

QNIDQPTEMTATEGAIVQINCTYQTSGFNGLFWYQQHAGEAPTFLSYNVLDGLEEKGRFSSFL
SKGYSYLLKELQMKDSASYLCAVMDSNYQLIWGAGTKLIIPDIQNPDPAVYQLRDSKSSDKS
LFTDFDSQTNVSQSKDSDVYITDKCVLDMRSMDFKNSAVAWSNKSDFACANAFNNSIIPEDT
PSPIAGITQAPTSQILAAGRRTLRCTQDMRHNAMEWYRQDLGLGLRLIHYSNTAGTTGKGEV
GYSVSRANTDDFPLTLASAVPSQTSVYFCASSEAGGNTGELFFGEGSRLTVLEDLKNVFPPEV
FEPSEAEISHTQKATLVCLATGFYPDHVELWWVNGKEVHSGVCTDPQPLKEQPALNDSRYAL
RLRVSATFWQNPRNHFRCQVQFYGLSENDEWTQDRAKPVTQIVSAEAWGRAMRTHSLRYFR
VSDPIHGVPFISVGVDHSHPITTYDSVTRQKEPRAPWMAENLAPDHWEERYTQLLRGWQQMF
ELKRLQRHYNHSGSHTYQRMIGCELLEDGSTTGFLQYAYDGQDFLIFNKDTLSWLAVDNVAHT
AWEANQHELLYQKNWLEEECIAWLKRFLEYGKDQLQRTEPPLVRVNRKETFPGBTALFCKAHG
PPEIYMTWMKNGEEIVQEIDYGDILPSGDGTYQAWASIELDPQSSNLYSCHVEHSGVHMVLQV
QLVMHVGSHDEVHCSYLNSSQPDLEISAQAQYTGDGSPLIDGYDIEQVIEEGNKMWTMYIEPPY
HAKCFLATVGPFTEKRNVRVLPPETRQLTDKGYELFRKLWAICEEELWNKQYLLEHQNAEWAC
HAVNDVALWSLTDKNFILFDQGDYAYQLFGTTSGDELLECGIMRQYTHSGSHNYHRQLRKLEV
MQQWGRLLQTYREWHDPALNEAMWPARPEKQRTVSDYTTIPHSDVYGVSI FEPVGHIPDSVG
FYRLSHTRMARGWAEASVIQTVPKARDQTWEDNESLGYFQVQCRFHNRPNQWFTASVRLRS
AYRSDNLAPQEKLQPQDTCVGSHVEKGNVVWSLEVHDPYFGTALCVLTAKQTHSIEAESPEFV

TRain: T-cell Receptor Automated Immunoinformatics – User's Guide

Austin Seamann & Dario Ghersi



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<https://github.com/Aseamann/TRain>

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Overview of TRain

Input
Modeling
Pairing
Docking
Analysis

1 — Introduction

1.1 Overview of TRain

The TRain suite provides the accumulation and connection of several in-house and well cited bioinformatic tools to gain further utilization of T-cell receptor sequences and structures. TRain consists of five main components: input, modeling, pairing, docking, and analysis (see Figure 1.1). These components are interchangeable with different modeling and docking tools as the connection between each step is through common file types such as FASTA and PDB files.

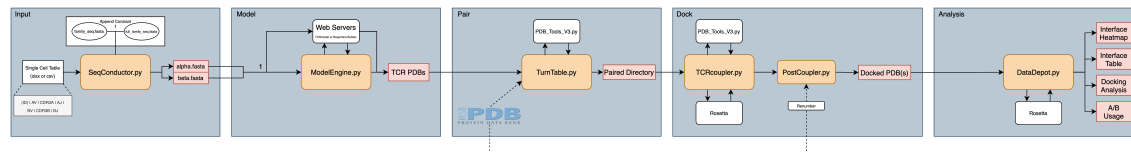


Figure 1.1: **Flowchart for the TRain suite.** General flow of TRain input to analysis. TRain takes input of single cell T-cell receptor sequencing data to produce full TCR sequences. Full TCR sequences are then submitted to modeling webserver of choice. Resulting PDB file containing TCR 3D structure then gets paired with a MHC and bound peptide to be docked. Docking is conducted with a prepared pipeline of Rosetta programs. Resulting TCR/pMHC complexes then can be further investigated with analytical tools.

1.2 Input

TRain is capable of handling single cell T-cell receptor sequencing output from sources such as immunoSeq. This introduces the first Python program SeqConductor. SeqConductor reads in either a xlsx or csv file containing TCR AV, AJ, BV, and BJ gene segments and CDR3 α and CDR3 β amino acid sequences. This constructs the amino acid sequences for the variable regions of each TCR.

The resulting variable region sequences can then be concatenated with a constant TCR region to produce a full TCR sequence. The primary goal of TRain is to gain further understanding of

TCR/pMHC interfaces. Thus, the constant region is not an extremely valuable portion but is needed for modeling purposes to avoid alignment errors. The full TCR sequences are then prepared to be passed to the following modeling steps.

Resulting fasta files are produced and can either be submitted directly to the suggested modeling suites or automated in the Modeling step.

1.3 Modeling

Currently, four modeling pipelines are readily available through web servers to produce 3D structures of TCRs from submitted alpha and beta chain sequences. These four services being LYRA, TCRmodel, Repertoire Builder, and TCRBuilder. TCRmodel has an advantage that it is also available as a local program through the Rosetta suite. However, this is not implemented into the pipeline.

Based on a leave-one-out approach in comparison of the modeling services, Repertoire Builder showed the best results through our internal testing. However, methods for submission are available for each platform. Including web automation to submit several sequences to web servers that don't support multi-sequence input. For Repertoire Builder and TCRmodel, which both allow for batch submission, ModelEngine is not needed.

1.4 Pairing

Resulting modeled TCRs become beneficial when you are able to analyze their theoretical biological function. To do this, the first step is pairing it with a potential antigen. Unlike antibodies, TCRs are specific to a peptide presenting structure known as the major histocompatibility complex (MHC), also known as human leukocyte antigen (HLA). As this tool is capable of analysis of TCRs non-exclusively to homo sapiens, the term MHC will be utilized.

The program TurnTable has several functions to easily pair TCR structure(s) to one or several pMHCs. First, TurnTable centers the TCR to coordinates 0,0,0 using PCA analysis and ensures receptor binding domain is facing down. Then TurnTable positions the TCR's interface in-line but at a distance to the pMHC interface as needed according to the RosettaDock 4.0 input desires. Along with repositioning the TCR and pMHC, TurnTable trims the TCR chains to just the variable region, removes the beta-2 microglobulin, and trims the MHC. This trimming process helps reduce run-time of docking steps in the next component of TRain. As a final step, TurnTable renames the resulting PDB atoms and residues as desired by RosettaDock 4.0. This renaming can be corrected for in the PostCoupler program.

1.5 Docking

TCRcoupler is where TCRs are docked to the paired pMHC antigen. TCRcoupler can be performed on a single PDB file or a directory of PDB files. Following RosettaDock 4.0 procedure, flexible side chain and flexible back bone settings are allowed for the TCR chains. However, less time-demanding but also less refined docking can be done with rigid settings.

Resulting docked TCR/pMHC complexes can be viewed using any protein visualization program. However, to make for better analysis, a few steps should be complete before continuing past docking. PostCoupler takes in resulting PDB files and cleans them of any residual information and most importantly, rematches the numbering of the native PDB files.

1.6 Analysis

With complete crystal modeled docked TCR/pMHC structures DataDepot provides several analytical figures and tables to gain insight to the binding/non-binding interface. This step is not all encompassing of analytical features but can provide quick insight into the docked structures produced. Analysis can be done to look closely at a single interface or compare across several with different tools.

2 — Installing the TRain suite

TRain code is available through GitHub at: <https://github.com/Aseemann/TRain>. TRain has several Python3 package and other dependencies. To allow for the flexible of use of TRain , packages and dependencies are laid out for each step. Use of a package manager such as Conda (<https://www.anaconda.com/products/individual>) may be beneficial.

2.1 All

BioPython is necessary for almost all portions of TRain and can be assumed it's installed.

```
1 pip install biopython
```

2.2 Input

SeqConductor executable code can be found in the folder 01_Input. Python packages required for use can be installed as followed:

```
1 pip install pandas
2 pip install openpyxl
```

For instructions on how to run SeqConductor and for a list of command line options, see chapter 3.

2.3 Model

ModelEngine can be run on a Linux, Mac, or Windows machine as long as it has the ability to run Chrome web browser. To run ModelEngine all packages besides Selenium are default packages. Selenium is a package that allows for website testing and automation. Selenium's python package can be installed using pip.

```
1 pip install selenium
```

Drivers are also needed to run Selenium. The Chrome driver is needed for the current version of ModelEngine and is available at <https://sites.google.com/chromium.org/driver/>. Be sure to remember where the chromedriver is installed as it'll be needed for the configuration file for ModelEngine.

To modify the configuration file, use your preferred text editor and modify line 2 writing the location of the chrome driver in the quotations.

2.4 Pair

The only non-standard dependencies of TurnTable are BioPython, numpy, SciPy, and sklearn. These are needed for the use of PDB_Tools_V3.py, which TurnTable utilizes.

```
1 pip install numpy  
2 pip install scipy  
3 pip install -U scikit-learn
```

2.5 Dock

Installed on server under "/opt/rosetta_src_2021.16.61629_bundle".

For docking, a local installation of Rosetta needs to be installed. Installation instructions can be found at https://new.rosettacommons.org/docs/latest/getting_started/Getting-Started and Rosetta must be built with MPI executable and instructions can be found at https://new.rosettacommons.org/docs/latest/build_documentation/Build-Documentation.

If MPI is not installed on the computer you can install using the following (apt installer):

```
1 sudo apt-get install openmpi openmpi-de
```

or following instructions at <https://www.open-mpi.org/faq/?category=building>.

The configuration file for TCRcoupler is inside directory 04_Dock. Modify the configuration file with your preferred text editor writing the location of Rosetta install in line 2 inside the quotations.

2.6 Analysis

Several dependencies are needed for DataDepot and can be installed using pip.

```
1 pip install pandas  
2 pip install matplotlib  
3 pip install seaborn
```

Using the SeqConductor program

- Input files
- Parameters
- Output

Under the hood

3 — The Input Step

The input step is handled by the SeqConductor program. The SeqConductor program takes in input from single cell sequencing data of T-cell receptor gene segments names and CDR3 amino acid sequences. The inputted segments are then aligned with the CDR3 amino acid sequence taking priority in final output. Output can be specified for an information table or an alpha and beta fasta file for direct submission to Repertoire Builder or TCRmodel. If other modeling suite required, refer to chapter 4, modeling (LYRA & TCRBuilder).

3.1 Using the SeqConductor program

3.1.1 Input files

SeqConductor requires a standard input and an optional input. Standard input for SeqConductor is the table of single cell sequencing data in the form of TRAV, CDR3 α sequence, TRAJ, TRBV, CDR3 β sequence, TRBJ. By default, TRAV, CDR3 α , TRAJ, TRBV, CDR3 β , TRBJ are respective to columns 1, 2, 3, 4, 5 and 6. While column 0 is reserved for sequence ID. This can be altered with parameters listed below. These tables can be in the format of xlsx or csv files. The program automatically detects based on the file's handle. Regarding xlsx files, sheets names can also be specified if not the primary sheet. The optional input is the reference fasta file of gene family segments. By default, the fasta file "family_seq.fasta" is utilized. This fasta file is pulled results from IMGT/GENE-DB specified for the TR molecular components of Homo sapiens. While full support isn't provided for other organism, "full_family_seq.fasta" contains the rest of the organisms that IMGT/GENE-DB has data for.

3.1.2 Parameters

The SeqConductor program accepts the following parameters:

Input:

Single Cell Table xlsx or csv of single cell t-cell data (Required)

- s Parameter allows for selection of an alternative sheet name from a submitted xlsx file
- a Append each chain with constant region
- c List of numbers correlating to the columns containing each segment. Can exclude Seq ID
eg. Seq ID, TRAV, CDR3 α , TRAJ, TRBV, CDR3 β , TRBJ = 0,1,2,3,4,5,6
- o Parameter allows you to change organism in use, requires "full_family_seq.fasta" gene family file
- genefamily Parameter allows for the selection an alternative gene family fasta file

Output:

- i (Default) Parameter outputs a table containing prominent information along with the resulting $\alpha\&\beta$ chain sequences.
- f Parameter provides fasta files with resulting $\alpha\&\beta$ chains separated to two fasta files.
- omission Characters to avoid in header ID for fasta file submission | comma sep.

Other:

- v Verbose: Show alignments being produced
- silent Silent: Mute even poor alignments

An example of a typical run:

```

1 python3 SeqConductor.py Sample_Files/Test_Table.xlsx -f -a
2     or
3 python3 SeqConductor.py Sample_Files/Test_Table.xlsx -f -a -s Sheet2
4     -c 3,4,5,7,8,9,11

```

3.1.3 Output

Sample outputs can be found in the sub-directory `Sample_Output/`.

1. alpha.fasta
2. beta.fasta
3. alpha_full.fasta (Appended constant region)
4. beta_full.fasta (Appended constant region)
5. alpha_full_rep.fasta (Removed "+" from Seq. IDs to avoid Repertoire Builder Error)
6. beta_full_rep.fasta (Removed "+" from Seq. IDs to avoid Repertoire Builder Error)
7. information_table.csv

3.2 Under the hood

The main steps carried out by SeqConductor:

1. Construct dictionary of gene family segments
2. Construct dictionary containing information from submitted table
3. Create full TCR sequences
4. Output:
 - (a) Information Table
 - (b) Fasta files: alpha.fasta & beta.fasta

Alignment of V-gene segment + CDR3 + J-gene segment is conducted in step 3. The CDR region, which from single cell sequencing is in full, is always retained while the start/end of each gene segment can be trimmed. All alignments are done with scores of Match = 2.5, Mismatch = -1, Gap = -1.5, and Extend Gap = -0.5. First, the program aligns the TRAV segment with CDR3 α . To perform the alignment, the V-segment sequence is trimmed to the final amino acids with a length of the CDR3 sequence minus 5. In the case that a CDR3 sequence is shorter than 7 amino acids, the last amino acids from the V-segments are pulled equal to the length of the CDR3 sequence. It is common to see very short overlaps in the V-segment + CDR3 overlap. To avoid poor alignments, where the CDR3 is placed in front of the V-segment, a sliding window approach is then used. In the case of a sliding window alignment, the V-segment is shrunk by one amino acid (from the head of seq.) until a good alignment is found or the alignment is less than 2 amino acids long. In the case the alignment is less than 2 amino acids long, the CDR is directly appended to the V-segment and a cation message is sent to the console. The V-segment/CDR3 + J-segment overlap is slightly different. In this case, larger overlap is more common. So a single alignment is completed between the V-segment/CDR3 + J-segment with no additional sliding window option. For this alignment, the V-segment/CDR3 is trimmed to the length of the J-segment length. In the case of a poor alignment with several gaps in this method, the J-segment is directly append to the V-segment/CDR3 sequence and a message is sent to the console again.

Using the ModelEngine program

Configuration File

Input files

Parameters

Output

Under the hood

4 — The Modeling Step

Many TCR modeling services are already developed and easily available to produce theoretical TCR $\alpha\&\beta$ chain pairs. These services include LYRA, TCRmodel, TCRBuilder, and Repertoire Builder. Several of these services offer batch submission which is helpful for generating models of TCR sequences collected. However, a few don't support batch submission and TCRmodel allows batch submission but without the function of a blacklist for batch submission. ModelEngine provides an automation of these web-tools, however it is greatly encouraged to visit the webpages before use to be sure to properly cite these tools and refer to their publications. Repertoire Builder allows for easy batch submission with attached blacklists so it's not included in this tool but previous testing has shown it to be one of the most reliable.

4.1 Using the ModelEngine program

4.1.1 Configuration File

Config.ini file must be prepared. Point to the location of chromedriver installation (refer to chapter 2).

4.1.2 Input files

Two separate fasta files containing alpha chains in one file and beta chains in the other can be submitted. Headers must match up between alpha and beta chains. SeqConductor will produce these fasta files.

To provide a blacklist, in which homologs are restricted to not include provided PDB id, follow the same practice as Repertoire Builder. An example is provided below. Be sure that both chain's headers contain the identical list.

```
1 >4jff ((3vxt,3pwp,4jff))
2 KQEVEQNSGPLSVPEGAIASLNCTYSFLGSQSFFWYRQYSGKSPELIMFTYREGDKEDGRF
3 TAQLNKASQHVSLLIRDSQPSDSATYLCAVNDGGRLTFGDGTLTVKPNIQNPDPAVYQLR
4 DSKSSDKSVCLTDFDSQTNVSQSKDSDVYITDKCVLDMRSMDFKNSAVAWSNKSDFAC
```

5 ANAFNNSIIPEDTFF

4.1.3 Parameters

The ModelEngine program accepts the following parameters:

- a Fasta file containing alpha chain sequences
- b Fasta file containing alpha chain sequences
- l Automate LYRA submission
- m Automate TCRmodel submission
- t Automate TCRBuilder submission
- s Optional starting position in fasta file. Provide header.
- d Download directory folder name in reference to program location default=Models
- long Scalar for wait times for elements to appear and page to load

An example of a typical run:

```
1 python3 ModelEngine.py -a Sample_Files/alpha_full.fasta  
2   -b Sample_Files/beta_full.fasta -l  
3   or  
4 python3 ModelEngine.py -a Sample_Files/alpha_full.fasta  
5   -b Sample_Files/beta_full.fasta -t -d Results
```

4.1.4 Output

Resulting structures that were successful will now be in the download directory `Models` or in the download directory file name provided. For sequences that didn't produce a successful model, an error list will be printed after completion of run.

4.2 Under the hood

ModelEngine utilizes the python package Selenium to automate the webtools. This allows for a test environment in a Chrome browser window. This tool works by pointing to HTML and CSS elements and providing input. For LYRA, this is straight forward as it submits the chain sequences and accompanying blacklist, then hitting submit. The tool waits for the results page to show and then downloads the file to the desired directory and then followed by renaming the file to the header provided in the fasta file. This is an identical procedure for TCRmodel and TCRBuilder. However, it is considerably faster to do batch submission for TCRmodel if no blacklist is needed. This method is highly reliant on the future modifications made to these webservers. If ever a tool is non-functional, please inform the contact references provided.

Timing:

Each web page has slightly different wait times for elements to appear on screen. For LYRA, we initially wait 10 seconds for the web page to load after it's first called. Then we wait until the chain 1 text entry box to appear for a maximum of 10 seconds. Then we input the blacklist, alpha chain, beta chain, and then hit submit. Then we wait for the results page to appear based on its "jsmol" element for a maximum of 200 seconds. This is typically well within the time it takes for LYRA to return a

result. We then wait a few seconds before clicking the download button for the PDB to ensure it's loaded. Once downloaded, ModelEngine renames the PDB file to the corresponding header. If the TCR model failed to produce, then it's appended to an error list and presented once the program is finished running. This is identical for TCRmodel. However, for TCRBuilder a maximum wait time for a result is 1800 seconds (20 minutes) as it typically takes longer for a result.

If for any reason that the timings are too short and resulting with errors the command line option "--long" can be included with your scalar. A scalar of 1.5 or 2 may be appropriate to initially try.

Using the TurnTable program

- Input files
- Parameters
- Output

Under the hood

5 — The Pairing Step

After constructing a modeled TCR structure, it then needs to get paired to a pMHC before the docking steps. TurnTable allows for submission of single or multiple TCR and pMHC PDB files. It then pairs each from the collection. To assist in the ability of utilizing crystal structures versus the modelled TCRs, full TCRpMHC complexes can be submitted and the TCR or pMHC extracted. To help with efficient docking, TCR chains are trimmed if from a crystal structure, the MHC is trimmed, and the beta-2 microglobulin is removed. Following trimming, the PDB(s) are then renumbered and relabeled as needed for TCRcoupler and this can be corrected for after docking with PostCoupler.py if you want to keep the native numbering.

5.1 Using the TurnTable program

5.1.1 Input files

There are several input methods for TurnTable . The standard is to supply one TCR and one pMHC, either can be stripped from a complete complex, and pairing will occur between those two structures. Submission of multiple TCRs or pMHC is possible with a folder containing the PDB files. A single directory of TCR/pMHC structures can also be submitted and for every TCR and every pMHC to be swapped around resulting the squared number of inputs. If submitting TCR crystal structure, ensure you utilize -trimA and -trimB command line options.

5.1.2 Parameters

The TurnTable program accepts the following parameters:

- t TCR pdb file or folder of TCR PDBs
- m pMHC pdb file or folder of pMHC PDBs
- p Folder of PDBs and all TCRs and pMHCs will be swapped
- trimA Trim TCR alpha, needed if full crystal structure
- trimB Trim TCR beta, needed if full crystal structure

-trimM Prevent trimming of MHC

An example of a typical run:

```
1 python3 TurnTable.py -t Sample_Files/2nx5.pdb -m Sample_Files/1qrn.pdb --trimA  
2   --trimB  
3     or  
4 python3 TurnTable.py -t Sample_Files/ -m Sample_Files/. --trimA --trimB  
5     or  
6 python3 TurnTable.py -p Sample_Files/ --trimA --trimB
```

5.1.3 Output

Resulting output will create a Paired/ sub-directory containing updated TCR/pMHC PDB files. PDB files are named as "TCR_pMHC.pdb". These file are now in a state prepared for docing, see chapter 6.

5.2 Under the hood

The main steps carried out by TurnTable:

1. Determine files versus folders.
2. Create temporary TCR PDB.
3. If crystal structure, clean PDB and split off TCR.
4. Update TCR labels to D and E for alpha and beta respectively.
5. If crystal structure, trim TCR.
6. Center TCR utilizing primary component axis (PCA).
7. Create temporary pMHC PDB.
8. Clean crystal structure and split off pMHC.
9. Superimpose reference to centered temporary TCR file.
10. Superimpose pMHC to reference pMHC.
11. Removed reference structure and save as "TCR Name"_"pMHC Name".pdb

The majority of TurnTable is handled by methods from PDB_Tools_V3.py .

Using the PrepCoupler program

Input files

Output

Example Run

Using the TCRcoupler program

Configuration File

Parameters

Flexible Docking

Rigid Docking

Using the PostCoupler program

Parameters

Input files

Output

Example Run

6 — The Docking Step

To gain a structural prediction a TCR/pMHC interface, two Rosetta docking protocols are part of the TRain suite. One or several PDBs can be provided for computational docking conducted following the RosettaDock 4.0 protocol [Mar+18] or alternatively a simplified rigid docking protocol following RosettaDock 3.2. Flexible docking allows for movement of sidechains and slight movement of the backbone of the TCR, this allows for potentially better predictions. Docking is a computationally taxing process; If a heavily multi-core computer is not available, restraining to rigid docking may be necessary. Optional parameters and alternative starting positions can allow for additional customization.

6.1 Using the PrepCoupler program

PrepCoupler is a useful program if TurnTable is not necessary for any reason. PrepCoupler prepares the PDB(s) for Rosetta. To do so, several modifications are made. The PDB is renumbered to a continuous count of amino acids versus a new count for each chain. As well as, the chains are labeled to standard TCR PDB chain labeling of MHC: A, Peptide: C, Alpha:D, and Beta:E. With the removal of the beta-2 microglobulin (chain: B) for more efficient docking as it is not a component of the interface. Then to further enhance efficiency, the MHC chain is trimmed to just the two alpha helices and the back beta pleated sheet. The $\alpha\&\beta$ chains are also trimmed to contain only the variable region. This program however does not separate the TCR from the pMHC. This program should also be ran on native structure being submitted for docking analysis.

6.1.1 Input files

Provide any TCR PDB file or folder of TCR PDB files.

6.1.2 Output

TCR PDB file with updates to comply with Rosetta applications including: proper order of chains, updated numbering, updated chain labels, removal of unneeded data.

6.1.3 Example Run

```

1 python3 PrepCoupler.py Sample_Files/1ao7.pdb
2     or
3 python3 PrepCoupler.py Sample_Files/PDB_Dir/

```

6.2 Using the TCRcoupler program

TCRcoupler is an automation of several Rosetta applications to generate a docked TCR/pMHC complex. Depending on the computational resources available, use of optional parameters may be necessary. Things to reduce run time include: changing from flexible to rigid docking, decrease number of relaxation steps, decrease number of docking permutations, decrease number of refinement runs, or increase core count.

To ensure TCRcoupler run smoothly a few things should be ensured. TCRcoupler relies on the mpi complied version of Rosetta. This allows TCRcoupler to unitize multiple cores to increase efficiency of runs. By default, TCRcoupler will request all cores available on the machine but can be changed with command line argument (see section 6.1.2). Rosetta requires a specific numbering scheme for the structure and chain order within the PDB. To avoid numbering and chain order issues, utilize TurnTable (refer to chapter 5).

6.2.1 Configuration File

Config.ini file must be prepared. Point to the location of Rosetta installation (refer to chapter 2).

6.2.2 Parameters

The TCRcoupler program accepts the following parameters:

| | |
|-----|---|
| pdb | PDB file or folder of PDBs for docking |
| -l | Changes to Linux runnable program default |
| -m | Changes to MacOS runnable program |
| -f | Initializes flexible docking default |
| -r | Initializes rigid docking |
| -c | Max number of cpu cores provided default=all |
| -p | (Flex) Number of pmhc relax runs default=100 |
| -x | (Flex) Number of xml relax runs for TCR default=40 |
| -b | (Flex) Number of back bone rub runs for TCR default=30 |
| -s | (Flex) Number of fast relax runs for TCR default=30 |
| -a | (Rigid) Number of relax runs performed default=100 |
| -d | (Both) Number of docking runs performed default=10000 |
| -e | (Both) Number of refinement runs, post dock default=100 |
| -n | Native structure folder to run optional comparison to crystal structure, pdbs have to be in folder and match numbering of submitted pdbs. |

6.2.3 Flexible Docking

Input files

PDB(s) submitted for flexible docking must be numbered properly and the alpha and beta chain must come after the MHC and peptide or resulting errors may occur. If PDBs submitted have not been processed through TurnTable then PrepCoupler should be run. PDBs provided as native structures should also be run through PrepCoupler. Flexible docking also requires PDB_Tools_V3.py to be located in the same directory as TCRcoupler.

Output

Docking generates several output files including: several flag files, split TCR and pMHC PDB files, input_files and output_files directories and sub-directories, several intermediate PDB files, and ensemble list files. To help reduce storage utilization, TCRcoupler automatically deletes PDBs produced that are not the highest scoring. Automatic removal of PDBs can be inhibited by command line parameter. For results, enter into the output_files directory and within /Refine/ the remaining PDB will be the best scoring docked structure.

Example Run

Setup:

Before we start the run, use your preferred text editor and modify config.ini to your Rosetta installation.

```
1 (Driver)
2 rosetta_loc="/opt/rosetta_src_2021.16.61629_bundle"
```

For an example run, create a new directory in 04_Dock for our run. Then copy TCRcoupler.py, PDB_Tools_V3.py, config.ini, and Sample_Files/2nx5_2nx5.pdb to our new folder.

```
1 cd 04_Dock
2 mkdir Run_1
3 cp TCRcoupler.py Run_1/
4 cp PDB_Tools_V3.py Run_1/
5 cp config.ini Run_1/
6 cp Sample_Files/2nx5_2nx5.pdb Run_1/
7 cd Run_1
```

Now let's do a flexible docking run. For this run, we're going to allocate 100 CPU cores.

```
1 python3 TCRcoupler.py 2nx5_2nx5.pdb -f -c 100
```

As an alternative, let's submit a folder of PDBs. Here's the entire process. Nohup and allow us to run in background

```
1 cd 04_Dock
2 mkdir Run_1
3 cp TCRcoupler.py Run_2/
4 cp PDB_Tools_V3.py Run_2/
5 cp config.ini Run_2/
6 cp -r Sample_Files/Multi_Run Run_2/
7 cd Run_2
8 nohup python3 TCRcoupler.py Multi_Run -f -c 100 &
```

6.2.4 Rigid Docking

Input files

PDB(s) submitted for rigid docking must be numbered properly and the alpha and beta chain must come after the MHC and peptide or resulting errors may occur. If PDBs submitted have not been processed through TurnTable then PrepCoupler should be ran. PDBs provided as native structures should also be ran through PrepCoupler.

Output

Rigid docking will produce less resulting files and directories. The produced item will include: three flag files, a time.txt file, several intermediate PDB files, and output_files directory. Just like with flexible docking, intermediate PDB files will be removed unless requested by user not to. Results can also be found in the output_files directory and within /Refine/ the remaining PDB will be the best scoring docked structure.

Example Run

Rigid docking follows in line with flexible docking but with just the replacement parameter of -f with -r. Keep in mind, a directory of PDBs can also be submitted for rigid docking.

```

1 cd 04_Dock
2 mkdir Run_3
3 cp TCRcoupler.py Run_3/
4 cp config.ini Run_3/
5 cp -r Sample_Files/2nx5_2nx5.pdb
6 cd Run_3
7
8 python3 TCRcoupler.py Multi_Run -r -c 100

```

6.3 Using the PostCoupler program

6.3.1 Parameters

The PostCoupler program accepts the following parameters:

| | |
|----------|--|
| pdb | PDB file being renumbered |
| -r | Reference PDB file being used to match numbering |
| -n | (Optional) Customized name for output file |
| -c | Chains that will be renumbered from reference PDB default=all |
| --custom | Custom numbering for smallest chain e.g. 9,11,2,4,5 Has to match num of aa to numbers provided |

6.3.2 Input files

PDB being submitted would be the docked structure. This was renumbered to the required scheme for Rosetta applications. Assumes chain labeling is identical.

6.3.3 Output

Resulting PDB with either "_orignum.pdb" appending PDB file provided or with custom name provided.

6.3.4 Example Run

Let's try to examples. One with a redocked structure where both the TCR and pMHC can use the same reference. In the second, we'll used a TCR and pMHC from two seperate structures and only renumber from their respective reference.

```
1 python3 PostCoupler.py Sample_Files/1qrn_1qrn_docked.pdb
2     -r Sample_Files/1qrn.pdb -n 1qrn_1qrn_updated.pdb
3         or
4     python3 PostCoupler.py Sample_Files/1qrn_1qse.pdb -r 1qse.pdb -c AC
5         and
6     python3 PostCoupler.py Sample_Files/1qrn_1qse.pdb -r 1qrn.pdb -c DE
```


Using the DataDepot program

- Configuration File
- Energy Breakdown
- ABusage
- Native Structure Comparison

7 — The Analysis Step

The analysis step is an accumulation of Python methods that allow for quick visual insight. All of these methods are encompassed in DataDepot . Some of these methods still rely on Rosetta to collect even further information from the structures produced. Methods that require a Rosetta installation will be signified and the configuration file must be updated, see more information below.

7.1 Using the DataDepot program

DataDepot assumes that TCRs are labeled A = MHC, C = Peptide, D = Alpha, and E = Beta. If labeling does not match, PrepCoupler can be used from chapter 6.

7.1.1 Configuration File

Config.ini file must be prepared if conducting Energy Breakdown and Native Structure Comparison. Point to the location of Rosetta installation (refer to chapter 2).

7.1.2 Energy Breakdown

Rosetta's application Residue Energy Breakdown provides details of residue-residue interactions occurring within a PDB file. This information can provide more insight than just distance value cutoffs for interactions occurring at the interface of a TCR/pMHC complex. DataDepot provides two forms to view the interactions happening at the interface of the TCR/pMHC complex. These two forms are in a simple table and in a heatmap.

For both, you can submit either a PDB file or an already complete energy breakdown table. If a PDB is provided, Energy Breakdown is ran and the resulting .out file is converted to a .csv. If a energy breakdown table is submitted .out and .tsv files are converted to a .csv. The resulting .csv file of either is then processed to either produce the table or heatmap.

Table

The Energy Breakdown table provides a list of interactions occurring between the alpha and beta chains of the TCR to the MHC or peptide. Both attractive and repulsive energy values are shown.

Example Run

```

1  python3 DataDepot.py --table Sample_Files/sample.pdb
2  or
3  python3 DataDepot.py --table Sample_Files/energy_breakdown.out

```

Heatmap

The Energy Breakdown heatmap provides similar information as the table. However, the heatmap makes it easier to get a visual reference of interactions occurring at the interface. Only attractive energy values are displayed in the heatmap.

Example Run

```

1  python3 DataDepot.py --heatmap Sample_Files/sample.pdb
2  or
3  python3 DataDepot.py --heatmap Sample_Files/energy_breakdown.out

```

7.1.3 AUsage

AUsage calculates interface scores for Alpha to pMHC and Beta to pMHC utilizing the Rosetta application InterfaceAnalyzer. The program is ran twice, once for the alpha chain and another for the beta chain. From the output of InterfaceAnalyzer for both, the delta G separated value is pulled. This provides us with an energy based value for the interactions occurring between Alpha to the pMHC and Beta to the MHC. The output is AB_Usage.csv, which is a table containing all delta G separated scores for each Alpha and Beta chain.

Example Run

```

1  python3 DataDepot.py --ab Sample_Files/sample.pdb
2  or
3  python3 DatDepot.py --ab Sample_Files/Multi_PDBs

```

7.1.4 Native Structure Comparison

TCRcoupler allows you to submit a native crystal structure along with a PDB being submitted for docking. This can be useful to compare a crystal structure of a TCRpMHC complex to a re-docking attempt or compare against slightly mutated submitted chains. Submitting a native structures adds additional columns to the score files produced during docking and refinement.

Example Run

```

1  python3 DataDepot.py --sc Sample_Files/1qrn_1qrn.sc -x l_sc -y Fnat
2  or
3  python3 DataDepot.py --sc Sample_files/Score_Files -x motif_dock -y rms

```

Axis Options:

1. I_sc
2. motif_dock
3. total_score
4. Fnat
5. rms
6. cen_rms

Bibliography

- [Mar+18] Nicholas A. Marze et al. “Efficient flexible backbone protein-protein docking for challenging targets”. In: *Bioinformatics* 34.20 (2018). ISSN: 14602059. DOI: 10.1093/bioinformatics/bty355 (cited on page 21).