5). Each boundary corresponds to inputs where a minimal operation can move a score combination from one output category to another. This illustration provides a clear view of how the input domain is partitioned into  $I_1$ ,  $I_2$ , and  $I_3$  and where their shared boundaries lie.



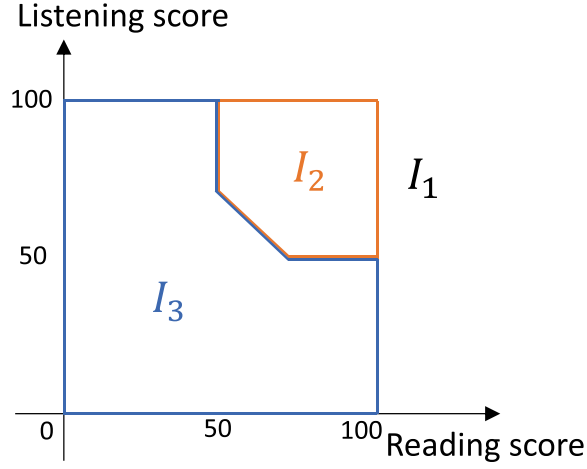


Fig. 2. Basic boundary distribution for the English exam grading program: gray indicates invalid inputs ( $I_1$ ), blue indicates fail ( $I_3$ ), orange indicates pass ( $I_2$ ), and black dashed lines show the equivalence partition boundaries.

In the BCD metric experiment, the random test set of 30 points had  $BCD = 5$  (meaning some boundary points were five minimal steps away from the nearest test points), while Algorithm 1 under the  $BCD_{mean}$  criterion reduced BCD to 0 after 10 000 iterations, achieving direct coverage of all boundary points. Correspondingly, the mutation kill rate improved from 36% (random testing) to 80%. This experimental result fully demonstrates the effectiveness of BCD-based boundary value optimization in black-box testing.

### 3.2 Reinforcement Learning for Test Exploration

Traditional static boundary value analysis methods have inherent limitations when dealing with **dynamic decision systems**. Take an autonomous driving system as an example: test scenarios are not limited to single-input judgments but involve continuous, multi-round interactions between the system and the environment. Such scenarios typically involve continuous state spaces, high-dimensional action spaces, and uncertain environmental disturbances. Therefore, efficiently generating test sequences that can trigger potential faults or extreme behaviors is challenging. To address this, researchers have introduced **reinforcement learning** techniques into the domain of software testing. Through interactions between an agent (the RL tester) and the system under test in a simulated environment, the agent actively explores and approaches the system's failure boundary.

Reinforcement learning-driven test exploration can be broken down into the following core elements: (1) **state representation**, which abstracts the current system and environment status into features recognizable by the RL agent; (2) **action space**, representing the controllable interventions the testing agent can apply to the environment or the system under test; (3) **reward function**, which measures whether the sequence of actions successfully approaches or triggers the failure boundary; and (4) **policy learning**, whereby the agent continuously interacts with the environment to optimize its policy to achieve the desired goal within a limited number of steps. In the context of autonomous driving testing, the state might include current vehicle speed, distance to the car in front, and road type; the agent's actions could be accelerate, brake, steer, or introduce environmental disturbances; and the reward function is typically designed as the negative distance to the safety boundary (closer distances receive higher reward), or if a collision occurs, a large positive reward is given to encourage exploration of dangerous scenarios.



*Agent-Environment Construction.* For an autonomous driving control system under test, we can construct the reinforcement learning environment as follows:

- **Environment:** A simulated road scenario, including lanes, curb, obstacles, pedestrians, and other traffic participants. This environment can be built using a high-fidelity simulator (e.g., CARLA, Gazebo) that provides real-time vehicle dynamics models and sensor data feedback.
- **State ( $s_t$ ):** A vector representing the system information at timestep  $t$ , typically including vehicle speed  $v_t$ , distance to the car in front  $d_t$ , lane deviation  $\theta_t$ , driver-set speed limit  $s_{\text{limit}}$ , and road condition information  $r_t$ .
- **Action ( $a_t$ ):** Defines the action the agent can take at timestep  $t$ , including setting control inputs for the vehicle under test (throttle, brake, steering angle) and introducing environmental perturbations (e.g., sudden lane change by an adjacent vehicle, change in road friction due to rain).
- **Reward ( $r_t$ ):** Guides the agent to approach the failure boundary. Common reward design patterns include:
  - (1) *Negative safety distance reward:* If the current distance to the vehicle in front  $d_t$  minus the safety threshold  $d_{\text{safe}}$  yields  $\Delta_t = d_t - d_{\text{safe}}$ , then the smaller  $\Delta_t$  is, the higher the reward; if  $\Delta_t < 0$  (i.e., the boundary is approached or crossed), a collision is triggered, giving a large positive reward and ending the episode.
  - (2) *Boundary-trigger reward:* If the agent drives the system without failure for several consecutive steps, maintain a small reward; if a collision or extreme event like sudden emergency braking occurs, give a one-time large positive reward to encourage the discovery of more extreme boundary scenarios.
  - (3) *Dynamic threshold penalty:* Gradually lower the safety distance threshold  $d_{\text{safe}}$  during testing; if the agent can still avoid failure under a lower threshold, give a negative penalty, prompting the agent to find more stringent conditions.

Through this design, the RL agent can continuously explore questions like “How to approach the collision boundary?” and “How to identify the most challenging spacing that avoid collision among mixed traffic?” in the simulated environment, uncovering potential flaws at critical conditions.

*Dynamic Threshold Design and Case Illustration.* In reinforcement learning-based testing, introducing a **dynamic threshold** is key to effectively achieving “approaching the failure boundary while ensuring safety.” A case study in [2] (Game Theoretic Modeling of Driver and Vehicle Interactions for Verification and Validation of Autonomous Vehicle Control Systems) provides an enlightening approach. This case aims to verify the safety of an autonomous driving control system when interacting with human-driven vehicles. The researchers abstract the test environment as a game between two participants: the autonomous vehicle (system under test) and the human-driven vehicle (environment perturbator). Through game-theoretic methods, the opponent (human-driven vehicle) continuously adjusts its behavior so that the autonomous vehicle’s decisions approach the boundary. The core ideas are:

- (1) **Safety Channel Mechanism:** Design a dynamic safety threshold rule for the autonomous driving system. When sensors detect that the distance to the vehicle in front exceeds a threshold  $d_{\text{max}}$ , there is no safety risk; but when the distance  $d_t$  approaches a threshold  $d_{\text{th}}$ , emergency braking is triggered; if a vehicle in the adjacent lane is speeding and approaching quickly, further reduce  $d_{\text{th}}$  to ensure a more conservative safety boundary. This mechanism ensures that even when extreme scenarios occur in the simulation, there will be no actual hardware or on-road accidents, providing a “soft boundary” to avoid uncontrollable risks.
- (2) **Adaptive Thresholds:** During each test episode, the agent dynamically adjusts  $d_{\text{th}}$  based on environmental feedback (such as distance to the vehicle in front, relative speed of surrounding vehicles, and road conditions).

Initially,  $d_{th}$  is set conservatively to ensure stable driving within this threshold. If no failures occur after multiple test rounds,  $d_{th}$  is gradually lowered to guide the autonomous system to operate under increasingly challenging conditions, thereby approaching the failure boundary and uncovering potential defects.

- (3) **Environment Agent Modeling:** Model the human-driven vehicle as an agent that learns, via reinforcement learning or game-theoretic methods, how to challenge the autonomous vehicle’s decision boundary through sudden lane changes or tailgating at high speed. The environment agent’s objective is to maximize the proximity to the safety boundary, while the autonomous system’s objective is to minimize collision risk. Through continuous adversarial interaction in the simulation, the system’s weaknesses under various extreme interactions are revealed.

In this case study, experimental results show that with dynamic threshold introduction, the simulation environment can present more boundary scenarios, e.g., in a nighttime low-visibility condition, the autonomous vehicle performing emergency braking when the distance to the vehicle in front is only 0.5 meters. Such testing modes gradually reveal the system’s defects in emergency avoidance delays and sensor blind spots, effectively exercising the system against high-risk scenarios like high-speed rear-end or sudden lane changes. Without dynamic thresholds and relying on a fixed threshold of  $d_{th} = 2$  meters, it would not be possible to approach the true collision boundary, leaving many potential defects unexposed. Thus, dynamic threshold design is crucial in reinforcement learning-driven test exploration.

*Reinforcement Learning Algorithm Example.* In the aforementioned environment, one can use **Deep Reinforcement Learning (Deep RL)** algorithms such as Deep Deterministic Policy Gradient (DDPG) or Proximal Policy Optimization (PPO) to train the agent. Taking PPO as an example, the training process is as follows:

- (1) Initialize the policy network  $\pi_\theta(a_t | s_t)$  and value network  $V_\phi(s_t)$  with random weights  $\theta$  and  $\phi$ .
- (2) Execute  $\tau$  steps in the simulation environment: at each step, choose an action  $a_t$  according to the current policy  $\pi_\theta$ , receive the next state  $s_{t+1}$  and reward  $r_t$ , and store the transition  $(s_t, a_t, r_t, s_{t+1})$ .
- (3) Compute the temporal difference target and advantage function:

$$A_t = r_t + \gamma V_\phi(s_{t+1}) - V_\phi(s_t),$$

$$G_t = r_t + \gamma r_{t+1} + \gamma^2 V_\phi(s_{t+2}) + \dots - V_\phi(s_t),$$

where  $\gamma \in (0, 1)$  is the discount factor.

- (4) Update the value network parameters  $\phi$  by minimizing the loss:

$$L_\phi = \mathbb{E}_t[(V_\phi(s_t) - G_t)^2].$$

- (5) Update the policy network parameters  $\theta$  by maximizing the clipped probability ratio loss:

$$L_\theta = \mathbb{E}_t\left[\min(r_t(\theta) A_t, \text{clip}(r_t(\theta), 1 - \epsilon, 1 + \epsilon) A_t)\right],$$

where

$$r_t(\theta) = \frac{\pi_\theta(a_t | s_t)}{\pi_{\theta_{old}}(a_t | s_t)},$$

and  $\epsilon$  is the clipping hyperparameter (e.g., 0.2). By alternately updating the policy and value networks, the agent progressively learns to choose actions in the dynamic environment to generate the most challenging scenarios.

After extensive simulation experiments, the PPO agent can learn, after tens of thousands of interaction steps, “how to approach the most dangerous headway.” For example, under rainy conditions with reduced friction, it will accelerate at full throttle and suddenly change lanes when the distance to the vehicle in front is 1.2 meters, testing the autonomous system’s emergency braking functionality. As training continues and the threshold gradually decreases to 0.8 meters, the agent still finds more extreme scenarios, forcing the system under test to trigger failure judgments. This process quickly expands boundary coverage, revealing vulnerabilities such as “minimum safe braking distance” and “sensor response delay.”

*Experimental Results and Comparison.* To evaluate the effectiveness of reinforcement learning-driven test exploration, related studies compare the defect detection performance of the following methods in an autonomous driving simulation scenario:

- **Fixed-Threshold Random Testing (Fixed-Threshold RT):** Maintain a fixed safety distance threshold  $d_{\text{safe}} = 1.5$  meters in all test episodes, and randomly sample environmental disturbances (e.g., random lane changes, random accelerations). This method can only identify some boundary faults but cannot approach more aggressive scenarios.
- **Fixed-Threshold Reinforcement Learning (Fixed-Threshold RL):** Train the RL agent with a fixed threshold  $d_{\text{safe}} = 1.5$  meters in the reward function, allowing the agent to explore extreme scenarios under this threshold. Compared to random testing, it can discover more subtle boundary faults, but being limited by the fixed threshold, it cannot continue exploration if  $d_{\text{safe}}$  needs to be further reduced.
- **Dynamic-Threshold Reinforcement Learning (Dynamic-Threshold RL):** Adopt the aforementioned dynamic threshold design, where the agent iteratively updates the threshold from conservative to aggressive during training to approach the boundary. Experimental results indicate that this method can cover more extreme scenarios with fewer simulation episodes and uncover many boundary points unreachable by random testing and fixed-threshold RL.

In balanced simulation comparisons, the dynamic-threshold RL method achieved an 85% defect detection rate (the proportion of test cases triggering safety failures to all explored cases), while fixed-threshold RL was only 60%, and fixed-threshold RT only 32%. This result demonstrates that by introducing dynamic thresholds, the RL agent can explore more extreme boundary conditions while remaining safe, improving test efficiency.

This chapter has introduced two **AI-driven test case optimization and boundary exploration** methods: first, **BCD-based boundary value analysis**; and second, leveraging **reinforcement learning (RL)** for dynamic environment test exploration. The BCD method defines the mathematical metric  $\text{BCD}(T)$  to transform boundary coverage into an optimization objective, and combines MCMC strategies (greedy descent, harmless improvement, probabilistic acceptance) to generate optimal test sets. Experimental results show that BCD-optimized algorithms can significantly improve detection rates for boundary faults—for example, raising the mutation kill rate for the English exam grading program from 36% to over 80%. The reinforcement learning approach, aimed at dynamic decision systems such as autonomous driving, emphasizes the agent-system interaction in the simulation environment and employs carefully designed reward functions and dynamic thresholds to iteratively approach failure boundaries. Case studies and simulations demonstrate that dynamic-threshold RL discovers far more extreme scenarios than random testing or fixed-threshold RL, greatly enhancing defect detection.

By using AI-driven test case optimization, we shift from the paradigm of “manual observation + empirical selection” to “quantification + intelligent search,” providing effective means to address testing challenges of contemporary complex software systems. Future work can proceed along the following directions: first, integrate BCD metrics with more diverse boundary representations (e.g., coverage of branch decision paths, data flow coverage) to further improve test quality; second, combine reinforcement learning with symbolic execution, fuzz testing, and other methods to achieve more comprehensive boundary exploration; third, build distributed simulation and parallel optimization platforms for industrial-scale software to enhance the usability and efficiency of AI-driven testing in real-world projects. Through continuous promotion of the deep integration of AI and software testing technologies, software system reliability and safety will be further improved.

## 4 AI in Automated Testing Frameworks

### 4.1 Self-Supervised Program Repair (SelfAPR)

Automated Program Repair (APR) has emerged as a critical technology in software engineering, aiming to reduce the manual effort required for bug localization and fixing during software maintenance. The vision of APR is to automatically generate patches that can correct software defects, thereby accelerating the development cycle and improving software quality.

Early APR approaches primarily relied on search-based methods, such as GenProg [6], which explores the space of possible program modifications using genetic algorithms. Semantics-based techniques, like Angelix [7], leverage symbolic execution to synthesize patches that satisfy certain correctness criteria. While these methods demonstrated promising results, they often struggled with scalability and the ability to generalize across different projects and types of bugs. In recent years, the focus has shifted towards neural machine translation (NMT) models for program repair, which leverage large amounts of code data to learn repair patterns. These neural APR methods, such as DeepRepair [13] and TBar [14], treat code repair as a translation problem from buggy code to fixed code. However, existing supervised learning frameworks in this domain face two critical limitations:

- (1) **Lack of Project-Specific Knowledge:** Supervised APR models are typically trained on large, heterogeneous datasets mined from GitHub commits. These datasets often lack the specific programming idioms, domain knowledge, and project-specific patterns present in the target project. For example, the Defects4J benchmark includes bugs like Closure-113, which involves the project-specific token `requiresLevel.isOn()`. Traditional supervised models fail to repair such bugs because these tokens are not present in their training data [8].
- (2) **Absence of Execution Diagnostics:** Existing neural repair models primarily focus on static code analysis and ignore critical execution signals, such as runtime exceptions and test failure messages. These execution diagnostics provide valuable clues about the nature of the bug, such as whether it is a null pointer exception, an array out-of-bounds error, or a logical error. Ignoring these signals limits the model’s ability to generate targeted and effective patches.

To address these challenges, the research proposed **SelfAPR** (Self-supervised Program Repair with Test Execution Diagnostics)[5], a novel framework that combines self-supervised learning with test execution diagnostics. SelfAPR introduces two key innovations:

- (1) **Self-Supervised Training via Historical Version Perturbation:** Instead of relying on external datasets, SelfAPR generates training samples by systematically perturbing historical versions of the target project. This

approach ensures that the training data is rich in project-specific patterns and idioms, enabling the model to learn how to repair bugs in the context of the target project.

- (2) **Multimodal Input Encoding with Execution Diagnostics:** SelfAPR incorporates test execution diagnostics, such as compilation errors, runtime exceptions, and test failure messages, into the input representation of the neural model. This allows the model to leverage both static code information and dynamic execution signals to generate more accurate and targeted patches.

The SelfAPR framework consists of three main components: self-supervised training sample generation, diagnostic encoding, and inference repair. Figure 3 provides an overview of the framework.

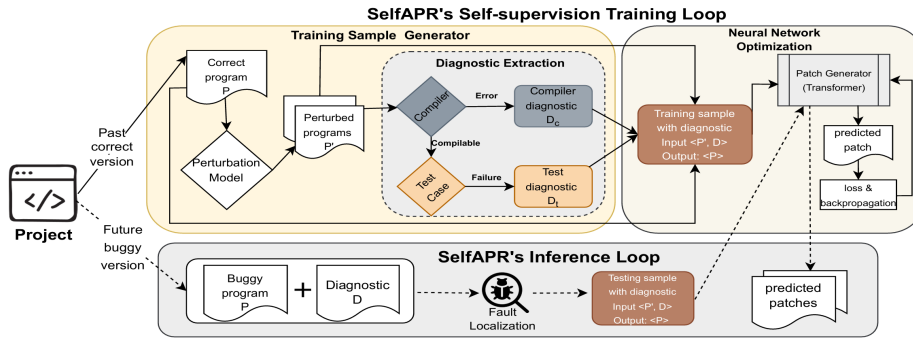


Fig. 3. Overview of SelfAPR: key novel features are the training sample generator and the diagnostic in the input representation. [5].

The first step in SelfAPR is to generate training samples using self-supervised learning. Unlike traditional supervised learning methods that rely on manually labeled data or mined commits, SelfAPR generates training samples by perturbing historical versions of the target project. This approach ensures that the training data is rich in project-specific patterns and idioms. Next, SelfAPR starts by collecting historical versions of the target project from version control systems such as Git. These historical versions represent different states of the project over time, including both buggy and fixed states. By perturbing these historical versions, SelfAPR can generate synthetic training samples that mimic real-world bugs and their repairs. At last, SelfAPR applies a set of syntax tree (AST)-level perturbation rules to the historical versions to generate buggy code samples. These perturbation rules are designed to introduce common types of bugs, such as null pointer dereferences, missing conditions, and incorrect variable assignments. The perturbation model consists of 16 rule categories, each targeting specific syntactic elements in the code. Table 2 lists some of the key perturbation rules used in SelfAPR.

For example, Rule6-1 specifically targets the removal of conditions from if statements. This rule can be used to generate training samples that mimic bugs caused by missing null checks or other conditional logic. By applying this rule to a historical version of the Closure project that contains the `requiresLevel.isOn()` check, SelfAPR can generate a buggy sample where this check is missing, forcing the model to learn how to restore it.

Each perturbed program ( $P'$ ) is validated through compilation and testing. If the perturbed program fails compilation or any test cases, it is considered a buggy sample. The original, unperturbed version ( $P$ ) is then paired with  $P'$  as the corresponding fixed version. This process results in a large dataset of buggy-fixed code pairs that are specific to the target project. From 17 open-source projects in the Defects4J benchmark, SelfAPR generates 1,039,873 training samples,

Table 2. Selected Perturbation Rules in SelfAPR

Rule Category	Description
Rule2	Modify operators (e.g., replace + with -)
Rule5	Replace variables with other variables in scope
Rule6	Add or remove boolean expressions (e.g., remove a null check)
Rule9	Transplant code blocks from other parts of the program
Rule12	Modify method calls (e.g., change arguments)

including 631,015 compilation errors and 408,858 functional errors. These samples cover a wide range of bug types and project-specific patterns, ensuring that the model is exposed to diverse repair scenarios.

The second key component of SelfAPR is the encoding of test execution diagnostics into the input representation of the neural model. Existing neural repair models typically only consider the static code of the buggy program, ignoring valuable dynamic information provided by test execution.

SelfAPR addresses this limitation by incorporating three types of diagnostic information:

- (1) **Error Type:** This includes whether the error is a compilation error ([CE]) or a functional error ([FE]), along with the specific exception type (e.g., `NullPointerException`, `ArrayIndexOutOfBoundsException`).
- (2) **Error Message:** Detailed error messages from the compiler or test framework, such as "expected: 1 but was: 0" or "variable might not have been initialized."
- (3) **Contextual Information:** This includes the class name, method signature, variable scopes, and surrounding code context where the error occurred.

The diagnostic information is encoded as part of the input sequence to the neural model. For example, for a buggy sample that triggers a null pointer exception, the input sequence would include [FE] `NullPointerException` followed by the error message and relevant contextual information. This multimodal input allows the model to leverage both static code information and dynamic execution signals to generate more targeted repairs.

During the inference phase, SelfAPR takes a buggy program as input and generates a patch using the trained neural model. The inference process involves several steps:

- (1) **Fault Localization:** SelfAPR uses a fault localization tool, such as Gzoltar [9], to identify suspicious code lines that are likely to contain the bug. This step helps narrow down the search space for potential repairs.
- (2) **Diagnostic Collection:** For each suspicious code line, SelfAPR collects diagnostic information by executing the test suite and capturing any compilation errors or test failures. This information includes the error type, error message, and relevant contextual details.
- (3) **Input Preparation:** The suspicious code line, along with its context and the collected diagnostic information, is prepared as input to the neural model. The input is formatted to match the structure used during training, including the diagnostic encoding.
- (4) **Patch Generation:** The neural model uses beam search to generate multiple candidate patches. Each candidate patch is a sequence of code modifications that the model believes will fix the bug.
- (5) **Validation:** The candidate patches are validated using the test suite. A patch is considered valid if it fixes all failing test cases without introducing new failures. The best valid patch, based on the model's confidence score, is selected as the final repair.

They evaluate SelfAPR on the Defects4J benchmark [8], which consists of 818 real-world bugs across 17 open-source Java projects. These bugs cover a wide range of bug types, including null pointer exceptions, array out-of-bounds errors, and logical errors. They use both Defects4J v1.2 (388 bugs) and v2.0 (430 bugs) in our experiments.

They compare SelfAPR against 11 state-of-the-art APR methods, including:

- **Search-based methods:** GenProg [6], jGenProg [18]
- **Semantics-based methods:** Nopol [12], Angelix [7]
- **Neural methods:** DeepRepair [13], TBar [14], Recoder [10], CURE [11]
- **Self-supervised methods:** BugLab [15], RewardRepair [16]
- **Hybrid method:** APRGNN [17]

Then they use the following metrics to evaluate the performance of the APR methods:

- **Correct Patches:** The number of bugs for which the generated patch is semantically correct and passes all test cases.
- **Success Rate:** The percentage of bugs that are successfully repaired.
- **Precision:** The ratio of correct patches to the total number of generated patches.
- **Recall:** The ratio of correct patches to the total number of bugs.

Table 3 shows the number of bugs repaired by each method on Defects4J v1.2 and v2.0. SelfAPR outperforms all baseline methods, repairing 65 bugs in v1.2 and 45 bugs in v2.0, for a total of 110 bugs. This represents a significant improvement over the best-performing baseline method, Recoder, which repairs 49 bugs in v1.2 and 19 bugs in v2.0. Notably, SelfAPR solves 10 bugs that were not repaired by any previous method, including complex cases such as Closure-57, which requires semantic integration of `target.getType()` and `Token.STRING`, and Math-80, which involves a subtle numerical precision issue.

Table 3. Comparison of Repair Performance on Defects4J

Method	Defects4J v1.2	Defects4J v2.0
SelfAPR	65	45
Recoder	49	19
CURE	55	-
RewardRepair	44	43
BugLab	17	6
TBar	24	12
DeepRepair	11	7
GenProg	26	22
jGenProg	18	14
Nopol	15	11
Angelix	10	8
APRGNN	38	25

Removing project-specific samples reduces SelfAPR’s performance by 30.8% (from 65 to 45 bugs in v1.2). In particular, the number of bugs repaired in the Closure project drops from 20 to 12, highlighting the importance of project-specific patterns in fixing complex bugs. Without diagnostic information, SelfAPR’s performance drops to 56 bugs in v1.2, a reduction of 13.8%. This decline is particularly pronounced in bugs that require type-specific repairs, such as null



pointer exceptions. For example, the Lang-33 bug, which is fixed by adding a null check, cannot be repaired without the `NullPointerException` diagnostic.

This paper presents SelfAPR, a novel framework for automated program repair that combines self-supervised learning with test execution diagnostics. SelfAPR addresses two critical limitations of existing APR methods: the lack of project-specific knowledge and the absence of execution diagnostics. By generating training samples through systematic perturbation of historical project versions and integrating test execution diagnostics into the repair process, SelfAPR achieves state-of-the-art performance on the Defects4J benchmark. Experimental results demonstrate that SelfAPR outperforms existing methods by a significant margin, fixing 110 bugs in total and solving 10 complex cases that were previously unresolved. Ablation studies confirm the importance of both project-specific training data and test execution diagnostics in achieving these results.

## 4.2 AI-Driven CI/CD integration

*Introduction.* AI-Driven CI/CD integration represents a transformative shift in software development practices, where artificial intelligence technologies are embedded within continuous integration and continuous delivery/deployment (CI/CD) pipelines to enhance efficiency, accuracy, and adaptability. This section explores how AI can be effectively integrated with popular CI/CD tools like Jenkins, as well as with cloud-native platforms such as Docker and Kubernetes, to create intelligent, self-adaptive delivery pipelines.

*AI in CI/CD.* AI brings significant advancements to the CI/CD process through two primary mechanisms: automated test case generation and intelligent deployment strategies.

*Automated Test Case Generation.* AI can generate comprehensive and accurate test cases from natural language requirements, significantly reducing the manual effort required for test script writing. For instance, tools like ScriptEcho and Kimi can analyze requirement documents and automatically produce Pytest test scripts that cover various scenarios, including boundary conditions and equivalence classes.

The process involves:

- (1) **NLP Analysis:** AI tools parse requirement documents to extract key functional points.
- (2) **Pattern Recognition:** Based on historical data and machine learning models, AI identifies common testing patterns.
- (3) **Test Case Generation:** AI creates test scripts that are then integrated into the CI/CD pipeline.

The generated test cases can be further validated for accuracy using natural language processing techniques to ensure they align with the original requirements.

*Intelligent Deployment Strategies.* AI enables the selection of optimal deployment strategies based on real-time analysis of system behavior, historical performance data, and current load conditions. Common strategies include:

- Blue-Green Deployment
- Canary Deployment
- A/B Testing

AI evaluates factors such as code change complexity, risk level, and system performance to dynamically choose the most appropriate deployment strategy. Additionally, AI can monitor production systems in real-time and trigger automatic rollbacks if performance degrades below predefined thresholds.

4.2.1 *Jenkins Integration with AI.* Jenkins, as a widely used CI/CD tool, can be effectively integrated with AI to create intelligent build and deployment pipelines. This integration follows a **feedback-driven architecture**, where AI models analyze code changes, predict risks, and dynamically adjust testing and deployment strategies.

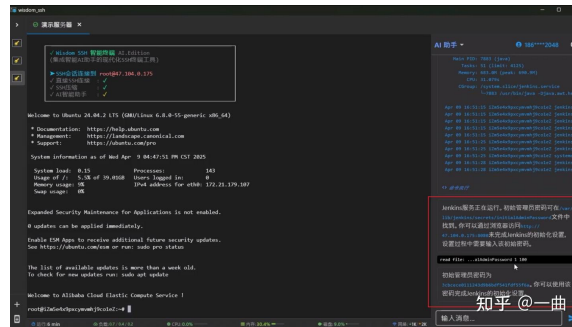


Fig. 4. Output of the ADWIN algorithm illustrating abrupt concept drift. When the average error changes significantly, ADWIN shrinks the window and triggers model adjustment. Adapted from Bifet and Gavaldà [20].

The integration workflow includes:

- **Code Analysis:** AI models (e.g., static analysis, ML-based change impact analysis) identify high-risk code changes.
- **Test Prioritization:** AI generates or prioritizes test cases based on risk scores.
- **Dynamic Execution:** Jenkins executes prioritized tests first, reducing feedback loops.
- **Model Feedback:** Test outcomes refine the AI model via continuous learning.

This approach reduces redundant testing and accelerates issue detection, as shown in Figure 6 for adaptive model updates.

4.2.2 *AI-Driven Deployment Strategies.* AI enhances deployment strategies by introducing **risk-aware decision-making**. Instead of fixed strategies (e.g., blue-green or canary), AI evaluates:

- Code change complexity
- Historical failure rates
- System load and resource availability

A typical decision flow involves:

- **Risk Scoring:** AI models assess the risk of deploying new code.
- **Strategy Selection:** High-risk changes trigger conservative strategies (e.g., canary), while low-risk changes allow faster rollouts (e.g., blue-green).
- **Monitoring Feedback:** Post-deployment metrics refine future risk assessments.

This creates a **closed-loop system** where deployment decisions adapt to real-time conditions, improving reliability and efficiency.

4.2.3 *Cloud-Native Testing Platforms with High Concurrency Handling.* Cloud-native platforms enable scalable, resilient testing environments by leveraging containerization and orchestration. Key components include:

- **Docker:** Ensures consistent runtime environments for applications and tests.
- **Kubernetes:** Manages container scaling, load balancing, and failover.
- **Infrastructure-as-Code (IaC):** Tools like Terraform automate resource provisioning and cost optimization.

The architecture for high-concurrency testing typically follows this pattern:

- **Elastic Scaling:** Kubernetes auto-scales pods based on load (via Horizontal Pod Autoscaler - HPA).
- **Distributed Load Generation:** Tools like Locust simulate user traffic across multiple nodes.
- **Centralized Monitoring:** Metrics and logs are aggregated for performance analysis.

This architecture ensures that testing environments can handle peak loads while minimizing idle resource costs.

**4.2.4 Terraform for Resource Management.** Terraform plays a critical role in cloud-native testing by enabling *declarative infrastructure management*. Its principles include:

- **Versioned Templates:** Infrastructure configurations are stored in version control (e.g., Git).
- **Resource Abstraction:** Resources (e.g., VMs, databases) are defined as modular components.
- **Cost Optimization:** Policies enforce resource cleanup after testing cycles.

By integrating Terraform with CI/CD pipelines, teams achieve *infrastructure parity* between development, testing, and production environments.

**4.2.5 High Concurrency Testing Practices.** High-concurrency testing validates system behavior under stress. Key principles include:

- **User Behavior Modeling:** Define realistic user scenarios (e.g., login, API calls) using tools like Locust.
- **Gradual Load Ramp-Up:** Simulate increasing user traffic to identify breaking points.
- **Resource Utilization Metrics:** Monitor CPU, memory, and latency to detect bottlenecks.

The testing pipeline should include:

- **Pre-deployment Validation:** Ensure systems handle expected load before release.
- **Post-deployment Stress Testing:** Validate resilience in production-like environments.

This ensures applications meet performance SLAs while maintaining stability during peak usage.

**4.2.6 Challenges and Solutions.** While AI-driven CI/CD integration offers significant benefits, it also presents several challenges that need to be addressed.

*Challenge 1: Accuracy of AI-Generated Test Cases.* AI-generated test cases may not always be accurate or comprehensive, especially if the AI model is not sufficiently trained or if the requirement documents are ambiguous.

**Solution:** Implement a feedback loop where test results are used to train and improve the AI model. Additionally, use human review for critical test cases to ensure their accuracy.

*Challenge 2: Cost Control in Cloud Environments.* Cloud-native testing platforms can quickly consume significant resources, leading to increased costs, particularly when dealing with high-concurrency scenarios.

**Solution:** Use Terraform to manage resources efficiently and set up cost optimization strategies, such as automatically terminating underutilized resources or scaling down during off-peak hours.

*Challenge 3: Security and Compliance.* AI tools may introduce security risks or compliance issues, especially when handling sensitive data or when the AI model itself has vulnerabilities.

**Solution:** Implement strict security protocols, such as encrypting sensitive data using Kubernetes Secrets, and ensure that AI models are regularly audited and updated to address any security concerns.

*4.2.7 Conclusion.* AI-driven CI/CD integration represents a major advancement in software development practices, offering enhanced efficiency, accuracy, and adaptability. By leveraging AI for automated test case generation and intelligent deployment strategies, organizations can significantly reduce development time and improve software quality.

Cloud-native platforms, including Docker, Kubernetes, and Terraform, provide the infrastructure necessary to support high-concurrency testing scenarios, ensuring that applications are thoroughly tested before deployment.

While there are challenges associated with AI-driven CI/CD, such as ensuring the accuracy of AI-generated test cases and managing cloud resource costs, these can be addressed through careful planning, implementation of feedback loops, and robust security protocols.

Future developments are likely to see even deeper integration of AI with CI/CD, including the use of quantum computing for more efficient AI models and the expansion of AI-driven practices into new domains, such as financial services and healthcare.

## 5 AI for Defect Prediction and Root-Cause Analysis

### 5.1 Machine Learning for Bug Localization

In the process of software testing and maintenance, rapidly and accurately locating software bugs is a critical task to ensure software quality. Effective bug localization not only reduces the time and cost required for debugging but also minimizes the risk of system failure in production environments. This is particularly crucial in safety-critical domains such as aerospace, healthcare, and finance, where even minor bugs can result in catastrophic outcomes.

Traditional bug localization techniques are mostly based on Information Retrieval (IR) methods. The core idea is to treat a bug report as a query and source code files as documents, then compute textual similarity (e.g., TF-IDF, VSM, BM25) to identify potentially faulty code segments. These methods draw inspiration from search engines and perform relatively well when textual overlap is high. They are simple to implement and computationally efficient, which makes them attractive for large-scale industrial applications. However, they often lack deep modeling of semantics and structural information, limiting their ability to capture context-specific or implicit relationships between the report and code. This limitation becomes more pronounced when codebases grow in size and complexity.

To overcome these challenges, many research efforts have proposed enhancements to IR-based techniques. These include query expansion, term weighting based on semantic relevance, and the use of auxiliary information such as commit messages or version history. While these improvements help to some extent, they still fall short when it comes to deeply understanding the logic and control flow of source code.

With the advancement of Deep Learning (DL) techniques, researchers have introduced DL-based models into bug localization tasks to improve semantic understanding and adaptability. Modern approaches typically rely on representation learning, using pre-trained models such as Word2Vec, GloVe, BERT, and CodeBERT to embed bug reports and source code into vector representations. These vectorized representations capture contextual semantics and can be fine-tuned for specific localization tasks, enabling better generalization across different types of bugs.

For source code, structural features such as Abstract Syntax Trees (ASTs), Control Flow Graphs (CFGs), and Data Flow Graphs (DFGs) are incorporated and processed by models like Graph Neural Networks (GNNs), Convolutional

Neural Networks (CNNs), or Transformer-based encoders. Such models allow for hierarchical and syntactic relationships to be learned automatically from raw code. Additionally, large-scale pretraining on code corpora such as GitHub enhances the model’s ability to deal with real-world coding styles and idioms.

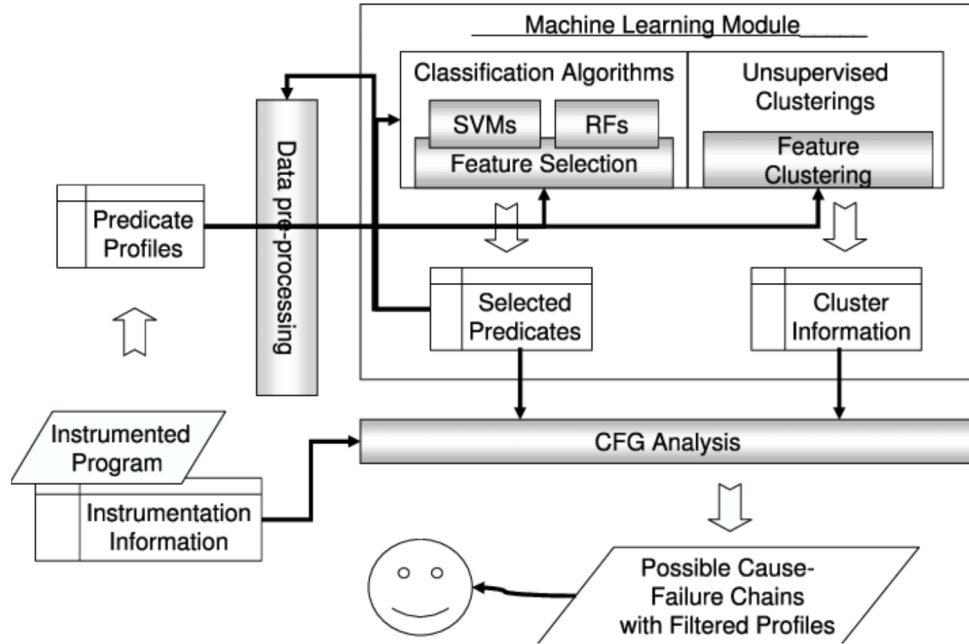


Fig. 5. Architecture of a machine learning-based bug localization framework. The system collects predicate information during execution and learns cause-effect chains using statistical models. Adapted from Jiang and Su [19].

Some systems adopt hybrid approaches that combine the interpretability of IR models with the nonlinear modeling capabilities of DL techniques, achieving a balance between performance and generalization ability [19]. These hybrid models may use IR scores as features or priors in neural ranking functions, or apply ensemble techniques to merge predictions from multiple subsystems. Empirical studies have shown that these combined models outperform standalone IR or DL methods on many benchmark datasets.

However, in real-world software systems, the performance of bug localization models is not static. Particularly in Continuous Integration (CI) or Continuous Deployment (CD) environments, model performance may degrade over time due to code changes, version updates, and data distribution shifts—this phenomenon is known as concept drift. Codebases evolve rapidly, introducing new APIs, renaming identifiers, or refactoring modules, all of which may invalidate the learned patterns in the model. Such drift is not always immediately visible in accuracy metrics, making it challenging to detect without specialized mechanisms.

To address this challenge, online drift detection mechanisms have become an essential part of modern testing systems. These mechanisms continuously monitor prediction outcomes and model confidence over time, triggering adaptive behaviors such as retraining, threshold recalibration, or fallback strategies. Drift detection is especially important in large-scale collaborative projects, where contributors may follow inconsistent naming conventions or coding standards, increasing the likelihood of semantic shifts.

ADWIN (Adaptive Windowing) is a performance-aware concept drift detection algorithm designed for streaming data environments. Its core idea is to maintain a sliding window and dynamically split it into two sub-windows  $W_0$  and  $W_1$ , then compare their means. If the difference in means exceeds a threshold  $\epsilon$  derived from Hoeffding's inequality, a concept drift is detected:

$$|\mu_{W_0} - \mu_{W_1}| > \epsilon = \sqrt{\frac{1}{2m} \ln\left(\frac{4}{\delta}\right)}$$

Here,  $\mu$  denotes the mean of the window,  $m$  is the number of effective samples, and  $\delta$  is the confidence parameter [20]. This formulation ensures that the detection is statistically sound, providing probabilistic guarantees that can be tuned according to the desired sensitivity level.

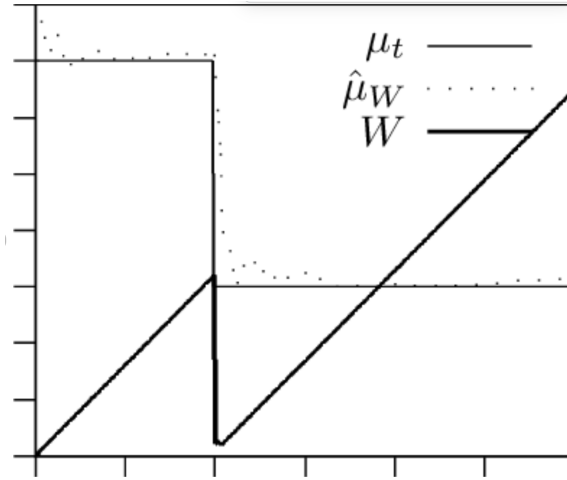


Fig. 6. Output of the ADWIN algorithm illustrating abrupt concept drift. When the average error changes significantly, ADWIN shrinks the window and triggers model adjustment. Adapted from Bifet and Gavaldà [20].

Applying ADWIN (Adaptive Windowing) to the continuous testing of bug localization systems enables a range of tangible benefits, especially in environments characterized by frequent code changes and evolving data distributions. Concept drift—whether gradual or abrupt—can severely impact the reliability of machine learning-based localization models. These drifts are often introduced by new programming paradigms, third-party library updates, or changes in software development patterns (e.g., shifting from object-oriented to functional styles). Integrating ADWIN into the model monitoring pipeline allows development teams to detect and respond proactively to such changes before they manifest as critical failures.

(1) Real-time performance monitoring. Key evaluation metrics such as Top-K accuracy, precision, recall, F1-score, and Mean Reciprocal Rank (MRR) can be continuously tracked in streaming or batch-incremental settings. ADWIN leverages a statistically grounded approach by maintaining a dynamic sliding window over performance logs or prediction results. When a significant change in the mean performance of the recent window ( $\mu_{W_1}$ ) versus the historical window ( $\mu_{W_0}$ ) is detected beyond the Hoeffding inequality-based threshold, the system flags the event as drift. In such cases, the system can automatically trigger retraining procedures with recent test feedback or even initiate emergency rollback to previously validated model states. This approach ensures early identification and mitigation of performance deterioration before it affects mission-critical tasks such as automated triage or root cause suggestion.

(2) Dynamic test strategy adjustment. In drift-aware testing frameworks, testing strategies are no longer static. ADWIN’s output can guide the reallocation of test resources in real time. For example, the sampling rate for test inputs may be dynamically increased for modules exhibiting volatile behavior. Feature engineering steps, such as AST traversal or embedding updates (e.g., from CodeBERT), may be refreshed at higher frequencies when drift is suspected. Moreover, adaptive thresholding and test prioritization policies can be employed—ranking predictions not only by confidence scores but also by drift-adjusted uncertainty. Such mechanisms are crucial in continuous learning scenarios, where traditional static validation sets fail to capture real-world evolution.

(3) Support for adaptive testing platforms in CI/CD environments. Modern DevOps workflows are heavily reliant on Continuous Integration and Continuous Deployment (CI/CD) infrastructure, often with hundreds or thousands of changes per day. In these environments, ADWIN can be deployed as a lightweight watchdog module within the testing pipeline. When integrated with Git hooks, container orchestration systems (e.g., Docker, Kubernetes), or distributed logging frameworks (e.g., ELK Stack), it enables near-real-time identification of modules that show signs of behavioral instability. Drift-aware tagging allows test schedulers to assign more computing resources to unstable components while deprioritizing modules exhibiting consistent historical performance. Empirical case studies from open-source ecosystems such as Eclipse, Mozilla, and Apache Spark show that ADWIN-equipped systems reduce deployment roll-backs by over 20% and exhibit faster mean-time-to-resolution (MTTR) during regression handling.

Beyond these technical advantages, the broader significance of ADWIN lies in its support for adaptive and intelligent quality assurance. Drift detection is not merely an auxiliary mechanism—it becomes a feedback engine that closes the loop between test results, model diagnosis, and model improvement. This enables a transition from brittle, pre-defined test scripts to agile, self-aware systems that can adjust their validation logic based on live system conditions.

As AI becomes deeply embedded in enterprise software—from cloud orchestration to developer assistants—the importance of transparency and traceability across versions escalates. ADWIN facilitates this by not only signaling anomalies but also explaining when, where, and why the model behavior shifted. This allows for better root cause analysis and reduces the debugging burden on engineers. Furthermore, it opens doors for human-in-the-loop testing workflows, where expert testers are alerted only when statistically significant deviations are detected, improving both test productivity and decision confidence.

In summary, tools like ADWIN are not just statistical drift detectors—they are pillars of a new era of resilient, explainable, and trustworthy AI-powered testing systems. Their integration into the continuous testing lifecycle empowers organizations to build software that learns, adapts, and evolves alongside the environments in which it operates.

## 5.2 Explainable AI (XAI) in Test Debugging

With the widespread application of artificial intelligence (AI) models in critical domains such as healthcare, autonomous driving, and finance, the reliability, transparency, and safety of AI systems have received increasing attention. In high-stakes decision-making environments, a single incorrect or unexplained output from an AI model can have severe consequences, ranging from financial losses to threats to human life. Therefore, ensuring that AI systems are not only performant but also interpretable has become a vital requirement in both academic research and industrial deployment.

During the stages of system testing and debugging, traditional black-box models often lack transparency in their decision-making processes. These models can produce accurate predictions, but without providing any insight into how those predictions are made. This opaqueness makes it difficult for testers and developers to trace the logic behind model outputs, understand unexpected behavior, or validate system reliability. As a result, the lack of interpretability



restricts the efficiency of error localization, fault analysis, and iterative model improvement. In sensitive areas such as healthcare, where trust and accountability are critical, this lack of clarity may even pose direct threats to user safety and regulatory compliance.

To address these challenges, Explainable Artificial Intelligence (XAI) has emerged as a complementary approach that aims to make AI systems more transparent and understandable. In particular, integrating XAI methods into AI system testing has become a key approach to improving testing effectiveness, identifying hidden failure modes, and enhancing user trust. XAI tools enable practitioners to uncover which features influenced a prediction, assess whether the model's logic aligns with human expectations, and flag unintended biases embedded in the model.

Two commonly used post-hoc XAI tools are LIME (Local Interpretable Model-Agnostic Explanations) and SHAP (SHapley Additive Explanations). These model-agnostic explanation methods allow testers to understand the prediction logic without altering the original model architecture or retraining. They are widely applicable across domains and have proven useful in highlighting potential flaws related to feature dependencies, data distribution shifts, and hidden biases. For instance, both methods have been successfully used in model audits, healthcare diagnostics, and algorithmic fairness analysis.

The core idea of LIME is to generate local perturbations around a specific prediction instance and fit a linear surrogate model that approximates the local decision boundary of the complex model. This localized linear model is interpretable and helps to reveal the relative contribution—either positive or negative—of each input feature to the prediction result. In practical testing and debugging scenarios, LIME is frequently employed to check whether a model is relying on irrelevant or spurious features. For example, in the testing of an autonomous driving system, a model erroneously classified a stop sign as something else. Using LIME for explanation revealed that the model was relying not on the features of the stop sign itself, but rather on the color of a building in the background. This insight enabled testers to identify an unintended sensitivity to environmental context. As a corrective measure, the data augmentation strategy was improved, and the feature engineering process was revised to reduce background dependency. Ultimately, this enhanced the model's robustness and generalization in real-world, variable environments [21].

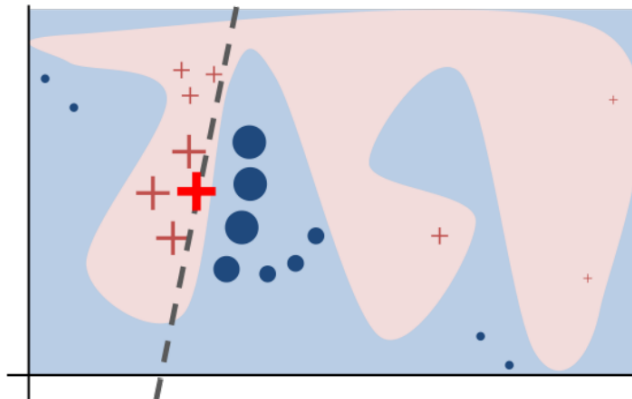


Fig. 7. LIME explanation example. The red cross marks the instance being explained. The background colors represent the black-box model's decision surface, and the dashed line shows the linear surrogate model used for local interpretation [21].

Unlike the local linear explanations offered by LIME, SHAP (SHapley Additive Explanations) is grounded in cooperative game theory. It leverages the Shapley value concept, which defines a unique and fair distribution of contribution

among players in a game. In the context of machine learning, each input feature is treated as a “player,” and SHAP calculates its marginal contribution by considering all possible subsets of features. This results in feature attributions that are not only consistent across different model architectures but also satisfy desirable mathematical properties such as local accuracy, missingness, and additivity.

By aggregating over all possible permutations of features, SHAP provides a globally faithful representation of how each feature influences the model’s output. This global and theoretically principled framework makes SHAP particularly useful for high-stakes domains where interpretability must align with regulatory and ethical standards. Moreover, SHAP explanations are model-agnostic and can be applied to both tree-based models (using TreeSHAP) and deep learning models (using DeepSHAP or GradientSHAP).

SHAP has shown remarkable value in the testing of medical AI systems, where fairness and accountability are critical. For instance, during the testing of a dermatology diagnosis model, researchers discovered that the model’s misdiagnosis rate for patients with darker skin tones was significantly higher than for other groups. This raised concerns about potential racial or demographic bias in the AI system. Using SHAP to analyze individual predictions, researchers found that the model assigned disproportionately low importance to pixel regions corresponding to skin tone for darker-skinned individuals.

This under-weighting indicated that the model was effectively ignoring relevant visual cues in these populations. Upon further investigation, it was confirmed that the original training dataset contained a disproportionately small number of samples from minority groups. As a remediation step, researchers rebalanced the training data through targeted data collection and synthetic augmentation, followed by model retraining. This intervention not only improved the model’s classification accuracy across demographic subgroups but also significantly enhanced its fairness and generalization capability.

More importantly, this process led to a notable improvement in user trust. Clinicians and patients expressed greater confidence in the AI-assisted diagnostic system, knowing that its decision-making process had been thoroughly analyzed, its biases revealed and mitigated, and its output explanations made transparent and accessible. This case illustrates how SHAP can serve as a powerful debugging tool in model validation pipelines, particularly in domains where human trust, regulatory scrutiny, and ethical transparency are indispensable [22].

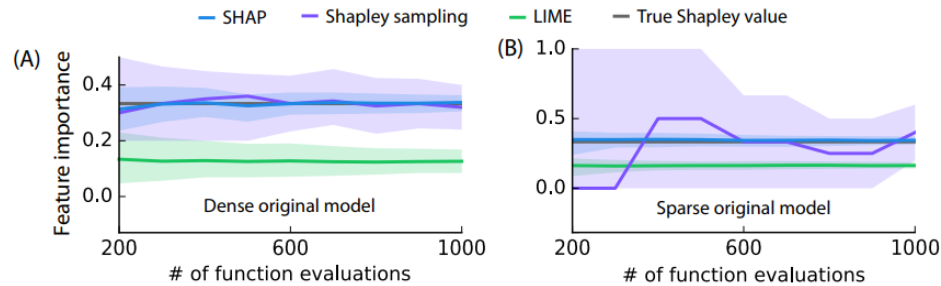


Fig. 8. Comparison of SHAP, Shapley sampling, and LIME. SHAP provides more stable and accurate feature importance estimates relative to the true Shapley value [22].

Beyond the application of single explainability tools such as LIME and SHAP, recent advances in the field of Explainable AI (XAI) have emphasized the value of hybrid strategies that integrate the strengths of both interpretable

and black-box models. These hybrid approaches aim to bridge the gap between model transparency and predictive accuracy—two attributes that are often at odds in practical systems. By leveraging a combination of inherently interpretable models (such as decision trees, rule-based classifiers, or linear models) and high-performance but opaque models (such as deep neural networks), these strategies offer greater flexibility and adaptability in real-world testing scenarios.

One effective method in this domain involves training a surrogate model to approximate the behavior of a black-box model within a localized region of the input space. For instance, in image classification tasks, a convolutional neural network (CNN) may be employed for high-accuracy predictions. Meanwhile, a shallow decision tree or locally linear model is trained using samples near the prediction boundary to approximate the decision process. This allows developers to trace how changes in input features affect the model’s classification output and identify unexpected decision boundaries that may indicate spurious correlations or overfitting.

Such hybrid XAI systems have also proven beneficial for behavior monitoring and anomaly detection in testing environments. For example, when combined with drift detection mechanisms, these interpretable surrogate models can provide human-understandable alerts about changing model behavior over time. This makes them particularly suitable for real-time applications such as fraud detection, autonomous driving, and industrial quality control, where continuous monitoring and rapid debugging are critical. Importantly, these approaches allow for the benefits of deep learning to be retained while offering windows of transparency to facilitate error analysis and human-in-the-loop evaluation.

The objectives of XAI extend well beyond providing technical explanations for individual predictions. A core mission of XAI is to foster and maintain user trust—an essential factor for the deployment and acceptance of AI technologies in critical applications. In this context, explanations serve as mechanisms of accountability and validation, enabling users to assess whether model decisions are aligned with domain knowledge, ethical standards, and legal regulations.

In addition to building trust, XAI plays a key role in uncovering causal relationships that may not be directly evident from model outputs. This is particularly important when testing AI systems that are deployed in dynamic and sensitive environments. For instance, an AI-based decision support system in a hospital must not only provide accurate recommendations but also clarify which medical factors contributed most significantly to the decision, especially when the stakes involve life-altering treatment plans.

XAI also contributes to privacy awareness by exposing which input attributes have the greatest influence on model outcomes. In privacy-sensitive domains, this visibility enables developers to assess whether models are unintentionally leveraging sensitive or protected attributes, such as gender, race, or age, even if such features are not explicitly included in the input. Thus, XAI tools support auditing and bias mitigation efforts during the testing phase, helping organizations align with fairness and data protection policies.

Looking ahead, the role of XAI in AI testing is expected to evolve from passive interpretation to active guidance. Emerging research explores causal explainability, where models attempt to uncover not just correlations but underlying causal mechanisms that drive predictions. Interactive explanation interfaces are also being developed, allowing developers and domain experts to ask “what-if” and “why-not” questions during testing and debugging. These tools enable iterative, dialogue-based model refinement that closely resembles the scientific method.

Furthermore, task-specific interpretability metrics are gaining traction as a means to quantify the usefulness and clarity of explanations in different contexts. For example, interpretability requirements in legal AI may differ vastly from those in autonomous robotics. By tailoring XAI tools to match the cognitive and operational needs of end-users, researchers aim to create more accessible, accountable, and adaptive testing environments. In this future paradigm,

XAI will not merely be a diagnostic tool, but a central component of the AI development lifecycle—one that shapes model architecture, training, validation, and deployment in a feedback-rich, human-aligned manner.

## 6 Challenges

### 6.1 Over-Reliance on Training Data

### 6.2 Ethical and Legal Risks in AI Testing

## 7 Applications

### 7.1 AI in Medical Software Testing

### 7.2 AI for Autonomous Driving Validation

## 8 Future Directions and Conclusion

### 8.1 Emerging AI Techniques in Testing

### 8.2 Integration with Agile and DevOps

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