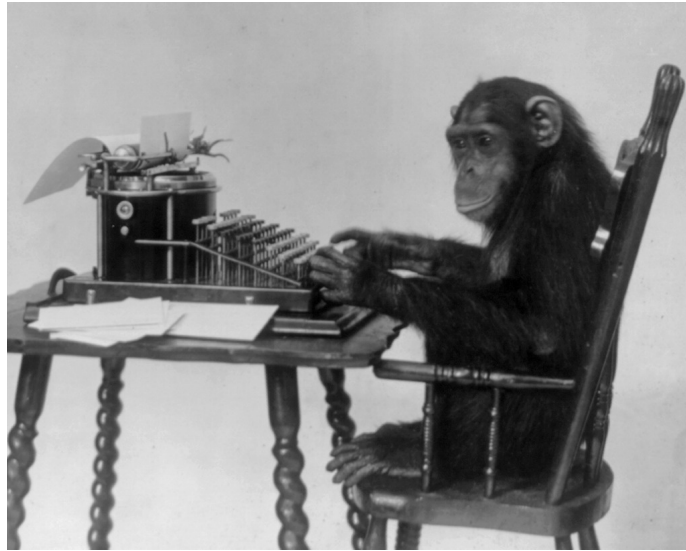


ECTA Homework 1  
Genetic Algorithms  
and  
Infinite Monkeys

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The infinite monkey theorem states that if a chimpanzee hits keys at random on a typewriter for an infinite amount of time, it will eventually type the complete works of William Shakespeare. Is this what evolution is doing? It has been argued that the genetic mutations required to move from a single cell to multicellular life are as unlikely as a monkey typing Hamlet's soliloquy. But is evolution just a monkey banging on a typewriter?

## 1 Assignment Description

1. Build a simple Genetic Algorithm and test the effect of each component
  - Mutation
  - Crossover
  - Elitism
2. Answer the question, "is evolution just a monkey banging on a typewriter?"
  - Grading Scheme
    - ☐ Code a GA (40 pts)
      - ☐ Selection (10pts)
      - ☐ Crossover (10pts)

- ☐ mutation (10pts)
- ☐ Elitism (10pts)
- ☐ GA Component Comparisons (40 pts)
  - ☐ Comparison vs. Standard Implementation (5 pts)
  - ☐ Comparison of Components (5 pts)
  - ☐ Short Answer #1 (10 pts)
  - ☐ Short Answer #2 (10 pts)
  - ☐ Short Answer #3 (10 pts)
- ☐ GA vs Monkey (20 pts)
  - ☐ Solve the soliloquy (10 pts)
  - ☐ Brute force calculation (5 pts)
  - ☐ Short Answer #4 (5 pts)

## 2 Submission Instructions

Follow along with the instructions in this PDF, filling in your own code, data, and observations as noted. Your own data should be inserted into the latex code of the PDF and recompiled. All code must be done in MATLAB. The basic structure of the code and fitness function are provided, but all code should be submitted as a separate zipped file in LEA. Relevant sections of code can be inserted directly into this document using the mcode latex package. This package is attached with documentation, and in this document I have provided usage examples.

To be perfectly clear we expect two submissions to LEA:

1. 1 PDF (report) – a modified version of this PDF, with your own code snippets, figures, and responses inserted
2. 1 ZIP (code and data) – a .zip file containing all code use to run experiments (.m files) *and* resulting data as a .mat file

### 3 Assignment Overview

#### the Task

Like our monkey, your Genetic Algorithm will be tested as to how closely it can reproduce Shakespeare. Two benchmarks are given: `hamletQuote` and `hamletSoliloquy`. These functions take one or more genes of length 18 for the quote or 1446 for the soliloquy and return a fitness value which corresponds to the number of letters that match the target text.

One gene is a number between 0 and 27, corresponding to a space (0), letters a-z (1-26), and a new line (27).

#### the Algorithm

I have created the basic structure of the GA for you. The magic happens in the loop here in `monkeyGa.m`:

```
%% Evolutionary Operators

% Selection -- Returns [MX2] indices of parents
parentIds = my.selection(fitness, p); % Returns indices of parents

% Crossover -- Returns children of selected parents
children = my.crossover(pop, parentIds, p);

% Mutation -- Applies mutation to newly created children
children = my.mutation(children, p);

% Elitism -- Select best individual(s) to continue unchanged
eliteIds = my.elitism(fitness, p);

% Create new population -- Combine new children and elite(s)
newPop = [pop(eliteIds,:); children];
pop = newPop(1:p.popSize,:); % Keep population size constant

% Evaluate new population
fitness = feval(p.task, pop);
```

It will be your job to implement each of the evolutionary operators and measure how they effect performance of the algorithm on the `hamletQuote` task.

## Running the Algorithm

To run the algorithm and view the results, you can use the snippet provided at the start of `monkeyExperiment.m`:

```
%% Run the algorithm once
clear;
p = monkeyGa('hamletQuote');           % Set hyperparameters
output = monkeyGa('hamletQuote',p); % Run with hyperparameters

% View Result
gene2text(output.best(:,end))
plot([output.fitMax; output.fitMed], 'LineWidth', 3);
legend('Max Fitness', 'Median Fitness', 'Location', 'NorthWest');
xlabel('Generations'); ylabel('Fitness'); set(gca, 'FontSize', 16);
title('Performance on Hamlet Task')
```

To run a section in matlab (a code block marked by `%%`, with the cursor inside the code block click the ‘Run Section’ button in the editor portion of the ribbon, or more simply hit ‘CTRL + Enter’). Run it a few times. As the only operator which is implemented is initialization, it will give you a pretty terrible result.

## Comparing Algorithms

As evolutionary algorithms are based on stochastic processes, they will not perform the same every time. Whenever a comparison between two algorithms or algorithm settings is made, it *must* be a comparison over several runs. Comparisons between runs must take into account the effect of randomness, including significance of results (how likely the result is to be because of chance).

## 4 The Assignment

### 4.1 Coding a simple GA

Begin by implementing the four given genetic operators, replacing the filler code with your own. The expected inputs and outputs, as well as hints as how to perform each operation are included within the code. Please put your code in the report here using the given ‘firstline/lastline’ syntax in the L<sup>A</sup>T<sub>E</sub>X.

*Don’t overthink it! Each of these can be done in less than 10 lines!*

#### 1. Tournament Selection

```
%----- BEGIN CODE -----

%% This is 'random' selection of parent pairs, can you do better?
parentIds = randi(p.popSize, [p.popSize 2]);

%% TOURNAMENT SELECTION SOLUTION
for child.count = 1:p.popSize
    group = randi(p.popSize, [p.sp,2]); % Get two sets of ...
        random individuals
    [~, winner_index] = max(fitness(group)); % Get a parent ...
        from each set based on highest fitness

    first.parent_index = group(winner_index(1,1), 1); % Get ...
        the first parent
    second.parent_index = group(winner_index(1,2), 2); % Get ...
        the second parent

    parentIds(child.count, :) = [first.parent_index ...
        second.parent_index];
end
%----- END OF CODE -----
```

#### 2. Crossover

```
%----- BEGIN CODE -----

%% No crossover happening, can you do better?
children = pop( parentIds(:,1) ,:);

%% ONE POINT Crossover SOLUTION
for index=1:size(pop,1)
    doCrossOver = rand(1) < p.crossProb; % Determine ...
        whether to do cross over or not

    if(doCrossOver)
```

```

        crossOverPoint = randi(size(pop,2));    % Define cross ...
        over point in case of cross over

        first_parent_index = parentIds(index,1);
        first_part = pop(first_parent_index,1:crossOverPoint); ...
        % Take first part from the first parent

        second_parent_index = parentIds(index,2);
        second_part = pop(second_parent_index, ...
        crossOverPoint+1: size(pop,2)); % Take second ...
        part from second parent

        children(index,:) = [first_part second_part];    % ...
        combine the two parts
    else
        children(index, :) = pop(parentIds(index,1), :);    % ...
        Take first parent in case of no cross over
    end
end
%----- END OF CODE -----

```

### 3. Mutation

```

%----- BEGIN CODE -----

%% No mutation happening, can you do better?
children = children;

%% POINT MUTATION SOLUTION
for childIndex=1:p.popSize
    genes = children(childIndex, :);

    doMutationFlag = rand(1, p.nGenes) < p.mutProb;
    doMutationInverseFlag = ~doMutationFlag;

    genes = genes .* doMutationInverseFlag; % Set 0 for the ...
    genes to change
    newGenes = randi([0,27],[1,p.nGenes]) .* doMutationFlag; % ...
    Set 0 for the genes not to change
    genes = genes + newGenes; % Combine the original genes ...
    with new values

    children(childIndex, :) = genes;
end
%----- END OF CODE -----

```

### 4. Elitism

```

%% Here we just keep the first individual as an elite, can you ...
do better?
eliteIds = 1;

%% ELITISM SOLUTION
[~,sorted_population_indices]= sort(fitness,'descend');

```



```

eliteIds = sorted(population.indices(1:ceil(p.popSize * ...
    p.elitePerc))); % Take population with highest fitness
%----- END OF CODE -----

```

## 4.2 Ablation Study

One common technique for better understanding an algorithm is remove each component and see the result. What happens when we don't use elitism or we skip crossover? In this section we test a few combinations.

### 4.2.1 Comparing Algorithms

Provided are versions of each operator with the prefix 'adam' instead of 'my'. These can be used to validate your own results. I included a version which uses them in the file 'adamGa', which is exactly the same except this part:

```

% Selection -- Returns [MX2] indices of parents
parentIds = adam_selection(fitness, p); % Returns indices of ...
    parents

% Crossover -- Returns children of selected parents
children = adam_crossover(pop, parentIds, p);

% Mutation -- Applies mutation to newly created children
children = adam_mutation(children, p);

% Elitism -- Select best individual(s) to continue unchanged
eliteIds = adam_elitism(fitness, p);

```

As this is a stochastic algorithm to get a fair comparison we should run the algorithm multiple times and compare statistically. Lets use all the cores on your computer to do this as fast as possible using a `parfor` loop. This is just like a `for` loop, except it runs each iteration on a different core. Get the result of 20 runs of your code and mine and save it to disk:

```

% changing 'for' to 'parfor'.
clear; p = monkeyGa('hamletQuote');
parfor iExp = 1:20
    output = adamGa('hamletQuote',p);
    fitness(iExp,:) = output.fitMax;
end

```

```

standardResult = fitness;

parfor iExp = 1:20
    output = monkeyGa('hamletQuote',p);
    fitness(iExp,:) = output.fitMax;
end
myResult = fitness;
save('runData.mat','standardResult','myResult')

```

With this data saved you can compare the two algorithms and compute the significance of the comparison. I have given you a few helper functions:

```

load('runData.mat')
gens = 1:length(standardResult);

% Get Significance of comparisons
fit1 = standardResult; fit2 = myResult;
[p,h] = sigPerGen(fit1,fit2);

% Plot results at every generation
figure(2); clf; hold on; C = parula(8); % Create figures and color map

% Plot Significance at every generations
hS1 = scatter(gens(~h),ones(1,sum(~h))*19,20,C(1,:), 'filled','s');
hS2 = scatter(gens(h),ones(1,sum(h))*19,20,C(7,:), 'filled','s');

% Plot median and percentiles
[hLine(1), hFill(1)] = percPlot(gens,fit1 ,C(2,:));
[hLine(3), hFill(2)] = percPlot(gens,fit2,C(5,:));

% Label and make pretty
hLeg = legend([hFill hS1 hS2], 'Baseline', 'Full', 'p > 0.05', 'p < ...
    0.05', 'Location', 'SouthEast');

```

This plot shows the median performance at each generation (dashed lines) of each algorithm along with their upper and lower quartiles. Indicated at the top is the probability that the two algorithms are the same. Unsurprisingly, both runs of the same algorithm are statistically the same. Replace this plot with one of your own creation, comparing my code with your own implementation, to ensure that your code is working.

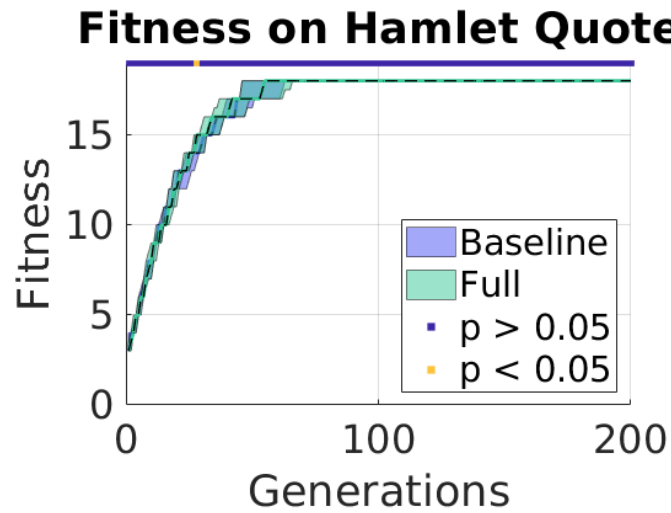


Figure 1: Baseline vs Full

Perform the following comparisons of your algorithm with various components removed and replace the plots with your own, this can be done by replacing the functions in the code and saving the result (e.g. replacing the `my_crossover` function with the `no_crossover` function: (1) Your full implementation vs. No crossover, (2) Your full implementation vs. No mutation, (3) No crossover vs. No crossover AND no elitism, and (4) No mutation vs. No mutation AND no elitism.

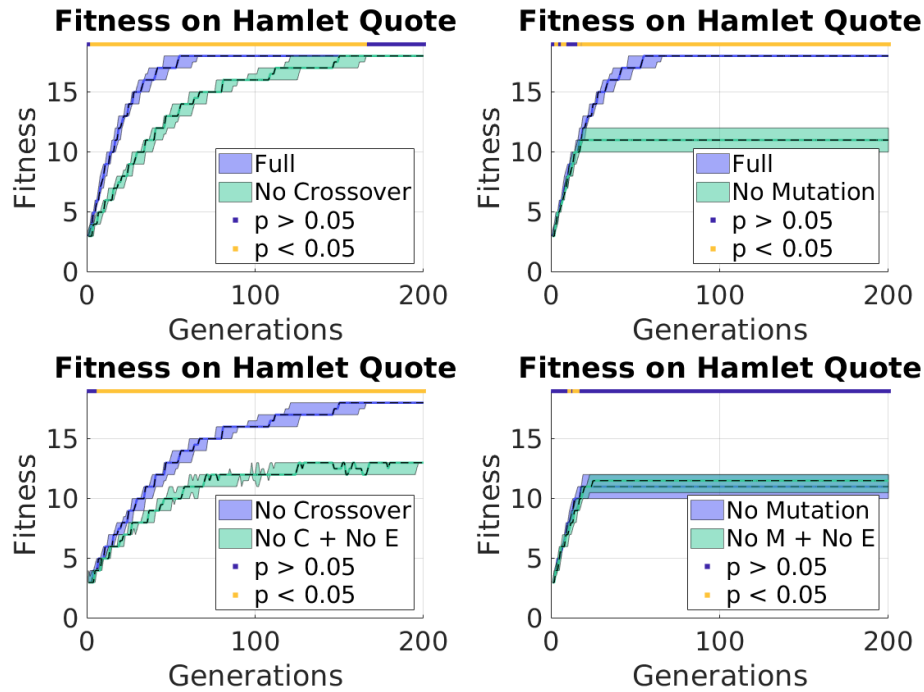


Figure 2: **GA performance when operators are removed**

Top Left: No Crossover, Top Right: No Mutation, Bottom Left: No Crossover vs. No Crossover and No Elitism, Bottom Right: No Mutation and No Elitism

#### 4.2.2 Analyzing the Results

1. Describe the main purposes of crossover and of mutation, how do your results illustrate their operation?

**Crossover:**

Crossover combines existing solutions to form new solutions. It allows genes with higher fitness to be maintained and spreaded in following generations. The graph (1) shows that the fitness grows slowly without crossover compared to full implementation.

**Mutation:**

Mutation introduces new information and building blocks in the genes. The comparison (2) shows that, without mutation, the fitness value reaches its limit very quickly and the limit is quite less compared to the full implementation.

2. When only crossover is used the problem is not typically solved. Why? Could you devise an experiment that would support your explanation? One in which crossover could solve the problem

every time?

*WRITE SOMETHING HERE*

3. Describe the benefits of elitism in the crossover and mutation only cases.

*Elitism is used to prevent losing progress over generations. The best solutions are copied into the next population without any change.*

*WRITE SOMETHING HERE*

## 4.3 Monkeys on a Typewriter

### 4.3.1 Using the GA

Now time to test your algorithm on the entire soliloquy. Is it really better than just banging on a typewriter? Switch out the fitness function and give the whole speech a try. It might take a little time, you may have to increase the number of generations to get to 100%, for this purpose don't worry about replicates:

```
%% The whole monologue
clear; p = monkeyGa('hamletSoliloquy');
p.maxGen = 10000;           % Increase the number of generations
tic;                       % Start the timer
output = monkeyGa('hamletSoliloquy',p);
gene2text(output.best(:,end)) % Show the found text
percentCorrect = (output.fitMax(end)/1446);
timeToComplete = toc;      % End the timer
disp([num2str(100*percentCorrect) '% correct in ' ...
      num2str(timeToComplete) ' seconds'])
```

By using the `tic` and `toc` commands we can time how long a program takes to execute. How long did it take your algorithm to find the whole speech?

*99.5159% correct in 89.0095 seconds*

### 4.3.2 Brute force

How long would it take to find the same solution by a monkey on a type writer, i.e. by brute force? The average and worst case for a brute force algorithm can be easily calculated by counting the possible states. Let's be charitable and say this is a particularly clever monkey, who is systematic and never typing the same thing twice. Lets be even more charitable and say that this clever monkey also has a MATLAB license and has created a program to do the typing for him. How many possible states are there? How long will it take this MATLAB monkey to explore them all? Please show your work and use appropriate units for your answer.

(hint: to time a very fast piece of code, repeat in many times and take the average time, like this: )

```
% Timing a single evaluation
aWholeBunchOfTimes = 100000;
test = randi([0 27], [aWholeBunchOfTimes, p.nGenes]);
tic; hamletSoliloquy(test); tEnd = toc;
oneEval = tEnd/aWholeBunchOfTimes;
disp(['One evaluation in ' num2str(oneEval) ' seconds'])
```

- Time required for one evaluation:  $2.8245 \times 10^{-06}$  seconds.
- Now, number of possible states can be  $28^{nGenes}$
- So time required to compute this would be  $2.8245 \times 10^{-06} \times 28^{nGenes}$
- For hamletSoliloquy ( $nGenes = 1446$ ), the time required will tend to infinity.
- For hamletQuote( $nGenes = 18$ ), the time required is  $9.0890 \times 10^{12}$  years.

How comparable are these methods? Is random search comparable to evolutionary search?

*For evolutionary search, as we have seen before, the time required for hamletSoliloquy is around 89 seconds, whereas for random search will need infinite time to come up with a similar solution. Thus, we can conclude that these approaches are not comparable.*

## 5 Inserting MATLAB code into LATEX — 3 ways

1) This inline demo `for i=1:3, disp('cool'); end;` uses the `\mcode{}` command.<sup>1</sup>

2) The following is a block using the `lstlisting` environment.

```
for i = 1:3
    if i ≥ 5 && a ≠ b           % literate programming replacement
        disp('cool');          % comment with some  $\TeX$  in it:  $\pi x^2$ 
    end
    [:,ind] = max(vec);
    x_last = x(1,end) - 1;
    v(end);
    really really long really really long really really long really ...
        really long really really long line % blaaaaaaaaa
    ylabel('Voltage ( $\mu$ V)');
end
```

Note: Here, the package was loaded with the `framed`, `numbered`, `autolinebreaks` and `useliterate` options. **Please see the top of `mcode.sty` for a detailed explanation of these options.**

3) Finally, you can also directly include an external m-file from somewhere on your hard drive (the very code you use in MATLAB, if you want) using the `\lstinputlisting{/SOME/PATH/FILENAME.M}` command. If you only want to include certain lines from that file (for instance to skip a header), you can use `\lstinputlisting[firstline=6, lastline=15]{/SOME/PATH/FILENAME.M}`.

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<sup>1</sup>Works also in footnotes: `for i=1:3, disp('cool'); end;`