

## Homework Assignment - 6

## Indraprastha Insitute of Information Technology, Delhi

COMPUTER SCIENCE AND APPLIED MATHEMATICS

## Introduction to Quantitative Biology (BIO213)

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## 1 Solution

In our solution we have displayed the following on the console:

- Number of molecules: The number of molecules participating in the random walk.
- Size of the lattice: The size of the lattice in which the molecules are placed.
- Number of iterations: The number of iterations for which we perform the walk.
- Mean X displacement: The mean displacement along the X direction.
- Mean Y displacement: The mean displacement along the Y direction.
- Mean displacement vector: A vector denoting the mean displacement.
- Magnitude of mean displacement ( $\langle r \rangle$ ): Magnitude of the mean displacement vector.
- Mean square displacement ( $< r^2 >$ ): Magnitude of the mean square displacement. As can be seen in the output attached, the magnitude of the mean displacement, < r > should be close to 0 and in our case, it is 0.9014, which is approximately 0. Hence, the simulation was successful.

```
% BIO213 - IQB
% Homework Assignment 6
% Brihi Joshi (2016142)
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analysis(100, 1000, 50);
%This function initialises a list of x-coordinates and y-coordinates of
% molecules in the lattice.
function [x_coords, y_coords] = initialize_coords(n, size)
  x coords = zeros(1, n);
  y coords = zeros(1, n);
  i = 1;
  while i < (n + 1)
     x = randi(size, 1);
     y = randi(size, 1);
     flag = 0;
     for j = 1 : i-1
        if x_coords(j) == x && y_coords(j) == y
          flag = 1;
           break;
        end
     end
     if flag == 1
       continue;
     else
        x coords(i) = x;
        y_{coords(i)} = y;
        i = i + 1;
     end
  end
end
% The function moves molecules left, right, up or down, depending upon the
% direction code provided. The direction codes are as follows:
% 0 -> Up
% 1 -> Right
% 2 -> Down
% 3 -> Left
% In case the destination in a particular direction is already occupied by
% another molecule or the molecule is at the boundary of the lattice, then
% it is not moved.
```

```
function [x, y] = update_coords(xi, yi, dir, x_coords, ...
    y coords, size, index)
   x = xi;
    y = yi;
    if dir == 0
        if yi == size
           return;
        else
            for i = 1 : length(x_coords)
               if (x_coords(i) == xi && y_coords(i) == yi + 1)
            end
            x = xi;
            y = yi + 1;
            y_{coords(index)} = y;
    elseif dir == 1
        if xi == size
            return;
        else
            for i = 1 : length(y_coords)
               if (x_coords(i) == xi + 1 && y_coords(i) == yi)
                   return;
               end
            end
            x = xi + 1;
            y = yi;
            x coords(index) = x;
         end
    elseif dir == 2
        if yi == 1
            return;
        else
            for i = 1 : length(y_coords)
               if (x_coords(i) == xi && y_coords(i) == yi-1)
                   return;
               end
            end
            x = xi;
            y = yi - 1;
            y_coords(index) = y;
    elseif dir == 3
        if xi == 1
            return;
        else
            for i = 1 : length(y_coords)
               if (x_coords(i) == xi - 1 && y_coords(i) == yi)
                   return;
               end
            end
            x = xi - 1;
            y = yi;
            x coords(index) = x;
         end
    end
end
```

```
% This function simulates a random walk from the number of molecules,
% number of iterations and the size of the lattice
function [copy_x, copy_y, xs, ys] = random_walk(n, iter, size)
   [x coords, y coords] = initialize coords(n, size);
   copy_x = x_coords;
   copy_y = y_coords;
   for i = 1 : iter
       for p = 1 : n
          dir = 0;
          chance = rand();
           if chance < 0.25
              dir = 0;
           elseif chance < 0.50</pre>
              dir = 1;
           elseif chance < 0.75</pre>
              dir = 2;
           elseif chance < 1.00
              dir = 3;
           end
           [a, b] = update_coords(x_coords(p), y_coords(p), dir, ...
              x coords, y coords, size, p);
           x_{coords}(p) = a;
           y coords(p) = b;
       end
   end
   xs = x_coords;
   ys = y_coords;
end
function [rx, ry, r2] = analysis(n, iter, size)
   [xi, yi, xf, yf] = random walk(n, iter, size);
   rx = mean(xf - xi);
   ry = mean(yf - yi);
   r = sqrt(rx^2 + ry^2);
   r2 = mean((xf-xi).^2) + mean((yf-yi).^2);
   fprintf("Following is the analysis of the random walk simulation:\n\n");
   fprintf("1. Number of molecules = %d\n", n);
   fprintf("2. Size of lattice = %d\n", size);
   fprintf("3. Number of iterations = %d\n", iter);
   fprintf("4. Mean X displacement = %.4f\n", rx);
   fprintf("5. Mean Y displacement = %.4f\n", ry);
   fprintf("6. Mean displacement vector = %.4fî + %.4fĵ\n", rx, ry);
   fprintf("7. Magnitude of mean displacement = %.4f\n", r);
   fprintf("8. Mean square displacement = %.4f\n\n", r2);
```

Following is the analysis of the random walk simulation:

- 1. Number of molecules = 100
- 2. Size of lattice = 50
- 3. Number of iterations = 1000
- 4. Mean X displacement = -0.5800
- 5. Mean Y displacement = -0.6900
- 6. Mean displacement vector =  $-0.5800\hat{1} + -0.6900\hat{j}$
- 7. Magnitude of mean displacement = 0.9014
- 8. Mean square displacement = 399.9100

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