IT-314 Software Engineering Lab 9

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Topic: Mutation Testing

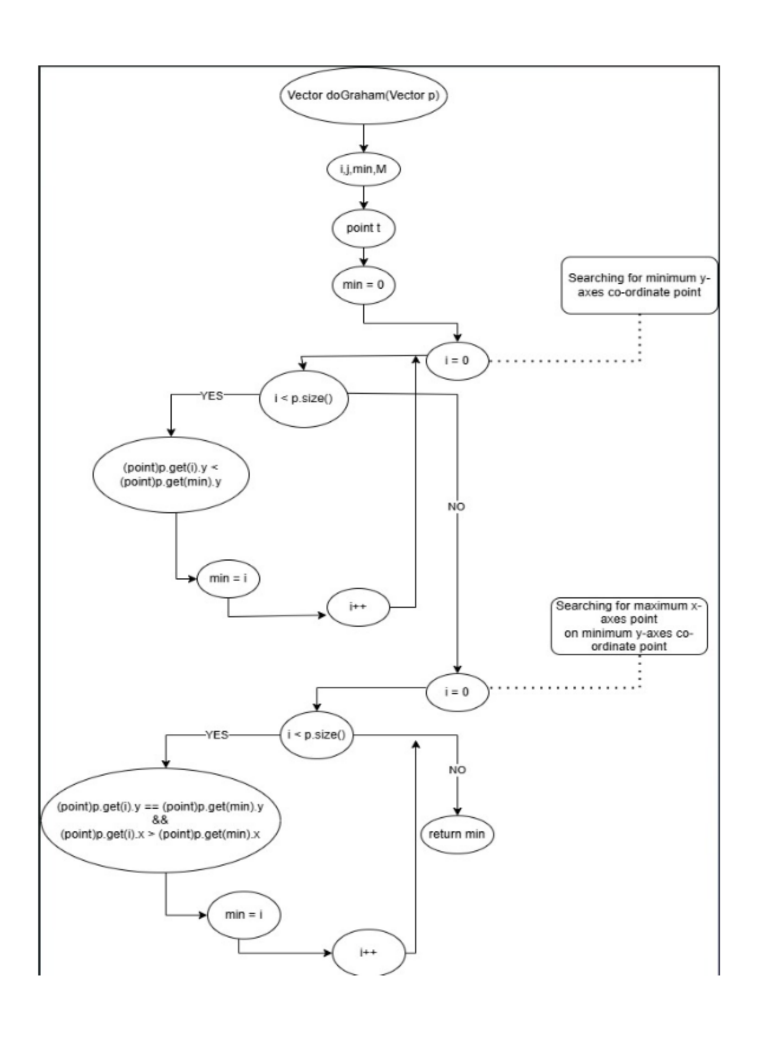
Q.1. The code below is part of a method in the ConvexHull class in the VMAP system. The following is a small fragment of a method in the ConvexHull class. For the purposes of this exercise, you do not need to know the intended function of the method. The parameter p is a Vector of Point objects, p.size() is the size of the vector p, (p.get(i)).x is the x component of the ith point appearing in p, similarly for (p.get(i)).y. This exercise is concerned with structural testing of code, so the focus is on creating test sets that satisfy some particular coverage criteria.

```
Vector doGraham(Vector p) {
       int i, j, min, M;
       Point t;
        min = 0;
        // search for minimum:
        for(i=1; i < p.size(); ++i) {
            if( ((Point) p.get(i)).y <
                         ((Point) p.get(min)).y)
               min = i;
            }
        }
        // continue along the values with same y component
        for(i=0; i < p.size(); ++i) {
            if(( ((Point) p.get(i)).y ==
                          ((Point) p.get(min)).y ) &&
                 (((Point) p.get(i)).x >
                          ((Point) p.get(min)).x ))
                 min = i;
            }
        }
```

❖ Executable Code in Java:-

```
import java.util.Vector;
class Point {
    public Point(int x, int y) {
       this.y = y;
    public String toString() {
public class ConvexHull {
    public static void doGraham(Vector<Point> p) {
        min = 0;
        System.out.println("Searching for the minimum
y-coordinate...");
        for (i = 1; i < p.size(); ++i) {</pre>
            System.out.println("Comparing " + p.get(i) + " with " +
p.get(min));
            if (p.get(i).y < p.get(min).y) {</pre>
                System.out.println("New minimum found: " +
p.get(min));
        System.out.println("Searching for the leftmost point with
the same minimum y-coordinate...");
        for (i = 0; i < p.size(); ++i) {</pre>
            System.out.println("Checking if " + p.get(i) + " has the
same y as " + p.get(min) +
```

❖ Control Flow Diagram



Construct test sets for your flow graph that are adequate for the following criteria:

- a. Statement Coverage.
- **b.** Branch Coverage.
- c. Basic Condition Coverage.

a. Statement Coverage

Objective: Verify that every line of code in the flow graph is executed at least once.

Test Set

- 1. Test Case 1:
 - Inputs: A list p with more than one point (e.g., [(0, 1), (1, 2), (2, 0)]).
 - Purpose: This case traverses the entire code, covering the logic for finding both the minimum y-coordinate and the leftmost point with that y-coordinate.
- 2. Test Case 2:
 - o **Inputs**: [(2, 2), (2, 2), (3, 3)].
 - Purpose: Tests cases where points share the same y-coordinate, ensuring the code correctly identifies the leftmost point.

b. Branch Coverage

Objective: Ensure that every possible branch (true/false outcome) from each decision point is taken at least once.

Test Set:

- 1. Test Case 1:
 - o **Inputs**: [(0, 1), (1, 2), (2, 0)].
 - o **Purpose**: Covers the true branch when searching for the minimum y-coordinate.
- 2. Test Case 2:
 - o **Inputs**: [(2, 2), (2, 2), (3, 3)].
 - Purpose: Checks the branch logic where y-coordinates are equal, ensuring the code proceeds to compare x-coordinates.
- 3. Test Case 3:
 - o **Inputs**: [(1, 2), (1, 1), (2, 3)].
 - Purpose: Ensures that the flow takes the false branch when evaluating new minimum y-coordinates, and also tests the leftmost point check.

c. Basic Condition Coverage

Objective: Validate that each condition within a decision point can independently produce both true and false outcomes.

Test Set

1. Test Case 1:

- o **Inputs**: [(1, 1), (2, 2), (3, 3)].
- Purpose: Ensures both true and false evaluations for y-coordinate comparisons.

2. Test Case 2:

- o **Inputs**: [(1, 1), (1, 1), (1, 2)].
- **Purpose**: Examines cases with identical y-coordinates, testing the x-coordinate comparison logic.

3. Test Case 3:

- o **Inputs**: [(3, 1), (2, 2), (1, 3)].
- Purpose: Confirms that each condition in the loop can be evaluated, verifying the robustness of the function's logic.
- For the test set you have just checked can you find a mutation of the code (i.e. the deletion, change or insertion of some code) that will result in failure but is not detected by your test set. You have to use the mutation testing tool.

Types of Possible Mutations

We can apply typical mutation types, including:

- Relational Operator Changes: Modify <=to <or ==to !=in the conditions.
- Logic Changes: Remove or invert a branch in an if-statement.
- Statement Changes: Modify assignments or statements to see if the effect goes undetected.

Potential Mutations and Their Effects

- 1. Changing the Comparison for Leftmost Point:
 - Mutation: In the second loop, change p.get(i).x < p.get(min).xto p.get(i).x <= p.get(min).x.</p>
 - Effect: This would cause the function to select points with the same xcoordinate as the leftmost, potentially breaking the uniqueness of the minimum point.
 - Undetected by Current Test: The current tests do not cover the case where multiple points have the same y and x values, which would reveal if the function mistakenly allows such points as the leftmost.

2. Altering the y-Coordinate Comparison to <=in the First Loop:

- Mutation: Change p.get(i).y < p.get(min).yto p.get(i).y <= p.get(min).yin the first loop.</p>
- Effect: This would allow points with the same y-coordinate but different x-coordinates to overwrite min, potentially selecting a non-leftmost minimum point.
- Undetected by Current Tests: The current test set lacks cases where several points have the same y-coordinate, and this mutation would go undetected. To reveal this, we would need a test where multiple points have the same y and different x coordinates.

3. Removing the Check for x-coordinate in the Second Loop:

- Mutation: Remove the condition p.get(i).x < p.get(min).xin the second loop.
- Effect: This would cause the function to select any point with the same minimum y-coordinate as the "leftmost," regardless of its x-coordinate.
- Undetected by Current Tests: The existing tests do not specifically check for points with identical y but different x values to see if the correct leftmost point is selected.

Additional Test Cases to Detect These Mutations

To detect these mutations, we can add the following test cases:

1. Detect Mutation 1:

- Test Case: [(0, 1), (0, 1), (1, 1)]
- Expected Result: The leftmost minimum should still be (0, 1)despite having duplicates.
- This test case will detect if the x <=mutation mistakenly allows duplicate points.

2. Detect Mutation 2:

- Test Case: [(1, 2), (0, 2), (3, 1)]
- Expected Result: The function should select (3, 1) as the minimum point based on the y-coordinate.
- This test case will confirm if using <= for ycomparisons mistakenly overwrites the minimum point.

3. Detect Mutation 3:

- Test Case: [(2, 1), (1, 1), (0, 1)]
- **Expected Result**: The leftmost point (0, 1) should be chosen.
- This will reveal if the x-coordinate check was mistakenly removed.

These additional test cases would help ensure that any such mutations do not survive undetected by the test suite, strengthening the coverage.

Python Code for Mutation:-

```
from math import atan2
class Point:
   def __init__(self, x, y):
       self.x = x
       self.y = y
    def __repr__(self):
        return f"({self.x}, {self.y})"
def orientation(p, q, r):
    # Cross product to find orientation
   val = (q.y - p.y) * (r.x - q.x) - (q.x - p.x) * (r.y - q.y)
   if val == 0:
        return 0 # Collinear
    elif val > 0:
        return 1 # Clockwise
    else:
       return 2 # Counterclockwise
def distance_squared(p1, p2):
```

```
return (p1.x - p2.x) ** 2 + (p1.y - p2.y) ** 2
def do_graham(points):
     # Step 1: Find the bottom-most point (or leftmost in case of a tie)
     n = len(points)
     min_y_index = 0
     for i in range(1, n):
                      if (points[i].y < points[min_y_index].y) or \</pre>
                (points[i].y == points[min_y_index].y and points[i].x < points[min_y_index].x):
                 min_y_index = i
     points[0], points[min_y_index] = points[min_y_index], points[0] p0 = points[0]
     # Step 2: Sort the points based on polar angle with respect to
p0
     points[1:] = sorted(points[1:], key=lambda p: (atan2(p.y - p0.y, p.x - p0.x),
distance_squared(p0, p)))
     # Step 3: Initialize the convex hull with the first three points hull = [points[0], points[1],
     points[2]]
     # Step 4: Process the remaining points for i in range(3,
     n):
```

```
# Mutation introduced here: instead of checking `!= 2`, we
incorrectly use `== 1`
        while len(hull) > 1 and orientation(hull[-2], hull[-1],
points[i]) == 1:
            hull.pop()
        hull.append(points[i])
    return hull
# Sample test to observe behavior with the mutation
points = [Point(0, 3), Point(1, 1), Point(2, 2), Point(4, 4),
          Point(0, 0), Point(1, 2), Point(3, 1), Point(3, 3)]
hull = do graham(points)
print("Convex Hull:", hull)
```