Supplementary Materials And Methods

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1. Functions defined for ceRNA models and workflow of method

We defined the functions that can be used with R programming. Briefly, these functions process a given miRNA:gene dataset and convert to graph object. All values that are significant in miRNA:target interactions are stored in edge variables and processed with formulations that are given in previous section. The functions and steps of approach are explained as following (Figure S1 in Supplementary Figures):

Conversion of dataset: priming_graph() function processes the given dataset that includes competing elements in first variable and repressive element in second variable. If the affinity and/or degradation factors are specified in the function, factors are taken into account, are processed with defaults in vice versa. The formulations that are given in equations (1-4) are performed in this function. This step gives the graph object which contains efficiency values of miRNA:competing target pairs in steady-state in terms of amount. It is assumed that the initial target amounts in the dataset is observed after the repressive activity of miRNAs in steady-state.

Transition of variables in graph: In the previous step, the calculations are performed in the edge variables of the graph object. However, the graph object allows to use node variables, while the node features are handled to the graph. In this direction, update_nodes function carries the amount values to node variables. This step must be applied with "once" option because it is primary process.

Trigger change in graph: The dataset are assumed as steady-state in previous step and the efficiency coefficients are calculated according to this acceptance. In the network that is found in steady-state conditions, the change is applied to the graph object for distribution of steady-state. To provide the distribution in the network the workflow offer two methods: update_variables and update_how. The first, a new dataset that is contained competing and repressive element names and current values of these can be processed with update_variables. The second option, the amount of the given node name in update_how function can be changed according to "how" argument.

Updating current values of variables: After variables updating in edge variables, these are carried to node variables. Current and previous values of variables are stored as node variables with update_variables function.

Simulation of competing behavior of targets: After the change in the steady-state conditions, the network elements try to gain steady-state again. This process progresses as repeating of regulations after the spreading the changes in the network. In this step, simulation of regulations according to given cycle count in simulate function is applied. After each simulation cycle, the miRNA repression values are re-calculated and the current values of competing elements are found and saved. The process is performed in the edge data and at the same time outputs of the calculations are carried from edge to node data.

The node elements in the dataset are handled as two type; repressive (miRNAs) and competing (targets). It is assumed in approach that while targets are degrading or inhibiting by miRNAs continuously, miRNAs reversibly used. If the trigger of the network is a miRNA, it maintains the current value of amount that provides by user. On the contrary, it tries to help this process to provide steady-state through the regulations on its amount, if a competing element is used as a trigger. The functions that are used in the approach are developed with R programming so as can be used with other packages. These are can be found in the Github repository ceRNAnetsim Github page and improved with contributions from others.

```
# install.packages('deutools')
# deutools::install_github('selcenari/ceRNAnetsim')
library(ceRNAnetsim)
```

• load minsamp data

```
data("minsamp")
minsamp
     competing miRNA Competing_expression miRNA_expression seed_type region
##
## 1
         Gene1 Mir1
                                    10000
                                                       1000
                                                                 0.43
                                                                        0.30
## 2
         Gene2 Mir1
                                    10000
                                                       1000
                                                                 0.43
                                                                        0.01
         Gene3 Mir1
## 3
                                     5000
                                                       1000
                                                                 0.32
                                                                        0.40
## 4
        Gene4 Mir1
                                    10000
                                                       1000
                                                                 0.23
                                                                        0.50
                                                                 0.35
                                                                        0.90
## 5
        Gene4 Mir2
                                    10000
                                                       2000
## 6
        Gene5 Mir2
                                     5000
                                                       2000
                                                                 0.05
                                                                        0.40
## 7
         Gene6 Mir2
                                     10000
                                                       2000
                                                                 0.01
                                                                        0.80
##
     energy
## 1
        -20
## 2
        -15
## 3
        -14
## 4
        -10
## 5
        -12
```

See Table S1 in Supplementary Tables file.

6

7

-11

-25

minsamp dataset analysis in lack of interaction factors.

Firstly, we have analysed minimal data without interaction factors between miRNA:target.

• 1. We have evaluated graph in the steady state conditions as followings (Figure S2 in Supplementary Figures):

```
priming_graph(minsamp, competing_count = Competing_expression,
    miRNA_count = miRNA_expression)
```

```
## # A tbl_graph: 8 nodes and 7 edges
## #
## # A rooted tree
## #
## # Node Data: 8 x 7 (active)
     name type node_id initial_count count_pre count_current
##
     <chr> <chr>
                   <int>
                                  <dbl>
                                            <dbl>
                                                          <dbl>
## 1 Gene1 Comp~
                       1
                                  10000
                                            10000
                                                           10000
## 2 Gene2 Comp~
                       2
                                  10000
                                            10000
                                                          10000
## 3 Gene3 Comp~
                       3
                                  5000
                                            5000
                                                           5000
## 4 Gene4 Comp~
                       4
                                  10000
                                            10000
                                                          10000
```

```
## 5 Gene5 Comp~
                       5
                                   5000
                                              5000
                                                            5000
                                            10000
                       6
                                  10000
                                                           10000
## 6 Gene6 Comp~
## # ... with 2 more rows, and 1 more variable: changes_variable <chr>
## #
## # Edge Data: 7 x 19
##
      from
              to Competing name miRNA name Competing expre~ miRNA expression
##
     <int> <int> <chr>
                                 <chr>>
                                                        <dbl>
                                                                          <dbl>
               7 Gene1
## 1
         1
                                 Mir1
                                                        10000
                                                                           1000
## 2
         2
               7 Gene2
                                 Mir1
                                                        10000
                                                                           1000
## 3
         3
                                                                           1000
               7 Gene3
                                 Mir1
                                                         5000
    ... with 4 more rows, and 13 more variables: dummy <dbl>,
## #
       afff_factor <dbl>, degg_factor <dbl>, comp_count_list <list>,
## #
       comp_count_pre <dbl>, comp_count_current <dbl>,
## #
       mirna_count_list <list>, mirna_count_pre <dbl>,
## #
       mirna_count_current <dbl>, mirna_count_per_dep <dbl>,
## #
       effect_current <dbl>, effect_pre <dbl>, effect_list <list>
```

• 2. We have obtained graph after change on Gene2 expression as following (Figure S3 in Supplementary Figures):

```
priming_graph(minsamp, competing_count = Competing_expression,
    miRNA_count = miRNA_expression) %>% update_how("Gene2",
    2)
```

```
## # A tbl_graph: 8 nodes and 7 edges
## # A rooted tree
## #
## # Node Data: 8 x 7 (active)
     name type node_id initial_count count_pre count_current
##
                    <int>
                                  <dbl>
                                             <dbl>
     <chr> <chr>
                                                           <dbl>
## 1 Gene1 Comp~
                                  10000
                                             10000
                                                           10000
                        1
                        2
                                             10000
## 2 Gene2 Comp~
                                  10000
                                                           20000
## 3 Gene3 Comp~
                        3
                                   5000
                                              5000
                                                            5000
                        4
                                             10000
                                                           10000
## 4 Gene4 Comp~
                                  10000
## 5 Gene5 Comp~
                        5
                                   5000
                                              5000
                                                            5000
                        6
                                  10000
## 6 Gene6 Comp~
                                             10000
                                                           10000
## # ... with 2 more rows, and 1 more variable: changes_variable <chr>
## #
## # Edge Data: 7 x 19
##
              to Competing_name miRNA_name Competing_expre~ miRNA_expression
##
     <int> <int> <chr>
                                 <chr>>
                                                        <dbl>
                                                                          <dbl>
## 1
         1
               7 Gene1
                                 Mir1
                                                        10000
                                                                           1000
## 2
         2
                                                        10000
               7 Gene2
                                 Mir1
                                                                           1000
## 3
         3
               7 Gene3
                                 Mir1
                                                         5000
                                                                           1000
     ... with 4 more rows, and 13 more variables: dummy <dbl>,
       afff_factor <dbl>, degg_factor <dbl>, comp_count_list <list>,
## #
       comp_count_pre <dbl>, comp_count_current <dbl>,
## #
       mirna_count_list <list>, mirna_count_pre <dbl>,
## #
       mirna_count_current <dbl>, mirna_count_per_dep <dbl>,
## #
       effect_current <dbl>, effect_pre <dbl>, effect_list <list>
```

• 3. We have determined regulations after Gene2 upregulation (Figure S4 in Supplementary Figures):

```
priming_graph(minsamp, competing_count = Competing_expression,
    miRNA_count = miRNA_expression) %>% update_how("Gene2",
    2) %>% simulate(cycle = 2)
```

```
## # A tbl_graph: 8 nodes and 7 edges
## #
## # A rooted tree
## #
## # Node Data: 8 x 7 (active)
    name type node_id initial_count count_pre count_current
     <chr> <chr>
                   <int>
                                  <dbl>
                                            <dbl>
                                                           <dbl>
                                           10063.
                                                          10062.
## 1 Gene1 Comp~
                       1
                                  10000
## 2 Gene2 Comp~
                       2
                                  10000
                                           19841.
                                                          19845.
## 3 Gene3 Comp~
                       3
                                   5000
                                            5032.
                                                          5031.
## 4 Gene4 Comp~
                       4
                                  10000
                                           10063.
                                                          10059.
                       5
## 5 Gene5 Comp~
                                   5000
                                            5000
                                                           5001.
                       6
                                  10000
                                           10000
                                                          10002.
## 6 Gene6 Comp~
## # ... with 2 more rows, and 1 more variable: changes_variable <chr>
## #
## # Edge Data: 7 x 20
##
      from
              to Competing_name miRNA_name Competing_expre~ miRNA_expression
##
     <int> <int> <chr>
                                 <chr>>
                                                        <dbl>
                                                                         <dbl>
## 1
         1
               7 Gene1
                                 Mir1
                                                        10000
                                                                           1000
## 2
         2
               7 Gene2
                                 Mir1
                                                        10000
                                                                           1000
## 3
         3
               7 Gene3
                                                                          1000
                                 Mir1
                                                         5000
## # ... with 4 more rows, and 14 more variables: dummy <dbl>,
       afff_factor <dbl>, degg_factor <dbl>, comp_count_list <list>,
       comp_count_pre <dbl>, comp_count_current <dbl>,
## #
       mirna_count_list <list>, mirna_count_pre <dbl>,
## #
       mirna_count_current <dbl>, mirna_count_per_dep <dbl>,
## #
       effect_current <dbl>, effect_pre <dbl>, effect_list <list>,
## #
       mirna_count_per_comp <dbl>
```

Note that the regulations are colored according to expression changes of present and a previous value. So, it can be observed that whole gene expressions increase in comparison of initial steady-state. The overall regulations of gene expressions are as followings:

```
priming_graph(minsamp, competing_count = Competing_expression,
    miRNA_count = miRNA_expression) %>% update_how("Gene2",
    2) %>% simulate(2) %>% activate(edges) %>% as_tibble() %>%
    select(Competing_name, comp_count_list, effect_list) %>%
    unnest()
```

```
## # A tibble: 21 x 3
##
      Competing_name comp_count_list effect_list
##
                                <dbl>
## 1 Gene1
                               10000
                                              286.
                               10063.
    2 Gene1
                                              222.
## 3 Gene1
                               10062.
                                              224.
## 4 Gene2
                               10000
                                              286.
## 5 Gene2
                               19841.
                                              444.
## 6 Gene2
                               19845.
                                              441.
```

```
## 7 Gene3 5000 143.
## 8 Gene3 5032. 111.
## 9 Gene3 5031. 112.
## 10 Gene4 10000 286.
## # ... with 11 more rows
```

A tbl_graph: 8 nodes and 7 edges

A rooted tree

minsamp dataset analysis with interaction factors.

We have made the same analysis in presence of interaction factors (Sequentially shown at Figure S5-7 in Supplementary Figures).

```
priming_graph(minsamp, competing_count = Competing_expression,
   miRNA count = miRNA expression, aff factor = c(energy,
        seed_type), deg_factor = region)
## # A tbl graph: 8 nodes and 7 edges
## #
## # A rooted tree
## #
## # Node Data: 8 x 7 (active)
##
    name type node_id initial_count count_pre count_current
##
     <chr> <chr>
                  <int>
                                 <dbl>
                                           <dbl>
                                                          <dbl>
## 1 Gene1 Comp~
                      1
                                 10000
                                           10000
                                                          10000
## 2 Gene2 Comp~
                       2
                                 10000
                                           10000
                                                          10000
## 3 Gene3 Comp~
                       3
                                                          5000
                                  5000
                                            5000
## 4 Gene4 Comp~
                       4
                                 10000
                                           10000
                                                          10000
                       5
                                                          5000
## 5 Gene5 Comp~
                                  5000
                                            5000
## 6 Gene6 Comp~
                       6
                                 10000
                                           10000
                                                          10000
## # ... with 2 more rows, and 1 more variable: changes_variable <chr>
## #
## # Edge Data: 7 x 22
              to Competing name miRNA name Competing expre~ miRNA expression
##
     <int> <int> <chr>
                                <chr>
                                                       <dbl>
                                                                        <dbl>
               7 Gene1
                                                       10000
                                                                         1000
## 1
         1
                                Mir1
## 2
               7 Gene2
                                Mir1
                                                       10000
                                                                         1000
## 3
         3
               7 Gene3
                                Mir1
                                                        5000
                                                                         1000
## # ... with 4 more rows, and 16 more variables: energy <dbl>,
       seed_type <dbl>, region <dbl>, dummy <dbl>, afff_factor <dbl>,
       degg_factor <dbl>, comp_count_list <list>, comp_count_pre <dbl>,
## #
       comp_count_current <dbl>, mirna_count_list <list>,
## #
## #
       mirna_count_pre <dbl>, mirna_count_current <dbl>,
       mirna_count_per_dep <dbl>, effect_current <dbl>, effect_pre <dbl>,
## #
## #
       effect_list <list>
priming_graph(minsamp, competing_count = Competing_expression,
   miRNA_count = miRNA_expression, aff_factor = c(energy,
        seed_type), deg_factor = region) %>% update_how("Gene2",
   2)
```

```
## #
## # Node Data: 8 x 7 (active)
    name type node id initial count count pre count current
     <chr> <chr>
##
                   <int>
                                 <dbl>
                                            <dbl>
## 1 Gene1 Comp~
                       1
                                 10000
                                            10000
                                                          10000
                       2
                                            10000
                                                          20000
## 2 Gene2 Comp~
                                 10000
## 3 Gene3 Comp~
                       3
                                  5000
                                             5000
                                                           5000
## 4 Gene4 Comp~
                       4
                                  10000
                                            10000
                                                          10000
## 5 Gene5 Comp~
                       5
                                  5000
                                             5000
                                                           5000
## 6 Gene6 Comp~
                       6
                                  10000
                                            10000
                                                          10000
## # ... with 2 more rows, and 1 more variable: changes_variable <chr>
## #
## # Edge Data: 7 x 22
              to Competing_name miRNA_name Competing_expre~ miRNA_expression
##
     <int> <int> <chr>
                                                       <dbl>
                                <chr>
## 1
         1
               7 Gene1
                                 Mir1
                                                        10000
                                                                          1000
## 2
         2
               7 Gene2
                                                       10000
                                                                          1000
                                Mir1
## 3
         3
               7 Gene3
                                Mir1
                                                        5000
                                                                          1000
## # ... with 4 more rows, and 16 more variables: energy <dbl>,
       seed type <dbl>, region <dbl>, dummy <dbl>, afff factor <dbl>,
## #
       degg_factor <dbl>, comp_count_list <list>, comp_count_pre <dbl>,
       comp count current <dbl>, mirna count list <list>,
       mirna_count_pre <dbl>, mirna_count_current <dbl>,
## #
       mirna count per dep <dbl>, effect current <dbl>, effect pre <dbl>,
## #
       effect list <list>
priming_graph(minsamp, competing_count = Competing_expression,
    miRNA_count = miRNA_expression, aff_factor = c(energy,
        seed_type), deg_factor = region) %>% update_how("Gene2",
    2) %>% simulate(cycle = 2)
## # A tbl_graph: 8 nodes and 7 edges
## #
## # A rooted tree
## #
## # Node Data: 8 x 7 (active)
   name type node_id initial_count count_pre count_current
     <chr> <chr>
                   <int>
                                  <dbl>
                                            <dbl>
                                                          <dbl>
## 1 Gene1 Comp~
                                  10000
                                           10065.
                                                         10064.
                       1
## 2 Gene2 Comp~
                       2
                                 10000
                                           19997.
                                                         19997.
## 3 Gene3 Comp~
                       3
                                  5000
                                           5023.
                                                          5023.
## 4 Gene4 Comp~
                       4
                                  10000
                                           10029.
                                                         10028.
## 5 Gene5 Comp~
                       5
                                  5000
                                            5000
                                                          5000.
## 6 Gene6 Comp~
                       6
                                  10000
                                           10000
                                                         10000.
## # ... with 2 more rows, and 1 more variable: changes_variable <chr>
## # Edge Data: 7 x 23
##
              to Competing_name miRNA_name Competing_expre~ miRNA_expression
      from
     <int> <int> <chr>
                                                                         <dbl>
                                 <chr>>
                                                       <dbl>
                                                       10000
## 1
               7 Gene1
         1
                                Mir1
                                                                          1000
## 2
         2
               7 Gene2
                                Mir1
                                                       10000
                                                                          1000
               7 Gene3
         3
                                Mir1
                                                        5000
                                                                          1000
## # ... with 4 more rows, and 17 more variables: energy <dbl>,
## # seed_type <dbl>, region <dbl>, dummy <dbl>, afff_factor <dbl>,
```

```
## # degg_factor <dbl>, comp_count_list <list>, comp_count_pre <dbl>,
## # comp_count_current <dbl>, mirna_count_list <list>,
## # mirna_count_pre <dbl>, mirna_count_current <dbl>,
## # mirna_count_per_dep <dbl>, effect_current <dbl>, effect_pre <dbl>,
## # effect_list <list>, mirna_count_per_comp <dbl>
```

When the graphs which were resulted from analyses were examined, it was observed that behaviors were same. But, when the results were analysed in terms of expression values, the regulation differences can be observed.

```
priming_graph(minsamp, competing_count = Competing_expression,
    miRNA_count = miRNA_expression, aff_factor = c(energy,
        seed_type), deg_factor = region) %>% update_how("Gene2",
    2) %>% simulate(3) %>% activate(edges) %>% as_tibble() %>%
    select(Competing_name, comp_count_list, effect_list) %>%
    unnest()
```

```
## # A tibble: 28 x 3
      Competing_name comp_count_list effect_list
##
      <chr>
                                <dbl>
                                            <dbl>
##
   1 Gene1
                               10000
                                           263.
##
  2 Gene1
                               10065.
                                           198.
##
  3 Gene1
                               10064.
                                           199.
##
   4 Gene1
                               10064.
                                           199.
## 5 Gene2
                               10000
                                             6.58
##
  6 Gene2
                               19997.
                                             9.91
  7 Gene2
                                             9.88
##
                               19997.
                                             9.88
## 8 Gene2
                               19997.
## 9 Gene3
                                            91.5
                                5000
## 10 Gene3
                                            68.8
                                5023.
## # ... with 18 more rows
```

Common target perturbation in *minsamp* dataset.

Genes targeted by multiple miRNAs (referred to as "common target") are of special interest since they are subject to cooperative effect. Also, they perturb more than one neighborhood. In our small dataset, minsamp, Gene4 is regulated by two miRNAs. Let's simulate perturbation effects triggered by Gene4 (Shown at Figure S8 in Supplementary Figures .

```
## # A tbl_graph: 8 nodes and 7 edges
## #
## # A rooted tree
## #
## # Node Data: 8 x 7 (active)
##
    name type node_id initial_count count_pre count_current
                   <int>
     <chr> <chr>
                                 <dbl>
                                           <dbl>
                                                         <dbl>
## 1 Gene1 Comp~
                                 10000
                                          10028.
                                                       10027.
                       1
```

```
10001.
## 2 Gene2 Comp~
                                  10000
                                           10001.
## 3 Gene3 Comp~
                       3
                                  5000
                                            5010.
                                                          5009.
## 4 Gene4 Comp~
                       4
                                           19803.
                                  10000
                                                         19806.
                       5
                                                          5024.
## 5 Gene5 Comp~
                                   5000
                                            5024.
## 6 Gene6 Comp~
                       6
                                  10000
                                           10044.
                                                          10044.
## # ... with 2 more rows, and 1 more variable: changes variable <chr>
## # Edge Data: 7 x 23
##
      from
              to Competing_name miRNA_name Competing_expre~ miRNA_expression
     <int> <int> <chr>
                                <chr>
                                                       <dbl>
##
                                                                         <dbl>
         1
               7 Gene1
                                Mir1
                                                       10000
                                                                          1000
         2
               7 Gene2
                                                       10000
                                                                          1000
## 2
                                Mir1
               7 Gene3
                                                                          1000
## 3
                                Mir1
                                                        5000
## # ... with 4 more rows, and 17 more variables: energy <dbl>,
       seed_type <dbl>, region <dbl>, dummy <dbl>, afff_factor <dbl>,
       degg_factor <dbl>, comp_count_list <list>, comp_count_pre <dbl>,
## #
## #
       comp_count_current <dbl>, mirna_count_list <list>,
       mirna count pre <dbl>, mirna count current <dbl>,
## #
      mirna_count_per_dep <dbl>, effect_current <dbl>, effect_pre <dbl>,
## #
       effect_list <list>, mirna_count_per_comp <dbl>
```

The common target perturbation (increasing to two fold at Gene4 expression in presence of interaction factors) resulted in more prominent efficiency at the same conditions (shown in following).

```
priming_graph(minsamp, competing_count = Competing_expression,
    miRNA_count = miRNA_expression, aff_factor = c(energy,
        seed_type), deg_factor = region) %>% update_how("Gene4",
    2) %>% simulate(3) %>% activate(edges) %>% as_tibble() %>%
    select(Competing_name, comp_count_list, effect_list) %>%
    unnest()
```

```
## # A tibble: 28 x 3
##
      Competing_name comp_count_list effect_list
                                            <dbl>
      <chr>
##
                                <dbl>
##
  1 Gene1
                               10000
                                           263.
## 2 Gene1
                               10028.
                                           236.
## 3 Gene1
                               10027.
                                           237.
## 4 Gene1
                               10027.
                                           237.
## 5 Gene2
                               10000
                                             6.58
## 6 Gene2
                                             5.89
                               10001.
## 7 Gene2
                                             5.90
                               10001.
## 8 Gene2
                               10001.
                                             5.90
## 9 Gene3
                                5000
                                            91.5
## 10 Gene3
                                5010.
                                            81.9
## # ... with 18 more rows
```

Determination of perturbation efficiencies of elements in system.

```
find_node_perturbation(sample_graph, how = 2, cycle = 3,
    limit = 0.1)
## # A tibble: 8 x 9
##
     name type node_id initial_count count_pre count_current
##
     <chr> <chr>
                   <int>
                                  <dbl>
                                             <dbl>
                                                           <dbl>
                                  10000
                                            10000
                                                           10000
## 1 Gene1 Comp~
                       1
## 2 Gene2 Comp~
                       2
                                  10000
                                             10000
                                                           10000
## 3 Gene3 Comp~
                       3
                                   5000
                                             5000
                                                            5000
## 4 Gene4 Comp~
                                  10000
                                            10000
                                                           10000
## 5 Gene5 Comp~
                       5
                                   5000
                                             5000
                                                            5000
## 6 Gene6 Comp~
                       6
                                  10000
                                                           10000
                                            10000
## 7 Mir1 miRNA
                       7
                                   1000
                                             1000
                                                            1000
## 8 Mir2 miRNA
                       8
                                   2000
                                             2000
                                                            2000
## # ... with 3 more variables: changes_variable <chr>,
      perturbation_efficiency <dbl>, perturbed_count <dbl>
```

2. Obtaining breast cancer dataset and integration

This section describes how to apply ceRNAnetsim package on a breast cancer patient miRNA:target interaction dataset. Before the approach, we obtained three datasets and combined them.

2.1 How to get gene expression counts of TCGA-E9-A1N5 patient.

We have obtained the gene expression values of patient using TCGAbiolinks package from Bioconductor. For this process, we have followed the instructions of the package. TCGAbiolinks package provides to obtain data for whole number of given barcode(s) at once. But, we preferred to download them separately to show datasets.

• Obtain to gene expression counts of tumor tissue.

```
BCP_tumor <- GDCquery(project = "TCGA-BRCA", data.category = "Transcriptome Profiling",
    data.type = "Gene Expression Quantification", workflow.type = "HTSeq - Counts",
    barcode = "TCGA-E9-A1N5-01A-11R-A14D-07")

GDCdownload(BCP_tumor)

BCPGE_tumor <- GDCprepare(BCP_tumor)

TCGA_E9_A1N5_tumor <- as.data.frame(assay(BCPGE_tumor)) %>%
    mutate(ensembl_gene_id = rownames(.)) %>% dplyr::inner_join(as.data.frame(rowData(BCPGE_tumor)),
    by = "ensembl_gene_id") %>% dplyr::select(ensembl_gene_id,
    external_gene_name, 1)

colnames(TCGA_E9_A1N5_tumor)[3] <- "GE_tumor"

head(TCGA_E9_A1N5_tumor)</pre>
```

• Obtain to gene expression counts of normal tissue.

```
BCP_normal <- GDCquery(project = "TCGA-BRCA", data.category = "Transcriptome Profiling",
    data.type = "Gene Expression Quantification", workflow.type = "HTSeq - Counts",
    barcode = "TCGA-E9-A1N5-11A-41R-A14D-07")
GDCdownload(BCP_normal)

BCPGE_normal <- GDCprepare(BCP_normal)

TCGA_E9_A1N5_normal <- as.data.frame(assay(BCPGE_normal)) %>%
    mutate(ensembl_gene_id = rownames(.)) %>% dplyr::inner_join(as.data.frame(rowData(BCPGE_normal)),
    by = "ensembl_gene_id") %>% dplyr::select(ensembl_gene_id,
    external_gene_name, 1)

colnames(TCGA_E9_A1N5_normal)[3] <- "GE_normal"

head(TCGA_E9_A1N5_normal)</pre>
```

2.2 How to get miRNA expression counts of TCGA-E9-A1N5 patient.

We have used TCGAbiolinks package to obtain miRNA expression quantification. The query gives read count of miRNA as isoform chromosome coordination. The data also contains mature miRNA information. So, we processed data to attain -5p -3p isoform information using mirbase release21 dataset.

• Get the mirbase id of mature miRNA:

We downloaded the mirbase release 21 dataset from mirbase and processed the patient mirna expression datasets as following:

```
mirbase_id_conv <- read_tsv("hsa_mirna.txt", comment = "#",
    col_names = FALSE) %>% dplyr::select(mirna_type = X3,
    definition = X9) %>% filter(!endsWith(mirna_type,
        "primary_transcript")) %>% tidyr::separate(definition,
    c("ID", "Alias", "Name", "Derivated"), sep = ";") %>%
    dplyr::select(Alias, Name) %>% tidyr::separate(Alias,
    c("trash1", "ID"), sep = "=") %>% tidyr::separate(Name,
    c("trash2", "Name"), sep = "=") %>% dplyr::select(-trash1,
    -trash2)
```

```
## # A tibble: 6 x 2
## ID Name
## <a href="mailto:chr">chr</a> <chr>
## 1 MIMAT0027618 hsa-miR-6859-5p
## 2 MIMAT0027619 hsa-miR-6859-3p
## 3 MIMAT0005890 hsa-miR-1302
## 4 MIMAT0027618 hsa-miR-6859-5p
## 5 MIMAT0027619 hsa-miR-6859-3p
## 6 MIMAT0049032 hsa-miR-12136
```

• Obtain the miRNA expression of tumor tissue of patient:

```
BCP_mirnatumor <- GDCquery(project = "TCGA-BRCA", data.category = "Transcriptome Profiling",
    data.type = "Isoform Expression Quantification",
    workflow.type = "BCGSC miRNA Profiling", barcode = "TCGA-E9-A1N5-01A-11R-A14C-13")
GDCdownload(BCP_mirnatumor)
BCPME_mirnatumor <- GDCprepare(BCP_mirnatumor) %>%
    as.data.frame() %>% dplyr::select(miRNA ID, read count,
   reads_per_million_miRNA_mapped, miRNA_region) %>%
   dplyr::filter(startsWith(miRNA_region, "mature")) %>%
    dplyr::mutate(mirbase_id = str_remove(miRNA_region,
        "mature,")) %>% dplyr::select(-miRNA_region) %>%
   dplyr::inner_join(mirbase_id_conv, by = c(mirbase_id = "ID")) %>%
   dplyr::select(miRNA_name = Name, read_count, reads_per_million_miRNA_mapped) %>%
    dplyr::group_by(miRNA_name) %>% mutate(read_count = sum(read_count),
   reads_per_million_miRNA_mapped = sum(reads_per_million_miRNA_mapped)) %>%
    dplyr::ungroup() %>% distinct()
head(BCPME_mirnatumor)
```

• Obtain the miRNA expression of normal tissue of patient:

```
BCP_mirnanormal <- GDCquery(project = "TCGA-BRCA",
    data.category = "Transcriptome Profiling", data.type = "Isoform Expression Quantification",
    workflow.type = "BCGSC miRNA Profiling", barcode = "TCGA-E9-A1N5-11A-41R-A14C-13")
GDCdownload(BCP mirnanormal)
\# a616435d-0b69-48ac-813d-5d75ad9b85eb.mirbase21.isoforms.quantification.txt
BCPME_mirnanormal <- GDCprepare(BCP_mirnanormal) %>%
    as.data.frame() %>% dplyr::select(miRNA_ID, read_count,
    reads_per_million_miRNA_mapped, miRNA_region) %>%
   dplyr::filter(startsWith(miRNA_region, "mature")) %>%
    dplyr::mutate(mirbase_id = str_remove(miRNA_region,
        "mature,")) %>% dplyr::select(-miRNA_region) %>%
    dplyr::inner_join(mirbase_id_conv, by = c(mirbase_id = "ID")) %>%
   dplyr::select(miRNA_name = Name, read_count, reads_per_million_miRNA_mapped) %>%
    dplyr::group_by(miRNA_name) %>% mutate(read_count = sum(read_count),
   reads_per_million_miRNA_mapped = sum(reads_per_million_miRNA_mapped)) %>%
    dplyr::ungroup() %>% distinct()
head(BCPME_mirnanormal)
```

2.3 Get the high-throughput experimental miRNA:target dataset.

There are various datasets about miRNA:target pairs such as miRTarBase, DianaTools, miRecords, miRWalk etc. Some of these present the experimentally supported miRNA target pairs or only predicted ones. The experimentally supported datasets generally provides weak evidence for interactions. For these reasons, we obtained the high-throughput experimental miRNA:target dataset from two studies performed by Helwak et al. and Moore et al. These steps were not handle in this file because they contain many processes.

Briefly these datasets contain various common information about miRNA:target interactions such as the miRNA name, miRNAsequence, target name, target sequence, their chromosomal locations, binding location

on the target sequence, binding free energy, seed structure. But these datasets provides the informations with different data structures. So we followed the steps:

- The datasets were directly downloaded from supplementary data files of the studies.
- It was provided that the datasets are converted to same human genome build.
- The seed type information was organized as the same style.
- The datasets were combined.
- We committed the interaction factors as numeric values according to previous studies. (We added the interaction factors and their numeric values at Supplementary files, Process on miRNA:target pairs dataset for process codes and Supplementary Tables for used numeric values)

Finally, we have obtained the experimentally supported miRNA:target dataset.

```
experimentalmirnagene <- readRDS("data/experimentalmirnagene.RDS")
head(experimentalmirnagene)</pre>
```

```
## # A tibble: 6 x 18
##
     cluster chromosome start_position end_position strand hgnc_symbol
                                               <int> <chr>
##
     <chr>
            <chr>
                                  <int>
                                                            <chr>>
## 1 0727A-~ chr5
                             162864575
                                           162873157 1
                                                            CCNG1
## 2 L1HS-1~ chr14
                              95552565
                                            95624347 -1
                                                            DICER1
## 3 L2HS-8~ chr6
                             109307640
                                           109416022 -1
                                                            SESN1
## 4 L2HS-1~ chr5
                              36876861
                                            37066515 1
                                                            NIPBL
## 5 L2-407~ chr4
                             106603784
                                           106817143 -1
                                                            INTS12
## 6 L1HS-7~ chr5
                             130977407
                                           131132710 -1
                                                            FNIP1
## # ... with 12 more variables: Ensembl_Gene_Id <chr>,
       Ensembl_Transcript_Id <chr>, target_seq <chr>, miRNA <chr>,
       miR_seq <chr>, seed_type <chr>, Energy <dbl>, HG38build_loc <chr>,
       Genom_build <chr>, region <chr>, region_effect <dbl>,
## #
       seed_type_effect <dbl>
```

The methods about miRNA:target interactions are based a basic principle that is reading after isolation of miRNA:target chimerics. The datasets contain all the chimeric miRNA:target structures found in the medium during the experiment. On the other hand, it could be said that the reading is performed as snapshot.Because of that, the methods can provide different chimeric interactions the same miRNA:target pair. We have preferred to select most effective interaction parameters for the same miRNA:target pairs that can exhibit various interactions. The step is performed as:

```
tocombine_mirnagene <- experimentalmirnagene %>% dplyr::select(miRNA,
    Ensembl_Gene_Id, hgnc_symbol, Energy, seed_type_effect,
    region_effect) %>% distinct() %>% group_by(Ensembl_Gene_Id,
    miRNA) %>% mutate(seed_type_effect = ifelse(seed_type_effect ==
    max(seed_type_effect), seed_type_effect, max(seed_type_effect)),
    Energy = ifelse(Energy == min(Energy), Energy,
        min(Energy)), region_effect = ifelse(region_effect ==
        max(region_effect), region_effect, max(region_effect))) %>%
    distinct()

head(tocombine_mirnagene)
```

2.4 Combine the dataset

```
E9GE_mirnagenenormal <- BCPME_mirnanormal %>% dplyr::inner_join(tocombine_mirnagene,
   by = c(miRNA_name = "miRNA")) %>% dplyr::inner_join(TCGA_E9_A1N5_normal,
   by = c(Ensembl_Gene_Id = "ensembl_gene_id", hgnc_symbol = "external_gene_name")) %>%
   distinct() %>% dplyr::select(hgnc_symbol, miRNA_name,
   mirna_RPM = reads_per_million_miRNA_mapped, GE_normal,
   Energy, seed_type_effect, region_effect)
# saveRDS(E9GE_mirnagenenormal,
# 'data/E9GE mirnagenenormal.rda')
E9GE mirnagenenormal <- readRDS("data/E9GE mirnagenenormal.rda")
head(E9GE_mirnagenenormal)
## # A tibble: 6 x 7
    hgnc_symbol miRNA_name mirna_RPM GE_normal Energy seed_type_effect
##
                             <dbl> <dbl> <dbl>
## 1 CCNG1
                hsa-let-7~ 111204.
                                        5245 -25.1
                                                                 0.05
                hsa-let-7~ 111204.
## 2 DICER1
                                         3285 -24.4
                                                                 0.43
## 3 SESN1
                hsa-let-7~ 111204.
                                        1179 -22.2
                                                                 0.05
## 4 NIPBL
                hsa-let-7~ 111204.
                                         4503 -22.1
                                                                 0.05
## 5 INTS12
                hsa-let-7~
                                          600 -21.9
                            111204.
                                                                 0.05
## 6 FNIP1
                hsa-let-7~
                            111204.
                                         1248 -21.8
                                                                 0.43
## # ... with 1 more variable: region_effect <dbl>
E9GE_mirnagenetumor <- BCPME_mirnatumor %>% dplyr::inner_join(tocombine_mirnagene,
   by = c(miRNA name = "miRNA")) %>% dplyr::inner join(TCGA E9 A1N5 tumor,
   by = c(Ensembl_Gene_Id = "ensembl_gene_id", hgnc_symbol = "external_gene_name")) %>%
   distinct() %>% dplyr::select(hgnc_symbol, miRNA_name,
   mirna_RPM = reads_per_million_miRNA_mapped, GE_tumor,
   Energy, seed_type_effect, region_effect)
# saveRDS(E9GE_mirnagenetumor,
# 'data/E9GE_mirnagenetumor.rda')
E9GE_mirnagenetumor <- readRDS("data/E9GE_mirnagenetumor.rda")</pre>
head(E9GE_mirnagenetumor)
## # A tibble: 6 x 7
    hgnc_symbol miRNA_name mirna_RPM GE_tumor Energy seed_type_effect
##
    <chr>
                <chr>
                       <dbl>
                                       <dbl> <dbl>
                                                               <dbl>
## 1 CCNG1
                hsa-let-7~
                             62406.
                                        2467 -25.1
                                                                0.05
## 2 DICER1
                hsa-let-7~ 62406.
                                        5023 -24.4
                                                                0.43
                                         829 -22.2
## 3 SESN1
                hsa-let-7~ 62406.
                                                               0.05
                hsa-let-7~
## 4 NIPBL
                             62406.
                                        5126 -22.1
                                                               0.05
                                                               0.05
## 5 INTS12
                hsa-let-7~ 62406.
                                        1009 -21.9
## 6 FNIP1
              hsa-let-7~
                             62406.
                                        2144 -21.8
                                                               0.43
## # ... with 1 more variable: region_effect <dbl>
```

2.5 Selection of trigger node

We have determined the most important nodes of network. We applied find_node_perturbation function. We only defined nodes that affect the other nodes more than 1.05 fold change with 10 iteration when they increase 3 fold.

This dataset, perturbation of network, includes totally 423 effective nodes. We selected SERPINE2 gene to perturb the network because it is the most efficient node gene in network.

2.5 Approach of Method into Combined Datasets

We selected SERPINE2 gene for simulation of regulation on network.

2.5.1 Find iteration of simulation

The node amount of changed gene on the system in terms of percentage were shown at Figure S9 in Supplementary Figures. As seen, firstly, the changed gene count increase. The system which contains the hundreds of miRNAs and thousands of genes can slowly gain the steady-state again. At first glance, it can be assumed that when all nodes in the system are reached, stable state will be provided. However, although all nodes are reached, the nodes competing with each other cause the edits to continue for a while.

The dynamics of the approach are shown in package vignettes.

A directed acyclic simple graph with 8 components

So, we offered an approach about to find iteration. find_iteration function does not give the iteration to gain steady-state, but it gives the iteration which has maximum affected node counts. The function is applied as following:

```
as.data.frame(E9GE_mirnagenenormal) %>% priming_graph(competing_count = GE_normal,
    miRNA_count = mirna_RPM, aff_factor = c(Energy,
        seed_type_effect), deg_factor = region_effect) %>%
    update_how("SERPINE2", 2.75) %>% simulate(50) %>%
    find_iteration(limit = 1, plot = FALSE)
```

```
## Warning in priming_graph(., competing_count = GE_normal, miRNA_count = mirna_RPM, : First column is ;
## [1] 31
```

2.5.2 Simulation of dataset

We tried to apply two fold of the point that SERPINE2 has maximum affected genes on network.

```
## Warning in priming_graph(., competing_count = GE_normal, miRNA_count = mirna_RPM, : First column is ]
## # A tbl_graph: 8217 nodes and 25614 edges
## #
```

```
## #
## # Node Data: 8,217 x 7 (active)
    name type node_id initial_count count_pre count_current
     <chr> <chr>
##
                  <int>
                              <dbl>
                                          <dbl>
                                                     <dbl>
## 1 CCNG1 Comp~
                      1
                                 5245
                                          5249.
                                                         5250.
## 2 DICE~ Comp~
                      2
                                 3285
                                          3285.
                                                        3290.
## 3 SESN1 Comp~
                      3
                                 1179
                                         1179.
                                                        1179.
## 4 NIPBL Comp~
                      4
                                 4503
                                          4503.
                                                        4504.
## 5 INTS~ Comp~
                      5
                                  600
                                           600.
                                                         600.
                      6
                                 1248
                                                        1252.
## 6 FNIP1 Comp~
                                          1247.
## # ... with 8,211 more rows, and 1 more variable: changes_variable <chr>
## #
## # Edge Data: 25,614 x 23
             to Competing_name miRNA_name GE_normal mirna_RPM Energy
##
     <int> <int> <chr>
                               <chr>
                                             <dbl>
                                                        <dbl>
        1 7873 CCNG1
## 1
                               hsa-let-7~
                                               5245
                                                       111204.
                                                               -25.1
## 2
        2 7873 DICER1
                               hsa-let-7~
                                                3285
                                                       111204. -24.4
## 3
        3 7873 SESN1
                               hsa-let-7~
                                                1179
                                                      111204. -22.2
## # ... with 2.561e+04 more rows, and 16 more variables:
      seed_type_effect <dbl>, region_effect <dbl>, dummy <dbl>,
## #
      afff_factor <dbl>, degg_factor <dbl>, comp_count_list <list>,
      comp_count_pre <dbl>, comp_count_current <dbl>,
      mirna_count_list <list>, mirna_count_pre <dbl>,
## #
      mirna_count_current <dbl>, mirna_count_per_dep <dbl>,
      effect_current <dbl>, effect_pre <dbl>, effect_list <list>,
## #
## #
      mirna_count_per_comp <dbl>
simulation_results <- as.data.frame(E9GE_mirnagenenormal) %>%
   priming_graph(competing_count = GE_normal, miRNA_count = mirna_RPM,
        aff_factor = c(Energy, seed_type_effect), deg_factor = region_effect) %>%
   update_how("SERPINE2", 2.75) %>% simulate(62) %>%
    as_tibble() %>% dplyr::select(name, initial_count,
    count_current)
```

Warning in priming_graph(., competing_count = GE_normal, miRNA_count = mirna_RPM, : First column is

2.5.3 Comparison of simulation results and tumor tissue expression values

4503

4 NIPBL

5126

```
E9GE_mirnagenetumor %>% dplyr::select(hgnc_symbol,
    GE_tumor) %>% dplyr::inner_join((E9GE_mirnagenenormal %>%
    dplyr::select(hgnc_symbol, GE_normal)), by = "hgnc_symbol") %>%
    inner_join(simulation_results, by = c(hgnc_symbol = "name")) %>%
    distinct()
## # A tibble: 7,813 x 5
##
     hgnc_symbol GE_tumor GE_normal initial_count count_current
##
      <chr>
                    <dbl>
                              <dbl>
                                             <dbl>
## 1 CCNG1
                                5245
                     2467
                                              5245
                                                           5250.
## 2 DICER1
                     5023
                                3285
                                              3285
                                                           3290.
## 3 SESN1
                     829
                                1179
                                              1179
                                                           1179.
```

4503

4504.

##	5	INTS12	1009	600	600	600.
##	6	FNIP1	2144	1248	1248	1252.
##	7	ACAD8	860	1249	1249	1249.
##	8	CCNB2	749	690	690	690.
##	9	ZNF260	1808	1067	1067	1067.
##	10	SYVN1	2565	2300	2300	2303.
##	#	with	7,803 more rows			

Actually, we have developed to provide a new approach mirna mediated regulation networks. This approach may not explain the whole regulation behaviors between miRNAs and targets but can be first step to more detailed and coherent miRNA:target regulation approach.