CPBS 7711 MODULE 4 DAY 3 ASSIGNMENT: Aishwarya Mandava

MOTIVATION:

Fanconi Anemia (FA) is a rare genetic disorder inherited in an autosomal recessive pattern and is characterized by physical abnormalities, bone marrow failure, and increased risk of malignancy. Approximately 90% of the individuals with FA have impaired bone marrow function leading to decrease in the production of Red blood cells, White blood cells, and Platelets. Previous studies have found 12 genes associated with FA, including BRCA and FANC genes that play a role in DNA damage response and repair mechanisms. The construction of functional networks involving these FA-associated genes could offer valuable insights into novel molecular mechanisms and pathways. Such analyses could prove valuable in identifying sets of mutually functionally related genes within the loci linked to FA, contributing to a deeper understanding of the disease and offering insights for potential therapeutic interventions.

COMPUTATIONAL PROBLEM:

We have a protein-protein interactions (PPI) network representing various interactions between the nodes/genes and a set of loci associated with FA disease. The computational objective is to prioritize groups of candidate genes from FA-associated loci on the basis of mutual functional relatedness, i.e., mutually connected gene groups. To address this task, an initial population of 5000 random FA associated subnetworks are subjected to a genetic algorithm to enrich dense subnetworks. The resultant optimized subnetwork is compared with non-informative subnetworks, similarly optimized using the genetic algorithm, to test for statistical significance. The final optimized FA associated population is used for computing the gene scores and subsequent ranking of the candidate genes. Visualizing and analyzing the top dense subnetworks would facilitate exploring dynamic interactions and provide insights into potential signatures within the protein-protein network within the specific context of FA disease-associated loci.

SPECIFIC APPROACH:

Given that we have the disease (FA) associated genes and the protein-protein interactions (PPI), the approach involves the initial generation of a population comprising 5000 random subnetworks. Each subnetwork is constructed to include a randomly chosen representative gene from each locus. The optimization of this initial population employs a Genetic Algorithm (GA) comprising mutation and mating steps.

In the mutation step, representative genes in each locus are swapped with other genes from the same locus with a 5% probability (with replacement) uniformly at random. The computation involves determining normalized weighted edge density for each subnetwork i denoted as s_i that is derived from cubically transforming edge weight d_i^3 of the subnetwork.

$$S_i^* = \frac{S_i}{5000}$$
$$\sum_{j=1}^{S} S_j$$

The mating step involves sampling pairs of subnetworks with replacement, with the probability of selecting a parent subnetwork i equal to s_i^* . This results in child subnetworks by randomly selecting each gene in a locus from either parent. The GA optimization terminates when the average density of the child subnetworks fails to improve the average density of the parent population by more than 0.5%.

To test the statistical significance of the Genetic Algorithm-optimized population, a null case is generated through quantile-based binning. In this null case, FA genes within each locus are replaced with a non-FA gene from the same bin. The null case population undergoes optimization using the genetic algorithm, and after repeating this process 1000 times, the p-value is computed by comparing the test statistic in the FA-optimized population with that of the 1000 non-FA optimized populations and is equal to the number of times the null case test statistic is as extreme or more extreme as the test statistic of the optimized FA population.

We then score each candidate gene within the FA population by replacing the representative gene with every other gene in that locus, and subsequently computing the edge weights within each subnetwork. We then compute the "empty case" score by completely ignoring that specific locus. The difference between the candidate gene score and the empty case indicates the contribution of the candidate gene to the rest of the subnetwork. The final average gene scores are computed by taking an average of these candidate gene scores across the 5000 subnetworks.

SPECIFIC IMPLEMENTATION:

A) Data Description:

The protein-protein interactions (PPI) network was retrieved from the STRING database [2] in the tab-delimited format. This file has 1,972,248 interactions across various genes including both FA genes and non-FA genes. The FA disease genes were retrieved from the OMIM [3] database in the Gene Map Table (GMT) format. This file has 12 loci, and 584 FA disease genes. However, not all 584 interact with other genes, only 329 disease associated genes interact with other disease associated genes. **Table 1** shows disease associated genes within loci that are present in the input file and in the string file.

Locus	0	1	2	3	4	5	6	7	8	9	10	11
All FA genes (input file)	46	50	79	50	12	50	50	50	50	48	50	50
FA genes from STRING file (FA gene connected to FA gene)	28	27	54	31	4	27	29	27	28	24	25	26

Table1: Number of disease associated genes

B) Generating 5000 subnetworks with FA genes:

This step involves selecting a random gene from each loci to form a subnetwork. This is repeated 5000 times resulting in the generation of distinct subnetworks. There are 4.67e+16 potential ways of creating a

subnetwork by randomly selecting one gene per locus (note - these 12 genes may or may not have interactions with the other genes in that subnetwork) and each subnetwork could have at most 12C2 or (12*11)/2 or 66 edges. **Figure1** shows 4 loci and illustrates the 5000 random subnetworks generated from the 12 loci. The script uses min_edges parameter to select the minimum number of edges for the subnetwork to be considered for the population. Additionally, the script ensures to create unique subnetworks. By default it is set to 0, i.e., zero or more edges per subnetwork. Setting this higher ensures denser initial population, but the run time would be longer.

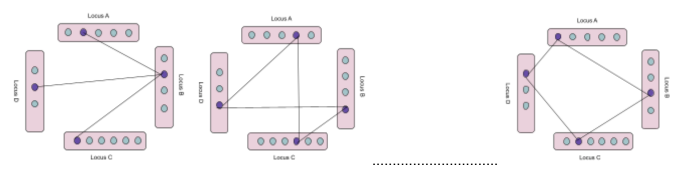


Figure1: 5000 random FA subnetworks

C) Genetic Algorithm:

Mutation: Within the mutation step, there is a 5% probability of swapping the representative genes in each subnetwork. This is achieved by generating a random number between 1 and 20 for each gene in the subnetwork and if the generated number is 1, the corresponding gene undergoes mutation. Additionally, the script ensures that the representative gene is not replaced by itself. The normalized edge weights and average edge weights are computed. **Figure 2** illustrates the mutation step with genes in yellow mutated with 5% probability.

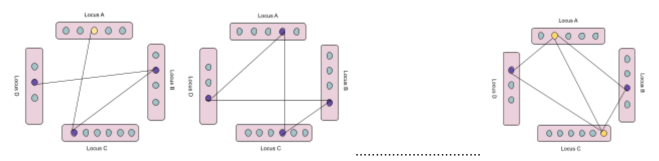


Figure 2: Mutation (Genes in Yellow are mutated with 5% probability)

Mating: Within this step, pairs of subnetworks are sampled from the population resulting in the preceding mutation step, and the probability of selection is determined by their normalized edge weights. From these selected pairs, a child subnetwork is generated by randomly selecting a gene for each locus from either parent. After 5000 such iterations, mean percent difference is calculated as:

((mean average density of the new generation - mean average density of the previous generation)*100)/(mean average density of the previous generation)

Figure 3 illustrates the mating step

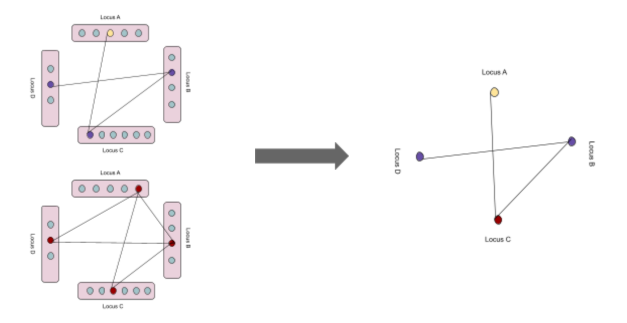


Figure3: Mating step with a pair of subnetworks sampled and a child subnetwork by selecting a gene from each loci from either parent (purple genes are from parent 1, yellow gene is mutated in the previous step, and red genes are from parent 2)

Optimization: The Genetic algorithm terminates when the mean percent difference between the mean population density from the current generation and the mean population density from the previous generation is less than 0.5%. The mating population from the previous generation is used as the final population of disease associated genes.

D) Generating a null case:

The initial population of 5000 random FA subnetworks were used to create the null case. This involved replacing each FA disease-associated gene with a non-FA gene within a similar range of edges. Quantile-based binning was used to categorize both FA and non-FA genes into 100 bins based on their edge numbers, with a minimum frequency of 0, a maximum frequency of 10101, and a bin width of 101. While using average edge weights per gene is more reasonable than the number of edges, this approach led to numerous bins containing 0 non-FA genes. This presented a challenge as replacing an FA gene within such bins is impractical, for instance, 13 bins out of 50 bins do not have non-FA genes.

The population of substituted non-FA subnetworks underwent optimization through the genetic algorithm. In the mutation step, non-FA genes were randomly assigned to 12 loci in each iteration of the null case. The final optimized generation, comprising 5000 subnetworks in each iteration, was used to compute the test statistic. The test statistic, represented by the average edge weight, was compared for each subnetwork of the optimized FA population against the test statistics of 1000 null case populations. The p-value is determined as the number of times the null case test statistic was as extreme or more extreme than that of the FA subnetwork. **Figure 4** illustrates the null case step.

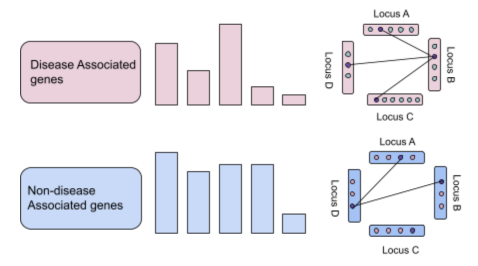


Figure 4: Generating a null case and subjecting it to the genetic algorithm (assigning random loci numbers to non-disease associated genes)

E) Gene Scoring:

In the final optimized population of FA subnetworks, wherein each subnetwork is generated by randomly selecting a representative gene from each locus, the first step computes the gene score equivalent to the number of edges in that specific subnetwork. Subsequently, each representative gene is replaced with every other gene within that specific locus and re-compute the gene score based on the edge counts. For instance, for loci L, comprising of Gi genes (g1, g2, g3,, gn), the representative gene g* is replaced with g1 through gn and re-compute their associated edge counts. For the empty case, the edge counts were computed by completely eliminating the loci L. The final gene score for this subnetwork is the difference between the candidate gene scores and edge density of the empty locus case. Genes with higher gene scores imply high connectivity to the other loci in that subnetwork, while genes with lower scores imply lower connectivity to the other loci in that subnetwork.

The final gene scores are applied across all the 5000 subnetworks in the final population and an average gene score is computed for each gene. In cases where a gene at a locus is not found in the network file, an "NA" score is assigned, implying its absence in the network.

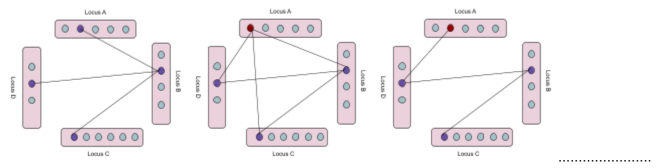


Figure 5: Candidate gene scores

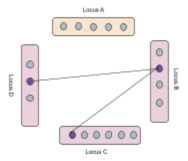


Figure 6: Empty locus case

RESULTS:

A) 5000 subnetworks:

There are 1024 edges that connect FA gene to FA gene. The 5000 random subnetwork generation uses a constraint of 0, i.e., zero or more edges in a subnetwork for it to be considered for the initial population.

B) Genetic Algorithm:

The initial population had a mean edge density of 1.006. The genetic algorithm ran for 13 iterations with the final iteration less than 0.5%. The optimized mean population density is 8.785. Table 2 shows mean percent population density, mean current population density, mean difference, and percentage difference. The optimized subnetwork had a maximum of 20 edges and a minimum of 2 edges in the subnetworks. Table 3 shows the top ten subnetworks arranged in descending order of edge numbers. Figure 7 shows distribution of the first generation.

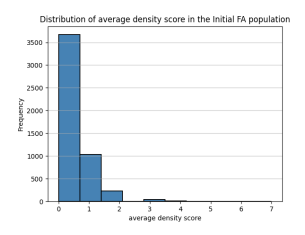
Generation	Mean parent population density	Mean current population density	Mean difference	% difference
1	1.006	0.889	-0.117	-11.630
2	0.889	1.945	1.055	118.673
3	1.945	3.88	1.937	99.598
4	3.88	5.843	1.960	50.480
5	5.84	7.146	1.303	22.311
6	7.146	7.952	0.805	11.268
7	7.952	8.342	0.390	4.910
8	8.342	8.482	0.140	1.681
9	8.482	8.590	0.107	1.265
10	8.590	8.662	0.071	0.837
11	8.662	8.751	0.089	1.036

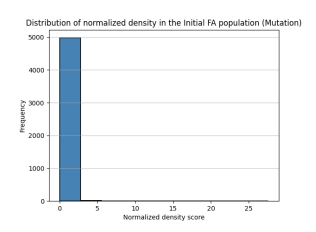
12 8.751 8.785 0.033	0.381
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 Table 2: Genetic Algorithm optimization (13 generations)

Subnetwork	Number of edges
1830	20
783	18
142	18
896	17
3980	17
3723	17
3002	17
734	16
669	16
58	16

Table 3: Top 10 subnetworks sorted by descending order of number of edges





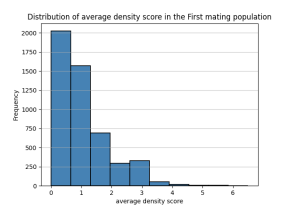


Figure 7: a) Average density of the initial FA population (top left) b) Normalized weights for the first FA population (top right) c) Average density score for the mating step in the first generation (bottom left)

C) Gene Scores: Figures 8 and 9 show the distribution of gene scores across loci.

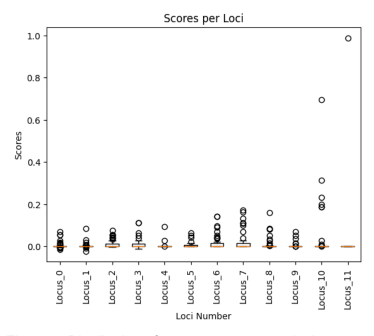


Figure 8: Distribution of gene scores across loci

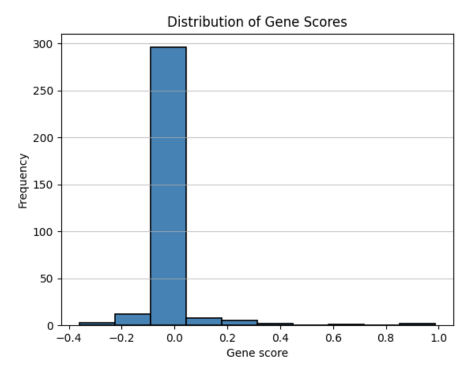


Figure 9: Distribution of gene scores

DISCUSSION:

Table 2 shows results from the genetic algorithm. As we would expect, It is observed that mean population density increases with every generation of the genetic algorithm. Figures in 7 show the increase in mean density within the first generation from mutation step to mating step.

For the gene scores, genes in loci 10 and 11 have the highest gene scores. Genes with high connectivity to other loci in the subnetwork have high scores, and similarly, genes with low connectivity to other loci in the subnetwork have low scores.

LIMITATIONS:

- 1. The null case does not use average edge weights for generating bins. While using average edge weights per gene is more reasonable than the number of edges, this approach led to numerous bins containing 0 non-FA genes. This presented a challenge as replacing an FA gene within such bins is impractical, for instance, 13 bins out of 50 bins do not have non-FA genes.
- 2. The optimization of nonFA networks takes several hours to complete.

PSEUDOCODE:

// Function to create dictionary of input file (loci_diction) and FA network (fa_diction), sets of all fa genes (fa_set) i.e., 584 genes; FA genes that are in the STRING file (string_set), i.e., 508 genes; and FA genes that are only connected to other FA genes (fa string set) i.e., 329 genes

```
FUNCTION data_to_diction(input_path,string_path):

OPEN input_path as input:

CREATE loci_diction as loci ID as key and genes as values

CREATE fa_set with all the genes

INITIALIZE fa_diction empty dictionary

INITIALIZE fa_string_set empty set

INITIALIZE string_set empty list

INITIALIZE all_genes_set

OPEN string_path as string_file:
```

SORT columns of each row

FOR edge in string file:

IF gene in first column and second column are in fa set:

UPDATE genes to fa string set

IF genes in first column is in fa_diction.keys():

IF genes in two columns not in fa diction inner keys:

UPDATE inner dictionary to the key

ELIF first column not in fa diction.keys():

ADD inner dictionary to the key

ELIF gene in first column or second column are in fa set:

IF gene in first column is in fa set:

APPEND gene in first column to string_set list

IF gene in second column is in fa_set:

APPEND gene in second column to string_set list

ADD gene in first column to all_genes_set

ADD gene in second column to all_genes_set

ADD fa_set to all_genes_set

CONVERT string_set list to a set

RETURN loci_diction, fa_set,fa_diction,fa_string_set,string_set,all_genes_set

// Function to create dictionary string file dictionary

FUNCTION string_to_diction(string_path):

INITIALIZE string_diction empty dictionary

INITIALIZE string_gene_set empty set

WITH open(string_path,'r') as string_file:

FOR edge in string_file:

ASSIGN sorted(edge.strip().split('\t')) to genes

UPDATE genes[:2] to string_gene_set

IF first column gene IN string diction:

```
IF ((second column gene NOT IN (string_diction[genes[0]].keys()) AND
string_diction[genes[0]].keys()):
                                 ASSIGN edge weight to string diction[genes[0]][genes[1]]
                   ELIF first column genes NOT IN string diction:
                          ASSIGN {genes[1]:genes[2]} to string_diction[genes[0]]
     RETURN string gene set, string diction
// Function to create the first parent population with 5000 random subnetworks
FUNCTION first parent population(loci diction,fa diction,num iterations=5000,min edges=0):
     INITIALIZE subnetwork diction empty dictionary
     INITIALIZE subnetwork diction edge counts empty dictionary
     INITIALIZE first_parent_subnetwork empty dictionary
     INITIALIZE i to 0
     SET random seed to 123
     WHILE len(subnetwork.keys()) LESS THAN num_iterations:
            INITIALIZE iteration fa genes empty list
            INITIALIZE random index set empty list
            FOR key, value in loci diction.items():
                   ASSIGN random index between 0 and len(value)-1 to random index
                   APPEND value[random index] to iteration fa genes
                   APPEND random index to random index set
            IF NOT ANY(iteration == values FOR values in first_parent_subnetwork.values()):
                   ASSIGN iterations fa genes to first parent subnetwork['subnetwork i']
            INITIALIZE temp list empty list
            FOR outer g, inner d in fa diction.items():
                   FOR inner g,edge value in inner d.items():
                          IF outer g AND inner g IN first parent subnetwork['subnetwork i']:
                                 APPEND([outer g,inner g,edge value]) TO temp list
                          IF len(temp list) >= min edges:
                                 ASSIGN len(temp_list) to subnetwork_diction_edge_counts[subnetwork_i]
                                 ASSIGN temp_list to subnetwork_diction['subnetwork_i']
            i+=1
```

RETURN first parent subnetwork, subnetwork diction

```
// Function to generate a random between a range excluding a specific number
FUNCTION generate random excluding(range start, range end, exclude number):
     FOR num in range(range start,range end+1):
            IF num NOT EQUAL TO exclude number:
                  ASSIGN num to valid numbers
     RETURN random.choice(valid numbers)
// Function to mutate the parent subnetwork
FUNCTION ga mutation(loci diction,fa diction,parent subnetwork,num iterations=5000):
     INITIALIZE subnetwork_diction_updated empty dictionary
     INITIALIZE subnetwork mutated weighted empty dictionary
     INITIALIZE subnetwork_normalized_weighted empty dictionary
     SET random seed to 123
     FOR subnetwork index, subnet genes list IN parent subnetwork.items():
            FOR locus, mut genes in enumerate (subnet genes list):
                  ASSIGN random integer between 1 and 20 to prob_5
                  IF prob 5 EQUAL TO 1:
                         ASSIGN index of mut genes in locus to mutate gene
                         ASSIGN random number excluding mutate gene to rand replace
                         ASSIGN rand replace index gene to parent subnetwork['subnetwork i']
     FOR subnetwork index,genes list IN parent subnetwork.items():
            ASSIGN temp_list_2 empty list
            FOR outer gene, inner diction IN fa diction.items():
                  FOR inner gene, edge value IN inner diction.items():
                              ALL(current fa gene
                                                     IN
                                                           genes list
                                                                        FOR
                                                                                current fa gene
                                                                                                   IN
(outer gene,inner gene)):
                                IF(outer gene IN genes list) AND (inner gene IN genes list):
                                       APPEND([outer_g,inner_g,edge_value]) TO temp_list_2
            ASSIGN temp_list_2 TO subnetwork_diction_updated
            ASSIGN sum of edge weights in temp list 2 to subnetwork mutated weighted
            ASSIGN cubic sum of edge weights in temp list 2 to subnetwork normalized weighted
```

ASSIGN sum of subnetwork normalized weighted.values() to sum norm weights parent

ASSIGN normalized subnetwork mutated weighted to normalized weights parent

RETURN normalized_weights_parent, subnetwork_diction_updated, subnetwork_mutated_weighted, parent subnetwork

// Function for mating a mutated subnetwork

FUNCTION

ga_mating(fa_diction,normalized_weights_parent,subnetwork_mutated_weighted,num_iterations=5000,min_edges=0):

SET random seed to 123

INITIALIZE mate population empty dictionary

INITIALIZE mate_weighted empty dictionary

INITIALIZE iteration to 0

INITIALIZE mate_parent_subnetwork empty dictionary

WHILE len(mate_population.keys()) LESS THAN num_iterations:

ASSIGN random indices to subnet_pair //Based on weights or probabilities INITIALIZE sub_parent empty dictionary

INITIALIZE mate_parent_subnetwork['subnetwork_i'] empty list

FOR r in range(12):

ASSIGN random integer 1 or 2 to pair random

APPEND parent subnetwork[subnet pair[pair random-1][r]]

mate parent subnetwork['subnetwork i']

INITIALIZE temp list 3 empty list

FOR outer_gene,inner_diction in fa_diction.items():

FOR inner gene, edge value in inner diction.items():

IF outer_gene in mate_parent_subnetwork['subnetwork_i'] AND inner_gene in mate_parent_subnetwork['subnetwork_i']:

APPEND outer_gene,inner_gene,edge_value to temp_list_3

to

ASSIGN temp_list_3 to mate_population['subnetwork_i']
ASSIGN sum of edge weights in temp_list_3 to mate_weighted

INCREMENT iteration by 1

ASSIGN sum of mate_weighted.values() to tot_mate_weighted

ASSIGN average mate_weighted.values() to mean_current_weights

```
ASSIGN average subnetwork mutated weighted.values() to mean parent weights
    ASSIGN difference between mean current weights and mean parent weights to mean delta
    ASSIGN percentage difference to mean delta perc
    RETURN mate population, mate parent subnetwork, mate weighted, mean delta, mean delta perc
//Function to a dictionary of gene frequencies
Function gene frequency(string diction, string gene set)
    INITIALIZE gene_freq dictionary
    FOR outer key,inner diction in string diction.items()
           FOR inner key, inner value in inner diction. items()
                  IF outer key is in string gene set
                         INCREMENT gene_freq[outer_key] by 1
                  IF inner key is in string gene set
                         INCREMENT gene_freq[inner_key] by 1
    RETURN gene_freq
//Function to create a nested dictionary of bin sizes for fa genes and non fa genes
Function bin dictionary(gene freq,num bins,filter bin data)
    Bin_width = (maximum(gene_freq.values)-minimum(gene_freq.values))/num_bins
    INITIALIZE an empty dictionary bin diction
    FOR i in range of num bins
           Lower bound = minimum(gene freq.values)+i*bin width
           Upper_bound=maximum(gene_freq.values)+i*bin_width
           INITIALIZE outer keys in bin diction
           INITIALIZE inner keys in bin diction
           FOR gene, freq in gene_freq.items
                  IF freq >= lower_bound and freq <= upper_bound
                         IF gene is in fa set
                                APPEND gene to bin_diction fa_genes
                         ELSE
                                APPEND gene to bin diction non fa genes
// Function to match the bins for a given FA gene
Function match_fa_nonfa_by_bucket(nested_dict,disease_gene)
    FOR out g, inner dict in nested dict.items()
```

CALL match fa nonfa by bucket(inner gene, disease gene)

ASSIGN out g to bin cat

IF inner dict is dictionary

IF result is not NONE

RETURN out_g

```
//Function to generate nonFA subnetwork from FA
FUNCTION generate nonfa from fa(optimized FA subnetwork,random seed,bin diction):
    INITIALIZE nonFA parent empty dictionary
    INITIALIZE nonFA parent size empty dictionary
    INITIALIZE iteration to 0
    INITIALIZE nonFA parent genes empty dictionary
    INITIALIZE all nonFA null empty set
    SET random seed to random seed
    FOR subnet index, list genes in optimized FA subnetwork. items():
           INITIALIZE iteration_nonfa_genes empty list
           INITIALIZE nonFA parent genes['subnetwork '+str(iteration+1)] empty list
           FOR genes in list genes:
                  CALL match fa nonfa by bucket(bin diction,genes) and ASSIGN to bin
                  ASSIGN bin_diction[bin]['non_fa_genes'] to non_fa_values
                  ASSIGN random integer to random index nonfa
                  APPEND non_fa_values[random_index_nonfa] to iteration_nonfa_genes
                  ADD non fa values[random index nonfa] to all nonFA null
           ASSIGN iteration nonfa genes to nonFA parent genes['subnetwork '+str(iteration+1)]
           INITIALIZE temp list 4 empty list
           FOR outer gene, inner dict in bin diction. items():
                  IF outer_gene IN iteration_nonfa_genes:
                         FOR inner gene, inner edge IN inner diction. items():
                                IF inner gene IN iteration nonfa genes:
                                       APPEND outer_gene, inner_gene, inner_edge to temp_list_4
           INCREMENT iteration by 1
           ASSIGN temp_list_4 to nonFA_parent['subnetwork '+str(iteration+1)]
           ASSIGN len(temp_list_4) to nonFA_parent_size['subnetwork_'+str(iteration+1)]
      RETURN nonFA_parent_nonFA_parent_size,nonFA_parent_genes
// Function to create random loci for nonFA genes
FUNCTION generate random loci NonFA(nonFA parent genes):
```

ASSIGN {'Locus_{}'.format(I): [] for I in range(12)} to non_fa_loci INITIALIZE all_nonfa_subnet_genes empty set

FOR nonfa_sub,non_fa in nonFA_parent_genes.items():

FOR nonfa_locus,non_fa_g in enumerate(non_fa):

ADD non_fa_g to all_nonfa_subnet_genes

ASSIGN non_fa.index(non_fa_g) to nonfa_index

APPEND non_fa_g to non_fa_loci['Locus_{}'.format(nonfa_index)]

FOR i in all_genes_set:

IF i not in all_nonfa_subnet_genes:

ASSIGN a random integer between 0,11 to rand_nonfa_loci

RETURN non_fa_loci