



INDRAPRASTHA INSTITUTE of
INFORMATION TECHNOLOGY
DELHI

HOMework ASSIGNMENT - 6

INDRAPRASTHA INSTITUTE 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 ($\langle r^2 \rangle$): Magnitude of the mean square displacement.

As can be seen in the output attached, the magnitude of the mean displacement, $\langle r \rangle$ should be close to 0 and in our case, it is 0.9014, which is approximately 0. Hence, the simulation was successful.


```

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)
                return;
            end
        end
        x = xi;
        y = yi + 1;
        y_coords(index) = y;
    end

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;
    end

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
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
                dir = 1;
            elseif chance < 0.75
                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 = %.4fi + %.4fj\n", rx, ry);
    fprintf("7. Magnitude of mean displacement = %.4f\n", r);
    fprintf("8. Mean square displacement = %.4f\n\n", r2);

end

```

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 = $-0.5800\hat{i} + -0.6900\hat{j}$
7. Magnitude of mean displacement = 0.9014
8. Mean square displacement = 399.9100