NETISCE Manual and Tutorials

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About

Welcome to the NETISCE manual and tutorials.

NETISCE is a workflow that identified control node perturbations for cellular reprogramming. This manual contains directions for intial setup, as well as tutorials for reproducing NETISCE cell reprogramming results in developmental, stem cell, and cancer biology.

Installation and Usage

2.1 Download NETISCE

NETISCE pipelines can be downloaded from our github repository: https://github.com/veraliconaresearchgroup/netisce

We recommend that you run NETISCE on a high-performance cluster (hpc), as you may generate files that are quite large, or run computations that may take a long time. However, we provide two Nextflow pipelines, one designed for hpcs (NETISCE_hpc), and another for running NETISCE on a local machine (NETISCE_local).

2.2 Install Nextflow

Nextflow is required to run the NETISCE pipeline. Please follow the instructions from https://www.nextflow.io/ (see 'Getting Started' steps 1 & 2) to install Nextflow in the appropriate NETSICE folder (_local or _hpc).

2.3 Docker Image

docker image provided here TBD

2.4 Prerequisuites

If you are not using the Docker image, the following packages will need to be installed:

- scipy
- pandas
- sklearn
- vellowbrick

2.5 Parameters and Configuration

Whether on your local machine or hpc, to run NETISCE you must specify the files and parameters within the .nf file

- params.expressions: csv file containing normalized expression data for network nodes in different samples
- params.network: network file (sif format)
- params.samples: text file specifying the phenotype for each sample in params.expressions file (tab delimited)
- params.internal_control: text file containing a list of nodes to be used as internal marker nodes
- params.alpha: alpha parameter for signal flow analysis (default =0.9)
- params.undesired: string of the undesired phenotype (as labeled in the params.samples file)
- params.desired: string of the desired phenotype (as labeled in the params.samples file)
- params.filter: filtering parameter for criterion 2 ("strict" or "relaxed")
- params.kmeans_min_val: minimum k-means value for clustering (default=2)
- params.kmeans_max_val: maximum k-means value for clustering (default=10)
- params.num_nodes: number of nodes in network for which normalized expression data exists (within the params.expressions file)
- params.num_states: number of randomly generated initial states (default=100000, or 3^n where n is the number of network nodes and 3^n is less than 100000)

Please see the **input_data** folder for examples of files to match the formatting.

2.5.1 NETISCE mutations.nf

If you are interested in including mutational information, please use the NETISCE_mutations.nf pipeline. You must additionally specify params.mutations: a csv file containing mutational configuration for network nodes (0 for loss of function, 1 for gain of function). Please see example in input data for formatting.

2.5.2 nextflow.config

If you are running nextflow on an hpc, please specify your executor, and clusterOptions within the nextflow.config file. Please see https://www.nextflow.io/docs/latest/config.html for more information regarding your executor.

2.6 Running NETISCE

Once you have specified the parameters, run NETSICE using the following command:

./nextflow run NETISCE.nf -resume ##or NETISCE_mutations.nf if including mutational data

We recommend using the -resume flag in the case that you change a file or parameter within your pipeline. This way, nextflow caches results that remain unchanged, preventing pipeline steps from being re-run.

NETSICE output

After the NETISCE computations are complete, the output files will be located in the results folder. Please note that we have included in this folder the most relevant output files that you may want to use for further analysis. However, you can explore all outputs by checking within each step of the pipeline's work folder.

The contents of each file are briefly described below. For more details and to see example outputs, please see the Toy Network Examples.

exp_internalmarkers.txt

This file contains the resultant steady state values for the internal marker nodes for the provided experimental samples (those specified in samples.txt) from Signal Flow Analysis.

$experimental_internal markers.pdf$

This pdf is a figure of the steady state values for the internal marker nodes for the provided experimental samples. This can be used to verify the validity of the internal marker nodes.

elbow.png

A graph of the elbow metric for determining the optimal k for k-means.

silhouette.pdf

A graph of the silhouette metric for determining the optimal k for k-means.

fvs.txt

This file contains the node names for the FVS used as control nodes for the NETISCE run.

crit1perts.txt

This file contains a list of IDs for the control node perturbations that passed criterion 1.

$pert1_internal_markers.txt$

This file contains a table of the internal marker node steady state values from control node perturbations whose associated attractors passed the first filtering criterion.

$successful_controlnode_perturbations.txt$

This file contains a table of the control node perturbations that pass both the 1st and 2nd filtering criteria. it also contains the number of upregulation,downregulations, and total number of nodes perturbed for each perturbation set.

Toy Network Examples

Here, we will walk through a brief tutorial of a NETISCE run. The files necessary to complete the tutorial are within the input data folder of both NETISCE_local and NETISCE_hpc. The results from these Toy examples can be found in the toy_example_results folder of the main github repository.

4.1 Overview

We will use a simple toy network of 6 nodes and 9 edges.



Figure 4.1: Simple Toy Network

4.2 Data

You can find the relevant data files in the input_data folder.

In this example, we have 2 samples, A and B, with three replicates each (A_1,A_2,A_3, etc). The normalized expression data is housed in expressions.csv, and contains normalized expression values for 4 of the network nodes (in this case, node C was not found to be expressed in the samples).

X	A_1	A_2	A_3	B_1	B_2	B_3
A	2	1	2	6	7	6
В	6	7	6	2	1	1
\overline{D}	-1	-1	-2	6	7	6
\overline{E}	2	1	2	6	7	6

The samples.txt file specifies that A is associated to a treatment sensitive phenotype, while B is associated to a resistance phenotype.

name	phenotype
A_1	sensitive
	sensitive
A_3	sensitive
B_1	resistant
B_2	resistant
B_3	resistant

Note that you can use any term to describe the phenotypes. Just be sure to be consistent with the param.desried and param.undesired variables within the Nextflow .nf file.

Lastly, we need to include a list of internal marker nodes. This list is in internal_marker.txt. For our small network, the internal marker node is C.

С

4.3 NETISCE run configuration

With all your input data files loaded, next we configure the nextflow run. Within either NETISCE_local or NETSICE_hpc (Note: while we do recommend you run NETISCE on a hpc, this example is small enough to run locally).

Open up NETISCE.nf. Here, you need to specify the parameters for the Nextflow run on lines 3-19. Please refer to section 2.5 for parameter definitions.

For this example, your parameters should look like:

```
params.expressions = "$baseDir/input_data/expressions.csv"
params.network = "$baseDir/input_data/network.sif"
params.samples = "$baseDir/input_data/samples.txt"
params.internal_control="$baseDir/input_data/internal_marker.txt"
params.alpha = 0.9
params.undesired = 'resistant'
params.desired = 'sensitive'
params.filter ="strict"

params.kmeans_min_val = 2
params.kmeans_max_val = 10

params.num_nodes = 4 // that have expression data
params.num_states = 1000
```

Some Notes: make sure to include \$baseDir before pointing to the folder containing your input data. Also, be sure that params.num_nodes is the number of nodes where there exists normalized expression data within expression.csv. Finally, in NETISCE.nf, mutations are not considered, so that like is commented out.

4.4 Run NETSICE

In your terminal/command prompt, navigate to the appropriate NETISCE folder _hpc or local. To start your run, enter ./nextflow run NETISCE.nf -resume. While NETISCE is running, your terminal should look like this, where you can see the progress on each step of the pipeline:

Once the run has successfully completed, the process will end and the following will be displayed:

4.5 NETSICE Results

Let's take a look at the results of our NETISCE run, where the goal was to shift the system from the undesired state B, and towards the desired state A. These results can be found in the toy_example_1 subfolder of the toy_example_results folder of the main github repository.

Figure 4.2: Terminal when running NETISCE

```
Completed at: 02-Dec-2021 10:19:50
Duration : 4m 29s
CPU hours : 0.1
Succeeded : 16
```

Figure 4.3: Terminal when running NETISCE

$exp_internalmarkers.txt$

Our internal marker node was node C. In this file we see the steady state values of the node in the A and B sample replicates (the output values from SFA).

name	C
1	0.4278056
	0.4802943
A_3	0.4361991
B_1	0.1590962
B_2	0.0982107
B_3	0.0935476

experimental_internalmarkers.pdf

The above numbers may be a little challenging to read! So, we have included a plot of the values in the experimental_internalmarkers.pdf:

On this histogram, we see bars for each of the samples and their replicates. The A (sensitive) samples are marked by a blue vertical line at their steady state value, while the B (resistant) samples are marked by a red vertical line at their steady state value. Here, we see that the values of node C are well separated between the two phenotypes (all of the A values are greater than all of the B



Figure 4.4: experimental marker node steady state values

values). We will assume that this also aligns with the biological knowledge of the system.

In this example, since there are only 4 network nodes that have normalized expression values, NETISCE generates the maximum number of random initial states, 3^4 , or 81.

After estimating attractors for the experimental and randomly generated initial states, the resultant attractors were clustered using k-means clustering. The elbow and silhouette metrics are used to determine the optimal number k.

elbow.png

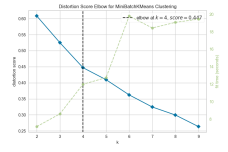


Figure 4.5: elbow metric for optimal k

The elbow metric found the optimal number of k clusters to be k=4.

silhouette.pdf

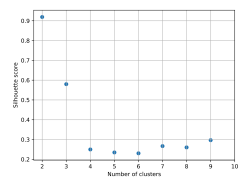


Figure 4.6: silhouette metric for optimal k

The silhouette metric found the optimal number of k clusters to be k=2.

Since the optimal ks identified by the silhouette metric and the elbow metric do not match, NETISCE chooses the smaller k, as long as the phenotypes remain separate (NETISCE checks to make sure this is true).

fvs.txt

This file contains the node names that were identified by the FVS finding algorithm.

name
D
В

The FVS finding algorithm identified nodes B and D to be the minimal FVS control nodes in the toy network. Since the FVS control node set contained 2 nodes, 9 combinations of perturbations were performed on the control node sets.

crit1perts.txt

This file contains a list of IDs for the perturbations to FVS control nodes that passed criterion 1.

V1	
$pert_{_}$	_0
$pert_{_}$	_1
pert_{-}	_2
pert_	_4
pert_{-}	_5
pert_{-}	_6
pert_{-}	_7
pert_{-}	_8

8 out of the 9 pertrubations passed the machine learning filtering criterion.

pert1_internal_markers.txt

This file contains a table of the internal marker node state values from control node perturbations whose associated attractors passed the first filtering criterion.

name	С
pert_0	-2.7000000
pert_1	1.0209997
pert_2	6.3000000
pert_4	-0.0427445
pert_5	6.3000000
pert_6	-2.7000000
pert_7	-2.8528056
pert_8	6.3000000

$successful_controlnode_perturbations.txt$

This file contains a table of the perturbations on FVS control nodes that passed both the 1st and 2nd filtering criteria. it also contains the number of upregulation, downregulations, and total number of nodes perturbed for each perturbation set.

	D	В	up	down	total
pert_1	down	nochange	0	1	1
pert_5	nochange	up	1	0	1
pert_2	down	up	1	1	2
pert_8	up	up	2	0	2

Here, we see that four perturbations that passed both filtering criteria.

Let's take a quick look at the steady state values for these perturbations, and the attractors generated from the experimental data:

	name	C
1	A_1	0.4278056
2	A_2	0.4802943
3	A_3	0.4361991
4	B_1	0.1590962
5	B_2	0.0982107
6	B_3	0.0935476
21	pert_1	1.0209997
31	pert_2	6.3000000
51	pert_5	6.3000000
8	pert_8	6.3000000

Indeed, we see that the steady-state expression values of node C in the attractors generated by peturbations to the FVS control nodes are all are greater than the steady-state expression values of node C in the attractors generated from the sensitive A sample. A successful reprogramming from resistant (B) to sensitive (A) cells has occurred!

4.6 Toy Example with mutations

Let's say that in our system, gene A exhibits a loss of function mutation in the sensitive phenotype (A samples). If we want to include this in our simulations, we will use the NETISCE_mutations.nf pipeline.

First, we must add to our input_data folder a .csv file containing the mutational profile. Let's call this file mutations.csv:

X	A
A_1	0
	0
3	0
B_1	NA
B_2	NA
B_3	NA

The loss of function mutation is encoded with 0 (gain-of-function mutations can be encoded with "1").

Next, we make sure that the parameters in NETISCE_mutations.nf on lines 3-19 are set correctly for the conditions

For this example, your parameters should look like:

```
params.expressions = "$baseDir/input_data/expressions.csv"
params.network = "$baseDir/input_data/network.sif"
params.samples = "$baseDir/input_data/samples.txt"
```

```
params.internal_control="$baseDir/input_data/internal_marker.txt"
params.mutations="$baseDir/input_data/mutations.csv"
params.alpha = 0.9
params.undesired = 'resistant'
params.desired = 'sensitive'
params.filter ="strict"

params.kmeans_min_val = 2
params.kmeans_max_val = 10

params.num_nodes = 4 // that have expression data
params.num_states = 1000
```

Note, the additional parameter params.mutations that points to the mutations.csv.

As above, to run Netisce, enter ./nextflow run NETISCE.nf -resume.

Results

By including mutational information, the results of NETISCE have changed. These results can be found in the toy_example_2 subfolder of the toy_example_results folder of the main github repository. Now, our successful_controlnode_perturbations.txt file contains pert_0 instead of pert_8

	В	D	up	down	total
pert_1	down	nochange	0	1	1
pert_5	nochange	up	1	0	1
pert_0	down	down	0	2	2
pert_2	down	up	1	1	2

Let's take a look at the steady-state expression values of node C in the attractors generated from the successful perturbations and the experimental initial states when mutational information is included.

name	C		
A_1	-0.6621166		
B_1	0.1590962		
pert_0	-2.7000000		
pert_1	-2.7000000		
pert_2	-2.7000000		
pert_5	-2.8528056		

Though the values are different in this system with mutations, we still see that the steady-state expression values of node C in the attractors generated by peturbations to the FVS control nodes are all are greater than the steady-state expression values of node C in the attractors generated from the sensitive A sample. A successful reprogramming from resistant (B) to sensitive (A) cells has occurred!

Cell Fate Specification in Ascidian Embryo

This section contains instructions to reproduce the results of simulating FVS control node perturbations in a model of ascidian embryo cell specification. You can read the original report here: link

The input data, nextflow pipeline, and results of this simulation can be found in the ascidian embryo folder in the github repository

These simulations were run on a high performance cluster that uses a SLURM executor. Although we recommend that you run NETISCE on an hpc, this simulation is small enough that it can be run on a local machine. If you choose to run it locally, then remove the nextflow.config file from the directory.

5.1 Input Data

The goal of this simulation was to reproduce the results of experimental perturbations to the FVS nodes of the cell fate specification GRN for ascidian embryos using Signal Flow Analysis. Therefore, we use a modified version of the NETISCE pipeline to simulate these specific perturbations. We are only interested in performing the 7 perturbations to the 6 FVS control nodes that were experimentally verified to induce cell tissue fates.

expression.csv contains the initial activities for the unperturbed state and the 7 FVS control node perturbations. Here, all simulations have Gata.a and Zic-r.a=1, as the activation of these two genes is required for normal embryonic development.

perturbations.csv contains the specified perturbations for each FVS node in the appropriate perturbation simulation. 0 denotes downregulation, wherease 1 encodes upregulation. If no value is set, then there is no fixed perturbation to the FVS node, as in the unperturbed case.

internal-marker-nodes.txt contains the 7 internal marker nodes used to verify if the specified cell reprogramming had been successfully simulated.

5.2 Run the simulation

To run the simulation, simply execute the ascidian-embryo.nf file using the following command: ./nextflow run ascidian-embryo.nf -resume

5.3 Results

The nextflow pipeline generates 1 result file exp_internalmarkers.txt, which contains the steady state values of the internal-marker nodes for the unperturbed attractor, and the attractors generated from the perturbations on FVS control nodes.

knitr::kable(read.delim('ascidian_embryo/results/exp_internalmarkers.txt',sep="\t"))

name	Alp	Bco	Celf3.a	Epi1	Fli.l
unperturbed	-0.0000341	0.0018729	0.0018729	0.0217995	0.001
Adentz (Endoderm perturbation)	0.1179951	-0.4252100	-0.4252100	0.2707276	-0.183
adentZ (brain+pan-neural perturbation)	-0.3845691	0.4252100	0.4252100	0.2707276	-0.237
adeNtz (pan-neural perturbation)	-0.5243074	-0.4252100	-0.4252100	0.2707276	-0.300
adEntZ (mesenchyme perturbation)	-0.2630083	0.4252100	0.4252100	-0.1563935	0.284
adentz (epidermis perturbation)	-0.5243074	-0.4252100	-0.4252100	0.2707276	-0.300
adenTz (muscle perturbation)	-0.5243074	-0.4252100	-0.4252100	0.2707276	-0.300
aDentz (notochord perturbation)	-0.3792942	-0.4252100	-0.4252100	0.1988199	-0.396

5.4 Visualizing Results

We can use radar plots to visualize the results of the SFA simulations of perturbations to the FVS control nodes. This can help us identify which perturbations successfully induced the appropriate tissue fate. In this context, a perturbation is considered successful if the internal-marker node in the attractor generated from the perturbed FVS control nodes has a larger steady-state value than that in the unperturbed attractor.

The following code for generating radar plots was adapted from datanovia.com is used to generate the radar charts. **Note:** you may need to adjust the

formatting of exp_internalmarkers.txt so that the strings within quotations are placed into one column.

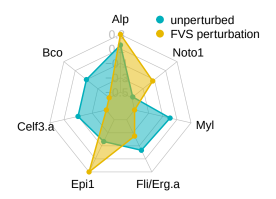
```
create_beautiful_radarchart <- function(data, color = "#00AFBB",</pre>
                                        vlabels = colnames(data), vlcex = 1,
                                         caxislabels = NULL, title =row.names(data)[4], ...){
  radarchart(
   data, axistype = 1,
    # Customize the polygon
   pcol = color, pfcol = scales::alpha(color, 0.5), plwd = 2, plty = 1,
    # Customize the grid
   cglcol = "grey", cglty = 1, cglwd = 0.8,
    # Customize the axis
   axislabcol = "grey",
    # Variable labels
   vlcex = vlcex, vlabels = vlabels,
   title = title,
   centerzero = F,
   caxislabels = caxislabels
  )
}
```

```
library(fmsb)
d1<-read.delim("ascidian_embryo/results/exp_internalmarkers.txt",sep="\t",row.names = 1,check.name
maxcol<-apply(d1, 2, max)
mincol<-apply(d1, 2, min)

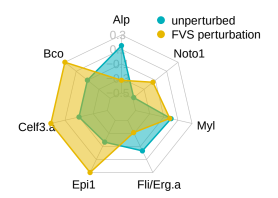
d2<-rbind(maxcol,mincol, d1)
rownames(d2)[1:2]<- c("Max", "Min")

# Define settings for plotting in a 3x4 grid, with appropriate margins:
for (i in 4:nrow(d2)) {
   create_beautiful_radarchart(d2[c(1:3, i), ],color = c("#00AFBB", "#E7B800"),caxislabels = seq(not beautiful_radarchart(d2]c(1:3, i), ],color = c("#00AFBB", "#E7B800"), caxislabels = seq(not beautiful_radarchart(d2]c(1:3, i), ],color = c("#00AFBB", "#E7B800"), caxislabels = seq(not beautiful_radarchart(d2]c(1:3, i), ],color = c("#00AFBB", "#E7B800"), caxislabels = seq(not beautiful_radarchart(d2]c(1:3, i), ],color = c("#00AFBB", "#E7B800"), caxislabels = seq(not beautiful_radarchart(d2]c(1:3, i), ],color = c("#00AFBB", "#E7B800"), caxislabels = seq(not beautiful_radarchart(d2]c(1:3, i), ],color = c("#00AFBB", "#E7B800"), caxislabels = seq(not beautiful_radarchart(d2]c(1:3, i), ],color = c("#00AFBB", "#E7B800"), caxislabels = seq(not beautiful_radarchart(d2]c(1:3, i), ],color = c("#00AFBB", "#E7B800"), caxislabels = seq(not beautiful_radarchart(d2]c(1:3, i), ],color = c("#00AFBB", i), bty = "n", pch=20, col=c("#00AFBB", i), bty = "n", pch
```

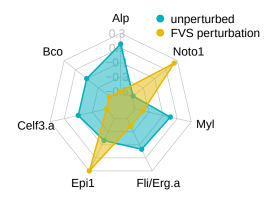
Adentz (Endoderm perturbation)



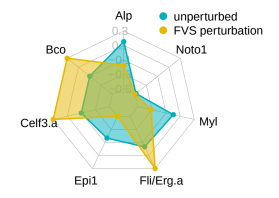
adentZ (brain+pan-neural perturbation)



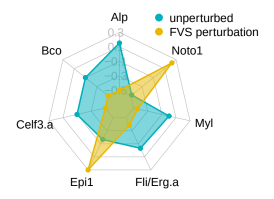
adeNtz (pan-neural perturbation)



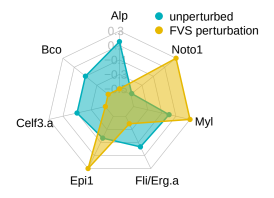
adEntZ (mesenchyme perturbation)



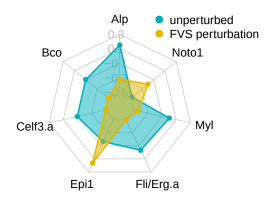
adentz (epidermis perturbation)



adenTz (muscle perturbation)



aDentz (notochord perturbation)



Pluripotent Stem Cell Example

Adaptive Resistance in Colorectal Cancer Example

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Footnotes and citations

8.1 Footnotes

Footnotes are put inside the square brackets after a caret ^[]. Like this one ¹.

8.2 Citations

Reference items in your bibliography file(s) using @key.

For example, we are using the **bookdown** package [Xie, 2021] (check out the last code chunk in index.Rmd to see how this citation key was added) in this sample book, which was built on top of R Markdown and **knitr** [Xie, 2015] (this citation was added manually in an external file book.bib). Note that the .bib files need to be listed in the index.Rmd with the YAML bibliography key.

The RStudio Visual Markdown Editor can also make it easier to insert citations: https://rstudio.github.io/visual-markdown-editing/#/citations

¹This is a footnote.

Blocks

9.1 Equations

Here is an equation.

$$f\left(k\right) = \binom{n}{k} p^{k} \left(1 - p\right)^{n - k} \tag{9.1}$$

You may refer to using \@ref(eq:binom), like see Equation (9.1).

9.2 Theorems and proofs

Labeled theorems can be referenced in text using \@ref(thm:tri), for example, check out this smart theorem 9.1.

Theorem 9.1. For a right triangle, if c denotes the length of the hypotenuse and a and b denote the lengths of the **other** two sides, we have

$$a^2 + b^2 = c^2$$

 $Read\ more\ here\ https://bookdown.org/yihui/bookdown/markdown-extensions-by-bookdown.html.$

9.3 Callout blocks

The R Markdown Cookbook provides more help on how to use custom blocks to design your own callouts: https://bookdown.org/yihui/rmarkdown-cookbook/custom-blocks.html

Sharing your book

10.1 Publishing

HTML books can be published online, see: https://bookdown.org/yihui/bookdown/publishing.html

10.2 404 pages

By default, users will be directed to a 404 page if they try to access a webpage that cannot be found. If you'd like to customize your 404 page instead of using the default, you may add either a _404.Rmd or _404.md file to your project root and use code and/or Markdown syntax.

10.3 Metadata for sharing

Bookdown HTML books will provide HTML metadata for social sharing on platforms like Twitter, Facebook, and LinkedIn, using information you provide in the index.Rmd YAML. To setup, set the url for your book and the path to your cover-image file. Your book's title and description are also used.

This gitbook uses the same social sharing data across all chapters in your bookall links shared will look the same.

Specify your book's source repository on GitHub using the edit key under the configuration options in the _output.yml file, which allows users to suggest an edit by linking to a chapter's source file.

Read more about the features of this output format here:

https://pkgs.rstudio.com/bookdown/reference/gitbook.html

Or use:

?bookdown::gitbook

Bibliography

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