SMORE

Synteny Modulator Of Repetitive Elements

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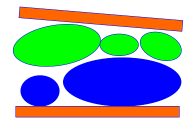
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1 Introduction

SMORE is a pipeline to detect evolutionary events along a phylogenetic tree based on a set of repetitive elements and multiple sequence alignments. This manual provides a more detailed description of the SMORE pipeline published in [1] and [2]. An example is shown in Section 6.

The pipeline is composed of two modular parts: (1) the inference of the orthology relation and (2) the quantitative analysis of the orthology relation, see Fig. 1. The first component identifies a map of genomic anchor points that are used to partition the annotated elements of interest into an initial set of candidate clusters. These are then processed to account for the most common artefacts in the input data and refined using information that is provided by analysing related but distinguishable sequence elements together. The second part of the pipeline is largely independent of the first and can also be employed using input data generated by other, third-party methods. With our pipeline, we provide an uninterrupted workflow that returns results based on input files and user-defined parameters. With the exception of breaks between subcommands indicated in Fig. 1 and where output data is provided for the user, the UNIX pipes utilizes to transfer data between software components.

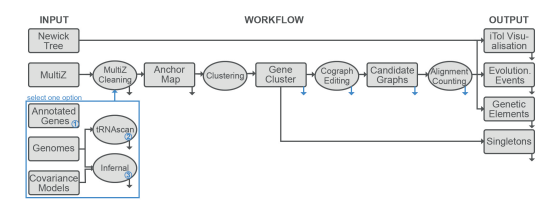


Figure 1: Summary of the computational workflow implemented in the SMORE pipeline for analyzing the evolution of mutlicopy genes. The compilation of orthology estimates and the quantitative analysis are logically separated and can also be used independent of each other. See text for details. Black arrows pointing into the direction of the pipeline show an uninterrupted workflow and hence no printing and reading of files in between single steps of the pipeline. Black arrows pointing downwards indicate output files that are always part of the output whereas blue arrows pointing downwards indicate the creation of temporary files and of optional output for the user, respectively.

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2 Quickstart

The program can be downloaded here https://github.com/AnneHoffmann/Smore. No further program installation is required to run the SMORE pipeline except when used with additional programs such as Infernal or tRNAscan-SE (see the following subsection for details). The pipeline is written in Perl and Python, hence interpreters for both languages are required as well as gzip. Further details on versions are listed in Sec. 2.1, specifying software requirements.

A usual pipeline run consists of two steps: i) retrieving the clusters of genetic elements from the input data and ii) analysing the clusters and counting evolutionary events. Parts of the pipeline can be used independently from each other in order to repeat some processing steps. Please see **Section 4** for further details. To start a complete run of the SMORE pipeline, use:

```
SMORE bake --out OUTPATH --ref REFERENCE_SPECIES
--maf PATH_TO_MAF_FILES --genomes PATH_TO_GENOMEFOLDER
--newick NEWICK_TREE <MODE-OPTIONS>
```

Upper case terms in the command need to be replaced by the specific information needed. MAF files are multiple sequence alignments in MultiZ format. All MAF files should be loacted in the same folder. Newick tree is a tree in newick format. Only species that occur in the tree can be analysed. Each MAF alignment is based on a reference species. Please give the species identifier of the MAF reference species in the –ref option. The mode-options are options corresponding on the mode chosen for retrieving the input data. Further details in modes ar explained in Section 2.2 and Section 3. Following modes are available: (a) gene list mode, (b) loci list mode, (c) tRNAscan-SE mode and (d) Infernal mode.

Note that species identifier have to be the same in the MAF files, mode-dependent input data, newick tree and reference species!

The output will be written to the specified output directory (--out option). The output contains a list of genetic clusters as well as a phylogenetic tree where nodes are labeled with numbers of evolutionary events. For more detailed information, see **Section 5**. When giving input files and folders as input to the pipeline, please specify the complete absolute path, as files cannot be found without. Additionally, pathes to installed programs such as Perl, Python, Infernal, tRNAscan-SE must not be specified except if they are not stored in the environment path variable.

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2.1 Software requirements

SMORE is programmed in Python and Perl, hence it requires Python3 and Perl5 and gzip in order to reduce the usage of memory. Additionally, it requires a current version of Infernal [3] and tRNAscan-SE [4] in case genetic elements are automatically detected using a covariance model or using tRNAscan-SE. A current version of tRNAscan-SE can be downloaded at http://eddylab.org/software.html and a current version of Infernal at http://eddylab.org/infernal/. Our pipeline only requires the subprogram cmsearch of the Infernal program suite.

2.2 Mode overview

The SMORE pipeline includes several subcommands, that can be used independently depending on the current data set. Additionally, the subcommands SMORE prep and SMORE bake can be applied using different modes of pre-processing. Fig. 2 gives an overview of available modes. The modes will be explained in detail in the next section, Sec. 3 which will used the subcommand SMORE bake in order to explain the modes. All other subcommands will be explained in in Section 4.

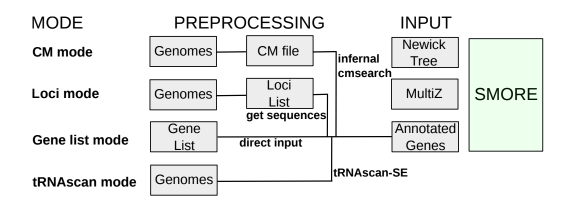


Figure 2: Overview of pre-processing modes of the SMORE pipeline. The pre-processing steps are part of the pipeline in order to retrieve the data needed to construct gene clusters. The construction of gene clusters will use the output of the pre-processing step which is automatically handed over.

Help pages for the pipeline can be called using one of the following commands:

```
SMORE --help
SMORE [subcommand] --help
```

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3 Input for pre-processing modes

The subcommands SMORE bake and SMORE prep can be used running different modes of how the input gene list is created, hence different sets of parameters are required for different modes of running. This section lists all the parameters sorted by usage mode.

3.1 General input

The following parameters are used in all modes and hence, needed for subcommands SMORE bake and SMORE prep:

option	description
obligatory parameter	
out -o OUTPATH	Directory where to write the output files. If the folder
	does not exist, it will be created.
optional parameter	
tool -t TOOLPATH	Directory where SMORE is located. This parameter can
	usually be omitted.
python PYTHON_PATH	Path to Python installation if the one specified in the
	path environment is not used.
perl PERL_PATH	Path to Perl installation if the one specified in the
	path environment is not used.
filter NUM	Percentage of low scoring MAF blocks to be discarded,
	NUM is between 0 and 1. This parameter is optional,
	default = 0.

3.2 Gene list mode

This mode does not run any pre-processing step but the user provided input will be directly used. This mode is used if only specific genes are included in the analysis that cannot be automatically retrieved by tRNAscan-SE or Infernal.

option	description
genes GENELIST	A list of genetic elements as input.

Table 3 specifies the format of the input gene list. The gene list should be tab-separated with one entry for each gene and columns should contain the information as shown in the

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second row of the table. The columns are specified by:

- 1. chr: chromosome,
- 2. start: start coordinate of genetic element in the chromosome,
- 3. end: end coordinate of genetic element in the chromosome,
- 4. spec: species identifier,
- 5. strand: + or -
- 6. type: type of genetic element e.g. Met when tRNA-Met,
- 7. pseudogene or not: true or false,
- 8. struc: secondary structure in dot-bracket notation,
- 9. seq: sequence,
- 10. comment: any comment by the user.

The third row of the table specifies if a column is obligatory (obl) or can be omitted (opt). If information in optional columns are omitted, the field has to be filled with 'NA'. For the case of optional sequence field in the gene list, please see the **Subsection 3.3**.

1	2	3	4	5	6	7	8	9	10
chr	start	end	spec	strand	type	pseudogene	struc	seq	comment
obl	obl	obl	obl	obl	opt	opt	obl	opt	opt

Table 3: This table defines the format of the gene list required as a user provided input when running SMORE in *gene list mode*. The second row provides information on which column is obligatory (obl) and which information is optional (opt). For optional information that is not provided, fields cannot be empty but have to be filled by 'NA'. Abbrevations are: chr: chromosome, spec: species, seq: sequence, struc: secondary structure.

Note that species identifier and chromosome identifier have to be the same in the MAF files, gene list, newick tree and reference species!

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3.3 Loci list mode

The *loci list mode* can be applied if only coordinates of elements are known. Given the corresponding genomes, SMORE will automatically retrieve the gene sequences that are needed for the analysis. Retrieving the sequences based on the provided coordinates and genomes is the only pre-processing done by the *loci list mode*.

option	description
genomes -g GENOMES	Folder with genomes of the species used to retrieve the
FOLDER	sequence of the genetic elements in the loci list based
	on their coordinates. Filenames should match species
	names in loci list files.
loci FILE	List with genetic elements without sequence.

The loci list is a tab-separated table with one entry for each gene. The format is the same as for the gene list but omits the last three columns, as shown in **Table 5**. The columns are specified as follows:

- 1. chr: chromosome,
- 2. start: start coordinate of genetic element in the chromosome,
- 3. end: end coordinate of genetic element in the chromosome,
- 4. spec: species identifier,
- 5. strand: + or -,
- 6. type: type of genetic element e.g. Met when tRNA-Met,
- 7. pseudogene or not: true or false.

The third row of the table specifies if a column is obligatory (obl) or can be omitted (opt). If information in optional columns are omitted, the field has to be filled with 'NA'.

Note that species identifier and chromosome identifier have to be the same in the MAF files, genome files, loci list, newick tree and reference species!

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1	2	3	4	5	6	7
chr	start	end	spec	strand	type	pseudogene
obl	obl	obl	obl	obl	opt	opt

Table 5: This table defines the format of the loci list required as a user provided input when running SMORE in *loci list mode*. The second row provides information on which column is obligatory (obl) and which information is optional (opt). For optional information that is not provided, fields cannot be empty but have to be filled by 'NA'. Abbrevations are: chr: chromosome, spec: species.

3.4 Infernal mode

The Infernal program suite provides several programs to search for genes with a specified covariance model (CM), to scan given genomes or create a covariance model for a set of training sequences. Our pipeline uses cmsearch in order to retrieve genetic events from a given genome based on a provided CM. Hence, Infernal can be used to create the CM in advance. For more details, we refer to the Infernal manual or publication.

option	description
cm -c CM	Covariance model file, input for Infernal.
genomes -g GENOMES	Folder with genomes of the species used as an input
FOLDER	to Infernal to scan the genomes for genetic elements
	specified by the CM. Filenames should match species
	names in MultiZ files.
incE NUM	Optional parameter for cmsearch (Infernal), e-value
	threshold.
incT NUM	Optional parameter for cmsearch (Infernal), bitscore
	threshold.
infernal PATH	Path to cmsearch, only needed if the one specified in
	the path environment is not used.
pseudo NUM	Optional parameter, bitscore threshold that defines
	pseudogenes.

Note that species identifier and chromosome identifier have to be the same in the MAF files, genome files, newick tree and reference species!

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$3.5 \quad tRNAscan-SE \ mode$

In bioinformatic applications, tRNAscan-SE [4] is a popular tool to retrieve tRNAs from a given genome. Hence, when in tRNAscan-SE mode, our pipeline automatically applies tRNAscan-SE on the given genomes. The tRNAscan-SE output files will be parsed and reformatted in order to retrieve all information needed to construct gene clusters. As mentioned in [2], tRNA genes are the best studied elements when working on concerted evolution. tRNA genes show a high turnover rate and frequent pseudogenization events.

option	description
genomes -g GENOMES	Folder with genomes of the species used as an input
FOLDER	to Infernal to scan the genomes for genetic elements
	specified by the CM. Filenames should match species
	names in MultiZ files.
trna	Option to activate the usage of tRNAscan-SE on the
	given genomes.
trnascan PATH	Path to tRNAscan-SE, only needed if it is not installed
	in the environment path variable.

Note that species identifier and chromosome identifier have to be the same in the MAF files, genome files, newick tree and reference species!

4 Subcommands

In order to repeat only some parts of the pipeline or have a more detailed look into intermediary files, parts of the pipeline can be used independently. This section gives an overview of all possible subcommands explaining input, output and options. **Fig. 3** gives a graphical overview of subcommands in comparison to the workflow.

The SMORE pipeline can be applied in one complete run but also splitted in several parts that allow the user to change and compare results created by different parameter sets. Additionally, the pipeline of subcommands consisting of SMORE prep, SMORE mix, SMORE roast and SMORE eat is able to process large amounts of data by splitting the set of clusters in disjoint subsets (SMORE mix), analyse each part individually (SMORE roast) and summarizing all results at the end (SMORE eat).

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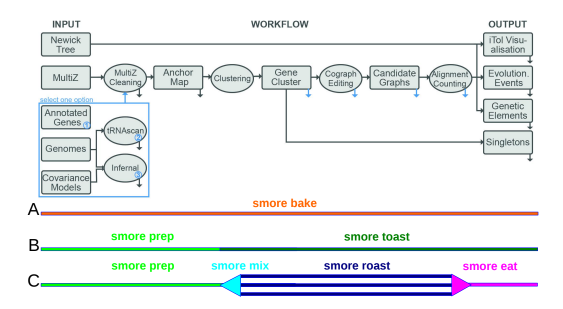


Figure 3: Overview of subcommands of the SMORE pipeline next to the workflow.

4.1 SMORE bake

The subcommand SMORE bake will run the complete pipeline for a given input. Different modes such as *gene list mode* or *Infernal mode* can be specified by the set of parameters and the input data. The output is saved in the specified output folder (option --out). In case the output folder does not exist, it will be created. As SMORE bake is a combination of the subcommands prep and toast, the parameter set is the same as for prep and toast. Thus, please see the tables in the **Subsection 4.2 and 4.3** for parameter descriptions. The help pages for SMORE bake is SMORE bake --help or SMORE bake -h.

4.2 SMORE prep

By applying SMORE prep only the first part of the pipeline will be executed. Hence, only input data concerning the creation of genetic clusters is required. This subcommand can be used to create a basic dataset that will be the input of following steps of the pipeline. Afterwards, succeeding subcommands can be applied with different parameter sets. Depending on the number of genetic elements and genome size, SMORE prep might take several a while to process the input data. The output will be saved in the specified output folder (option --out). The help page for SMORE prep is SMORE prep --help or SMORE prep -h.

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parameter	obl/opt	description
general		
out -o OUTPATH	obl	Directory where to write the output files. If the folder
		does not exist, it will be created.
maf MAF_FILES	obl	Directory where all MultiZ alignment files are located.
ref	obl	Name of the reference species as specified in the ${\tt MultiZ}$
REFERENCE_SPECIES		alignment files.
tool -t TOOLPATH	opt	Directory where SMORE is located. This parameter can
		usually be omitted.
python PYTHON_PATH	opt	Path to Python installation if the one specified in the
		path environment is not used.
perl PERL_PATH	opt	Path to Perl installation if the one specified in the
		path environment is not used.
filter NUM	opt	Percentage of low scoring MAF blocks to be discarded,
		NUM is between 0 and 1. This parameter is optional,
		default = 0.
gene list mode		
genes GENELIST	obl	A list of genetic elements as input.
Infernal mode		
cm -c CM	obl	Covariance model file, input for Infernal.
genomes -g	obl	Folder with genomes of the species used as an input
GENOMES FOLDER		to Infernal to scan the genomes for genetic elements
		specified by the CM. Filenames should match species
		names in MultiZ files.
incE NUM	opt	Parameter for cmsearch (Infernal), e-value thresh-
		old.
incT NUM	opt	Parameter for cmsearch (Infernal), bitscore thresh-
		old.
infernal PATH	opt	Path to cmsearch, only needed if the one specified in
		the path environment is not used.
pseudo NUM	opt	Bitscore threshold that defines pseudogenes.

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$tRNAscan ext{-}SE\ mode$			
genomes -g	obl	Folder with genomes of the species used as an input	
GENOMES FOLDER		to Infernal to scan the genomes for genetic elements	
		specified by the CM. Filenames should match species	
		names in MultiZ files.	
trna	obl	Option to activate the usage of tRNAscan-SE on the	
		given genomes.	
trnascan PATH	opt	Path to tRNAscan-SE, only needed if it is not installed	
		in the environment path variable.	
loci list mode			
genomes -g	obl	Folder with genomes of the species used to retrieve the	
GENOMES FOLDER		sequence of the genetic elements in the loci list based	
		on their coordinates. Filenames should match species	
		names in loci list files.	
loci FILE	obl	List with genetic elements without sequence.	

4.3 SMORE toast

The subcommand SMORE toast can be applied after SMORE prep. It will take SMORE prep output (option --prep) and analyse the given genetic clusters. The output will be saved in the specified output folder (option --out). As SMORE toast can be applied using different sets of parameters, applying it several times on the same input data can help estimating the best parameter set e.g. joining of clusters (--join) or similarity thresholds (--seq, --struc). By omitting verbose output, SMORE toast is able to process given input data very fast. Verbose output (if --verbose) or any combination of them (if --clus, --graph, --aln) will help to get a deeper insight into intermediary steps and how the clusters are composed, but will take a lot more time to process the data and more space capacities as all intermediary files are stored in the output folder. The help page for SMORE toast is SMORE toast --help.

parameter	obl/opt	description
out -o OUTPATH	obl	Directory where to write the output files. If the folder
		does not exist, it will be created.

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prep PATH	obl	Directory where SMORE prep output files are located.
newick NEWICK_FILE	obl	Phylogenetic tree in newick format including all species
		contained in the analysis. For automatic conversion of
		species identifiers, use parameter –id.
id ID_FILE	opt	In case species identifier in the tree and MultiZ files
		do not match, this file can be used to automatically
		convert species identifier. Format of the table: cur-
		rent_name_in_tree (tab) name_to_translate_to
join LEVEL	opt	Level of how to join the clusters. Levels are none, strict
		and relaxed. Default: relaxed. The level refers to the
		adjacency constraints of gene clusters, see Section 7
		for further details.
seqsim NUM	opt	Percentage of sequence similarity in order to define
		two sequences as orthologs. Number between 0 and 1,
		default: 0.9. Set to -1 in case the sequence similarity
		should be omitted for the analysis (hence, only check
		structure similarity).
strucsim NUM	opt	Percentage of structure similarity in order to define
		two structures as orthologs. Number between 0 and 1,
		default: 0.9 Set to -1 in case the structure similarity
		should be omitted for the analysis (hence, only check
		sequence similarity).
nomiss	opt	Using this option, the pipeline will omit the check
		for missing data. Hence, deletions in the tree won't
		distinguish between a deletion because of missing data
		or a 'real' deletion. Only useful when running on a
		large data set.
verbose	opt	Using this option, the pipeline will print all intermedi-
		ary files such as clusters, graphs and alignments. This
		will slow down the running time of the program but
		will give deeper insights into the data.
clus	opt	Using this option, the pipeline will print all intermedi-
		ary cluster files.

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graph	opt	Using this option, the pipeline will print all intermedi-
		ary graph files.
aln	opt	Using this option, the pipeline will print all intermedi-
		ary alignment files.
tool -t TOOLPATH	opt	Directory where SMORE is located. This parameter can
		usually be omitted.
python PYTHON_PATH	opt	Path to Python installation if the one specified in the
		path environment is not used.
perl PERL_PATH	opt	Path to Perl installation if the one specified in the
		path environment is not used.

4.4 SMORE mix

SMORE mix is designed to only run the joining of the clusters based on given parameters and SMORE prep output. It will return a list of the original clusters and another file listing clusters after joining. Additionally, SMORE mix can be used to handle large amounts of data. With option --max the maximal number of clusters can be specified that should be included in the pipeline's next processing step. SMORE mix will hence split the resulting clusters in several files that do not exceed the specified number. In order to help the user proceed to the next step of the pipeline (SMORE roast), the output will contain a command list, specifying the commands needed to call the next step with each part of the cluster list. The user can easily copy the commands and call the next step or modify the commands by adding parameters. Verbose output can be used with SMORE mix but might slow down running time of the program. The possible parameters for SMORE mix include parameters for SMORE roast, too, as the parameters will be used to create the command list. The help page for SMORE mix is SMORE mix --help.

parameter	obl/opt	description
out -o OUTPATH	obl	Directory where to write the output files. If the folder
		does not exist, it will be created.
prep PATH	obl	Directory where SMORE prep output files are located.

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species FILE	obl	File listing all species included in the analysis. Thus,
Species 1122		the file includes the species identifier, one per line. The
		file is created by SMORE prep and thus will be located
		in the SMORE prep output folder, named specieslist.
newick NEWICK_FILE	obl	Phylogenetic tree in newick format including all species
		contained in the analysis. For automatic conversion of
		species identifiers, use parameter –id.
id ID_FILE	opt	In case species identifier in the tree and MultiZ files
		do not match, this file can be used to automatically
		convert species identifier. Format of the table: cur-
		rent_name_in_tree (tab) name_to_translate_to
join LEVEL	opt	Level of how to join the clusters. Levels are none, strict
		and relaxed. Default: relaxed. The level refers to the
		adjacency constraints of gene clusters, see Section 7
		for further details.
max NUM	opt	This parameter can be used to specify the maximal
		number of clusters used in following steps of the analy-
		sis. If there are more clusters, the program will auto-
		matically split the data set and create a command list
		in order to run the next steps in parallel. The default
		value is 50000.
seqsim NUM	opt	Percentage of sequence similarity in order to define
•	•	two sequences as orthologs. Number between 0 and 1,
		default: 0.9. Set to -1 in case the sequence similarity
		should be omitted for the analysis (hence, only check
		structure similarity).
strucsim NUM	opt	Percentage of structure similarity in order to define
SOLUCSIM MON	opι	two structures as orthologs. Number between 0 and 1,
		,
		default: 0.9. Set to -1 in case the structure similarity
		should be omitted for the analysis (hence, only check
		sequence similarity).

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nomiss	opt	Using this option, the pipeline will omit the check
		for missing data. Hence, deletions in the tree won't
		distinguish between a deletion because of missing data
		or a 'real' deletion. Only useful when running on a
		large data set.
verbose	opt	Using this option, the pipeline will print all intermedi-
		ary files such as clusters, graphs and alignments. This
		will slow down the running time of the program but
		will give deeper insights into the data.
clus	opt	Using this option, the pipeline will print all intermedi-
		ary cluster files.
tool -t TOOLPATH	opt	Directory where SMORE is located. This parameter can
		usually be omitted.
python PYTHON_PATH	opt	Path to Python installation if the one specified in the
		path environment is not used.
perl PERL_PATH	opt	Path to Perl installation if the one specified in the
		path environment is not used.

4.5 SMORE roast

The subcommand SMORE roast is applied on the SMORE mix output. The program SMORE mix will output a list of commands on how to proceed. Thus SMORE roast is called based on these commands. However the user can change or add some parameters to the preprinted commands in order to adjust the program call. If there is more than one commands, it is important that all commands are executed with the same parameter set in order to obtain a homogenuous result at the end. If some of the commands are omitted, the final result will only include the data that was analysed. The next step of the pipeline SMORE eat will summarize the SMORE roast output into final results of the pipeline run. The resulting files of SMORE roast are given specified names to make sure that the next step of the pipeline will find all input data. Thus, file names should not be changed and files not be moved to another folder. Verbose output can be used with SMORE roast, also for just some of the commands but might slow down the running time of the program. The help page for SMORE roast is SMORE roast --help.

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parameter	obl/opt	description
out -o OUTPATH	obl	Directory where to write the output files. If the folder
		does not exist, it will be created.
in -i FILE	obl	Output file of SMORE mix including the list of clusters.
species FILE	obl	File listing all species included in the analysis. Thus,
		the file includes the species names as given in the
		newick tree and prep output, one species identifier per
		line.
newick NEWICK_FILE	obl	Phylogenetic tree in newick format including all species
		contained in the analysis. For automatic conversion of
		species identifiers, use parameter –id.
id ID_FILE	opt	In case species identifier in the tree and MultiZ files
		do not match, this file can be used to automatically
		convert species identifier. Format of the table: cur-
		rent_name_in_tree (tab) name_to_translate_to
seqsim NUM	opt	Percentage of sequence similarity in order to define
		two sequences as orthologs. Number between 0 and 1 ,
		default: 0.9. Set to -1 in case the sequence similarity
		should be omitted for the analysis (hence, only check
		structure similarity).
strucsim NUM	opt	Percentage of structure similarity in order to define
		two structures as orthologs. Number between 0 and 1 ,
		default: 0.9. Set to -1 in case the structure similarity
		should be omitted for the analysis (hence, only check
		sequence similarity).
nomiss	opt	Using this option, the pipeline will omit the check
		for missing data. Hence, deletions in the tree won't
		distinguish between a deletion because of missing data
		or a 'real' deletion. Only useful when running on a
		large data set.

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verbose	opt	Using this option, the pipeline will print all intermedi-
		ary files such as clusters, graphs and alignments. This
		will slow down the running time of the program but
		will give deeper insights into the data.
graph	opt	Using this option, the pipeline will print all intermedi-
		ary graph files.
aln	opt	Using this option, the pipeline will print all intermedi-
		ary duplication alignment files.
tool -t TOOLPATH	opt	Directory where SMORE is located. This parameter can
		usually be omitted.
python PYTHON_PATH	opt	Path to Python installation if the one specified in the
		path environment is not used.
perl PERL_PATH	opt	Path to Perl installation if the one specified in the
		path environment is not used.

4.6 SMORE eat

The subcommand SMORE eat is applied on output of SMORE roast. Here, the output folder of SMORE roast is the input to SMORE eat which will take into account all files in the folder with having specific names as given by SMORE roast. This last part of the program will then summarize all the information such that evolutionary events can be counted and numbers added to the tree. The help page for SMORE eat is SMORE eat --help.

parameter	obl/opt	description
out -o OUTPATH	obl	Directory where to write the output files. If the folder
		does not exist, it will be created.
prep PATH	obl	Directory where SMORE prep output files are located.
mix PATH	obl	Directory where SMORE mix output files are located.
roast PATH	obl	Directory where SMORE roast output files are located.
newick NEWICK_FILE	obl	Phylogenetic tree in newick format including all species
		contained in the analysis. For automatic conversion of
		species identifiers, use parameter –id.

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id ID_FILE	opt	In case species identifier in the tree and MultiZ files
		do not match, this file can be used to automatically
		convert species identifier. Format of the table: cur-
		rent_name_in_tree (tab) name_to_translate_to.
nomiss	opt	Using this option, the pipeline will omit the check
		for missing data. Hence, deletions in the tree won't
		distinguish between a deletion because of missing data
		or a 'real' deletion. Only useful when running on a
		large data set.
tool -t TOOLPATH	opt	Directory where SMORE is located. This parameter can
		usually be omitted.
python PYTHON_PATH	opt	Path to Python installation if the one specified in the
		path environment is not used.
perl PERL_PATH	opt	Path to Perl installation if the one specified in the
		path environment is not used.

5 Output

All output files are located in the specified folder, given with option --out or -o. The output consists of multiple files and depends on the subcommand(s) used. The following subsections describe the output files in more details. For an example, see **Section 6**.

5.1 General output

Each of the SMORE subcommands will provide an output file that summarizes the current run and provides information about the status of the program during runtime. Additionally, each subpart of the pipeline will create a file where all errors are collected. If the file is empty, no errors occured.

output file	description
Summary.txt	This file gives a short overview of the program's run
	specifying parameter and running time. This file can
	also be used to check the status of the program while
	running as it gives information based on the current
	steps of the pipeline.

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$errors_SUBCOMMAND$	For each part of the SMORE pipeline, there is a file
	giving errors that happened during the run. If no
	errors occured, the file is empty.

5.2 Output of completed pipeline run

As specified in **Section 2.2** by **Fig. 2**, there are several ways on how to do a complete run of the SMORE pipeline. The options are:

- 1. SMORE bake
- 2. SMORE prep and SMORE toast
- 3. SMORE prep and SMORE mix and SMORE roast and SMORE eat

This section will describe the output after running one of the three options mentioned above. For output of intermediary steps, please see one of the following subsections, depending on the subcommand. The table lists the output files including final results. For intermediary files, see following tables.

output file	description
OutTree.txt	The resulting tree in newick format with numbers at
	the nodes given in brackets. This format can be used
	to visualize the tree with newick compatible programs.
geneticEvents.txt	File listing all genetic events counted during the anal-
	ysis. The numbers are sorted by event and node of
	the tree. The file includes a event called 'Other'. This
	will give the difference of genetic elements between the
	total amount and the elements used in the analysis.
	For a successful run of the pipeline, the numbers should
	be 0.

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$data_iTOL$	Folder giving files that can be uploaded to <i>itol.embl.de</i> [5]. The file called F0tree.txt is uploaded at http://itol.embl.de/upload.cgi. The remaining files can be added using drag and drop into the browser window. This will result in a interactive visualization of the resulting tree. A legend is added automatically. All nodes in the tree will have unique names, thus some nodes might have names such as 'innerNode0' because it was added automatically. For an explanation of how	
	to visualize the final output, see Section 5.7 .	
$allClusters_original.txt,$	These two files contain lists of clusters showing which el-	
allClusters_joined.txt	ements are contained together in one cluster before and	
	after joining. A line starting with > is followed by the	
	cluster number. Each line after the cluster ID contains	
	one genetic element of the cluster having the following	
	format: chromosome species_elementID start	
	end strand left_block right_block structure	
	sequence type pseudogene comment. In case some	
	information is not provided, the field is filled with	
	'NA'.	
$list_cographs.txt,$	These files contain statistics about graphs that were	
$list_noncographs.txt$	cographs from the beginning or had to be edited in	
	order to become a cograph. The tables list number	
	of nodes, number of edges, number of corrected edges	
	(see Section. 7) and density of the graphs. Entries	
	with 0 are based on graphs that have no nodes or not	
	more than 3 nodes, as all graphs below 4 nodes are	
	cographs and need not to be checked.	
remoldings.txt,	These files contain genetic elements that (a) have highly	
in remoldings.txt	similar sequences but different types or (b) have the	
	same types but clearly distinct sequences.	

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all Types.txt,	These two files list the different types of genetic ele-	
all PseudoTypes.txt	ments for all species and functional or pseudogenized	
	genes. This can be used to analyse the distribution of	
	different types of genetic elements.	

5.3 Verbose output

The verbose version of SMORE toast will output three additional files (if --verbose) or any combination of them (if --clus, --graph, --aln) for each cluster, named with left and right anchor numbers to match all three files. They will be in three different folders: cluster, graph and duplication_alignment. The files contain the specific structures of the cluster in each step of the analysis and can be used to gain a deeper insight into the data.

output file	description	
GENECLUSTER.clus	The files consist of a list of genetic elements being	
	part of this cluster. There is one entry for each	
	genetic element in the cluster. The format for	
	an entry is a tab-separated line with following	
	fields: chromosome species_elementID start	
	end strand left_block right_block structure	
	sequence type pseudogene comment. In case some	
	information is not provided, the field is filled with	
	'NA'.	
GENECLUSTER.edli	Files showing the graph structure of a gene cluster using	
	a weighted edge list. The graphs are complete, thus	
	there exists an edge between each pair of nodes. Hence,	
	there is one entry for each edge in the graph. The	
	entry is a tab-separated line with format: node1 node2	
	seq_sim struc_sim. The nodes are genetic elements	
	and the weights are the sequence similarity score and	
	structure similarity score (if structure is available).	

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The alignment files show the results of the duplication alignments for each cluster. Hence, for each species in the cluster, there is a sequence of genetic elements. Those elements are represented by single letters. If the elements similarity is above the thresholds, they share the same letter. The duplication alignment are a Needleman-Wunsch alignment including the possibility to detect duplications. Duplications are represented by '~', gaps by '-'. For each pair of species in a cluster, there is an entry consisting of the alignment and the

5.4 Output SMORE prep

The subcommand SMORE prep is the first part of the pipeline, sorting genetic elements in between genomic anchors in order to create gene clusters. Hence, SMORE prep will output data in three different subfolders in the output folder:

alignment score.

- 1. genes: This data is needed to continue the analysis of gene clusters. The folder contains a file in bed format for each of the species. The files contain a tabsaparated entry line for each genetic element. The format is as follows: chromosome species_elementID start end strand left_anchorID right_anchorID structure sequence type pseudogene comment. In case some information is not provided, the field is filled with 'NA'. When running Infernal as a pre-processing step, the comment section in the table will contain the Infernal score for each entry.
- 2. bed: This data is not used any further in the analysis but might be useful when having a deeper look into the results. The folder contains a gzipped file for each species. The files contain information about the genomic anchors retrieved from the MultiZ files. For each genomic anchor, there is one tab-separated line with the following format: chromosome species_anchorID start end strand ID_of_leftadjacent_anchor ID_of_rightadjacent_anchor. In case no anchor could be detected, the field is filled with 'None'.
- 3. **temp**: The files in this folder are needed in a later step of the analysis when deletions in the tree are clustered by cause, hence either missing data or deletion of the genetic

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elements. The folder contains a gzipped file for each species containing one entry for each genomic anchor. The information differs from the one in the bed folder as it contains scores for each of the anchors. The format is a tab-separated line for each anchor with chromosome species_anchorID anchorStart anchorLength strand refSpecies anchorScore. The 'refSpecies' field will contain 'True' if the species is the reference species, and 'False' if not.

5.5 Output SMORE mix

The subcommand SMORE mix reads SMORE prep output as its input and summarizes gene clusters. The program will join clusters based on the specified parameter (--join) and in case the number of cluster is higher than the specified maximal number (--max), it will split the set of clusters. In this way, the next step of the pipeline can be started on several subsets of clusters which will speed up the overall running time. Hence, SMORE mix will create a folder called clusterlists containing list of gene clusters. If all the lists are combined, the original set of clusters is retrieved. Additionally, SMORE mix will output a file called commandlist.txt. This file consists of a list of commands calling SMORE roast for each of the cluster lists. Hence, the user only needs to copy-paste the commands to execute or add or edit some parameters before running the commands. Additionally, SMORE mix creates output files listed in the following table.

output file	description	
species list	This files contains the species identifier as they appear	
	in the analysis, one line for each species.	
nones.txt, pseunones.txt	Each genetic element that cannot be set in between	
	genomic anchors, the anchor identifier is specified as	
	'None'. After starting the pipeline, all genetic elements	
	with 'None'-anchors will be excluded from the analsis	
	and written in a separate file.	

5.6 Output SMORE roast

The subcommand SMORE roast will use SMORE mix output, hence a cluster list. In case, SMORE roast is applied in parallel to several subdivided parts of the original list of clusters, SMORE roast will output intermediary files for each run that can be summarized running SMORE eat. Hence, SMORE roast output files should stay in the specified folder and not

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be renamed or moved. The following table specifies the output files. The name of the corresponding cluster list will be added to the filenames in order to have unique names in case SMORE roast is started several times. For further explanations on event counts, see Section 7.

output file	description	
singletons, pseusingletons	Lists counts for singletons or pseudogene singletons (if	
	present) for each species.	
matches, pseudomatches	Lists counts for matches or pseudogene matches (if	
	present) for each species.	
insertions, pseudoinsertions	Lists counts for insertions or pseudogene insertions (if	
	present) for each species.	
duplications	Lists counts for duplications for each species.	
delCheck, pseudelCheck	Lists counts for deletions or pseudogene deletions (if	
	present) for each species. The deletions might be	
	caused due to missing data or evolutionary deletion of	
	genes. This will be checked during SMORE eat.	
remoldings, inremoldings	Lists counts for remoldings or inremoldings (if present)	
	for each species.	
errors	For each run of SMORE roast there will be a file listing	
	errors occured during the run. If the file is empty, no	
	errors occured.	
$list_cographs,\ list_noncographs$	Lists cographs and non-cographs with identifier, node	
	number, edge number, corrected edge number and	
	density for each graph.	

5.7 Output visualization

After a complete successful run of the pipeline, the specified output folder will contain a subfolder called $data_iTOL$. This folder contains files that are formatted in a way such that after uploading them on the iTOL webpage, the resulting phylogenetic tree will be displayed in an interactive environment. Hence, the tree can be edited further and downloaded in several formats. The filenames are numbered, F0 to F8, which shows the recommended order in which they are uploaded. Changing the order does not cause any problems, except for file F0, which needs to be uploaded first, as it contains the underlying tree structure.

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The following steps have to be done to visualize the tree:

- After the folder data_iTOL is created and contains files F0 to F8, open the iTOL webpage in a browser, hence either open http://itol.embl.de and click on Annotate or directly open http://itol.embl.de/upload.cgi.
- 2. Click on *Browse* and choose the file *F0tree.txt* from the folder *data_iTOL*. After clicking on *Upload* the tree structure is displayed in the browser.
- 3. The remaining files from the folder data_iTOL can be added to the tree by using drag and drop on the files. Following the order specified by the numbering of the files is recommended but not mandatory.
- 4. Using the interactive browser tool iTOL, colors and labels can be changed, as well as sizes of branches and distances between nodes. The tool provides several formats for downloading the tree.
- 5. Numbers in the tree are either written at the leaves or at inner edges of the tree. Numbers at the leaves are specific for the species whereas numbers at inner edges refer to all species that are 'below' of 'after' that edge on the way from the root to the leaves. Numbers are color-coded and a legend is provided. Colors can be changed using iTOL.
- 6. In order to display the tree correctly, every node will have a label. If the label was not present in the original tree, the nodes will receive a name during the pipelines run. Hence, some nodes will have identifier such as 'InnerNode0'. These can of course be deleted.
- 7. For examples on how the trees look like, please see **Section 6**.

6 Examples

This section will show some examples of how to start the pipeline and how to read and visualize the output files.

6.1 SMORE bake with tRNAscan-SE

The SMORE pipeline will be started using six primate species and tRNAscan-SE in order to obtain counts of evolutionary events for tRNAs withing primate species. The species

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minoidae) Catarrhini;.

identifier have to be the same for the genomes, MultiZ files, reference species and in the newick tree. The data of this example was published in [2]. The input data such as MultiZ files were downloaded from http://hgdownload.cse.ucsc.edu/downloads.html. Genomes in fasta format and phylogenetic tree in newick format can be downloaded from UCSC or NCBI. The following species are included in the analysis.

species	abbreviation
Homo sapiens	hg38
Pan troglodytes	panTro4
Gorilla gorilla	gorGor3
Pongo abelii	$\mathrm{ponAbe2}$
Nomascus leucogenys	nomLeu3
Macaca mulatta	rheMac3

When giving input files and folders as input to the pipeline, please specify the complete absolute path, as files cannot be found without. Additionally, pathes to installed programs such as Perl, Python, Infernal, tRNAscan-SE must not be specified except if they are not stored in the environment path variable. The pipeline is called as follows:

```
SMORE bake --out OUTPATH/Output_smorebake --ref hg38
--maf PATH_TO_MAF_FILES --genomes PATH_TO_GENOMEFOLDER
--newick tree_6primates.newick --trna
```

Uppercase names have to be changed based on the system where the pipeline is executed. Fig. 4 shows the tree given as an input to SMORE bake displayed using iTOL. The newick

As described in **Section 5.7**, the interactive tool iTOL can be used to edit the tree and adapt features such as size, colors, labels. The resulting slightly edited tree of the SMORE bake run on six primate species and tRNAs is displayed in **Fig. 5**.

Further output files are a file called Summary.txt which includes status reports about the program's execution and statistics about the number of clusters, the size of clusters and information about graph structures and the number of corrected graphs.

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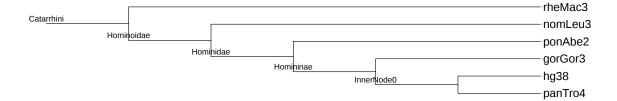


Figure 4: Phylogenetic tree displayed by iTOL as given as an input to SMORE bake. As for the analysis, all nodes have to have labels, the program will automatically name nodes without label. Hence, one of the inner nodes in the tree is called *InnerNode0*.

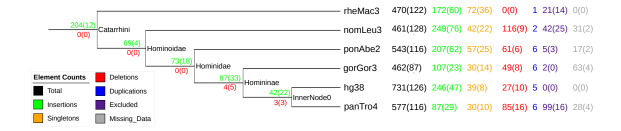


Figure 5: Resulting phylogenetic tree from the SMORE bake run on six primate species using tRNAscan-SE displayed by iTOL.

For the current example we get the following statistics:

```
Number of species: 6
Number of different element types: 24
Number of original clusters: 1926
Average number of elements of original clusters: 1.93
Number of joined clusters: 1038
Average number of elements per cluster in joined cluster: 3.58

Information on graph structures and corrections:
Number of cographs 650, with on average
3.74 nodes, 4.78 edges and a density of 0.82.
Number of non-cographs 5, with on average
8.40 nodes, 21.80 edges and a density of 0.28.
All non-cographs were corrected to obtain a cograph structure.
```

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Further output files, called allTypes.txt and allPseudoTypes.txt can be used to visualize the distribution of element types and species in the data set. For each pair of species and element type, there is one entry in the files giving the number of elements with this combinations. Fig. 6 shows the distribution of types within active and pseudogenized tRNA genes.

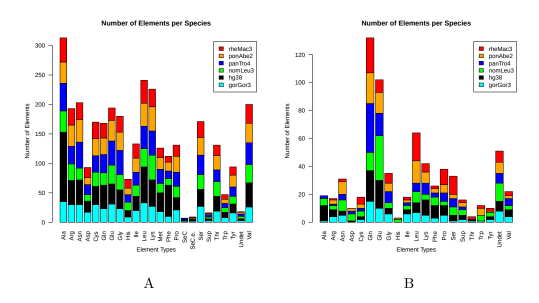


Figure 6: The plots show the distribution of tRNA types in the analysis based on six primate species. (A) shows the distribution of active tRNA genes whereas (B) shows the distribution of pseudogene types. The types of genes are identified using tRNAscan-SE.

6.2 SMORE bake with loci list and gene list

The calls for SMORE bake with loci list and gene list are very similar, and thus explained both here. A loci list is a list which contains information about genetic elements such as chromosome, species, strand, start and end coordinates. In order to apply similarity thresholds, our pipeline needs the genomic sequences and/or secondary structures of the elements. Hence, the loci list will be extended by the sequences taken from the species' genomes.

The call for running SMORE bake with a loci list as input looks as follows:

```
SMORE bake --out OUTPATH/Output_smorebake --ref hg38

--maf MAF_FILES_FOLDER --genomes GENOMEFOLDER

--newick tree_6primates.newick --loci locilist.tsv
```

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Calling SMORE bake with a gene list has a similar call but the genomes are not needed:

```
SMORE bake --out OUTPATH/Output_smorebake --ref hg38
--maf MAF_FILES_FOLDER
--newick tree_6primates.newick --gene genelist.tsv
```

As an example, the loci list with tRNAs in hg38 looks as follows:

```
chr1 16725566 16725638 hg38 + Val T
chr1 16727285 16727355 hg38 + Gly F
chr1 16860198 16860270 hg38 + Val T
chr1 16861921 16861991 hg38 + Gly F
chr1 16872583 16872654 hg38 + Glu F
```

The first column specifies the chromosome, the next two columns are start and end coordinates of the gene. The fourth and fifth columns specify the species and the strand. The last two columns show the type of the gene and if the gene is considered to be a pseudogene (T) or not (F).

The following shows the same entries as before, but here as a **gene list** input (sequences and structures shortened):

The gene list additionally includes a column specifying the secondary structure in dotbracket notation and a column for the sequence. The last column of the gene list is a comment column which might be useful for user-created list that only contain very specific genes.

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6.3 SMORE bake with Infernal

In order to apply SMORE bake using the Infernal option, a covariance model has to be specified using option --cm. The call for running SMORE bake with a loci list as input looks as follows:

```
SMORE bake --out OUTPATH/Output_smorebake --ref hg38

--maf MAF_FILES_FOLDER --genomes GENOMEFOLDER

--newick tree_6primates.newick --cm trna.cm
```

Ready-to-use covariance models can be downloaded at http://rfam.xfam.org/. Otherwise, covariance models can be created based on sample sequences using Infernal.

6.4 SMORE prep and ..

As depicted in Fig. 3 showing the different combinations of subcommands, it is possible to run parts of the pipeline independently from each other. The first part, including the pre-processing step, sorts the genetic elements inbetween the genomic anchors and thus, defines gene clusters. This part is executed when calling SMORE prep. As explained in Section 5.4, SMORE prep output consists of three different folders. This data forms the basis for further analysis steps using SMORE. Hence, subsequent steps of the pipeline can be called several times with different parameters in order to have a comparison when using distinct similarity thresholds or levels of joining clusters. Calling SMORE prep is similar as calling SMORE bake except that the option for the phylogenetic tree is not needed within SMORE prep. A call for SMORE prep with tRNAscan-SE is written as:

```
SMORE prep --out OUTPATH/Output_smoreprep --ref hg38 --trna
--maf PATH_TO_MAF_FILES --genomes PATH_TO_GENOMEFOLDER
```

For further details on SMORE prep parameter, see Section 4.2.

6.4.1 .. SMORE toast

Given SMORE prep output, the subcommand SMORE toast can be started easily by referring to SMORE prep output with --prep option. Hence, a simple SMORE toast call will look like:

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```
SMORE toast --out OUTPATH/Output_smoretoast
--prep OUTPATH/Output_smoreprep
--newick tree_6primates.newick
```

The SMORE toast has several additional optional parameter, that will change the outcome of the analysis, such as similarity thresholds and the level of joining clusters. Other options will increase the number of output files such that the user can have a closer look into the intermediary data. While more verbose output will increase running time of the program, other options such as skipping checks for deletions will decrease the running time. Please see Section 4.3 for more details on parameters.

6.4.2 .. SMORE mix, roast, eat

This combination of subcommands is mostly for larger data sets or if the user wants to run several test runs with different combinations of parameters. Hereby, the user can check intermediary files without running the whole pipeline and repeat steps directly. In case of a large data set, SMORE mix can split the data. This will be shown in the following toy example based on the tRNA data of six primate species as above.

Given the SMORE prep output, we now start SMORE mix. In order to reduce the running time of the next step, we want to have at most 400 clusters in each instance of execution. Hence, we call SMORE mix with the following options:

```
SMORE mix --out OUTPATH/Output_smoremix --prep OUTPATH/Output_smoreprep
--newick tree_6primates.newick
--max 400 --species OUTPATH/Output_smoreprep/specieslist
```

The parameter --species requires a file with species identifiers, one in each line. This file is created by the SMORE prep run and will be located in SMORE prep output files.

SMORE mix output include a file called commandlist and a subfolder called clusterlists which contains the complete cluster list divided into several files. The commands in the command list can be copied and directly used to call the next step of the pipeline SMORE roast. For our example, the list of clusters was divided into three sublists, hence the command list lists the following commands:

```
(1)SMORE roast --tool TOOLPATH/SMORE --out OUTPATH/Output_smoreroast
--python /usr/bin --perl /usr/bin -s 0.8 -p 0.8
--in OUTPATH/Output_smoremix/clusterlists/clusList_part0.txt
```

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```
--newick tree6species.newick
--species OUTPATH/Output_smoreprep/specieslist

(2)SMORE roast --tool TOOLPATH/SMORE --out OUTPATH/Output_smoreroast
--python /usr/bin --perl /usr/bin -s 0.8 -p 0.8
--in OUTPATH/Output_smoremix/clusterlists/clusList_part1.txt
--newick tree6species.newick
--species OUTPATH/Output_smoreprep/specieslist

(3)SMORE roast --tool TOOLPATH/SMORE --out OUTPATH/Output_smoreroast
--python /usr/bin --perl /usr/bin -s 0.8 -p 0.8
--in OUTPATH/Output_smoremix/clusterlists/clusList_part2.txt
--newick tree6species.newick
```

As it can be seen, the calls are exactly the same except for the cluster list file. Even though the output folder is the same, output files will have unique names by including the name of the corresponding cluster list. It is important, that output files of all SMORE roast runs that belong to the same original cluster list are located in the same folder as the next step of the pipeline, SMORE eat will summarize all those files. Hint: The output folders of SMORE roast and SMORE mix can be the same as the next step, SMORE eat will need the locations of the folders.

After running SMORE roast for all clusters lists, SMORE eat will summarize all output files and output the final results such as the phylogenetic tree with event counts, see:

```
(1)SMORE eat --out OUTPATH/Output_smoreeat

--prep OUTPATH/Output_smoreprep

--mix OUTPATH/Output_smoremix

--roast OUTPATH/Output_smoreroast

--newick tree6species.newick

--species OUTPATH/Output_smoreprep/specieslist
```

As the example data is the same as in all other examples above, the output, e.g. the phylogenetic tree, will be the same as above.

7 Theory

This section will explain some theoretical parts that might be needed to understand the internal processes of SMORE. For further details and references, we refer to the publication [2].

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7.1 Graph representation

As explained in the corresponding publication [2], graph structures are used to represent orthology relations. Hence, nodes in the graphs are genetic elements. There is an edge between two nodes if the corresponding genes are orthologs (i.e. fulfill the similarity thresholds and genes are from different species, hence orthologs). I was shown that a valid orthology relation should have a cograph structure [6] i.e., it must not include a path P_4 on four vertices as an induced subgraph. As input data might contain errors or there are some outliers in the distribution of similarity thresholds, the cograph structure might not always be given. In order to obtain a valid orthology relation, non-cographs are corrected by editing as less as possible edges. See **Fig. 7** for an example.

Additionally, it is useful to know how graph structures look like on average. Hence, number of nodes and edges as well as densities are collected and an averaged value is contained in the output. The density of a graph usually acts as an indicator for the number of edges in comparison to the number of nodes. Given an orthology graph with only edges from the same species (paralogs), no edges will be drawn. Hence, even though genes are very similar, the density of the graph will be zero. Hence, for density calculations, we use a *corrected* number of edges, which only takes into account similarity thresholds and not the fact if genes are from the same species or not. This corrected edge number is included in cograph and non-cograph list to give a better intuition about the graph structure.

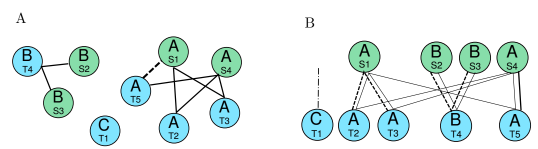


Figure 7: Example of the graph G for a cluster consisting of two groups of orthlogous elements in two species S and T (A). Thick edges indicate above-threshold sequence similarity. The dashed edge, which was included initially must be inserted to correct G: otherwise T5-S4-T3-S1 would form a P_4 . Modified Needleman-Wunsch alignment for graph G (B). The inserted edge to correct for a cograph is now part of the thick edges showing the orthology relation. The alignment will remove crossing edges of the orthology graph and detect duplications (dashed edges). The edge attached to node T_1 indicates a deletion in species S as there is no target node for this edge.

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After correction graphs for cograph structure, clusters undergo the duplication alignment. Here, a sequence of genetic elements for each species in the cluster is created. Each genetic element is represented by one letter. If two elements fulfill the similarity thresholds, they are assigned the same letter, for all of the species and inside a species, too. The order of elements in a cluster sequence depends on the order of genes in the corresponding genome. Given two cluster sequences from different species, an alignment is created such that insertions and deletions can be detected. Additionally, the modified alignment algorithm allows to align one element in the first species to more than one element in the other species if all element are of the same type and adjacent to each other when in the same species. This is shown in Fig. 7 B.

7.2 Counting

Given a set of genetic elements per species, they will be assigned to an evolutionary event in the phylogenetic tree. After collecting the information of all genetic elements, the assignments will be counted and the corresponding evolutionary event together with the number of elements will be inserted in the tree. The following evolutionary events can be found in the resulting tree (for a figure showing the events, we refer to **Section 6**):

- *Total*: This shows the total number of elements that were given as an input or contained in the resulting gene list based on the covariance model.
- Excluded: These genetic elements could not be assigned in between two genetic anchors. Hence, they are excluded from the analysis. In order to fit the total number of elements per species, they are still included in the resulting tree.
- Singleton: These elements appear as the only elements in the defined cluster. Additionally, no orthologuous cluster was detected. Hence, it is not possible to set those elements in the evolutionary context, so they are counted for each species separately at the leaves of the tree.
- Duplication: Duplications are detected by applying the modified list alignments. Thus, they are in a cluster together with a second highly similar element (or copy) in the same species. Other species might include only one copy of this element. Thus, the additional element is counted as a duplication.
- Insertion: Given orthologuous elements in several species, the element is assumed to have appeared in a common ancestor.

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- Deletion: An element is said to be deleted if several related species have an orthologuous version of it but it cannot be found in the orthologuous cluster of the current species.
- Missing Data: In case of a deletion but the absence of orthologuous anchors in the
 current species, this is count as missing data as this could also be induced by low
 quality of the multiple sequence alignments or underlying sequences.
- Pseudogene: Pseudogenizations are not explicitly stated in the tree but within each evolutionary event, we give a number (in parentheses) that gives the number of pseudogenes counted at this node of the tree. Hence, pseudogenes are included in the analysis but counted separately.

8 Help pages

The SMORE pipeline can be started using general options with:

```
SMORE [general options]
```

or:

```
SMORE <subcommand> [options]
```

For both cases, calling --help as option, the help pages will be displayed. Additionally, there is an extra help page explaining output files that can be called using --helpout.

9 Contact

When using SMORE, please cite

```
SMORE: Synteny Modulator Of Repetitive Elements.

Berkemer, S.J. and Hoffmann, A. and Murray, C.R. and Stadler, P.F.

MDPI Life, 2017 (submitted)
```

In case of further questions, please contact:

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