User Guide

for

T3SEpp

v1.0

02/03/2020

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1. Introduction

This manual was prepared for the standalone version of T3SEpp (version 1.0) and the related modules. For usage of the T3SEpp webserver, please refer to the HELP page of the T3SEpp website. The link is: http://www.szu-bioinf.org/T3SEpp.

This manual demonstrated the usage of T3SEpp and the modules with detailed examples, but did not introduce too much on methodology. For methodological details, parameter optimization and model performance please refer to http://www.szu-bioinf.org/T3SEpp/modules.html.

T3SEpp could be run as a whole pipeline. Alternatively, the modules could be used independently. The manual illustrated the usage of T3SEpp in Section 2, and the modules or related models in the remaining sections.

For any technical question, please feel free to contact anyone of us:

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Chen, 531634084@qq.com

Please cite the following reference when you use T3SEpp or the modules.

- 1. Xinjie Hui, Zewei Chen, Mingxiong Lin, Yueming Hu, Yingying Zeng, Xi Cheng, Le Ou-Yang, Ming-an Sun, Aaron P. White, Yejun Wang. T3SEpp: An Integrated Prediction Pipeline for Bacterial Type III Secreted Effectors. *mSystems*. In revision.
- 2. Yueming Hu, He Huang, Xi Cheng, Xingsheng Shu, Aaron P White, John Stavrinides, Wolfgang Köster, Guoqiang Zhu, Zhendong Zhao, Yejun Wang. A global survey of bacterial type III secretion systems and their effectors. *Environ Microbiol.* 2017; 19(10):3879-3895.

2. Manual of T3SEpp

- 1) System requirement: Linux or Mac.
- 2) Software or package prerequisites: The following software or packages should be pre-installed and configured into environmental variable path. Note that in Linux system, the Tensorflow needs to be activated before running T3SEpp (source /root/tensorflow/bin/activate).

```
Perl 5
R version 3.3 or later

Python3
*GO (version 1, if source code compilation needed)

BLAST suite version 2.2.30+

HMMER3.1

Numpy 1.18.1

Pandas 0.25.3

Tensorflow 1.3.0

Keras 2.1.2

argparse
```

3) Download and installation of T3SEpp:

The T3SEpp package (T3SEpp.v1.0) for Linux or Mac system could be downloaded from the website: http://www.szu-bioinf.org/T3SEpp.

Decompress the "T3SEpp.tar.gz" and get into ~/T3SEpp/ from terminal. If the pre-compiled version does not work, try to re-compile all the GO scripts in the "codes" sub-folder of ~/T3SEpp/sigHMM/, ~/T3SEpp/cbdHMM/, ~/T3SEpp/effectHMM/, ~/T3SEpp/transHMM/, ~/T3SEpp/flBlast/, ~/T3SEpp/SeqAac/, ~/T3SEpp/SignalP/, ~/T3SEpp/PSORTb/ and ~/T3SEpp/TMHMM/, and replace the binary files in to the "bin" sub-folder of each module with the newly compiled

ones, respectively. Compiling GO scripts:

\$ cd ~/T3SEpp/MODULE_NAME/bin

\$ go build ../codes/xxx.go

"xxx.go" is the script in "codes" sub-folder of MODULE_NAME. Compile all the scripts. Also, compile "T3SEppPrep.go" and "T3SEpp.go" and put the binary result files in the same directory (~/T3SEpp/).

4) Input files:

There is a necessary protein sequence file and four optional other input files (promoter sequence file, PSORTb prediction file, SignalP prediction file and TMHMM prediction file). If there is more than one file, the protein or gene name for each file must be consistent, in terms of the total number and identity. The names should not contain a space or an illegal character such as '|', and it is suggested not to use pure numbers as the protein or gene name in case of unexpected errors in different systems. For the 'protein sequence file' and 'promoter sequence file', T3SEpp requires them to be FASTA-formatted, as exemplified by the demonstrated "test.fa" and "testPromoter.fa" files (Fig 1).

>T1 protein sequence file - FASTA format

VDCSIKNKKGKNTMEINPIPIGSINILTTQSLTESQSTEEAKIEQG
SYQAKRIISQEPRLSTEFDPSLFKNKTERYNARLLKTVFEDTPNTS
YQQLWLTPNKQHKLTEEELSTIISKEIKNILIKEQVIKELQTELD
>T2
MGSSHHHHHHSSGLVPRGSHMFLTFPNVAITRDNRIDKLSENDLEI
TFKVSFSTTDRAMFRERHIEWQGNAIRLERQLNTGLNVSRG
>T3
MLKPICHSGSIKVPEYLETDKEKNAGRTPLSSDIQQVRNVVEDVPI
YGRLFGKQVLAHIHSRCQRDADIIREKALRRISRECGAEIDCALLI
SYALGCRPGDLPAYNVGRDSVETKAFELEKLADSPYAPYGQTGGFS
>T4
MGHLIPIVLPPHRQLDSDVDMLEHIEPTPLFDRVTDSPTFLDGLT
MFSNVLSLFSGRALNQNGVICSKAVEQNLAALECGMINDFWVAES
>T5
MPPMNESLKSNFDTLHRQMRQMPLSHFTVEPNAPDYSGIRQSGFF/
HISVQEQQVAQAFQALSGLLFSEDSPIDKWKVTDMARVDQQSRVGG
RLSEQGIIPGRVPESDVHPDSWRYISYRNELRSERGGGEMQSQALF

promoter sequence file - FASTA format >T1 AGGAACGCTTACGGGCTGTCTTCTCATTTTTAAGCGGCTCTCGG GAGAGCGTGCTTGCGCTGGGGCGTCCCGTCGGCAGCAACTGAAA GCCGTCGCACCTCGCCGGCTCCTTGCTGCCCCTTCAGAAGAGAT >T2 TCCTGATTATTCAACCACTACAAAGTGTCCATGGCAGAAACGCG CGGTCTGTATGTGAGGCTTTAGCCTGTTCTGCGTCTGCCGGCGG TTCCAGTAGCCGTCGATTTTGTACCTCCGTTGATGCTCATGAAC >T3 TTGAAAAACGCCCTCAAACCAGTGTCTTTTTAATCGAAAAATGA AATGTCCACTAATCGATTTCCGGCACAGTACACCATTCCGGCAG GACATTGCCCGTGATTTTATGGAACCGCTGTACGGCCTACGCCA >T4 TTATTATATTCACCTTTGTGTAGCTCTTTTTTCTCTTAGATATC GACTTATGGGGCTAAAAAATTCACGGTAAAAGAACTTTCTTAAT AATTCCGATGTCCCCCCCTTTTTTTTAAACGCCCCTGTGAAGG AACAGGCGGTATTTATCCACTGAATTAATGGGAAAATTTCACA

Fig 1. Input sequence file format - FASTA, name consistency

The files 'PSORTb.out.txt', 'SignalP.out.txt' and 'TMHMM.out.txt' are optional and generated by submitting the 'protein sequence file' to PSORTb 3.0, SignalP 4.1 and TMHMM 2.0 webserver, respectively. The links for the three tools are listed as below:

PSORTb: https://www.psort.org/psortb/index.html

SignalP: https://services.healthtech.dtu.dk/service.php?SignalP-4.1

TMHMM: https://services.healthtech.dtu.dk/service.php?TMHMM-2.0

'Short Format (tab delimited)', 'Short – no graphics' and 'One line per protein' are selected as the output format of PSROTb, SignalP and TMHMM, respectively. The file format for 'PSORTb.out.txt', 'SignalP.out.txt' and 'TMHMM.out.txt' are shown in Fig 2. Once generated, the files are moved into ~/T3SEpp/. For this demonstration, "testPSORTb.txt", "testSignalP.txt" and "testTMHMM.txt" are the example files of 'PSORTb.out.txt', 'SignalP.out.txt' and 'TMHMM.out.txt' respectively.

	T1 C	ytoplasmic	8.96		
	T2 Ui	nknown 2.0	0		
		ytoplasmic			
		ytoplasmic			
		nknown 2.0			
		ytoplasmic			
		ignalP.out.txt			
T1 0.119 67 0.124	34 0.179	10 0.14	3 0.133 N	0.570	SignalP-noTM
T2 0.141 30 0.123	32 0.164			0.570	SignalP-noTM
T3 0.106 26 0.112	26 0.127			0.570	SignalP-noTM
T4 0.116 66 0.103	33 0.187			0.570	SignalP-noTM
T5 0.164 57 0.119	57 0.160			0.570	SignalP-noTM
T6 0.108 19 0.126	11 0.228	1 0.15	1 0.138 N	0.570	SignalP-noTM
		TMHMM.ou	it txt		
T1 len=343 ExpAA		rst60=0.1		1-0	Topology=o
•					
T2 len=121 ExpAA		rst60=0.1			Topology=o
T3 len=374 ExpAA	.=0.00 Fi	rst60=0.0	0 PredH	lel=0	Topology=o
T4 len=246 ExpAA	.=0.00 Fi	rst60=0.0	0 PredH	lel=0	Topology=o
T5 len=216 ExpAA	=0.00 Fi	rst60=0.0	0 PredH	lel=0	Topology=o
T6 len=523 ExpAA	=0.00 Fi	rst60=0.0	0 PredH	lel=0	Topology=o

Fig 2. Input file format of PSORTb, SignalP and TMHMM prediction

After the input files have been prepared, get into ~/T3SEpp/ through command line:

\$ cd ~/T3SEpp/

5) Running T3SEpp in one command line:

```
$ perl T3SEpp.pl -prot test.fa -prom testProm.fa -psortb testPSORTb.txt -signalp testSignalP.txt -tmhmm testTMHMM.txt -cutoff 0.2
```

The arguments are explained as below:

-prot	PRPTEIN_SEQUENCE_FILE_TO_BE_PREDICTED	Necessary
-prom	PROMOTER_SEQUENCE_FILE	Optional
-psortb	PSORTb_PREDICTION_FILE	Optional
-signalp	SIGNALP_PREDICTION_FILE	Optional
-tmhmm	TMHMM_PREDICTION_FILE	Optional
-cutoff	CUSTOMIZED_CUTOFF (DEFAULT:0.5)	Optional

6) Output file format:

The prediction results are listed in ~/T3SEpp/Results/. 'T3SEpp.out.txt' gives the overall prediction results of the integrated pipeline, while the other files provide the detailed results from different modules (Fig 3).

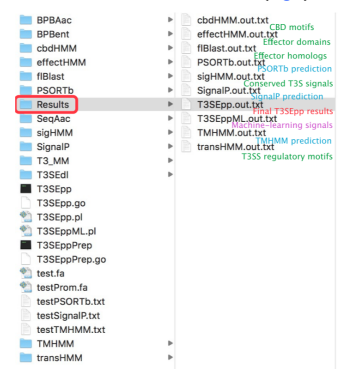


Fig 3. Prediction result files and their details

'T3SEpp.out.txt' summarizes the prediction results of different modules, the general prediction scores and the classification according to the customized or default (0.5) cutoff (Fig 4).

prot	T3SEppML	flBlast	sigHMM	cbdHMM	effectHMM	transHMM	TMHMM	PSORTb	SignalP	T3SEpp	Pred
T2	0	1	0	1	0	0	1	1	0	0.35	T3S
Т3	0	1	0	1	0	1	1	1	0	0.55	T3S
T4	0	1	0	0	0	0	1	1	0	0.28	T3S
T5	0	1	1	0	1	0	1	1	0	0.63	T3S
Т6	0	1	1	0	1	0	1	1	0	0.63	T3S
T1	0	1	0	1	0	1	1	1	0	0.55	T3S

Fig 4. The final prediction result of T3SEpp

3. Manual of T3SEppML

The module *T3SEppML* can also be run independently to screen the proteins with the atypical signal features of T3SEs.

- 1) System requirement: Linux or Mac.
- 2) Software or package prerequisites: The following software or packages should be pre-installed and configured into environmental variable path. Note that in Linux system, the Tensorflow needs to be activated before running T3SEpp (source /root/tensorflow/bin/activate).

```
Perl 5
R version 3.3 or later

Python3

*GO (version 1, if source code compilation needed)

Numpy 1.18.1

Pandas 0.25.3

Tensorflow 1.3.0

Keras 2.1.2

argparse
```

3) Download and installation of T3SEppML:

The *T3SEppML* module for Linux or Mac system could be downloaded from the website: http://www.szu-bioinf.org/T3SEpp.

Decompress the "T3SEppML.tar.gz" and get into ~/T3SEppML/ from terminal. If the pre-compiled version does not work, try to re-compile the GO scripts in ~/T3SEppML/SeqAac/codes/ and replace the binary files in ~/T3SEppML/SeqAac/bin with the newly compiled ones. Compiling GO scripts:

```
$ cd ~/T3SEppML/SeqAac/bin
$ go build ../codes/xxx.go
```

"xxx.go" is the script in ~/T3SEppML/SeqAac/codes/. Compile all the scripts.

4) Running flBlast in command lines:

```
$ cd ~/T3SEppML/
```

\$ perl T3SEppML.pl test.fa

Input file should be a FASTA-formatted protein sequence file, as exemplified by "test.fa". The protein name should neither be separated by space nor be with illegal characters such as "|" or purely numbers.

5) Output file format:

A single prediction result file will be generated named as "T3SEppML.out.txt". The format of the prediction result file is shown in Fig 5, containing 3 columns, protein ID, the scores of six models, and the classification with default cutoff of each model, where "1" means "T3S signal" and "0" means "non-T3S signal".

prot T1 T2 T3 T4	SeqAac BPBAac BPBent T3_MM T3SEdnn T3SErnn score 1 0.25 0.20 0.20 0.01 0.97 0 0.31 0.09 0.07 1.00 0.97 1 0.74 0.40 -0.01 1.00 0.21 1 0.23 0.34 -0.02 0.01 0.14	SeqAac BPBAac BPBent T3_MM T3SEdnn T3SErnn 1 1 1 1 0 1 0 1 1 1 11 1 1 1 0 1 0 1 1 1 0 0 0
T5	1 0.17 0.27 -0.01 0.01 0.04	1 1 1 0 0 0
T6	1 0.13 0.21 0.09 0.94 0.06	1 1 1 1 1 0

Fig 5. T3SEppML prediction result file format

Except for the T3SEdl, we do not provide independent manual for the other machine learning models in this version of UserGuide. However, they can still be run independently.

4. Manual of T3SEdl

The module *T3SEd1* can also be run independently to screen the proteins with the atypical signal features of T3SEs with two deep-learning models: *T3SEdnn* and *T3SErnn*.

- 1) System requirement: Linux / Mac / Windows.
- 2) Software or package prerequisites: The following software or packages should be pre-installed and configured into environmental variable path. Note that in Linux system, the Tensorflow needs to be activated before running T3SEpp (source /root/tensorflow/bin/activate).

Python3

Numpy 1.18.1

Pandas 0.25.3

Tensorflow 1.3.0

Keras 2.1.2

argparse

3) Download and installation of T3SEdl:

The *T3SEd1* module could be downloaded from the website: http://www.szu-bioinf.org/T3SEpp. Decompress the "T3SEdl.tar.gz" and get into ~/ T3SEdl/ from terminal.

- 4) Running T3SEdl in command lines:
- i) Predict with T3SEdnn:

```
$ python3 T3SEdl.py -t test.fa -m DNN
```

ii) Predict with T3SEdnn:

\$ python3 T3SEdl.py -t test.fa -m CNN_LSTM

Input file should be a FASTA-formatted protein sequence file, as exemplified by "test.fa", with each protein sequence 100-aa length (N-terminal 2-101 amino acids from each candidate protein) in one line. The protein name should neither be separated by space nor be with illegal characters such as "|" or purely numbers.

5) Output file format:

Either 'DNN' or 'CNN_LSTM' mode generates a "result.csv" file, curating the *T3SEdnn* or *T3SErnn* prediction results, respectively. The result file contains three columns: ID – the name of protein to be predicted, Score and Class ('1' – T3S signal or '0' – non-T3S signal). The default cutoff for classification is set as '0.5', i.e., >= 0.5 – T3S signal, < 0.5 – non-T3S signal.

5. Manual of flBlast

The module *flBlast* can also be run independently to screen the proteins with homology to validated T3SEs.

- 1) System requirement: Linux or Mac or Windows.
- 2) Software or package prerequisites: The following software or packages should be pre-installed and configured into environmental variable path.

*GO (version 1, if source code compilation needed)
BLAST suite version 2.2.30+

3) Download and installation of flBlast:

The *flBlast* module for Linux or Mac system could be downloaded from the website: http://www.szu-bioinf.org/T3SEpp.

Decompress the "flBlast.tar.gz" and get into $^{\sim}/flBlast/$ from terminal. There are three sub-folders and a testing file (Fig 6). The flBlast scripts have been precompiled and put in $^{\sim}/flBlast/bin/$. Users can also compile the source code in $^{\sim}/flBlast/codes/$ and place the binary file in $^{\sim}/flBlast/bin/$:

\$ cd ~/flBlast/bin \$ go build ../codes/xxx.go

"xxx.go" is the script in ~/flBlast/codes/. Compile all the scripts.

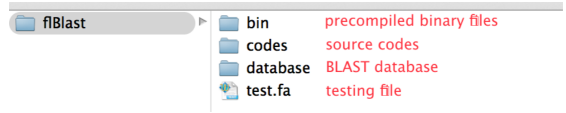


Fig 6. The folder organization of flBlast module

4) Running flBlast in command lines:

```
$ cd ~/flBlast/
```

\$ blastp -query test.fa
-db ./database/Validated_effectors_20160311 -evalue 1e-5
-outfmt 6 -out blast.1.txt

- \$./bin/bestpicker blast.1.txt >blast.2.txt
- \$./bin/simcovfilter ./database/Validated_effectors_20160311.length ./database/Validated_effectors_20160311.family ./database/Validated_effectors_20160311.function blast.2.txt

>flBlast.out.txt

Input file should be a FASTA-formatted protein sequence file, as exemplified by "test.fa". The protein name should neither be separated by space nor be with illegal characters such as "|" or purely numbers.

5) Output file format:

Three files were output and the "flBlast.out.txt" file designated by user contains the final prediction results. The result file has seven columns, representing candidate protein name, the most significantly homologous hit, effector family, functional group, length coverage and similarity for the covered regions (Fig 7).

Query	Subject	Subject_Acc	Family	Function	Coverage	Similarity
T5	Sal_SPI12_SpvC	CCF76774.1	0SPF	Substrate	1.00	0.73
T14	Pse_Pse2_HopAS1	AAZ37064.1	Pse_Pse2_HopAS1	Substrate	1.00	1.00
T15	Pse_Pse2_HopD1	NP_790715.1	X0PB	Substrate	1.00	0.94
T16	Xan_Xan_XopE2	CAJ23957.1	X0PE	Substrate	1.00	0.81
T17	Pse_Pse2_HopAF1	NP_791393.1	Pse_Pse2_HopAF1	Substrate	1.00	1.00
T18	Pse_Pse2_HopAK1	NP_793862.1	Pse_Pse2_HopAK1	Translocon	1.00	0.82
T22	Xan_Xan_XopAL1	AEL06398.1	X0PE	Substrate	0.85	0.35
T23	Erw_Hrp_AvrRpt2	CBA23177.1	AVRPT2	Substrate	1.00	0.99
T24	Bor_Bor_BspR	NP_888184.1	Bor_Bor_BspR	Substrate	0.85	0.98
T26	Shi_Shi_IpaH9.8	YP_406100.1	YOPM	Substrate	0.98	0.75
T27	Shi_Shi_IpaH7.8	YP_406039.1	YOPM	Substrate	1.00	0.63
T28	Bra_Bra_NopM	NP_768544.1	YOPM	Substrate	0.90	0.48
T29	Rhi_Rhi_NopT	NP_444174.1	Y0PT	Substrate	1.00	1.00

Fig 7. flBlast prediction result file format

6. Manual of sigHMM

The module *sigHMM* can also be run independently to screen the proteins with conserved T3S signal sequences.

- 1) System requirement: Linux or Mac.
- 2) Software or package prerequisites: The following software or packages should be pre-installed and configured into environmental variable path.

*GO (version 1, if source code compilation needed)
HMMER3.1

3) Download and installation of sigHMM:

The *sigHMM* module for Linux or Mac system could be downloaded from the website: http://www.szu-bioinf.org/T3SEpp.

Decompress the "sigHMM.tar.gz" and get into ~/sigHMM/ from terminal. There are four sub-folders and a testing file (Fig 8). The sigHMM script has been precompiled and put in ~/sigHMM/bin/. Users can also compile the source code in ~/sigHMM/codes/ and place the binary file in ~/sigHMM/bin/:

\$ cd ~/sigHMM/bin

\$ go build ../codes/xxx.go

"xxx.go" is the script in ~/sigHMM/codes/. Compile all the scripts.



Fig 8. The folder organization of sigHMM module

4) Running effectHMM in one command line:

\$ cd ~/sigHMM/

\$./bin/sigHMM hmmsearch ./hmmprofile test.fa out >test.out.txt
Input file should be a FASTA-formatted protein sequence file, as
exemplified by "test.fa". The protein name should neither be separated
by space nor be with illegal characters such as "|" or purely numbers.

5) Output file format:

The predictions were output into the result file with self-designated name, e.g., "test.out.txt" in this example (Fig 9, left), with two columns (Fig 9, right). The first column lists the names of proteins to be predicted, and the second column gives the prediction results: '-' represents no hit while 'SigFAM_NUMBER_N50' indicates the T3S signal family profile that the predicted protein contains.



Fig 9. sigHMM prediction results and output file format

7. Manual of effectHMM

The module *effectHMM* can also be run independently to screen the proteins with known effector domains.

- 1) System requirement: Linux or Mac.
- 2) Software or package prerequisites: The following software or packages should be pre-installed and configured into environmental variable path.

*GO (version 1, if source code compilation needed)
HMMER3.1

3) Download and installation of effectHMM:

The *effectHMM* module for Linux or Mac system could be downloaded from the website: http://www.szu-bioinf.org/T3SEpp.

Decompress the "effectHMM.tar.gz" and get into ~/effectHMM/ from terminal. There are four sub-folders and a testing file (Fig 10). The effectHMM script has been precompiled and put in ~/effectHMM/bin/. Users can also compile the source code in ~/effectHMM/codes/ and place the binary file in ~/effectHMM/bin/:

\$ cd ~/effectHMM/bin

\$ go build ../codes/xxx.go

"xxx.go" is the script in ~/effectHMM/codes/. Compile all the scripts.



Fig 10. The folder organization of effectHMM module

4) Running effectHMM in one command line:

\$ cd ~/effectHMM/

\$./bin/effectHMM hmmsearch ./hmmprofile test.fa out >test.out.txt
Input file should be a FASTA-formatted protein sequence file, as
exemplified by "test.fa". The protein name should neither be separated
by space nor be with illegal characters such as "|" or purely numbers.

5) Output file format:

The predictions were output into the result file with self-designated name, e.g., "test.out.txt" in this example (Fig 11, left), with two columns (Fig 11, right). The first column lists the names of proteins to be predicted, and the second column gives the prediction results: '-' represents no hit while 'Effector_FAM_NUMBER' indicates the effector domain family that the predicted protein belongs to.



Fig 11. effectHMM prediction results and output file format