sangeranalyseR

Oct 06, 2022

Contents

1	Why	sangera	nalyseR	1		
2	Main	feature	S	3		
3	What sangeranalyseR doesn't do					
4	User Manual					
5	User support					
6	Key	Key contributors				
7	Docu	mentati	on	13		
	7.1	Installa	tion	13		
		7.1.1	System requirements	13		
		7.1.2	Install from Bioconductor	13		
		7.1.3	Install the development version	14		
		7.1.4	Where to go from here?	14		
	7.2	Ouick S	Start Guide	15		
		7.2.1	Super-Quick Start (3 lines of code)	15		
		7.2.2	Step 1: Prepare your input files	15		
		7.2.3	Step 2: Load and analyse your data	15		
		7.2.4	Step 3 (optional): Explore your data	16		
		7.2.5	Step 4: Output your aligned contigs	16		
		7.2.6	Step 5 (optional): Generate an interactive report	16		
		7.2.7	A Reproducible Example	16		
	7.3		ers Guide	18		
		7.3.1	Step 1: Preparing your input files	19		
		7.3.2	Step 2: Loading and analysing your data	20		
		7.3.3	Step 3: Exploring your data with the Shiny app	20		
		7.3.4	Step 4: Outputting your aligned contigs	21		
		7.3.5	Step 5: Generating an interactive report	21		
		7.3.6	What's next ?	22		
	7.4		ced User Guide - SangerRead (AB1)	22		
	/	7.4.1	Preparing SangerRead AB1 input	23		
		7.4.2	Creating SangerRead instance from AB1	23		
		7.4.2	Visualizing SangerRead trimmed read	25 25		
		1.4.3		23		

	7.4.4	Updating SangerRead quality trimming parameters
	7.4.5	Writing SangerRead FASTA file (AB1) 25
	7.4.6	Generating SangerRead report (AB1) 26
	7.4.7	Code summary (<i>SangerRead</i> , ab1)
7.5	Advanc	ced User Guide - SangerContig (AB1)
	7.5.1	Preparing SangerContig AB1 inputs 28
	7.5.2	Creating SangerContig instance from AB1
	7.5.3	Updating <i>SangerContig</i> quality trimming parameters
	7.5.4	Launching SangerContig Shiny app
	7.5.5	Writing <i>SangerContig</i> FASTA files (AB1)
	7.5.6	Generating <i>SangerContig</i> report (AB1)
	7.5.7	Code summary (<i>SangerContig</i> , AB1)
7.6	Advanc	27 wed User Guide - SangerAlignment (AB1)
	7.6.1	Preparing SangerAlignment AB1 input
	7.6.2	Creating SangerAlignment instance from AB1
	7.6.3	Updating <i>SangerAlignment</i> quality trimming parameters
	7.6.4	Launching SangerAlignment Shiny app 56
	7.6.5	Writing SangerAlignment FASTA files (AB1)
	7.6.6	Generating SangerAlignment report (AB1)
	7.6.7	Code summary (SangerAlignment, AB1)
7.7	Advanc	xed User Guide - SangerRead (FASTA)
	7.7.1	Preparing SangerRead FASTA input
	7.7.2	Creating SangerRead instance from FASTA
	7.7.3	Writing SangerRead FASTA files (FASTA)
	7.7.4	Generating SangerRead report (FASTA)
	7.7.5	Code summary (<i>SangerRead</i> , fasta)
7.8		zed User Guide - SangerContig (FASTA)
	7.8.1	Preparing SangerContig FASTA input
	7.8.2	Creating <i>SangerContig</i> instance from FASTA
	7.8.3	Writing <i>SangerContig</i> FASTA files (FASTA)
	7.8.4	Generating <i>SangerContig</i> report (FASTA) 80
	7.8.5	Code summary (<i>SangerContig</i> , FASTA)
7.9		ced User Guide - SangerAlignment (FASTA)
,	7.9.1	Preparing SangerAlignment FASTA input
	7.9.2	Creating <i>SangerAlignment</i> instance from FASTA
	7.9.3	Writing SangerAlignment FASTA files (FASTA) 89
	7.9.4	Generating SangerAlignment report (FASTA)
	7.9.5	Code summary (<i>SangerAlignment</i> , FASTA)
7.10		92
	7.10.1	What is a regular expression?
	7.10.2	How to deal with secondary peaks
	7.10.3	How to work with FASTA files for input
7.11		fanual (functions)
/.11	7.11.1	SangerRead Constructor Parameters
	7.11.2	SangerContig Constructor Parameters
	7.11.3	SangerAlignment Constructor Parameters
7.12		ntly Asked Questions
	7.12.1	
7.13		sion
7.14		98
7.14		t
7.15		98
,.10	1	Inside help test
	1.10.1	

Why sangeranalyseR

sangeranalseR is an R package that provides fast, flexible, and reproducible workflows for assembling your sanger seuqencing data into contigs.

It adds to a list of already widely-used tools, like Geneious, CodonCode Aligner and Phred-Phrap-Consed;. What makes it different from these tools is that it's free, it's open source, and it's in R.

Main features

- **Pure R environment**: As far as we know, this is the first package that allows end-to-end analysis of Sanger sequencing data in a pure R environment.
- Automated data analysis: Given appropriately-named input files, a lot of the data analysis can be automated. Once you've set up an appropriate workflow for your data, you can run it again in seconds.
- Interactive Shiny apps: Local Shiny apps mean you visualize the data at many levels, view chromatograms, and adjust things like trimming parameters.
- Exporting and importing FASTA files: sangeranalyseR is primarily designed with loading raw ab1 files in mind, but it can also load sequences in FASTA format. Aligned results and trimmed reads can be written into FASTA file format.
- **Thorough report**: A single command creates a comprehensive interactive HTML report that provides a huge amount of detail on the analysis.

CHAPTER $\mathbf{3}$

What sangeranalyseR doesn't do

One really important feature that sangeranalyseR doesn't have is the ability to edit bases by hand. R is just not the right language for this. If you need to edit your reads by hand, we suggest doing that in another tool like Geneious, then exporting your reads as FASTA files and following the instructions for using sangeranalyseR with FASTA input.

User Manual

If you are already familiar with sangeranalyseR and want to have a quick look at function signatures, please refer to sangeranalyseR user manual

User support

Please go through the *Documentation* below first. If you have questions about using the package, a bug report, or a feature request, please use the GitHub issue tracker here:

https://github.com/roblanf/sangeranalyseR/issues

Key contributors

The first (and not very good) version of the package was written by Rob Lanfear (at ANU in Australia), in collaboration with Kirston Barton and Sarah Palmer (then both at the University of Sydney). The second and far far better version of the package was written by Kuan-Hao (Howard) Chao at ANU. (This section was written by Rob Lanfear, lest you think Howard wrote it!)

Documentation

7.1 Installation

7.1.1 System requirements

- R >= 4.0.0 (current)
- Rstudio (recommended)

7.1.2 Install from Bioconductor

sangeranalyseR is on Bioconductor 3.12 development now.

To install this package, start R (version "4.0") and enter:



Home Install

Home » Bioconductor 3.12 » Software Packages » sangeranalyseR (development version)

sangeranalyseR

Fig. 1: Figure 1. sangeranalyseR on Bioconductor 3.12 development.

7.1.3 Install the development version

If you haven't installed the devtools package before, please install it first:

install.packages("devtools")

Then run the following code in your R console to install the newest version from Github.

```
library(devtools)
## Install the release version
install_github("roblanf/sangeranalyseR", ref = "master")
## Install the development version
install_github("roblanf/sangeranalyseR", ref = "develop")
library(sangeranalyseR)
```

After installing sangeranalyseR, load it in R console.

library(sangeranalyseR)

Now, you are ready to go !

7.1.4 Where to go from here ?

Please continue to the Quick Start Guide or the more detailed Beginners Guide.

7.2 Quick Start Guide

This page provides simple quick-start information for using sangeranalyseR with AB1 files. Please read the *Beginners Guide* page for more details on each step.

If you haven't already, please follow the steps in the Installation page to install and load sangeranalyseR.

7.2.1 Super-Quick Start (3 lines of code)

The most minimal example gets the job done in three lines of code. More details below.

7.2.2 Step 1: Prepare your input files

Put all your AB1 files in a directory ./my_data/. The directory can be called anything.

Name your files according to the convention contig_index_direction.abl. E.g. Drosophila_COI_1_F.abl and Drosophila_COI_2_R.abl describes a forward and reverse read to assemble into one contig. You can have as many files and contigs as you like in one directory.

7.2.3 Step 2: Load and analyse your data

This command loads, trims, builds contigs, and aligns contigs. All of these are done with sensible default values, which can be changed. I

7.2.4 Step 3 (optional): Explore your data

launchApp(my_aligned_contigs)

This launches an interactive Shiny app where you can view your analysis, change the default settings, etc.

7.2.5 Step 4: Output your aligned contigs

writeFasta(my_aligned_contigs)

This will save your aligned contigs as a FASTA file.

7.2.6 Step 5 (optional): Generate an interactive report

generateReport (my_aligned_contigs)

This will save a detailed interactive HTML report that you can explore.

7.2.7 A Reproducible Example

If you are still confused about how to run sangeranalyseR and want to check whether it produces the results that you want, then check this section for more details. Here we demonstrate a simple and reproducible example for using sangeranalyseR to generate a consensus read from 8 sanger ab1 files (4 contigs and each includes a forward and a reverse read).

1. Prepare your input files & loading

The data of this example is in the sangeranalyseR package; thus, you can simply get its path from the library.

```
rawDataDir <- system.file("extdata", package = "sangeranalyseR")
parentDir <- file.path(rawDataDir, 'Allolobophora_chlorotica', 'ACHLO')</pre>
```

2. Load and analyse your data

Run the following on-liner to create the sanger alignment object.

```
ACHLO_contigs <- SangerAlignment(ABIF_Directory = parentDir,
REGEX_SuffixForward = "_[0-9]*_F.ab1$",
REGEX_SuffixReverse = "_[0-9]*_R.ab1$")
```

Following is the R shell output that you will get.

3. Explore your data

Launch the Shiny app to check the visualized results.

launchApp(ACHLO_contigs)

Following is the R shell output that you will get.

And a Shiny would popup as showed in Figure 1

sangeranalyseR	■ SangerAlignment Ov	erview Page	Save S4 instance	Close UI
SangerAlignment Overview Input Parameters:				
≣1 SangerContig 🗸				
 1 SangerContig Overview Forward SangerReads 		🖬 Re-calculate Contigs Alignment		
● Reverse SangerReads 🛛 🖌	Output Directory:	/var/folders/33/7v38jdjd2874jcxb6l71m00h0000gn/T//RtmpGkJZBm		
≣2 SangerContig <	Raw ABI Parent Directory:	/Library/Frameworks/R.framework/Versions/4.0/Resources/library/sangeranalyseR/ext	tdata/Allolobopho	ra_chloroti
	Trimming Method:	Method 1: 'Modified Mott Trimming'		
≣3 SangerContig <	Forward Suffix RegExp:	_[0-9]+_F+		
≡4 SangerContig <	Reverse Suffix RegExp:	_[0-9]+_R+		
	Contigs Number:	4		
	Alignment Parameters			-
	0.5 MinFractionCall	0.5 MaxFractionLost		
	Genetic Code Data Frame			_

Fig. 2: Figure 1. SangerAlignment Shiny dashboard.

4. Output your aligned contigs

Write each contig and the aligned consensus read into FASTA files.

writeFasta(ACHLO_contigs)

Following is the R shell output that you will get.

And you will get three FASTA files:

- (1) Sanger_all_trimmed_reads.fa
- (2) Sanger_contigs_alignment.fa
- (3) Sanger_contigs_unalignment.fa

5. Generate an interactive report

Last but not least, generate an Rmarkdown report to store all the sequence information.

generateReport(ACHLO_contigs)

For more detailed analysis steps, please choose one the following topics :

- Beginners Guide
- Advanced User Guide SangerRead (AB1)
- Advanced User Guide SangerContig (AB1)
- Advanced User Guide SangerAlignment (AB1)
- Advanced User Guide SangerRead (FASTA)
- Advanced User Guide SangerContig (FASTA)
- Advanced User Guide SangerAlignment (FASTA)

7.3 Beginners Guide

If you haven't already, please follow the steps in the Installation page to install and load sangeranalyseR.

This guide is for users who are starting with **AB1** (.ab1) files. If you are starting with **FASTA** (.fasta or .fa) files, please read through this guide then follow the slightly different path for those starting with FASTA data here: *Advanced User Guide - SangerAlignment (FASTA)*.

7.3.1 Step 1: Preparing your input files

sangeranalyseR takes as input a group of **AB1** files, which it then groups together into contigs. Once the individual contigs are built, all the contigs are aligned and a simple phylogenetic tree is made. This section explains how you should organize your files before running sangeranalyseR.

First, prepare a directory and put all your **AB1** files inside it (there can be other files in there too, sangeranalyseR will ignore anything without a *AB1* file extension). Files can be organised in as many sub-folders as you like. sangeranalyseR will recursively search all the directories inside <code>ABIF_Directory</code> and find all files that end with **AB1**.

Second, give sangeranalyseR the information it needs to group reads into contigs. To do this, sangeranalyseR needs two pieces of information about each read: the direction of the read (forward or reverse), and the contig that it should be grouped into. There are two ways you can give sangeranalyseR this information:

- using the file name itself
- using a three-column csv file

We'll cover both approaches using the following example. Imagine you have sequenced four contigs with a forward and reverse read, all from the same species, but from different locations. In this case you might have arranged your data something like *Figure_1*, below.

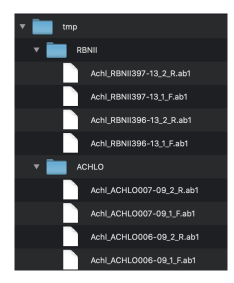


Fig. 3: Figure 1. Input ab1 files inside the parent directory, ./tmp/.

When using the filenames to group the reads, you'll need to specify three parameters: ABIF_Directory, REGEX_SuffixForward, and REGEX_SuffixReverse:

- ABIF_Directory: this is the directory that contains all the **AB1** files. In this example, the reads are in the /tmp/ directory, so for convenience we'll just say that ABIF_Directory should be /path/to/tmp/. In your case, it should be the absolute path to the folder that contains your reads.
- REGEX_SuffixForward: This is a regular expression (if you don't know what this is, don't panic it's just a way of recognising text that you will get the hang of fast), which tells sangeranalyseR how to use the end of a filename to determine a forward read. All the reads that are in forward direction have to contain this in their filename suffix. There are lots of ways to do this, but for this example, one uesful way to do it is _[0-9]*_F. ab1\$. This regular expression just says that the forward suffix is an underscore, followed at least one digit from 0-9, followed by another underscore then 'F', and ends with .ab1. The regex does not have to match to the end of the file name, but it's important to realise is that whatever comes before the part of the filename captured by this regex is by default the contig name. So in this case the regex also determines that the contig name for the first read is 'Achl_RBNII397-13'.

• REGEX_SuffixReverse: This is just the same as for the forward read, except that it determines the suffix for reverse reads. All the reads that are in reverse direction have to contain this in their filename suffix. In this example, its value is _[0-9]*_R.abl\$. I.e. all we've done is switch the 'F' in the forward read for an 'R' in the reverse read.

If you don't want to use the regex method, you can use the csv method instead. To use this method, just set processMethod parameter to csv and prepare an input .csv file with three columns:

- reads: the full file name (just the name, not the path) of the read to be grouped
- direction: "F" or "R" for forward and reverse reads, respectively
- contig: the name of the contig that reads should be grouped into

Following is an example of how you should organize your csv file in this example:

7.3.2 Step 2: Loading and analysing your data

After preparing the input files, you can create and align your contigs with just a single line of R code. In technical jargon, we are creating a *SangerAlignment* S4 instance.

It's important to note that this function is designed to be both *simple* and *flexible*. It's simple in that it has sensible defaults for all the usual things like trimming reads. But it's flexible in that you can change any and all of these defaults to suit your particular data and analyses. Here we just cover the simplest usage. The more flexible things are covered in the Advanced sections of the user guide.

So, let's create our contigs from our reads, and align them.

Here's how to do it using the regex method:

Here's how to do it using the csv file method

my_aligned_contigs is now a *SangerAlignment* S4 object which contains all of your reads, all the information on how they were trimmed, processed, and aligned, their chromatograms, and an alignment and phylogeny of all of your assembled contigs. The next section explains how to start digging into the details of that object.

7.3.3 Step 3: Exploring your data with the Shiny app

sangeranalseR includes a Shiny app that allows you to see, interact with, and adjust the parameters of your aligned contigs. For example, you can adjust things like the trimming parameters, and see how that changes your reads and your contigs.

To launch the interactive Shiny app use the launchApp function as follows

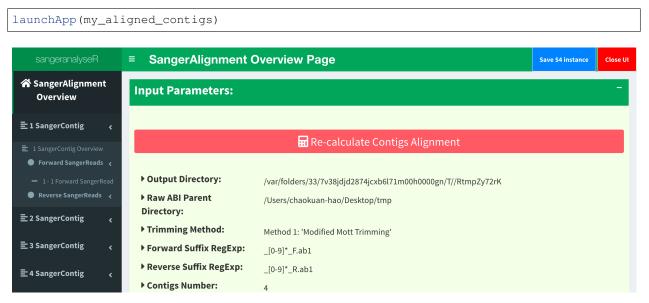


Fig. 4: Figure 2. SangerAlignment Shiny app user interface.

Figure_2 shows what the Shiny app looks like. On the left-hand side of *Figure_2*, there is a navigation menu that you can click to get more detail on every contig and every read. You can explore this app to get a lot more detail and make adjustments to your data. (Note that sangeranalyseR doesn't allow for editing individual bases of reads though - that's just not something that R is good for).

7.3.4 Step 4: Outputting your aligned contigs

Once you're happy with your aligned contigs, you'll want to save them somewhere.

The following function can write the *SangerAlignment* object into FASTA files. You just need to tell it where with the outputDir argument. Here we just wrote the alignment to the same folder that contains our reads.

writeFasta(my_aligned_contigs, outputDir = "/path/to/tmp/")

7.3.5 Step 5: Generating an interactive report

Last but not least, it is useful to store all the results in a report for future reference. You can generate a detailed report by running the following one-line function. *Figure_3* and *Figure_4*.

```
generateReport(my_aligned_contigs)
```

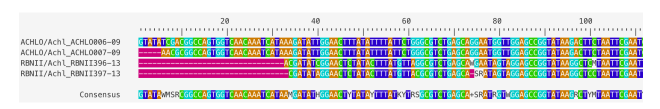


Fig. 5: Figure 3. An alignment of all contigs in the SangerAlignment object.

	RBNII/Achi RBNII397-13
	_
	RBNII/Achl RBNII396-13
—ACHLO/Achl ACHLO006-09	
ACHLO/Achi ACHLO007-09	

Fig. 6: Figure 4. A phylogenetic tree with contigs as the leaf nodes. This can help diagnose any issues with your contigs.

7.3.6 What's next?

Now you've finished the *Beginners Guide*, you should have a good overview of how to use the package. To dig a lot deeper into what you can do and why you might bother, there are also a set of advanced guides that focus on the three levels at which you can analyse Sanger data in the sangeranalyseR package. You can analyse individual reads with the *SangerRead* object, individual contigs with the *SangerContig* object, and alignments of two or more contigs (as we focussed on in this intro) with teh *SangerAlignment* object.

If you want to start the analysis from AB1 files, please choose the analysis level and read the following three links.

- Advanced User Guide SangerRead (AB1)
- Advanced User Guide SangerContig (AB1)
- Advanced User Guide SangerAlignment (AB1)

If you want to start the analysis from FASTA files, please choose the analysis level and read the following three links.

- Advanced User Guide SangerRead (FASTA)
- Advanced User Guide SangerContig (FASTA)
- Advanced User Guide SangerAlignment (FASTA)

7.4 Advanced User Guide - SangerRead (AB1)

SangerRead is in the bottommost level of sangeranalyseR (*Figure_1*), and each *SangerRead* object corresponds to a single read (one **AB1** file) in a Sanger sequencing experiment. *SangerRead* class extends *sangerseq* class from sangerseqR package and contains input parameters and results of quality trimming and chromatogram. In this section, we are going to go through detailed sangeranalyseR data analysis steps in *SangerRead level* with **AB1** file input.



Fig. 7: Figure 1. Hierarchy of classes in sangeranalyseR, SangerRead level.

7.4.1 Preparing SangerRead AB1 input

The main input file format to create *SangerRead* instance is **AB1**. Before starting the analysis, users need to prepare one target **AB1** file, and in this example, it is in the sangeranalyseR package; thus, you can simply get its path by running the following codes:

The only hard regulation of the filename, Achl_ACHLO006-09_1_F.ab1 in this example, is that the input file must have **.ab1** as its file extension. There are some suggestions about the filename in the note below:

Note:

- **AB1** file should be indexed for better consistency with file-naming regulation for *SangerContig* and *Sanger-Alignment*.
- Forward or reverse direction should be specified in the filename.

Figure_2 shows the suggested file-naming strategy. The filename should contain four main parts: "Contig name", "Index number", "Direction" and "ab1 file extension".

- "Contig name": Achl_RBNII397-13
- "Index number" : 1
- "Direction" : F
- "ab1 file extension" : . ab1

Achl_RBNII397-13_1_F.ab1

Fig. 8: Figure 2. SangerRead filename regulation.

In *SangerRead* section, it is not compulsory to follow the file-naming regulation because users can directly specify the filename in input (see *Creating SangerRead instance from AB1*); however, in the *SangerContig* and *SangerAlignment*, sangeranalyseR will automatically group files, so it is compulsory to have systematic file-naming strategy. For more details, please read *Advanced User Guide - SangerContig (AB1)* and *Advanced User Guide - SangerAlignment (AB1)*. *Figure_3* shows the suggested **AB1** file-naming regulation.

[Consensus Read Name] + _ + [index] + _ + [F,R] + .ab1

Fig. 9: Figure 3. Suggested AB1 file-naming regulation - SangerRead.

7.4.2 Creating SangerRead instance from AB1

After preparing the *SangerRead* input **AB1** file, A_chloroticaFFN, the next step is to create a *SangerRead* instance by running SangerRead constructor function or new method. The constructor function is a wrapper for the new method which makes instance creation more intuitive. The inputs include **Basic Parameters**, **Trimming Parameters**, and **Chromatogram Parameters**, and all of them have default values. In the example below, we show both *SangerRead* creation methods with important parameters.

<pre># using `constructor` function to create</pre>	
<pre>sangerReadF <- SangerRead(readFeature</pre>	= "Forward Read",
readFileName	= A_chloroticaFFN,
geneticCode	= GENETIC_CODE,
TrimmingMethod	= "M1",
MlTrimmingCuto	ff = 0.0001,
M2CutoffQualit	yScore = NULL,
M2SlidingWindo	wSize = NULL,
baseNumPerRow	= 100,
heightPerRow	= 200,
signalRatioCut	off = 0.33,
showTrimmed	= TRUE)
<pre># using `new` method to create SangerRea</pre>	d instance
<pre>sangerReadF <- new("SangerRead",</pre>	
readFeature	= "Forward Read",
readFileName	= A_chloroticaFFN,
geneticCode	
TrimmingMethod	= "M1",
M1TrimmingCutoff	
M2CutoffQualityScore	
M2SlidingWindowSize	
-	= 100,
heightPerRow	,
signalRatioCutoff	•
showTrimmed	= TRUE)
2110 M T T THUNGO	

The inputs of SangerRead constructor function and new method are the same. For more details about *SangerRead* inputs and slots definition, please refer to the sangeranalyseR reference manual. The created *SangerRead* instance, sangerReadF, is used as the input for the following functions.

Inside the R shell, you can run sangerReadF to get basic information of the instance or run sangerReadF@objectResults@readResultTable to check the creation result of every Sanger read after sangerReadF is successfully created.

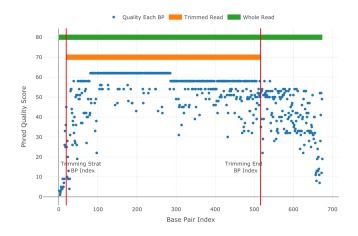
Here is the output of sangerReadF:

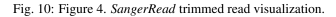
Here is the output of sangerReadF@objectResults@readResultTable:

readName	creationResult	errorType	errorMessage	inputSource	
<pre> →direction 1 Achl_ACHL0006-09_1_F.abl →Read </pre>	TRUE	None	None	ABIF	Forward

7.4.3 Visualizing SangerRead trimmed read

Before going to *Writing SangerRead FASTA file (AB1)* and *Generating SangerRead report (AB1)* pages, it is suggested to visualize the trimmed *SangerRead*. Run the qualityBasePlot function to get the result in *Figure_4*. It shows the quality score for each base pairs and the trimming start/end points of the sequence.





```
qualityBasePlot(sangerReadF)
```

7.4.4 Updating SangerRead quality trimming parameters

In the previous *Creating SangerRead instance from AB1* part, the constructor function applies the quality trimming parameters to the read. These parameters are not fixed. After instance creation, users can run updateQualityParam function which will change the *QualityReport* instance inside the *SangerRead* and update frameshift amino acid sequences.

TrimmingMethod = "M2",	d = "M2".	TrimmingMethod
	a 110 /	
MITTIMMINGCUTOII = NOLL,	off = NULL,	MlTrimmingCutoff
M2CutoffQualityScore = 29,	tyScore = 29,	M2CutoffQualitySco:
M2SlidingWindowSize = 15)	owSize = 15)	M2SlidingWindowSize

7.4.5 Writing SangerRead FASTA file (AB1)

After quality trimming, users can write <code>sangerReadF</code> into a **FASTA** file. Below is the one-liner that needs to be run. This function, writeFasta, mainly depends on writeXStringSet function in Biostrings R package. Users can

further set the compression level through it.

Users can download the output FASTA file of this example.

7.4.6 Generating SangerRead report (AB1)

Last but not least, users can save sangerReadF into a static HTML report by knitting Rmd files. In this example, tempdir function will generate a random path.

SangerRead_Report_ab1.html is the generated *SangerRead* report html of this example. Users can access to '*Basic Information*', '*DNA Sequence*', '*Amino Acids Sequence*', '*Quality Trimming*' and '*Chromatogram*' sections inside this report.

7.4.7 Code summary (SangerRead, ab1)

(1) Preparing SangerRead AB1 input

(2) Creating SangerRead instance from AB1

(continues on next page)

(continued from previous page)

readFeature	= "Forward Read",
readFileName	= A_chloroticaFFN)

Following is the R shell output that you will get.

(3) Visualizing SangerRead trimmed read

qualityBasePlot(sangerReadF)

(4) Writing SangerRead FASTA file (AB1)

writeFasta(sangerReadF)

Following is the R shell output that you will get.

And you will get one FASTA file:

(1) Achl_ACHL0006-09_1_F.fa

(5) Generating SangerRead report (AB1)

generateReport(sangerReadF)

You can check the html report of this SangerRead example (ABIF).

7.5 Advanced User Guide - SangerContig (AB1)

SangerContig is in the intermediate level of sangeranalyseR (*Figure_1*), and each *SangerContig* instance corresponds to a contig in a Sanger sequencing experiment. Among its slots, there are two lists, forward and reverse read list, storing *SangerRead* in the corresponding direction.

In this section, we are going to go through details about a reproducible *SangerContig* analysis example with the **AB1** file input in sangeranalyseR. By running the following example codes, you will get an end-to-end *SangerContig* analysis result.

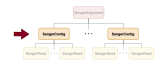


Fig. 11: Figure 1. Hierarchy of classes in sangeranalyseR, SangerContig level.

7.5.1 Preparing SangerContig AB1 inputs

The main input file format to create *SangerContig* instance is **AB1**. Before starting the analysis, users need to prepare one directory containing all **AB1** files, and all of them must be in the first layer of that directory. In other words, there should be no subdirectories. In this example, the data are in the sangeranalyseR package; thus, you can simply get its path by running the following codes:

```
rawDataDir <- system.file("extdata", package = "sangeranalyseR")
parentDir <- file.path(rawDataDir, "Allolobophora_chlorotica", "RBNII")</pre>
```

The value of parentDir is where all **AB1** files are placed. If your operating system is macOS, then its value should look like this:

And we showed the files under parentDir in Figure_2:



Fig. 12: Figure 2. SangerContig filename regulation.

Figure_2 shows the file-naming regulation and hierarchy. In this example, RBNII is the parent directory, and all **AB1** files must be under its first layer. There are two ways for users to group their **AB1** files which are **"regular expression matching"** and **"CSV file matching"**, and following are instructions of how to prepare and name your **AB1** input files.

(1) "regular expression matching" SangerContig inputs (AB1)

For regular expression matching method, sangeranalyseR will group **AB1** files based on their contig names and read directions in their filenames automatically; therefore, users have to follow the file-naming regulations below:

Note:

- All input files must have **.ab1** as its file extension
- All input files must have the same contig name in their filenames.
- Forward or reverse direction has to be specified in the filename.

There are four parameters, ABIF_Directory, contigName, REGEX_SuffixForward, and REGEX_SuffixReverse, that define the grouping rule to let sangeranalyseR automatically match correct **AB1** files and divide them into forward and reverse directions.

Note:

- ABIF_Directory: this is the directory that contains all **AB1** files, and it can be either an absolute or relative path. We suggest users to put only target **AB1** files inside this directory and do not include any other unrelated files.
- contigName: this is a regular expression that matches filenames that are going to be included in the *Sanger-Contig* analysis. grepl function in R is used.
- REGEX_SuffixForward: this is a regular expression that matches all filenames in forward direction. grepl function in R is used.
- REGEX_SuffixReverse: this is a regular expression that matches all filenames in reverse direction. grepl function in R is used.

If you don't know what regular expression is, don't panic - it's just a way of recognising text. Please refer to *What is a regular expression?* for more details. Here is an example of how it works in sangeranalseR:

So how sangeranalyseR works is that it first matches the contigName to exclude unrelated files and then separate the forward and reverse reads by matching REGEX_SuffixForward and REGEX_SuffixReverse. Therefore, it is important to make sure that all target **AB1** files share the same contigName and carefully select your REGEX_SuffixForward and REGEX_SuffixReverse. The bad file-naming and wrong regex matching might accidentally include reverse reads into the forward read list or vice versa, which will make the program generate wrong results. Therefore, it is important to have a consistent naming strategy. So, how should we systematically name **AB1** files? We suggest users to follow the file-naming regulation in *Figure_3*.

[Consensus Read Name] + _ + [index] + _ + [F,R] + .ab1

Fig. 13: Figure 3. Suggested AB1 file-naming regulation - SangerContig.

As you can see, the first part of the regulation is a consensus read name (or contig name), which narrows down the scope of **AB1** files to those we are going to examine. The second part of the regulation is an index. Since there might be more than one read that is in the forward or reverse direction, we recommend you to number your reads in the same

contig group. The third part is a direction which is either 'F' (forward) or 'R' (reverse). Last but not least, files have to end with **.ab1** file extension.

To make it more specific, let's go back to the true example. In *Figure_2*, there are a lot of **AB1** files from different contigs in RBNII (ABIF_Directory). First, we set contigName to "Achl_RBNII384-13" to reduce candidates from eight to two **AB1** files, Achl_RBNII384-13_1_F.ab1 and Achl_RBNII384-13_2_R.ab1. Then, we set REGEX_SuffixForward to " $[0-9] *_F.ab1$ " and REGEX_SuffixReverse to " $[0-9] *_R$. ab1\$" to let sangeranalyseR match and group forward and reverse reads automatically. By the regular expression rule, Achl_RBNII384-13_1_F.ab1 and Achl_RBNII384-13_2_R.ab1 will be categorized into "forward read list" and "reverse read list" respectively. The reason why we strongly recommend you to follow this file-naming regulation is that by doing so, you can directly adopt the example regular expression matching values, " $[0-9] *_F$. ab1\$" and " $[0-9] *_F$.ab1\$" and " $[0-9] *_F$.ab1\$

After understanding how parameters work, please refer to *Creating SangerContig instance from AB1* below to see how sangeranalseR creates 'Achl_RBNII384-13' *SangerContig* instance.

(2) "CSV file matching" SangerContig inputs (AB1)

For those who are not familiar with regular expression, we provide a second grouping approach, CSV file matching method. sangeranalyseR will group **AB1** files based on the information in a CSV file automatically; therefore, users have to follow the regulations below:

Note: Here is an example CSV file (*Figure_4*)

"reads","direction","contig" "Achl_RBNII384-13_1_F.ab1","F","Achl_RBNII384-13" "Achl_RBNII384-13_2_R.ab1","R","Achl_RBNII384-13"

Fig. 14: Figure 4. Example CSV file for SangerContig instance creation.

- There must be three columns, "reads", "direction", and "contig", in the CSV file.
- The "reads" column stores the filename of AB1 files that are going to be included in the analysis.
- The "direction" column stores the direction of the reads. It must be "F" (forward) or "R" (reverse).
- The "contig" column stores the contig name that each read blongs. Reads in the same contig have to have the same contig name, and they will be grouped into the same *SangerContig* instance.

There are three parameters, ABIF_Directory, contigName, and CSV_NamesConversion, that define the grouping rule to help sangeranalseR to automatically match correct **AB1** files and divide them into forward and reverse directions.

Note:

- ABIF_Directory: this is the directory that contains all **AB1** files, and it can be either an absolute or relative path. We suggest users to put only target AB1 files inside this directory and do not include any other unrelated files.
- contigName: this is a regular expression that matches filenames that are going to be included in the *Sanger-Contig* analysis. grepl function in R is used.
- CSV_NamesConversion: this is the path to the CSV file. It can be either an absolute or relative path.

The main difference between "CSV file matching" and "regular expression matching" is where the grouping rule is written. For "regular expression matching", rules are writtein in filenames, and thus more naming requirements are required. In contrast, rules of "CSV file matching" are written in an additional CSV file so it is more flexible on **AB1** file-naming.

So how sangeranalyseR works is that it first reads in the CSV file (with "reads", "direction", and "contig" columns), filter out rows whose "contig" is not the value of contigName parameter, find the names of **AB1** files listed in "reads", and assign directions to them based on "direction".

To make it more specific, let's go back to the true example. First, we prepare a CSV file (CSV_NamesConversion) and a file directory like *Figure_2* (ABIF_Directory) with some **AB1** files from different contigs. In the CSV file, both rows have the contig name "Achl_RBNII384-13", which is what we need to assign to the contigName parameter. sangeranalyseR then checks and matches "*reads*" of these two rows, "Achl_RBNII384-13_1_F.ab1" and "Achl_RBNII384-13_2_R.ab1", in RBNII directory and reduce candidates from eight to two **AB1** files. Last, these two reads are assigned into "forward read list" and "reverse read list" respectively by the "*direction*" column.

After understanding how parameters work, please refer to *Creating SangerContig instance from AB1* below to see how sangeranalseR creates 'Achl_RBNII384-13' *SangerContig* instance.

7.5.2 Creating SangerContig instance from AB1

After preparing the input directory, we can create a *SangerContig* instance by running SangerContig constructor function or new method. The constructor function is a wrapper for new method and it makes instance creation more intuitive. Their input parameters are same, and all of them have their default values. For more details about *SangerContig* inputs and slots definition, please refer to sangeranalyseR reference manual. We will explain two *SangerContig* instance creation methods, "regular expression matching" and "CSV file matching".

(1) "regular expression matching" SangerContig creation (AB1)

The consturctor function and new method below contain four parameters, ABIF_Directory, contigName, REGEX_SuffixForward, and REGEX_SuffixReverse, that we mentioned in the previous section. It also includes important parameters like quality trimming, chromatogram visualization, consensus alignment, and so on. Run the following code and create my_sangerContig instance.

<pre># using `constructor` function to create Sa</pre>	ngerContig instance
<pre>my_sangerContig <- SangerContig(inputSource</pre>	= "ABIF",
processMeth	.od = "REGEX",
ABIF_Direct	ory = parentDir,
contigName	= "Achl_RBNII384-13",
REGEX_Suffi	$xForward = "[0-9] *_F.ab1$",$
REGEX_Suffi	xReverse = "_[0-9]*_R.ab1\$",
TrimmingMet	hod = "M1",
M1TrimmingC	utoff = 0.0001,
M2CutoffQua	lityScore = NULL ,
M2SlidingWi	ndowSize = NULL ,
baseNumPerR	.ow = 100,
heightPerRo	w = 200,
signalRatio	Cutoff = 0.33,
showTrimmed	= TRUE,

(continues on next page)

(continued from previous page)

	(continued from previous page)
refAminoAcid	lSeq =
	IPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNNI
\hookrightarrow ",	
minReadsNum	= 2,
minReadLengt	h = 20,
minFractionC	Call = 0.5,
maxFractionI	lost = 0.5,
geneticCode	= GENETIC_CODE,
acceptStopCo	odons = TRUE ,
readingFrame	e = 1,
processorsNu	im = 1)
<pre># using `new` method to create SangerContig</pre>	instance
<pre>my_sangerContig <- new("SangerContig",</pre>	
inputSource	= "ABIF",
processMethod	= "REGEX",
ABIF_Directory	<pre>= parentDir, = "Achl_RBNII384-13",</pre>
contigName	= "Achl_RBNII384-13",
REGEX_SuffixForward	= "_[0-9]*_F.ab1\$",
REGEX_SuffixReverse	= "_[0-9]*_R.ab1\$",
TrimmingMethod	= " <u>M1</u> ",
M1TrimmingCutoff	= 0.0001,
M2CutoffQualityScore	= NULL,
M2SlidingWindowSize	
baseNumPerRow	= 100,
heightPerRow	= 200,
signalRatioCutoff	= 0.33,
showTrimmed	= TRUE,
refAminoAcidSeq	=
	IPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNNI
\hookrightarrow ",	
minReadsNum	= 2,
minReadLength	= 20,
minFractionCall	
maxFractionLost	
	= GENETIC_CODE,
	= TRUE,
readingFrame	= 1,
processorsNum	= 1)

In this example, contigName is set to Achl_RBNII384-13, so only Achl_RBNII384-13_1_F.abl and Achl_RBNII384-13_2_R.abl are selected. Moreover, by regular expression pattern matching, Achl_RBNII384-13_1_F.abl is categorized into the forward list, and Achl_RBNII384-13_2_R.abl is categorized into the reverse read. Both reads are aligned into a contig, my_sangerContig, and it will be used as the input for the following functions.

Inside the R shell, you can run my_sangerContig to get basic information of the instance or run my_sangerContig@objectResults@readResultTable to check the creation result of every Sanger read after my_sangerContig is successfully created.

Here is the output of my_sangerContig:

```
SangerContig S4 instance
    Input Source : ABIF
    Process Method : REGEX
    ABIF Directory : /Library/Frameworks/R.framework/Versions/4.0/Resources/
    →library/sangeranalyseR/extdata/Allolobophora_chlorotica/RBNII
```

(continues on next page)

Here is the output of my_sangerContig@objectResults@readResultTable:

readName	creationResult	errorType	errorMessage	inputSource	
→direction					
1 Achl_RBNII384-13_1_F.ab1	TRUE	None	None	ABIF	Forward
⇔Read					
2 Achl_RBNII384-13_2_R.ab1	TRUE	None	None	ABIF	Reverse
⊶Read					

(2) "CSV file matching" SangerContig creation (AB1)

The consturctor function and new method below contain three parameters, ABIF_Directory, contigName, and CSV_NamesConversion, that we mentioned in the previous section. It also includes important parameters like quality trimming, chromatogram visualization, consensus alignment, and so on. Run the following code and create my_sangerContig instance.

```
csv_namesConversion <- file.path(rawDataDir, "ab1", "SangerContig", "names_conversion_
\rightarrow 2.csv")
# using `constructor` function to create SangerContig instance
my_sangerContig <- SangerContig(inputSource</pre>
                                                    = "ABIF",
                                                    = "CSV",
                               processMethod
                               ABIF_Directory
                                                    = parentDir,
                               contigName
                                                     = "Achl_RBNII384-13",
                               CSV_NamesConversion = csv_namesConversion,
                                                     = "M1",
                               TrimmingMethod
                               M1TrimmingCutoff
                                                    = 0.0001,
                               M2CutoffQualityScore = NULL,
                               M2SlidingWindowSize = NULL,
                                                     = 100,
                               baseNumPerRow
                                                    = 200,
                               heightPerRow
                               signalRatioCutoff
                                                    = 0.33,
                                                      = TRUE,
                               showTrimmed
                               refAminoAcidSeq
                                                     =
↔ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNNI
∽",
                               minReadsNum
                                                      = 2,
                               minReadLength
                                                     = 20,
                                                     = 0.5,
                               minFractionCall
                                                     = 0.5,
                               maxFractionLost
                                                                       (continues on next page)
```

		(continued from previo	sus puge)
geneticCod	= GE	ENETIC_CODE,	
acceptStop	codons = TR	RUE,	
readingFra	ne = 1,		
processors	1um = 1)		
<pre># using `new` method to create SangerConti</pre>	r instance		
<pre>my_sangerContig <- new("SangerContig",</pre>			
Ĩ	= "ABIF",		
processMethod			
ABIF_Directory			
contigName	= "Achl_RBNII3	384-13",	
CSV_NamesConversion	= csv_namesCon	nversion,	
TrimmingMethod	= "M1",		
MlTrimmingCutoff	= 0.0001,		
M2CutoffQualityScor	= NULL,		
M2SlidingWindowSize	= NULL,		
baseNumPerRow	= 100,		
heightPerRow	= 200,		
signalRatioCutoff	= 0.33,		
showTrimmed	= TRUE,		
refAminoAcidSeq =			
↔"SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAEL	HPGALIGDDQIYNVIV	/TAHAFIMIFFMVMPIMIGGFGNW	VLVPLMLGAPDMAFPRMNN
⇔",			
minReadsNum	= 2,		
minReadLength	= 20,		
minFractionCall	= 0.5,		
maxFractionLost	= 0.5,		
geneticCode	= GENETIC_COD	DE,	
acceptStopCodons	= TRUE,		
readingFrame	= 1,		
processorsNum	= 1)		
-			

First, you need to load the CSV file into the R environment. If you are still don't know how to prepare it, please check (2) "CSV file matching" SangerContig inputs (AB1). Then, it will follow rules in the CSV file and create my_sangerContig. After it's created, inside the R shell, you can run my_sangerContig to get basic information of the instance or run my_sangerContig@objectResults@readResultTable to check the creation result of every Sanger read after my_sangerContig is successfully created.

Here is the output of my_sangerContig:

```
SangerContig S4 instance
         Input Source : ABIF
        Process Method : CSV
        ABIF Directory : /Library/Frameworks/R.framework/Versions/4.0/Resources/
→library/sangeranalyseR/extdata/Allolobophora_chlorotica/RBNII
  CSV Names Conversion : /Library/Frameworks/R.framework/Versions/4.0/Resources/
-library/sangeranalyseR/extdata/ab1/SangerContig/names_conversion_2.csv
            Contig Name : Achl_RBNII384-13
         'minReadsNum' : 2
      'minReadLength': 20
      'minFractionCall' : 0.5
      'maxFractionLost': 0.5
   'acceptStopCodons' : TRUE
         'readingFrame' : 1
     Contig Sequence :

ightarrow AGCAGGATAGTAGGGGCTGGTATAAGACTCCTAATTCGAATTGAGCTAAGACAGCCGGGAGCATTTCTAGGAAGGGATCAACTCTATAACACTATTGT
ightarrow
```

```
(continues on next page)
```

```
Forward reads in the contig >> 1
Reverse reads in the contig >> 1
SUCCESS [2021-12-07 17:11:48] 'Achl_RBNII384-13' is successfully created!
```

Here is the output of my_sangerContig@objectResults@readResultTable:

readName	creationResult	errorType	errorMessage	inputSource	L.
→direction					
1 Achl_RBNII384-13_1_F.ab1	TRUE	None	None	ABIF	Forward
⇔Read					
2 Achl_RBNII384-13_2_R.ab1	TRUE	None	None	ABIF	Reverse
⇔Read					

7.5.3 Updating SangerContig quality trimming parameters

In the previous *Creating SangerContig instance from AB1* part, the constructor function will apply the quality trimming parameters to all reads. After creating a *SangerContig* instance, users can change the trimming parameters by running updateQualityParam function which will update all reads with the new trimming parameters and redo reads alignment. If users want to do quality trimming read by read instead of all at once, please move on to the next section, *Launching SangerContig Shiny app* page.

```
newSangerContig <- updateQualityParam(my_sangerContig,
TrimmingMethod = "M2",
M1TrimmingCutoff = NULL,
M2CutoffQualityScore = 20,
M2SlidingWindowSize = 15)
```

7.5.4 Launching SangerContig Shiny app

We create an interactive local Shiny app for users to go into each *SangerRead* in *SangerContig* instance. Users only need to run one function, launchApp, with previously created instance as input and the *SangerContig* Shiny app will pop up. Here, we will go through *SangerRead* and *SangerContig* pages.

launchApp(my_sangerContig)

SangerContig page (SC app)

SangerContig page is the initial page of *SangerContig* Shiny app. *Figure 5* shows the overview page of the contig. Notice that there is a red "Re-calculate Contig" button. Users need to click the button after changing the quality trimming parameters in order to get the updated information. In SangerContig page, there are two expendable tabs, "Forward Reads" and "Reverse Reads" storing the corresponding reads on the left-hand side navigation panel in *Figure 5*. See *SangerRead page (SC app)* for more details of the subpage.



Fig. 15: Figure 5. SangerContig Shiny app initial page - SangerContig page.

The information provided in this page are input parameters and contig results including "genetic code table", "reference amino acid sequence", "reads alignment", "difference data frame", "dendrogram", "sample distance heatmap", "indels data frame", and "stop codons data frame".

Figure 6 shows reads alignment result and difference data frame. The alignment is generated by AlignSeqs or AlignTranslation function in DECIPHER package.

ignı	ment					
	1_Read_Achl_ACHL0006-09_1_F.ab1 2_Read_Achl_ACHL0006-09_2_R.ab1 Consensus Consensus	TATATCGACGGCCAG TATATCGACGGCCAG	GGTCAACAAATCA	1 1	TATATITTATICIG TATATITTATICIG	GGCGTCTGAGCAGGAA GGCGTCTGAGCAGGAA GGCGTCTGAGCAGGAA
	ences Data frame				Search:	
		¢		pairwise.diffs.to.co	1	unused.char
	10 ¢ entries	¢		pairwise.diffs.to.co	1	unused.char

Fig. 16: Figure 6. SangerContig page - reads alignment and difference data frame.

Figure 7 shows dendrogram result in both plot and in data frame. The results are generated by TreeLine function in DECIPHER package.

Figure 8 shows distance between **AB1** files. The results are generated by DistanceMatrix function in DECIPHER package. The heatmap is generated by plot_ly function in plotly package.

Figure 9 shows insertions, deletions and stop codons data frame.

SangerRead page (SC app)

Now, let's go to the next level which is also the lowest level, *SangerRead* page. *SangerRead* page contains all details of a read including its trimming and chromatogram inputs and results. All reads are in "forward" or "reverse" direction. In this example, there is one read in each direction and *Figure 10* shows "1 Forward Read" page. This page provides basic

endrogram					
1_Read_Achl_ACHLO006-09_1_F.ab1 -					7
2_Read_Achl_ACHL0006-09_2_R.ab1 - 0.0000		0.0005	0.0010	0.0015	
Show 10 🗘 entries				Search:	luster 🕯
1_Read_Achl_ACHLO006-09_1_F	.ab1				2
2_Read_Achl_ACHLO006-09_2_R	.ab1				1
2_Read_Achl_ACHLO006-09_2_R Showing 1 to 2 of 2 entries	.ab1			Previous 1	

Fig. 17: Figure 7. SangerContig page - dendrogram.

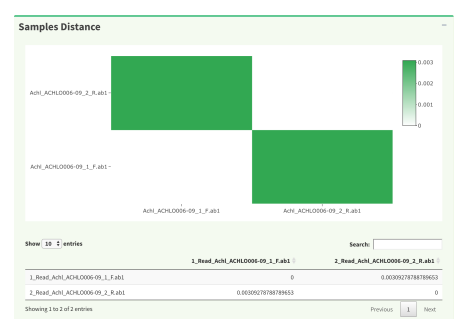


Fig. 18: Figure 8. SangerContig page - samples distance.

nde	ls Data frame				
Show	10 ¢ entries			Search:	
	read	÷	insertions 🔶	deletions 🝦	distance
1	/Users/chaokuan-hao/Desktop/tmp/ACHLO//Achl_ACHLO006- 09_1_F.ab1		3	0	0.280172413793103
2	/Users/chaokuan-hao/Desktop/tmp/ACHLO//Achl_ACHLO006- 09_2_R.ab1		4	0	0.25974025974026
Show	ing 1 to 2 of 2 entries			Previ	ous 1 Next
Stop	Codons Data frame				
Show	10 🜩 entries			Search:	
	read			÷	stop.codons
1	/Users/chaokuan-hao/Desktop/tmp/ACHLO//Achl_ACHLO006-0	9_1_F.a	ib1		8
2	/Users/chaokuan-hao/Desktop/tmp/ACHLO//Achl_ACHLO006-0	9_2_R.;	ab1		5
Show	ing 1 to 2 of 2 entries			Previ	ous 1 Next

Fig. 19: Figure 9. SangerContig page - indels and stop codons data frame.

information, quality trimming inputs, chromatogram plotting inputs etc. Primary/secondary sequences and quality Phred scores table in this figure are dynamic based on the signalRatioCutoff value for base calling and the length of them are always same. Another thing to mention is that primary/secondary sequences and the sequences in the chromatogram in *Figure 15* below will always be same after trimming and their color codings for A/T/C/G are same as well.



Fig. 20: Figure 10. SangerContig Shiny app - SangerRead page

In quality trimming steps, we removes fragment at both ends of sequencing reads with low quality score. It is important because trimmed reads will improves alignment results. *Figure 11* shows the UI for Trimming Method 1 (M1): 'Modified Mott Trimming'. This method is implemented in Phred. Users can change the cutoff score and click "Apply Trimming Parameters" button to update the UI. The value of input must be between 0 and 1. If the input is invalid, the cutoff score will be set to default 0.0001.

Figure 12 shows another quality trimming method for users to choose from, Trimming Method 2 (M2): 'Trimmomatics Sliding Window Trimming'. This method is implemented in Trimmomatics. Users can change the cutoff quality score as well as sliding window size and click "Apply Trimming Parameters" button to update the UI. The value of cutoff quality score must be between 0 and 60 (default 20); the value of sliding window size must be between 0 and 40

Quality Report:		
Trimming Paramet	ers Input	-
Your trimming method sele	tion : 'Modified Mott Trimming'	
1e-04 Cutoff Score	Ж	
Input Value		
1e-04		
	Apply Trimming Parameters	

Fig. 21: Figure 11. SangerRead page - Trimming Method 1 (M1): 'Modified Mott Trimming' UI.

(default 10). If the inputs are invalid, their values will be set to default.

rimming Daramotor	s Innut		
rimming Parameters	-	g Window Trimming'	
ar trimming method selection	in . Thinnonatics stium	g window miniming	
20	X	15	- 5 3
Contra ff Concelling Concerns			
Cutoff Quality Score		Sliding Window Size	
Cutoff Quality Score Input Value		Sliding Window Size	

Fig. 22: Figure 12. SangerRead page - Trimming Method 2 (M2): 'Trimmomatics Sliding Window Trimming' UI.

Figure 13 shows the quality report before and after trimming. After clicking the "Apply Trimming Parameters" button in *Figure 11* or *Figure 12*, the values of these information boxes will be updated to the latest values.

In *Figure 14*, the x-axis is the index of the base pairs; the y-axis is the Phred quality score. The green horizontal bar at the top of the plot is the raw read region and the orange horizontal bar represents the remaining read region. Both *Figure 14* trimming plot and *Figure 15* chromatogram will be updated once users change the quality trimming parameters and click the "Apply Trimming Parameters" button in *Figure 15*.

If we only see primary and secondary sequences in the table, we will loose some variations. Chromatogram is very helpful to check the peak resolution. *Figure 15* shows the panel of plotting chromatogram. Users can change four parameters: Base Number Per Row, Height Per Row, Signal Ratio Cutoff, and Show Trimmed Region. Among them, Signal Ratio Cutoff is a key parameter. If its value is default value 0.33, it indicates that the lower peak should be at least 1/3rd as high as the higher peak for it count as a secondary peak.

Here is an example of applying new chromatogram parameters. We click "Show Trimmed Region" to set its value from FALSE to TRUE and click the "Apply Chromatogram Parameters" button. *Figure 16* shows the loading notification popup during base calling and chromatogram plotting.

After replotting the chromatogram, we can see that trimmed region is showed in red striped region. *Figure 17* shows part of the chromatogram (1 bp ~ 240 bp). Moreover, chromatogram will be replotted when trimmed positions or

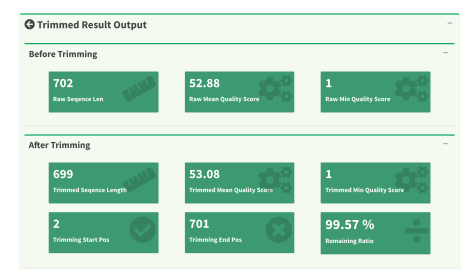


Fig. 23: Figure 13. SangerRead page - read quality report before / after trimming.



Fig. 24: Figure 14. SangerContig page - quality trimming plot.

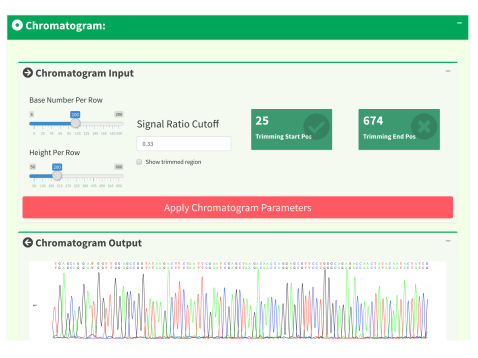


Fig. 25: Figure 15. SangerContig page - chromatogram panel.

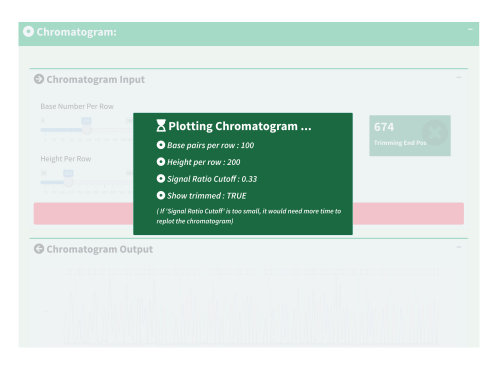


Fig. 26: Figure 16. SangerContig page - loading notification popup during replotting chromatogram.

chromatogram parameters are updated.

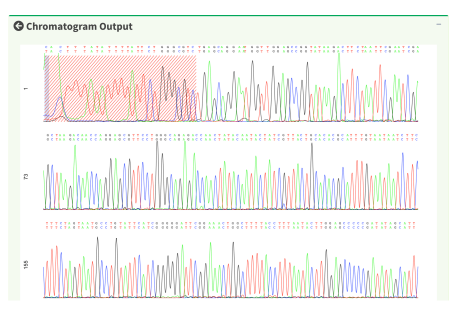


Fig. 27: Figure 17. SangerContig page - chromatogram with trimmed region showed.

To let users browse the trimmed primary/secondary sequences without finding "Trimming Start Point" and "Trimming End Point" by themselves, we provide the final trimmed primary/secondary sequences that will be used for reads alignment with quality scores in table format in *Figure 18*. Frameshift amino acid sequences are also provided.

Pr	im	ary	Seq	ue	۱ce																									
	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16 1	.7 1	19	20	21	22	23	24	25	26	27	28	29	30	31
1	с	Α	с	т	т	т	А	т	Α	т	т	т	т	Α	т	т	с т	G	G	G	с	G	т	С	т	G	Α	G	с	Α
E Se	col	ıda	ry S	eq	Jen	ce																								
Se			-				7	8	9	10	11	12	13	14	15	16 1	.7 1	19	20	21	22	23	24	25	26	27	28	29	30	31
5 Se			3	4	5	6		8 T							15 T		.7 1: C T	_		21 G				25 C	26 T	27 G	28 A		30 C	31 A
	1	2	3	4	5	6			_								_	_							26 T	_	_		30 C	31 A
1	1 T	2 A	3 C	4 T	5 T	6 T			_								_	_							26 T	_	_		30 C	31 A
1	1 T	2	3 C	4 T	5 T	6 T			_								_	_							26 T	_	_		30 C	31 A
1	1 T	2 A ty P	3 C	4 T	5 T	6 T			_	т		т	т		т	Т	_	G	G		С	G			26 T	_	_		30 C	31 A 31

Fig. 28: Figure 18. SangerContig page - trimmed primary/secondary sequences and Phred quality score in table format.

We have updated the trimming and chromatogram parameters for each read. Now, we need to click "Re-calculate contig" button to do alignment again. Last but not least, we can save all data into a new 'SangerContig' S4 instance by clicking "Save S4 Instance button". New S4 instance will be saved in **Rda** format. Users can run readRDS function to load it into current R environment. *Figure 19* shows some hints in the save notification popup.

sangeranalyseR	SangerContig Overview Page Save \$4 instance Close	e UI
☆ SangerContig Overview	• Basic Information:	
Forward SangerReads - 1 Forward SangerRead	🖬 Re-calculate Contig	
Reverse SangerReads	• Output Directory: • Raw ABI Parent Directory: • Contig Name: • Trimming Method: • Forward Suffix RegExp: • Forward Read Number: • Reverse Read Number: • Contig Parameters 2 MinReadsHum 2 MinReadsHum • Contig Parameters • Contig	

Fig. 29: Figure 19. SangerContig page - saving notification popup.

7.5.5 Writing SangerContig FASTA files (AB1)

Users can write the *SangerContig* instance, my_sangerContig, to **FASTA** files. There are four options for users to choose from in selection parameter.

- reads_unalignment: Writing reads into a single FASTA file (only trimmed without alignment).
- reads_alignment: Writing reads alignment and contig read to a single FASTA file.
- contig: Writing the contig to a single FASTA file.
- all: Writing reads, reads alignment, and the contig into three different files.

Below is the oneliner for writing out FASTA files. This function mainly depends on writeXStringSet function in Biostrings R package. Users can set the compression level through writeFasta function.

Users can download the output FASTA file of this example through the following three links:

- (1) Achl_RBNII384-13_reads_unalignment.fa
- (2) Achl_RBNII384-13_reads_alignment.fa
- (3) Achl_RBNII384-13_contig.fa

7.5.6 Generating SangerContig report (AB1)

Last but not least, users can save *SangerContig* instance, my_sangerContig, into a report after the analysis. The report will be generated in **HTML** by knitting **Rmd** files.

Users can set includeSangerRead parameter to decide to which level the *SangerContig* report will go. Moreover, after the reports are generated, users can easily navigate through reports in different levels within the **HTML** file.

One thing to pay attention to is that if users have many reads, it will take quite a long time to write out all reports. If users only want to generate the contig result, remember to set includeSangerRead to FALSE in order to save time.

Here is the generated SangerContig html report of this example (ABIF). Users can access to 'Basic Information', 'SangerContig Input Parameters', 'Contig Sequence' and 'Contig Results' sections inside it. Furthermore, users can also navigate through html reports of all forward and reverse SangerRead in this SangerContig report.

7.5.7 Code summary (SangerContig, AB1)

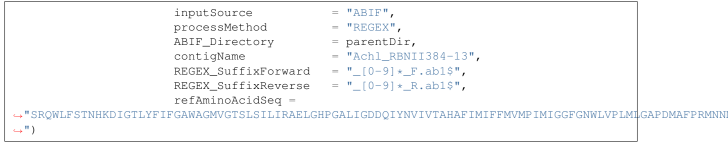
(1) Preparing SangerContig AB1 inputs

```
rawDataDir <- system.file("extdata", package = "sangeranalyseR")
parentDir <- file.path(rawDataDir, "Allolobophora_chlorotica", "RBNII")</pre>
```

(2) Creating SangerContig instance from AB1

(2.1) "Regular Expression Method" SangerContig creation (AB1)

(continues on next page)



Following is the R shell output that you will get.

(2.2) "CSV file matching" SangerContig creation (AB1)

```
csv_namesConversion <- file.path(rawDataDir, "ab1", "SangerContig", "names_conversion_
\leftrightarrow 2.csv")
# using `constructor` function to create SangerContig instance
my_sangerContig <- SangerContig(inputSource = "ABIF",</pre>
                                 processMethod
                                                       = "CSV",
                                 processmethod= "CSV",ABIF_Directory= parentDir,contigName= "Achl_RBNII384-13",
                                 CSV_NamesConversion = csv_namesConversion,
                                 refAminoAcidSeq =
↔ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNN
→ " )
# using `new` method to create SangerContig instance
my_sangerContig <- new("SangerContig",</pre>
                                              = "ABIF",
                       inputSource
                                            = "CSV",
                       processMethod
                       ABIF_Directory
                                             = parentDir,
                                             = "Achl_RBNII384-13",
                        contigName
                       CSV_NamesConversion = csv_namesConversion,
                        refAminoAcidSeg =
↔ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNNI
→ " )
```

Following is the R shell output that you will get.

(3) Updating *SangerContig* quality trimming parameters

(continues on next page)

```
M2CutoffQualityScore = 20,
M2SlidingWindowSize = 15)
```

(4) Launching SangerContig Shiny app

launchApp(my_sangerContig)

(5) Writing SangerContig FASTA files (AB1)

writeFasta(my_sangerContig)

Following is the R shell output that you will get.

You will get three FASTA files:

- (1) Achl_RBNII384-13_reads_unalignment.fa
- (2) Achl_RBNII384-13_reads_alignment.fa
- (3) Achl_RBNII384-13_contig.fa

(6) Generating SangerContig report (AB1)

generateReport(my_sangerContig)

You can check the html report of this SangerContig example (ABIF).

7.6 Advanced User Guide - SangerAlignment (AB1)

SangerAlignment is in the toppest level of sangeranalyseR (*Figure_1*), and each **SangerAlignment** instance corresponds to an alignment of contigs in a Sanger sequencing experiment. Among its slots, there is a *SangerContig* list which will be aligned into a consensus contig. Users can access to each *SangerContig* and *SangerRead* inside a *SangerAlignment* instance.

In this section, we are going to go through details about a reproducible *SangerAlignment* analysis example with the **AB1** file input in sangeranalyseR. By running the following example codes, you will get an end-to-end *SangerAlignment* analysis result.

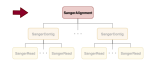


Fig. 30: Figure 1. Classes hierarchy in sangeranalyseR, SangerAlignment level.

7.6.1 Preparing SangerAlignment AB1 input

The main input file format to create *SangerAlignment* instance is **AB1**. Before starting the analysis, users need to prepare one directory containing all **AB1** files, and they can be either all placed in the first layer of that directory or be distributed in different subdirectories. In this example, the data are in the sangeranalyseR package; thus, you can simply get its path by running the following codes:

```
rawDataDir <- system.file("extdata", package = "sangeranalyseR")
parentDir <- file.path(rawDataDir, 'Allolobophora_chlorotica')</pre>
```

The value of parentDir is where all **AB1** files are placed. If your operating system is macOS, then its value should look like this:

And we showed the files under parentDir in *Figure_2*:

Figure_2 shows the file-naming regulation and hierarchy. In this example, Allolobophora_chlorotica is the parent directory, and **AB1** files are separated into ACHLO and RBNII directories. There are two ways for users to group their **AB1** files which are "**regular expression matching**" and "**CSV file matching**", and following are instructions of how to prepare and name your **AB1** input files.

(1) "regular expression matching" SangerAlignment inputs (AB1)

For regular expression matching method, sangeranalyseR will group **AB1** files based on their contig names and read directions in their filenames automatically; therefore, users have to follow the file-naming regulations below:

Note:

- All input files must have .ab1 as its file extension.
- Input files that are in the same contig group must have the same contig name in their filenames.
- Forward or reverse direction has to be specified in the filename.



Fig. 31: Figure 2. SangerAlignment filename regulation.

There are three parameters, ABIF_Directory, REGEX_SuffixForward, and REGEX_SuffixReverse, that define the grouping rule to let sangeranalyseR automatically match correct **AB1** files and divide them into forward and reverse directions.

Note:

- ABIF_Directory: this is the directory that contains all **AB1** files, and it can be either an absolute or relative path. We suggest users to put only target **AB1** files inside this directory and do not include any other unrelated files.
- REGEX_SuffixForward: this is a regular expression that matches all filenames in forward direction. grepl function in R is used.
- REGEX_SuffixReverse: this is a regular expression that matches all filenames in reverse direction. grepl function in R is used.

If you don't know what regular expression is, don't panic - it's just a way of recognising text. Please refer to *What is a regular expression?* for more details. Here is an example of how it works in sangeranalseR:

So how sangeranalyseR works is that it first matches the forward and reverse reads by matching REGEX_SuffixForward and REGEX_SuffixReverse. Then, sangeranalyseR uses the str_split function to split and vectorize their filenames into "contig name" and "direction-suffix" two parts. For those having the same "contig name" will be grouped into the same contig.

Therefore, it is important to have a consistent naming strategy. You need to make sure that **AB1** files in the same contig group share the same contig name and carefully select your REGEX_SuffixForward and REGEX_SuffixReverse. The bad file-naming and wrong regex matching might accidentally include reverse reads into the forward read list or vice versa, which will make the program generate wrong results. So, how should we systematically name **AB1** files? We suggest users to follow the file-naming regulation in *Figure_3*.

[Consensus Read Name] + + [index] + + [F,R] + .ab1

Fig. 32: Figure 3. Suggested AB1 file-naming regulation - SangerContig.

As you can see, the first part of the regulation is a consensus read name (or contig name), which helps sangeranalseR to identify which reads should be grouped into the same contig automatically. The second part of the regulation is an index; since there might be more than one read that is in the forward or reverse direction, we recommend you to number your reads in the same contig group. The third part is a direction which is either 'F' (forward) or 'R' (reverse). Last but not least, files have to end with **.ab1** file extension.

To make it more specific, let's go back to the true example. In *Figure_2*, there are two subdirectories, ACHLO and RBNII, containing lots of **AB1** files from different contigs in the root directory, Allolobophora_chlorotica (ABIF_Directory).

First, we set REGEX_SuffixForward to "_[0-9]*_F.ab1\$" and REGEX_SuffixReverse to "_[0-9]*_R.ab1\$" to let sangeranalyseR match and group forward and reverse reads automatically. By the regular expression rule, Ach1_ACHL0006-09_1_F.ab1, Ach1_ACHL0007-09_1_F.ab1, Ach1_ACHL0040-09_1_F.ab1, Ach1_ACHL0041-09_1_F.ab1, Ach1_RBNII384-13_1_F.ab1, Ach1_RBNII395-13_1_F.ab1, Ach1_RBNII396-13_1_F.ab1, and Ach1_RBNII397-13_1_F.ab1 are categorized into forward reads, and Ach1_ACHL0041-09_1_R.ab1, Ach1_ACHL0007-09_1_R. ab1, Ach1_ACHL0040-09_1_R.ab1, Ach1_ACHL0041-09_1_R.ab1, Ach1_RBNII395-13_1_R.ab1, Ach1_ACHL0041-09_1_R.ab1, Ach1_RBNII395-13_1_R.ab1, Ach1_RBNII395-13_1_R.ab1, Ach1_RBNII396-13_1_R.ab1, and Ach1_RBNII397-13_1_R. ab1 are categorized into reverse reads. Then, str_split function is used to split each file-name above into "contig name" and "direction-suffix". Eight contig names are detected in this example which are Ach1_ACHL0006-09, Ach1_ACHL0007-09, Ach1_ACHL0040-09, Ach1_ACHL0041-09,

Achl_RBNII384-13, Achl_RBNII395-13, Achl_RBNII396-13, and Achl_RBNII397-13. Last, a loop iterates through all contigs, and sangeranalseR creates each of them into a *SangerContig* instance. You can check *Advanced User Guide - SangerContig* (*AB1*) to see how sangeranalyseR creates a *SangerContig* instance.

The reason why we strongly recommend you to follow this file-naming regulation is that by doing so, you can directly adopt the example regular expression matching values, "[0-9]*F.ab1\$" and "[0-9]*R.ab1\$", to group reads and reduce chances of error. Everything mentioned above will be done automatically.

After understanding how parameters work, please refer to *Creating SangerAlignment instance from AB1* below to see how sangeranalseR creates *SangerAlignment* instance.

(2) "CSV file matching" SangerAlignment inputs (AB1)

For those who are not familiar with regular expression, we provide a second grouping approach, CSV file matching method. sangeranalyseR will group **AB1** files based on the information in a CSV file automatically. The note below shows the regulations:

Note: Here is an example CSV file (Figure 4)

"reads", "direction", "contig"

Fig. 33: Figure 4. Example CSV file for SangerAlignment instance creation.

- There must be three columns, "reads", "direction", and "contig", in the CSV file.
- The "reads" column stores the filename of AB1 files that are going to be included in the analysis.
- The "direction" column stores the direction of the reads. It must be "F" (forward) or "R" (reverse).

• The "contig" column stores the contig name that each read blongs. Reads in the same contig have to have the same contig name, and they will be grouped into the same contig.

There are two parameters, ABIF_Directory and CSV_NamesConversion, that define the grouping rule to help sangeranalseR to automatically match correct **AB1** files and divide them into forward and reverse directions.

Note:

- ABIF_Directory: this is the directory that contains all **AB1** files, and it can be either an absolute or relative path. We suggest users to put only target AB1 files inside this directory and do not include any other unrelated files.
- CSV_NamesConversion: this is the path to the CSV file. It can be either an absolute or relative path.

The main difference between "CSV file matching" and "regular expression matching" is where the grouping rule is written. For "regular expression matching", rules are writtein in filenames, and thus more naming requirements are required. In contrast, rules of "CSV file matching" are written in an additional CSV file so it is more flexible on **AB1** file-naming.

So how sangeranalyseR works is that it first reads in the CSV file (with "reads", "direction", and "contig" columns), find the names of **AB1** files listed in "reads", group them based on "contig", and assign directions to them based on "direction".

To make it more specific, let's go back to the true example. First, we prepare a CSV file (CSV_NamesConversion) and a file directory like Figure_2 (ABIF_Directory) with AB1 files from In the CSV file, there are 16 rows and 8 distinct contig names. different contigs. sangeranalyseR matches "reads" of these 16 rows to filenames in Allolobophora_chlorotica directory. Then sangeranalyseR groups all matched reads, Achl_ACHL0006-09_1_F.abl, Achl_ACHL0007-09_1_F. ab1, Achl_ACHL0040-09_1_F.ab1, Achl_ACHL0041-09_1_F.ab1, Achl_RBNII384-13_1_F. ab1, Achl RBNII395-13 1 F.ab1, Achl RBNII396-13 1 F.ab1, Achl RBNII397-13 1 F. ab1, Achl_ACHL0006-09_1_R.ab1, Achl_ACHL0007-09_1_R.ab1, Achl_ACHL0040-09_1_R. ab1, Achl_ACHLO041-09_1_R.ab1, Achl_RBNII384-13_1_R.ab1, Achl_RBNII395-13_1_R.ab1, Achl_RBNII396-13_1_R.abl, and Achl_RBNII397-13_1_R.abl, into 8 distinct contig names which are Achl_ACHL0006-09, Achl_ACHL0007-09, Achl_ACHL0040-09, Achl_ACHL0041-09, Achl_RBNII384-13, Achl_RBNII395-13, Achl_RBNII396-13, and Achl_RBNII397-13, by the "contig" column. Last, the directions of reads in each contig are assigned by the "direction" column. Take Ach1_ACHLO041-09 contig as an example. Its "forward read list" will include Ach1_ACHLO041-09_1_F. ab1, and its "reverse read list" will include Ach1 ACHLO041-09 1 R.ab1.

After understanding how parameters work, please refer to *Creating SangerAlignment instance from AB1* below to see how sangeranalseR creates *SangerAlignment* instance.

7.6.2 Creating SangerAlignment instance from AB1

After preparing the input directory, we can create a *SangerAlignment* instance by running *SangerAlignment* constructor function or new method. The constructor function is a wrapper for new method and it makes instance creation more intuitive. Their input parameters are same, and all of them have their default values. For more details about *SangerAlignment* inputs and slots definition, please refer to sangeranalyseR reference manual. We will explain two *SangerAlignment* instance creation methods, "regular expression matching" and "CSV file matching".

(1) "regular expression matching" SangerAlignment creation (AB1)

The consturctor function and new method below contain three parameters, ABIF_Directory, REGEX_SuffixForward, and REGEX_SuffixReverse, that we mentioned in the previous section. It also includes important parameters like quality trimming, chromatogram visualization, consensus alignment, contigs alignment, and so on. Run the following code and create my_sangerAlignment instance.



readingFrame	= 1,
processorsNum	= 2)

In this example, 16 reads are detected and 8 distinct *SangerContig* instances are created. These *SangerContig* instances are stored in a "contig list" in my_sangerAlignment, which will be used as the input for the following functions.

Inside the R shell, you can run my_sangerAlignment to get basic information of the instance or run my_sangerAlignment@objectResults@readResultTable to check the creation result of every Sanger read after my_sangerAlignment is successfully created.

Here is the output of my_sangerAlignment:

CATTCCTAGGAAGI
(

Here is the output of my_sangerAlignment@objectResults@readResultTable:

readName cr	reationResult erro	rType er	rorMessage input	Source	_
⇔direction					
1 Achl_ACHLO006-09_1_F.ab1	TRUE	None	None	ABIF	Forward
⇔Read					
2 Achl_ACHLO006-09_2_R.ab1	TRUE	None	None	ABIF	Reverse
⊶Read					
3 Achl_ACHLO007-09_1_F.ab1	TRUE	None	None	ABIF	Forward
⊶Read					
4 Achl_ACHLO007-09_2_R.ab1	TRUE	None	None	ABIF	Reverse
⇔Read					
5 Achl_ACHLO040-09_1_F.ab1	TRUE	None	None	ABIF	Forward
⇔Read					
6 Achl_ACHLO040-09_2_R.ab1	TRUE	None	None	ABIF	Reverse
→Read					
7 Achl_ACHLO041-09_1_F.ab1	TRUE	None	None	AB1F.	Forward
→Read				3010	D
8 Achl_ACHLO041-09_2_R.ab1 →Read	TRUE	None	None	ABIE	Reverse
9 Achl_RBNII384-13_1_F.ab1	TRUE	None	None	ADTE	Forward
→Read	INUL	None	None	ADIC	roiwaiu
10 Achl_RBNII384-13_2_R.ab1	TRUE	None	None	ABTE	Reverse,
→Read	INOL	None	None	ADIF	Neverse
11 Achl_RBNII395-13_1_F.ab1	TRUE	None	None	ABIE	Forward,
GRead	1100			11011	rorwara
12 Achl_RBNII395-13_2_R.ab1	TRUE	None	None	ABIF	Reverse,
GRead	11.01				
13 Achl_RBNII396-13_1_F.ab1	TRUE	None	None	ABIF	Forward,
→Read					
14 Achl_RBNII396-13_2_R.ab1	TRUE	None	None	ABIF	Reverse
→Read					_
15 Achl_RBNII397-13_1_F.ab1	TRUE	None	None	ABIF	Forward
⇔Read					_
				(es on next nage)

(continues on next page)

				(continued from previous page)
16 Achl_RBNII397-13_2_R.ab1	TRUE	None	None	ABIF Reverse
⇔Read				

(2) "CSV file matching" SangerAlignment creation (AB1)

The constructor function and new method below contain two parameters, ABIF_Directory, and CSV_NamesConversion, that we mentioned in the previous section. It also includes important parameters like quality trimming, chromatogram visualization, consensus alignment, contigs alignment, and so on. Run the following code and create my_sangerAlignment instance.

```
csv_namesConversion <- file.path(rawDataDir, "ab1", "SangerAlignment", "names_</pre>
↔ conversion_all.csv")
# using `constructor` function to create SangerAlignment instance
my_sangerAlignment <- SangerAlignment(inputSource = "ABIF",</pre>
                                                         = "CSV",
                                     processMethod
                                     ABIF_Directory = parentDir,
                                     CSV_NamesConversion = csv_namesConversion,
                                     TrimmingMethod = "M1",
                                     M1TrimmingCutoff = 0.0001,
                                     M2CutoffQualityScore = NULL,
                                     M2SlidingWindowSize = NULL,
                                     baseNumPerRow
                                                          = 100,
                                     heightPerRow
                                                          = 200,
                                     signalRatioCutoff
                                                          = 0.33,
                                     showTrimmed
                                                          = TRUE,
                                     refAminoAcidSeq
↔ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNNI
\rightarrow ",
                                     minReadsNum
                                                          = 2,
                                     minReadLength
                                                        = 20,
                                                        = 0.5,
                                     minFractionCall
                                     maxFractionLost
                                                        = 0.5,
                                                         = GENETIC_CODE,
                                     geneticCode
                                     geneticcode
acceptStopCodons = TR
= 1,
                                                          = TRUE,
                                                          = 1)
                                     processorsNum
# using `new` method to create SangerAlignment instance
my_sangerAlignment <- new("SangerAlignment",</pre>
                                             = "CSV",
                         processMethod
                         processMethod= "CSV",ABIF_Directory= parentDir,
                         CSV_NamesConversion = csv_namesConversion,
                         TrimmingMethod = "M1",
                         MlTrimmingCutoff = 0.0001,
                         M2CutoffQualityScore = NULL,
                         M2SlidingWindowSize = NULL,
                                             = 100,
                         baseNumPerRow
                         heightPerRow
                                              = 200,
                         signalRatioCutoff = 0.33,
                         showTrimmed
                                              = TRUE,
                         refAminoAcidSeq
                                             =
↔ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNNI
\rightarrow ",
```

(continues on next page)

minReadsNum minReadLength minFractionCall maxFractionLost geneticCode acceptStopCodons readingFrame processorsNum	<pre>= 2, = 20, = 0.5, = 0.5, = GENETIC_CODE, = TRUE, = 1, = 1)</pre>
processorsNum	= 1)

First, you need to load the CSV file into the R environment. If you are still don't know how to prepare it, please check (2) "CSV file matching" SangerAlignment inputs (AB1). Then, it will follow rules in the CSV file and create my_sangerAlignment. After it's created, inside the R shell, you can run my_sangerAlignment to get basic information of the instance or run my_sangerAlignment@objectResults@readResultTable to check the creation result of every Sanger read after my_sangerAlignment is successfully created.

Here is the output of my_sangerAlignment:

```
SangerAlignment S4 instance
Input Source : ABIF
Process Method : CSV
ABIF Directory : /Library/Frameworks/R.framework/Versions/4.0/Resources/
Ibrary/sangeranalyseR/extdata/Allolobophora_chlorotica
CSV Names Conversion : /Library/Frameworks/R.framework/Versions/4.0/Resources/
Ibrary/sangeranalyseR/extdata/abl/SangerAlignment/names_conversion_all.csv
Contigs Consensus : ______
TTATAYTTTATTYTRGGCGTCTGAAGCAGGATAGTAGGAGCYGGTATAAGACTCCTAATTCGAATTGAGCTAAGACARCCGGGAGCATTCCTAGGAAGI
SUCCESS [2021-14-07 01:48:28] 'SangerAlignment' is successfully created!
```

readName creatio	nResult err	orType erro	rMessage input	Source 🔒
⇔direction				
1 Achl_ACHLO006-09_1_F.ab1	TRUE	None	None	ABIF Forward
→Read				
2 Achl_ACHLO006-09_2_R.ab1	TRUE	None	None	ABIF Reverse
→Read				
3 Achl_ACHLO007-09_1_F.ab1	TRUE	None	None	ABIF Forward <mark>.</mark>
→Read				
4 Achl_ACHL0007-09_2_R.ab1	TRUE	None	None	ABIF Reverse
→Read				
5 Achl_ACHLO040-09_1_F.ab1	TRUE	None	None	ABIF Forward <mark>.</mark>
→Read				
6 Achl_ACHLO040-09_2_R.ab1	TRUE	None	None	ABIF Reverse
→Read				
7 Achl_ACHLO041-09_1_F.ab1	TRUE	None	None	ABIF Forward <mark>.</mark>
⇔Read				
8 Achl_ACHLO041-09_2_R.ab1	TRUE	None	None	ABIF Reverse
⇔Read				
9 Achl_RBNII384-13_1_F.ab1	TRUE	None	None	ABIF Forward <mark>.</mark>
⇔Read				
10 Achl_RBNII384-13_2_R.ab1	TRUE	None	None	ABIF Reverse
→Read				
11 Achl_RBNII395-13_1_F.ab1	TRUE	None	None	ABIF Forward
→Read	TDIT	Nege	Nege	
12 Achl_RBNII395-13_2_R.ab1	TRUE	None	None	ABIF Reverse
→Read	TRUE	None	None	ADTE Formation
13 Achl_RBNII396-13_1_F.ab1	IKUE	None	None	ABIF Forward_ (continues on next page)
⇔Read				(continues on next page)

Here is the output of my_sangerAlignment@objectResults@readResultTable:

				(continued from previous page)
14 Achl_RBNII396-13_2_R.ab1	TRUE	None	None	ABIF Reverse_
→Read				
15 Achl_RBNII397-13_1_F.ab1	TRUE	None	None	ABIF Forward
⇔Read				
16 Achl_RBNII397-13_2_R.ab1	TRUE	None	None	ABIF Reverse
⇔Read				_

7.6.3 Updating SangerAlignment quality trimming parameters

In the previous *Creating SangerAlignment instance from AB1* part, the constructor function will apply the quality trimming parameters to all reads. After creating a *SangerAlignment* S4 instance, users can change the trimming parameters by running updateQualityParam function which will update all reads with the new trimming parameters and redo reads alignment in *SangerContig* and contigs alignment in *SangerAlignment*. If users want to do quality trimming read by read instead all at once, please read *Launching SangerAlignment Shiny app*.

7.6.4 Launching SangerAlignment Shiny app

We create an interactive local Shiny app for users to go into each *SangerRead* and *SangerContig* in *SangerAlignment* instance. Users only need to run one function with previously created instance as input, my_sangerAlignment, and the *SangerAlignment* Shiny app will pop up. Here, we will go through pages in the three levels.

launchApp(my_sangerAlignment)

SangerAlignment page (SA app)

Figure 5 is the initial page and the toppest layer of *SangerAlignment* App. It provides basic parameters in *SangerAlignment* instance, contigs alignment result and phylogenetic tree etc. Before checking the results, users need to click "Re-calculate Contigs Alignment" button to do contigs alignment in order to get the updated results. From the left-hand side panel, we can clearly see the hierarchy of the *SangerAlignment* S4 instance and easily access to all reads and contigs in it.

Scroll down a bit, users can see the contigs alignment result generated by DECIPHER R package embedded in *Sanger-Alignment* page. *Figure 6* shows the contigs alignment result.

In *SangerAlignment* page, the phylogenetic tree result is provided as well (*Figure 7*). The tree is generated by ape R package which uses neighbor-joining algorithm.

sangeranalyseR	■ SangerAlignment	Overview Page	Save S4 instance	Close UI
☆ SangerAlignment Overview	Input Parameters:			-
≡1 SangerContig 🗸				
■ 1 SangerContig Overview		🖬 Re-calculate Contigs Alignment		
🕒 Forward SangerReads 🗸				
- 1 - 1 Forward SangerRead	Output Directory:	/var/folders/33/7v38jdjd2874jcxb6l71m00h0000gn/T//RtmpPG9k9i		
Reverse SangerReads	Raw ABI Parent	/Users/chaokuan-hao/Desktop/tmp		
≣ 2 SangerContig ∢	Directory:			
	Trimming Method:	Method 1: 'Modified Mott Trimming'		
≣3 SangerContig <	Forward Suffix RegExp:	_[0-9]*_F.ab1		
Ξ4 SangerContig ζ	Reverse Suffix RegExp:	_[0-9]*_R.ab1		
	Contigs Number:	13		
≣5 SangerContig <	Alignment Parameters			-
≣6 SangerContig <				
≣7 SangerContig ζ	0.5 MinFractionCall	0.5 MaxFractionLost		

Fig. 34: Figure 5. SangerAlignment Shiny app initial page - SangerAlignment Page.

Contigs Alignment	
	20 40
extdata/Allolobophora_chlorotica/ACHLO/Achl_ACHLO006-09	TATATCGACGGCCAGTGGTCAACAAATCATAAAGATATTGGAACTTTATATTTTA
extdata/Allolobophora_chlorotica/ACHLO/Achl_ACHL0007-09	CGCGGCCAG <mark>TGGTCAACAAATCATAAAG</mark> ATATT <mark>GG</mark> AACTTTATATTTTA
extdata/Allolobophora_chlorotica/ACHLO/Achl_ACHL0040-09	TAAACCGACGGACAGTGGTCAACAAATCATAAAGATATTGGAACTCTATACTTTA
extdata/Allolobophora_chlorotica/ACHLO/Achl_ACHL0041-09	ATCTGCGGCCAGTGGTCAACAAATCATAAAGATATCGGAACTCTATACTTTA
extdata/Allolobophora_chlorotica/RBNII/Achl_RBNII384-13 extdata/Allolobophora chlorotica/RBNII/Achl RBNII395-13	GATATT <mark>GG</mark> AACTCTATACTTTA GATATCAGACTCTATACTTTA
extdata/Allolobophora_chlorotica/RBNII/Achl RBNII396–13	GATATCGGAACTCTATACTTTA
extdata/Allolobophora_chlorotica/RBNII/Achl_RBNII397-13	GATATAGGAACTCTATACTTTA
extdata/Drosophila_melanogaster/Dmel_BBDCN941-10	<mark>G</mark> ATA <mark>TTGG</mark> ACCTTTATATTTTA
extdata/Drosophila_melanogaster/Dmel_BBDEE689–10	T <mark>GG</mark> TCAACAAATCATAAA <mark>G</mark> ATATT <mark>GG</mark> AACTTTATATTTTA
extdata/Drosophila_melanogaster/Dmel_PHDIP946-11	<mark>GTCCGCG</mark> ACCA <mark>GTGG</mark> TCAACAAATCATAAA <mark>G</mark> ATATTGGAACTITATATTTTA
extdata/Drosophila_melanogaster/Dmel_TDWGB557-10 extdata/Drosophila melanogaster/Dmel TDWGB669-10	AAGATATTGGAACTTTATATTTTA AAGATATTGGAACTTTATATTTTA

Fig. 35: Figure 6. SangerAlignment Page - contigs alignment result.

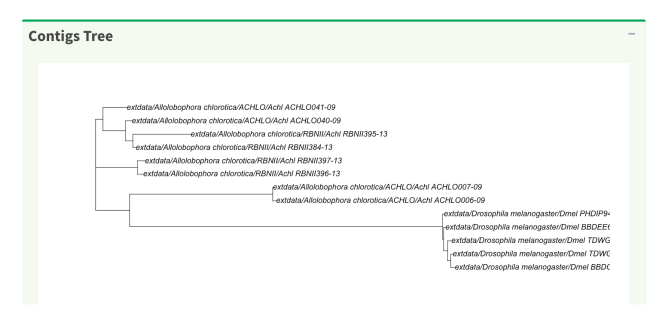


Fig. 36: Figure 7. SangerAlignment Page - phylogenetic tree result.

SangerContig page (SA app)

Now, let's go to the page in the next level, *SangerContig* page. Users can click into all contigs and check their results. *Figure 8* shows the overview page of Contig 1. Notice that there is a red "Re-calculate Contig" button. After changing the quality trimming parameters, users need to click the button before checking the results below in order to get the updated information.

The information provided in this page includes : "input parameters", "genetic code table", "reference amino acid sequence", "reads alignment", "difference data frame", "dendrogram", "sample distance heatmap", "indels data frame", "stop codons data frame". *Figure 9* and *Figure 10* show part of the results in the *SangerContig* page. The results are dynamic based on the trimming parameters from user inputs.

SangerRead page (SA app)

Now, let's go to the page in the lowest level, *SangerRead* page. *SangerRead* page contains all details of a read including its trimming and chromatogram inputs and results. All reads are in "forward" or "reverse" direction. Under "Contig Overview" tab (*SangerContig* page), there are two expendable tabs, "Forward Reads" and "Reverse Reads" storing corresponding reads on the left-hand side navigation panel in *Figure 11*. In this example, there are one read in each tab and *Figure 11* shows the "1 - 1 Forward Read" page. It provides basic information, quality trimming inputs, chromatogram plotting inputs etc. Primary/secondary sequences in this figure are dynamic based on the signalRatioCutoff value for base calling and the length of them are always same. Another thing to mention is that primary/secondary sequences and the sequences in the chromatogram in *Figure 16* below will always be same after trimming and their color codings for A/T/C/G are same as well.

In quality trimming steps, we removes fragment at both ends of sequencing reads with low quality score. It is important because trimmed reads will improves alignment results. *Figure 12* shows the UI for Trimming Method 1 (M1): 'Modified Mott Trimming'. This method is implemented in Phred. Users can change the cutoff score and click "Apply Trimming Parameters" button to update the UI. The value of input must be between 0 and 1. If the input is invalid, the cutoff score will be set to default 0.0001.

Figure 13 shows another quality trimming methods for users to choose from, Trimming Method 2 (M2): 'Trimmomatics Sliding Window Trimming'. This method is implemented in Trimmomatics. Users can change the cutoff quality score as well as sliding window size and click "Apply Trimming Parameters" button to update the UI. The value of

sangeranalyseR	■ 1 SangerContig O	verview Page	Save S4 instance	Close UI
☆ SangerAlignment Overview	• Basic Information:			-
≡1 SangerContig <				
1 SangerContig Overview		🖬 Re-calculate Contig		
● Forward SangerReads <				
Reverse SangerReads <	Output Directory:	/var/folders/33/7v38jdjd2874jcxb6l71m00h0000gn/T//RtmpPG9k9i		
≣2 SangerContig <	▶ Raw ABI Parent Directory:	/Users/chaokuan-hao/Desktop/tmp		
≣3 SangerContig <	▶ Consenesus Read Name:	Achl_ACHLO006-09		
≣4 SangerContig <	Trimming Method:	Method 1: 'Modified Mott Trimming'		
≣5 SangerContig <	Forward Suffix RegExp:	_[0-9]*_F.ab1		
	Forward Read Number:	1		
≣6 SangerContig <	Reverse Suffix RegExp:	_[0-9]*_R.ab1		
Ξ7 SangerContig <	▶ Reverse Read Number:	1		
≣8 SangerContig <	Contis Devendors			_

Fig. 37: Figure 8. SangerAlignment Shiny app - SangerContig page.

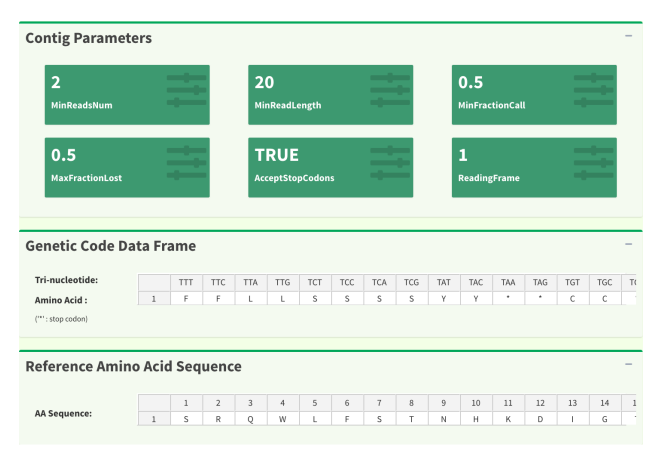
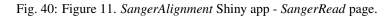


Fig. 38: Figure 9. SangerContig page - contig-related parameters, genetic code and reference amino acid sequence.

• Consensus Read Results: Alignment 20 40 60 111 1_Read_Achl_ACHL0006-09_1_F.ab1 CTC **GCGTCTGAGCAGG**A 2_Read_Achl_ACHL0006-09_2_R.ab1 GCGTCTGAGCAGGA TATATCGACGGCCACTGGTCAACAAATCATAAAGATATTGGAACTTTATATTTTATTCTG Consensus **GCGTCTG<mark>AG</mark>CA** GA TATATCGACGGCCAGTGGTCAACAAATCATAAAGATATTGGAACTTTATATTTTATTCTGGGCGTCTGAGCAGGA Consensus **Differences Data frame** Show 10 \$ entries Search: name pairwise.diffs.to.consensus unused.chars 1 1_Read_Achl_ACHLO006-09_1_F.ab1 0 0 0 2 2_Read_Achl_ACHLO006-09_2_R.ab1 0 Showing 1 to 2 of 2 entries Previous 1 Next

Fig. 39: Figure 10. SangerContig page - reads alignment and difference data frame.

sangeranalyseR	■ 1 SangerContig - 1 Forward SangerRead Page Save S4 instance Close U
☆ SangerAlignment Overview	• Raw File:
≣1 SangerContig <	Achl_ACHLO006-09_1_F.ab1
😑 1 SangerContig Overview	$(full\ path: /Users/chaokuan-hao/Desktop/tmp/extdata/Allolobophora_chlorotica/ACHLO/Achl_ACHLO006-09_1_F.ab1)$
🔵 Forward SangerReads 🔏	
 1 - 1 Forward SangerRead 	● Primary, Secondary DNA Sequences & Amino Acid Sequence (Before
Reverse SangerReads <	Trimming):
≣2 SangerContig <	
≣ 3 SangerContig <	≡ Primary Sequence
	1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25
Ξ4 SangerContig	1 C A C T T T A T A T T T A T C T G G G C G T C
≣5 SangerContig ζ	≡ Secondary Sequence
Ξ6 SangerContig ζ	1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25
$\equiv 6$ SangerContig \checkmark	1 TACTTTATATTTTATCTGGGCGTC



Quality Report:		
Trimming Paramet	ers Input	-
Your trimming method sele	ction : 'Modified Mott Trimming'	
1e-04 Cutoff Score	X	
Input Value		
1e-04		
	Apply Trimming Parameters	

Fig. 41: Figure 12. SangerRead page - Trimming Method 1 (M1): 'Modified Mott Trimming' UI.

cutoff quality score must be between 0 and 60 (default 20); the value of sliding window size must be between 0 and 40 (default 10). If the inputs are invalid, their values will be set to default.

rimming Parameters Input ur trimming method selection : 'Trimmomatics Sliding Window Trimming' 20 15 curation litra Series	ur trimming method selection : 'Trimmomatics Sliding Window Trimming'				
20 15	20 IS Stiding Window Size	rimming Paramete	ers Input		
	Cutoff Quality Score	ur trimming method selec	ction : 'Trimmomatics Slidin	ng Window Trimming'	
			Ж		5.3

Fig. 42: Figure 13. SangerRead page - Trimming Method 2 (M2): 'Trimmomatics Sliding Window Trimming' UI.

Figure 14 shows the quality report before and after trimming. After clicking the "Apply Trimming Parameters" button, the values of these information boxes will be updated to the latest values.

In *Figure 15*, the x-axis is the index of the base pairs; the y-axis is the Phred quality score. The green horizontal bar at the top of the plot is the raw read region and the orange horizontal bar represents the trimmed read region. Both *Figure 15* trimming plot and *Figure 16* chromatogram will be updated once users change the quality trimming parameters and click the "Apply Trimming Parameters" button in *Figure 16*.

If we only see primary and secondary sequences in the table, we will loose some variations. Chromatogram is very helpful to check the peak resolution. *Figure 16* shows the panel of plotting chromatogram. Users can change four parameters: Base Number Per Row, Height Per Row, Signal Ratio Cutoff, and Show Trimmed Region. Among them, Signal Ratio Cutoff is the key parameter. If its value is default value 0.33, it indicates that the lower peak should be at least 1/3rd as high as the higher peak for it count as a secondary peak.

Here is an example of applying new chromatogram parameters. We click "Show Trimmed Region" to set its value from FALSE to TRUE. *Figure 17* shows the loading notification popup during base calling and chromatogram plotting.

After replotting the chromatogram, trimmed region is showed in red striped region. *Figure 18* shows part of the the chromatogram (1 bp ~ 240 bp). Moreover, chromatogram will be replotted when trimmed positions or chromatogram

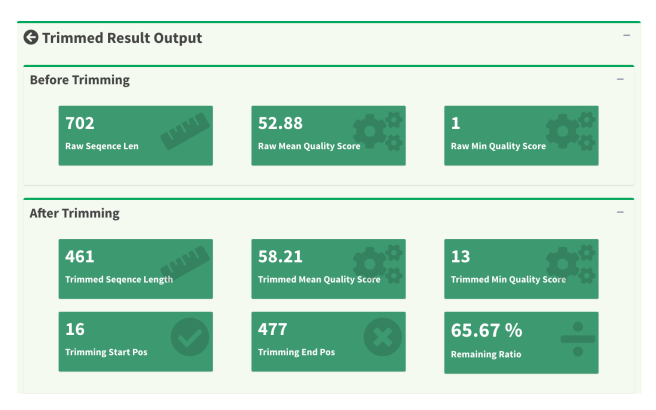


Fig. 43: Figure 14. SangerRead page - read quality report before / after trimming.



Fig. 44: Figure 15. SangerRead page - quality trimming plot.

Chromatogram Input			
Base Number Per Row			
5 80 20	。 ⊃ Signal Ratio Cutoff	21	466
5 25 45 65 85 105 125 145 165 185 200		Trimming Start Pos	Trimming End Pos
Height Per Row			
50 200 60	Whether show trimmed region		
50 105 160 215 270 325 380 435 490 545 60			
	Apply Chroma	atogram Parameters	
Chromatogram Output			
Chromatogram Output			

Fig. 45: Figure 16. SangerRead page - chromatogram panel.

Chromatogram:		
Chromatogram Input		
Base Number Per Row		
S 200 5 25 45 55 105 125 145 165 105 200 S 26 45 55 105 105 125 145 165 105 200	Z Plotting Chromatogram	466 Trimming End Pos
0. Height Per Row	Base pairs per row : 80	
50 200 600	• Height per row : 200	
50 105 160 215 270 325 380 435 490 545 600	Signal Ratio Cutoff : 0.33	
	• Show trimmed : TRUE	
	(If 'Signal Ratio Cutoff' is too small, it would need more time to replot the chromatogram)	
Chromatogram Output		

Fig. 46: Figure 17. SangerRead page - loading notification popup during replotting chromatogram.

parameters are updated.

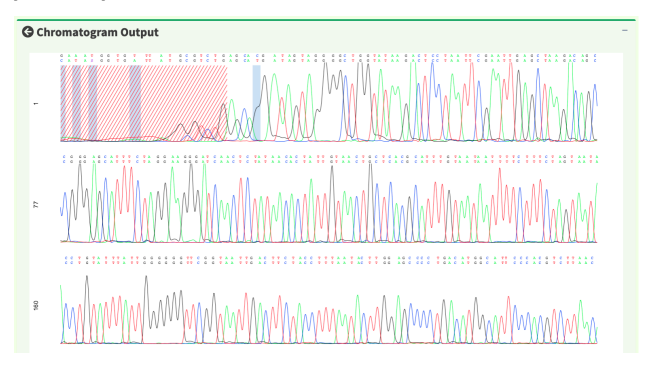


Fig. 47: Figure 18. SangerRead page - chromatogram with trimmed region showed.

To let users browse the trimmed primary/secondary sequences without finding "Trimming Start Point" and "Trimming End Point" by themselves, we provide the final trimmed primary/secondary sequences that will be used for reads alignment in table format with quality scores in *Figure 19*. Frameshift amino acid sequences are also provided.

We have updated the trimming and chromatogram parameters for each read. Now, we need to click "Re-calculate contig" button to do alignment again. Last but not least, we can save all data into a new 'SangerContig' S4 instance by clicking "Save S4 instance button". New S4 instance will be saved in **Rda** format. Users can run readRDS function to load it into current R environment. *Figure 20* shows some hints in the save notification popup.

7.6.5 Writing SangerAlignment FASTA files (AB1)

Users can write the *SangerAlignment* instance, my_sangerAlignment, to **FASTA** files. There are four options for users to choose from in selection parameter.

- contigs_unalignment: Writing contigs into a single FASTA file.
- contigs_alignment: Writing contigs alignment and contigs consensus read to a single FASTA file.
- all_reads: Writing all reads to a single FASTA file.
- all: Writing contigs, contigs alignment, and all reads into three different files.

Below is the oneliner for writing out FASTA files. This function mainly depends on writeXStringSet function in Biostrings R package. Users can set the compression level through writeFasta function.



Fig. 48: Figure 19. SangerRead page - trimmed primary/secondary sequences and Phred quality score in table format.

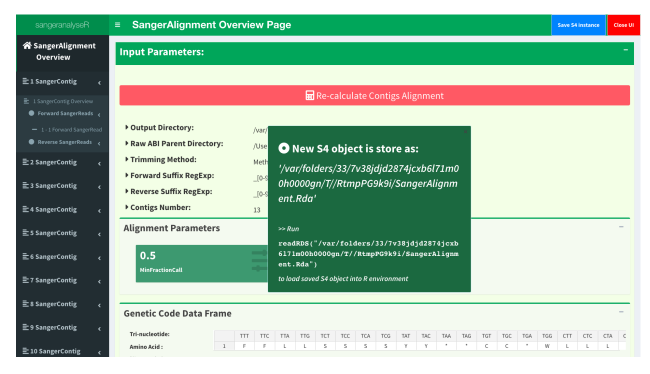


Fig. 49: Figure 20. SangerRead page - saving notification popup.

Users can download the output FASTA file of this example through the following three links:

- (1) Sanger_contigs_unalignment.fa
- (2) Sanger_contigs_alignment.fa
- (3) Sanger_all_trimmed_reads.fa

7.6.6 Generating SangerAlignment report (AB1)

Last but not least, users can save *SangerAlignment* instance, my_sangerAlignment, into a report after the analysis. The report will be generated in **HTML** by knitting **Rmd** files.

Users can set includeSangerContig and includeSangerRead parameters to decide to which level the *SangerAlignment* report will go. Moreover, after the reports are generated, users can easily navigate through reports in different levels within the **HTML** file.

One thing to pay attention to is that if users have many reads, it will take quite a long time to write out all reports. If users only want to generate the contig result, remember to set includeSangerRead and includeSangerContig to FALSE in order to save time.

Here is the generated SangerAlignment html report of this example (ABIF). Users can access to 'Basic Information', 'Contigs Consensus', 'Contigs Alignment', 'Contigs Tree', and 'Contig Reports' sections inside it. Furthermore, users can also navigate through html reports of all contigs and forward and reverse SangerRead in this SangerAlignment report.

7.6.7 Code summary (SangerAlignment, AB1)

(1) Preparing SangerAlignment AB1 inputs

```
rawDataDir <- system.file("extdata", package = "sangeranalyseR")
parentDir <- file.path(rawDataDir, 'Allolobophora_chlorotica')</pre>
```

(2) Creating SangerAlignment instance from AB1

(2.1) "Regular Expression Method" SangerAlignment creation (AB1)

```
# using `constructor` function to create SangerAlignment instance
my_sangerAlignment <- SangerAlignment(inputSource</pre>
                                                        = "ABIF",
                                                         = "REGEX",
                                     processMethod
                                     ABIF_Directory = parentDir,
                                     REGEX_SuffixForward = "[0-9] \star F.ab1\$",
                                     REGEX_SuffixReverse = "[0-9] *_R.ab1$",
                                     refAminoAcidSeq
↔ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNN
→ " )
# using `new` method to create SangerAlignment instance
my_sangerAlignment <- new("SangerAlignment",</pre>
                                              = "ABIF",
                         inputSource
                                             = "REGEX",
                         processMethod
                         ABIF_Directory = parentDir,
                         REGEX_SuffixForward = "[0-9] *_F.ab1$",
                         REGEX_SuffixReverse = "[0-9] *_R.ab1$",
                         refAminoAcidSeq
→ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNNI
→ " )
```

Following is the R shell output that you will get.

(2.2) "CSV file matching" SangerAlignment creation (AB1)

```
csv_namesConversion <- file.path(rawDataDir, "ab1", "SangerAlignment", "names_
⇔conversion_all.csv")
# using `constructor` function to create SangerAlignment instance
my_sangerAlignment <- SangerAlignment(inputSource = "ABIF",</pre>
                                                           = "CSV",
                                      processMethod
                                      ABIF_Directory
                                                           = parentDir,
                                      CSV_NamesConversion = csv_namesConversion,
                                      refAminoAcidSeq
→ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNN
→ " )
# using `new` method to create SangerAlignment instance
my_sangerAlignment <- new("SangerAlignment",</pre>
                                               = "CSV",
                          processMethod
                          ABIF_Directory
                                               = parentDir,
                          CSV_NamesConversion = csv_namesConversion,
                          refAminoAcidSeq
→ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNNI
→ " )
```

Following is the R shell output that you will get.

(3) Updating SangerAlignment quality trimming parameters (AB1)

(4) Launching SangerAlignment Shiny app (AB1)

launchApp(my_sangerAlignment)

(5) Writing SangerAlignment FASTA files (AB1)

writeFasta(my_sangerAlignment)

Following is the R shell output that you will get.

You will get three FASTA files:

- (1) Sanger_contigs_unalignment.fa
- (2) Sanger_contigs_alignment.fa
- (3) Sanger_all_trimmed_reads.fa

(6) Generating SangerAlignment report (AB1)

generateReport(my_sangerAlignment)

You can check the html report of this SangerAlignment example (ABIF).

7.7 Advanced User Guide - SangerRead (FASTA)

SangerRead is in the bottommost level of sangeranalyseR (*Figure_1*), and each SangerRead object corresponds to a single read in Sanger sequencing. In this section, we are going to go through detailed sangeranalyseR data analysis steps in SangerRead level with FASTA file input.



Fig. 50: Figure 1. Hierarchy of classes in sangeranalyseR, SangerRead level.

7.7.1 Preparing SangerRead FASTA input

The **FASTA** input method is designed for those who do not want to do quality trimming and base calling on their Sanger sequencing data; therefore, no quality trimming and chromatogram input parameters are needed. Before starting the analysis, users need to prepare a **FASTA** file, and in this example, it is in the sangeranalyseR package; thus, you can simply get its path by running the following codes:

The only hard regulation of the filename, $Achl_ACHLO006-09_1_F$. fa in this example, is that file extension must be .fasta or .fa.

7.7.2 Creating SangerRead instance from FASTA

After preparing an input **FASTA** file, the next step is to create a *SangerRead* instance by running SangerRead constructor function or new method. The constructor function is a wrapper for new method which makes instance creation more intuitive. All of the input parameters have their default values. We list important parameters in the two *SangerRead* creation methods below. readFileName stores the **FASTA** filename, and inside it, the string in the first line after ">" is the name of the read. Users need to assign the name of the read to fastaReadName which is used for read-matching. *Figure 2* is a valid **FASTA** file, Achl_ACHL0006-09_1_F.fa (example FASTA file), and the value of fastaReadName is Achl_ACHL0006-09_1_F.

Fig. 51: Figure 2. SangerRead FASTA input file.

The inputs of SangerRead constructor function and new method are the same. For more details about *SangerRead* inputs and slots definition, please refer to sangeranalyseR reference manual.

Inside the R shell, you can run sangerReadFfa to get basic information of the instance or run sangerReadFfa@objectResults@readResultTable to check the creation result of every Sanger read after sangerReadFfa is successfully created.

Here is the output of sangerReadFfa:

```
SangerRead S4 instance

Input Source : FASTA

Read Feature : Forward Read

Read FileName : Achl_ACHLO006-09_1_F.fa

Fasta Read Name : Achl_ACHLO006-09_1_F

Primary Sequence : ______

→CTGGGCGTCTGAGCAGGAATGGTTGGAGCCGGTATAAGACTTCTAATTCGAATCGAGCTAAGACAACCAGGAGCGTTCCTGGGCAGAACCAACTATAG

SUCCESS [2021-12-07 23:37:43] 'Achl_ACHLO006-09_1_F.fa' is successfully created!
```

Here is the output of sangerReadFfa@objectResults@readResultTable:

readName	creationResult	errorType	errorMessage	inputSource	direction
1 Achl_ACHL0006-09_1_1	F TR	UE Nor	Nor	E FASTA	Forward Read

7.7.3 Writing SangerRead FASTA files (FASTA)

Users can write sangerReadFfa to a FASTA file. Because the FASTA input method does not support quality trimming or base calling, in this example, the sequence of the output FASTA file will be the same as the input FASTA file. Moreover, users can set the compression level through the one-liner, writeFasta, which mainly depends on writeXStringSet function in Biostrings R package.

Users can download the Achl_ACHLO006-09_1_F.fa of this example.

7.7.4 Generating SangerRead report (FASTA)

Last but not least, users can save sangerReadFfa into a static HTML report by knitting Rmd files. In this example, tempdir function will generate a random path.

SangerRead_Report_fasta.html is the generated *SangerRead* report html of this example. Users can access to '*Basic Information*', '*DNA Sequence*' and '*Amino Acids Sequence*' sections inside this report.

7.7.5 Code summary (SangerRead, fasta)

(1) Preparing SangerRead FASTA input

(2) Creating SangerRead instance from FASTA

Following is the R shell output that you will get.

(3) Writing SangerRead FASTA files (FASTA)

writeFasta(sangerReadFfa)

Following is the R shell output that you will get.

And you will get one FASTA file:

(1) Achl_ACHL0006-09_1_F.fa

(4) Generating SangerRead report (FASTA)

generateReport(sangerReadFfa)

You can check the html report of this SangerRead example (FASTA).

7.8 Advanced User Guide - SangerContig (FASTA)

SangerContig is in the intermediate level of sangeranalyseR (*Figure_1*), and each *SangerContig* instance corresponds to a contig in a Sanger sequencing experiment. Among its slots, there are two lists, forward and reverse read list, storing *SangerRead* in the corresponding direction.

In this section, we are going to go through details about a reproducible *SangerContig* analysis example with the **FASTA** file input in sangeranalyseR. By running the following example codes, you will get an end-to-end SangerContig analysis result.

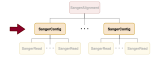


Fig. 52: Figure 1. Hierarchy of classes in sangeranalyseR, SangerContig level.

7.8.1 Preparing SangerContig FASTA input

In *Advanced User Guide - SangerContig (AB1)*, we demonstrated how to use **AB1** input files to create *SangerContig* instance. Here, we explain another input format - the **FASTA** input. Before starting the analysis, users need to prepare one **FASTA** file, which must end with **.fa** or **.fasta**, containing sequences of all reads. In this example, the **FASTA** file is in the sangeranalyseR package, and you can simply get its path by running the following codes:

```
rawDataDir <- system.file("extdata", package = "sangeranalyseR")
fastaFN <- file.path(rawDataDir, "fasta", "SangerContig", "Achl_ACHL0006-09.fa")</pre>
```

The value of fastaFN is where the **FASTA** file is placed. If your operating system is macOS, then its value should look like this:

And we showed the reads in fastaFN in Figure_2 (example FASTA file):

>Achl_ACHL0006-09_1_F
${\tt CTGGGCGTCTGAGCAGGAATGGTTGGAGCCGGTATAAGACTTCTAATTCGAATCGAGCTAAGACAACCAGGAGCGTTCCT}$
GGGCAGAGACCAACTATACAATACTATCGTTACTGCACACGCATTTGTAATAATCTTCTTTCT
${\tt TCGGGGGGATTCGGAAACTGGCTTTTACCTTTAATACTTGGAGCCCCCGATATAGCATTCCCTCGACTCAACAACATGAGA$
${\tt TTCTGACTACTTCCCCCATCACTGATCCTTTTAGTGTCCTCTGCGGCGGTAGAAAAAGGCGCTGGTACGGGGTGAACTGT}$
${\tt TTATCCGCCTCTAGCAAGAAATCTTGCCCACGCAGGCCCGTCTGTAGATTTAGCCATCTTTTCCCTTCATTTAGCGGGTG$
CGTCTTCTATTCTAGGGGGCTATTAATTTTATCACCACAGTTATTAATATGCGTTGAAGAGG
>Achl_ACHL0006-09_2_R
${\tt GAGGATGGGGTCTCCACCACCGGCAGGATCAAAGAATGAAGTATTGAGGTTTCGGTCGG$
${\tt CTGCTAGCACTGGTAAAGATAGAAGTAGAAGAAGAACAACTGTAATTAGCACAGCTCAGACAAACAGGGGAATTCGTTCAAGA}$
${\tt CGTAATCCTCTTCAACGCATATTAATAACTGTGGTGATAAAATTAATAGCCCCTAGAATAGAAGACGCACCCGCTAAATG}$
${\tt AAGGGAAAAGATGGCTAAATCTACAGACGGGCCTGCGTGGGCAAGATTTCTTGCTAGAGGCCGGATAAACAGTTCACCCCG}$
${\tt TACCAGCGCCTTTTTCTACCGCCGCAGAGGACACTAAAAGGATCAGTGATGGGGGAAGTAGTCAGAATCTCATGTTGTTG$
${\tt AGTCGAGGGAATGCTATATCGGGGGGCTCCAAGTATTAAAGGTAAAAGCCAGTTTCCGAATCCCCCGATGAATACAGGCAT}$
${\tt TACTAGAAAGAAGATTATTACAAATGCGTGTGCAGTAACGATAGTATTGTATAGTTGGTCTCTGCCCAGGAACGCTCCTG$
${\tt GTTGTCTTAGCTCGATTCGAATTAGAAGTCTTATACCGGCTCCAACCATTCCTGCTCAGACGCCCAGAATAAAATATAAA$

Fig. 53: Figure 2. SangerContig FASTA input file.

Inside the **FASTA** file (*Figure_2*; Ach1_ACHLO006-09.fa), the strings starting with ">" before each read are the read names. There are two ways of grouping reads which are "**regular expression matching**" and "**CSV file matching**", and following are instructions of how to prepare your **FASTA** input file.

(1) "regular expression matching" SangerContig inputs (FASTA)

For regular expression matching method, sangeranalyseR will group reads based on their contig name and read direction in their names automatically; therefore, users have to follow the read-naming regulations below:

Note:

- All reads in the same contig group must include the same contig name in their read names.
- Forward or reverse direction also has to be specified in their read names.

There are four parameters, FASTA_File, contigName, REGEX_SuffixForward and REGEX_SuffixReverse, that define the grouping rule to let sangeranalyseR automatically match correct reads in **FASTA** file and divide them into forward and reverse directions.

Note:

- FASTA_File: this is the path to **FASTA** file that contains all sequences of reads, and it can be either an absolute or relative path. We suggest users to include only target reads inside this **FASTA** file and do not include any other unrelated reads.
- contigName: this is a regular expression that matches read names that are going to be included in the *Sanger-Contig* analysis. grepl function in R is used.
- REGEX_SuffixForward: this is a regular expression that matches all read names in forward direction. grep1 function in R is used.
- REGEX_SuffixReverse: this is a regular expression that matches all read names in reverse direction. grep1 function in R is used.

If you don't know what regular expression is, don't panic - it's just a way of recognising text. Please refer to *What is a regular expression?* for more details. Here is an example of how it works in sangeranalseR:

So how sangeranalyseR works is that it first matches the contigName to exclude unrelated files and then separate the forward and reverse reads by matching REGEX_SuffixForward and REGEX_SuffixReverse. Therefore, it is important to make sure that all target reads in the **FASTA** file share the same contigName and carefully select your REGEX_SuffixForward and REGEX_SuffixReverse. The bad file-naming and wrong regex matching might accidentally include reverse reads into the forward read list or vice versa, which will make the program generate wrong results. Therefore, it is important to have a consistent naming strategy. So, how should we systematically name the reads? We suggest users to follow the file-naming regulation in *Figure_3*.

[Consensus Read Name] + + [index] + + [F,R]

Fig. 54: Figure 3. Suggested read naming regulation in FASTA file - SangerContig.

As you can see, the first part of the regulation is a consensus read name (or contig name), which narrows down the scope of reads to those we are going to examine. The second part of the regulation is an index. Since there might be more than one read that is in the forward or reverse direction, we recommend you to number your reads in the same contig group. The last part is a direction which is either 'F' (forward) or 'R' (reverse).

To make it more specific, let's go back to the true example. In *Figure_2*, there are two reads in the FASTA file (fasta_FN). First, we set contigName to "Achl_ACHL0006-09" to confirm that two of them, Achl_ACHL0006-09_1_F and Achl_ACHL0006-09_2_R, contain our target contigName and should be included. Then, we set REGEX_SuffixForward to "[0-9]*F\$" and REGEX_SuffixReverse to "[0-9]*R\$" to let sangeranalyseR match and group forward and reverse reads automatically. By the regular expression rule, Achl_ACHL0006-09_1_F and Achl_ACHL0006-09_2_R will be categorized into "forward read

list" and "reverse read list" respectively. The reason why we strongly recommend you to follow this file-naming regulation is that by doing so, you can directly adopt the example regular expression matching values, " $[0-9] \star F$ " and " $[0-9] \star R$ ", to group reads and reduce chances of error.

After understanding how parameters work, please refer to *Creating SangerContig instance from FASTA* below to see how sangeranalseR creates 'Achl_ACHLO006-09' *SangerContig* instance.

(2) "CSV file matching" SangerContig inputs (FASTA)

No doubt that read names in the original **FASTA** file do not follow the naming regulation, and you do not want to change the original **FASTA** file; thus, we provide a second grouping approach, CSV file matching method. sangeranalyseR will group reads in the **FASTA** file based on the information in a CSV file automatically, and users do not need to alter the read names in the **FASTA** file; therefore, users have to follow the regulations below:

Note: Here is an example CSV file (Figure_4)

```
"reads","direction","contig"
"Achl_ACHL0006-09_1_F","F","Achl_ACHL0006-09"
"Achl_ACHL0006-09_2_R","R","Achl_ACHL0006-09"
```

Fig. 55: Figure 4. Example CSV file for SangerContig instance creation.

- There must be three columns, "reads", "direction", and "contig", in the CSV file.
- The "reads" column stores the read names in the FASTA file that are going to be included in the analysis.
- The "direction" column stores the direction of the reads. It must be "F" (forward) or "R" (reverse).
- The "contig" column stores the contig name that each read blongs. Reads in the same contig have to have the same contig name, and they will be grouped into the same *SangerContig* instance.

There are three parameters, FASTA_File, contigName, and CSV_NamesConversion, that define the grouping rule to help sangeranalseR to automatically match correct reads in a **FASTA** file and divide them into forward and reverse directions.

Note:

- FASTA_File: this is the path to **FASTA** file that contains all sequences of reads, and it can be either an absolute or relative path. We suggest users to include only target reads inside this **FASTA** file and do not include any other unrelated reads.
- contigName: this is a regular expression that matches read names that are going to be included in the *Sanger-Contig* analysis. grepl function in R is used.
- CSV_NamesConversion: this is the path to the CSV file. It can be either an absolute or relative path.

The main difference between "CSV file matching" and "regular expression matching" is where the grouping rule is written. For "regular expression matching", rules are writtein in read names, and thus more naming requirements are required. In contrast, rules of "CSV file matching" are written in an additional CSV file so it is more flexible on naming reads.

So how sangeranalyseR works is that it first reads in the CSV file (with "*reads*", "*direction*", and "*contig*" columns), filter out rows whose "*contig*" is not the value of contigName parameter, find the read names in the **FASTA** file listed in "*reads*", and assign directions to them based on "*direction*".

To make it more specific, let's go back to the true example. First, we prepare a CSV file (CSV_NamesConversion) and a FASTA file (FASTA_File). In the CSV file, both rows have the contig name "Achl_ACHL0006-09", which is what we need to assign to the contigName parameter. sangeranalyseR then checks and matches "*reads*" of these two rows, "Achl_ACHL0006-09_1_F" and "Achl_ACHL0006-09_2_R". Last, these two reads are assigned into "forward read list" and "reverse read list" respectively by the "*direction*" column.

After understanding how parameters work, please refer to *Creating SangerContig instance from FASTA* below to see how sangeranalseR creates 'Achl_ACHLO006-09' *SangerContig* instance.

7.8.2 Creating SangerContig instance from FASTA

After preparing the input directory, we can create a *SangerContig* instance by running SangerContig constructor function or new method. The constructor function is a wrapper for new method and it makes instance creation more intuitive. Their input parameters are same, and all of them have their default values. For more details about *SangerContig* inputs and slots definition, please refer to sangeranalyseR reference manual. We will explain two *SangerContig* instance creation methods, "regular expression matching" and "CSV file matching".

(1) "regular expression matching" SangerContig creation (FASTA)

The consturctor function and new method below contain four parameters, FASTA_File, contigName, REGEX_SuffixForward, and REGEX_SuffixReverse, that we mentioned in the previous section. In contrast to **AB1** input method, it does not include quality trimming and chromatogram visualization parameters. Run the following code and create my_sangerContigFa instance.

```
# using `constructor` function to create SangerRead instance
my_sangerContigFa <- SangerContig(inputSource</pre>
                                                        = "FASTA",
                                                        = "REGEX",
                                  processMethod
                                  FASTA_File
contigName
                                                        = fastaFN,
                                                        = "Achl_ACHL0006-09",
                                  REGEX_SuffixForward = "[0-9] \star F \$",
                                  REGEX_SuffixReverse = "[0-9] * R;",
                                  refAminoAcidSeq
→ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNNI
\rightarrow ",
                                                        = 2,
                                  minReadsNum
                                  minReadLength
                                                        = 20,
                                  minFractionCall
                                                        = 0.5,
                                  maxFractionLost
                                                        = 0.5,
                                  geneticCode
                                                        = GENETIC_CODE,
                                  acceptStopCodons
                                                        = TRUE,
                                  readingFrame
                                                        = 1,
                                  processorsNum
                                                        = 1)
# using `new` method to create SangerRead instance
my_sangerContigFa <- new("SangerContig",</pre>
                         inputSource
processMethod
                                               = "FASTA",
                                             = "REGEX",
                         FASTA_File
contigName
                                              = fastaFN,
                                               = "Achl_ACHL0006-09",
                         REGEX_SuffixForward = "[0-9] \star F\$",
```

(continues on next page)

REGEX_SuffixReverse	= "_[0-9]*_R\$",	
refAminoAcidSeq	=	
→ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPG	GALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMLG.	APDMAFPRMNN
minReadsNum	= 2,	
minReadLength	= 20,	
minFractionCall	= 0.5,	
maxFractionLost	= 0.5,	
geneticCode	= GENETIC_CODE,	
acceptStopCodons	= TRUE,	
readingFrame	= 1,	
processorsNum	= 1)	

In this example, contigName is set to Achl_ACHL0006-09, so Achl_ACHL0006-09_1_F and Achl_ACHL0006-09_2_R are matched and selected. Moreover, by regular expression pattern matching, Achl_ACHL0006-09_1_F is categorized into the forward list, and Achl_ACHL0006-09_2_R is categorized into the reverse read. Both reads are aligned into a contig, my_sangerContigFa, and it will be used as the input for the following functions.

Inside the R shell, you can run my_sangerContigFa to get basic information of the instance or run my_sangerContigFa@objectResults@readResultTable to check the creation result of every Sanger read after my_sangerContigFa is successfully created.

Here is the output of my_sangerContigFa:

```
SangerContig S4 instance
         Input Source : FASTA
        Process Method : REGEX
      Fasta File Name : /Library/Frameworks/R.framework/Versions/4.0/Resources/
-library/sangeranalyseR/extdata/fasta/SangerContig/Achl_ACHL0006-09.fa
  REGEX Suffix Forward : _[0-9]*_F$
  REGEX Suffix Reverse : _[0-9]*_R$
            Contig Name : Achl_ACHLO006-09
         'minReadsNum' : 2
      'minReadLength': 20
      'minFractionCall' : 0.5
      'maxFractionLost' : 0.5
   'acceptStopCodons' : TRUE
         'readingFrame': 1
     Contig Sequence :
\rightarrow TTATATTTTATTCTGGGCGTCTGAGCAGGAATGGTTGGAGCCGGTATAAGACTTCTAATTCGAATCGAGCTAAGACAACCAGGAG\phiGTTCCTGGGCAGA
Forward reads in the contig >> 1
Reverse reads in the contig >> 1
SUCCESS [2021-13-07 11:52:40] 'Achl_ACHL0006-09' is successfully created!
```

Here is the output of my_sangerContigFa@objectResults@readResultTable:

readName c	reationResult errorTyp	e errorMessage	inputSource	direction
1 Achl_ACHL0006-09_1_F	TRUE N	one No	ne FASTA	Forward Read
2 Achl_ACHL0006-09_2_R	TRUE N	one No	ne FASTA	Reverse Read

(2) "CSV file matching" SangerContig creation (FASTA)

The consturctor function and new method below contain three parameters, FASTA_File, contigName, and CSV_NamesConversion, that we mentioned in the previous section. Run the following code and create my_sangerContigFa instance.

```
csv_namesConversion <- file.path(rawDataDir, "fasta", "SangerContig", "names_

→conversion_1.csv")

# using `constructor` function to create SangerRead instance
my_sangerContigFa <- SangerContig(inputSource = "FASTA",</pre>
                                 processMethod = "CSV",
FASTA_File = fastaFN,
                                  contigName = "Achl_ACHLO006-09",
                                  CSV_NamesConversion = csv_namesConversion,
                                  refAminoAcidSeq
→ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNN
\rightarrow ",
                                 minReadsNum = 2,
minReadLength = 20,
minFractionCall = 0.5,
maxFractionLost = 0.5,
                                 minReadsNum
                                                        = 2,
                                 maxFractionLost
                                 geneticCode
                                                      = GENETIC_CODE,
                                 acceptStopCodons = TRUE,
                                  readingFrame
                                                      = 1,
                                 processorsNum
                                                      = 1)
# using `new` method to create SangerRead instance
my_sangerContigFa <- new("SangerContig",</pre>
                                              = "FASTA",
                         inputSource
                         processMethod
                                             = "CSV",
                         FASTA_File
contigName
                                              = fastaFN,
                                              = "Achl_ACHL0006-09",
                         CSV_NamesConversion = csv_namesConversion,
                         refAminoAcidSeq
                                              =
↔ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNNI
∽",
                                             = 2,
                        minReadsNum
                        = GENETIC_CODE,
                         acceptStopCodons = TRUE,
                                              = 1,
                         readingFrame = 1,
processorsNum = 1)
```

First, you need to load the CSV file into the R environment. If you are still don't know how to prepare it, please check (2) "CSV file matching" SangerContig inputs (FASTA). Then, it will follow rules in the CSV file and create my_sangerContigFa. After it's created, inside the R shell, you can run my_sangerContigFa to get basic information of the instance or run my_sangerContigFa@objectResults@readResultTable to check the creation result of every Sanger read after my_sangerContigFa is successfully created.

Here is the output of my_sangerContigFa:

```
SangerContig S4 instance
Input Source : FASTA
Process Method : CSV
Fasta File Name : /Library/Frameworks/R.framework/Versions/4.0/Resources/
→library/sangeranalyseR/extdata/fasta/SangerContig/Achl_ACHL0006-09.fa
CSV Names Conversion : /Library/Frameworks/R.framework/Versions/4.0/Resources/
→library/sangeranalyseR/extdata/fasta/SangerContig/names_conversion_1.csv
Contig Name : Achl_ACHL0006-09
'minReadsNum' : 2
```

(continues on next page)

Here is the output of my_sangerContigFa@objectResults@readResultTable:

readName	creationResult	errorType	errorMessage	inputSource	directi	on
1 Achl_ACHL0006-09_1	_F TR	UE Noi	ne Nor	ne FASTA	Forward	Read
2 Achl_ACHL0006-09_2	_R TR	UE Noi	ne Nor	ne FASTA	Reverse	Read

7.8.3 Writing SangerContig FASTA files (FASTA)

Users can write the *SangerContig* instance, my_sangerContigFa, to **FASTA** files. There are four options for users to choose from in selection parameter.

- reads_unalignment: Writing reads into a single FASTA file (only trimmed without alignment).
- reads_alignment: Writing reads alignment and contig read to a single FASTA file.
- contig: Writing the contig to a single FASTA file.
- all: Writing reads, reads alignment, and the contig into three different files.

Below is the oneliner for writing out FASTA files. This function mainly depends on writeXStringSet function in Biostrings R package. Users can set the compression level through writeFasta function.

Users can download the output FASTA file of this example through the following three links:

- (1) Achl_ACHLO006-09_reads_unalignment.fa
- (2) Achl_ACHLO006-09_reads_alignment.fa
- (3) Achl_ACHL0006-09_contig.fa

7.8.4 Generating SangerContig report (FASTA)

Last but not least, users can save *SangerContig* instance, my_sangerContigFa, into a report after the analysis. The report will be generated in **HTML** by knitting **Rmd** files.

Users can set includeSangerRead parameter to decide to which level the *SangerContig* report will go. Moreover, after the reports are generated, users can easily navigate through reports in different levels within the **HTML** file.

One thing to pay attention to is that if users have many reads, it will take quite a long time to write out all reports. If users only want to generate the contig result, remember to set includeSangerRead to FALSE in order to save time.

Here is the generated SangerContig html report of this example (FASTA). Users can access to 'Basic Information', 'SangerContig Input Parameters', 'Contig Sequence' and 'Contig Results' sections inside it. Furthermore, users can also navigate through html reports of all forward and reverse SangerRead in this SangerContig report.

7.8.5 Code summary (SangerContig, FASTA)

1. Preparing SangerContig FASTA input

```
rawDataDir <- system.file("extdata", package = "sangeranalyseR")
fastaFN <- file.path(rawDataDir, "fasta", "SangerContig", "Achl_ACHL0006-09.fa")</pre>
```

2. Creating SangerContig instance from FASTA

```
# using `constructor` function to create SangerRead instance
my_sangerContigFa <- SangerContig(inputSource = "FASTA",</pre>
                                    processMethod
FASTA_File
contigName
                                                            = "REGEX",
                                                            = fastaFN,
                                                            = "Achl_ACHL0006-09",
                                    REGEX_SuffixForward = "[0-9] \star F$",
                                    REGEX_SuffixReverse = "[0-9] * R\$",
                                    refAminoAcidSeq
                                                            =
→ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNNI
→ " )
# using `new` method to create SangerRead instance
my_sangerContigFa <- new("SangerContig",</pre>
                                                  = "FASTA",
                           inputSource
                           inputSource = "FASTA",
processMethod = "REGEX",
                                                                              (continues on next page)
```

```
FASTA_File = fastaFN,

contigName = "Achl_ACHL0006-09",

REGEX_SuffixForward = "_[0-9]*_F$",

REGEX_SuffixReverse = "_[0-9]*_R$",

refAminoAcidSeq =

→ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMLGAPDMAFPRMNNM

→ ")
```

Following is the R shell output that you will get.

3. Writing SangerContig FASTA files (FASTA)

writeFasta(my_sangerContigFa)

Following is the R shell output that you will get.

And you will get three FASTA files:

- (1) Achl_ACHLO006-09_reads_unalignment.fa
- (2) Achl_ACHLO006-09_reads_alignment.fa
- (3) Achl_ACHL0006-09_contig.fa

4. Generating SangerContig report (FASTA)

generateReport(my_sangerContigFa)

You can check the html report of this SangerContig example (FASTA).

7.9 Advanced User Guide - SangerAlignment (FASTA)

SangerAlignment is in the toppest level of sangeranalyseR (*Figure_1*), and each **SangerAlignment** instance corresponds to an alignment of contigs in a Sanger sequencing experiment. Among its slots, there is a *SangerContig* list

which will be aligned into a consensus contig. Users can access to each *SangerContig* and *SangerRead* inside a *SangerAlignment* instance.

In this section, we are going to go through details about a reproducible *SangerAlignment* analysis example with the **FASTA** file input in sangeranalyseR. By running the following example codes, you will get an end-to-end *Sanger-Alignment* analysis result.

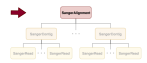


Fig. 56: Figure 1. Classes hierarchy in sangeranalyseR, SangerAlignment level.

7.9.1 Preparing SangerAlignment FASTA input

In *Advanced User Guide - SangerAlignment (AB1)*, we demonstrated how to use **AB1** input files to create *Sanger-Alignment* instance. Here, we explain another input format - the **FASTA** input. Before starting the analysis, users need to prepare one **FASTA** file, which must end with **.fa** or **.fasta**, containing sequences of all reads. In this example, the **FASTA** file is in the sangeranalyseR package, and you can simply get its path by running the following codes:

rawDataDir <- system.file("extdata", package = "sangeranalyseR")
fastaFN <- file.path(rawDataDir, "fasta", "SangerAlignment", "Sanger_all_reads.fa")</pre>

The value of fastaFN is where the **FASTA** file is placed. If your operating system is macOS, then its value should look like this:

And we showed the reads in fastaFN in Figure_2 (example FASTA file):

Inside the **FASTA** file (*Figure_2*; Sanger_all_reads.fa), the strings starting with ">" before each read are the read names. There are two ways of grouping reads which are "regular expression matching" and "CSV file matching", and following are instructions of how to prepare your **FASTA** input file.

(1) "regular expression matching" SangerAlignment inputs (FASTA)

For regular expression matching method, sangeranalyseR will group reads based on their contig name and read direction in their read names automatically; therefore, users have to follow the read-naming regulations below:

Note:

- All reads in the same contig group must include the same contig name in their read names.
- Forward or reverse direction also has to be specified in their read names.

There are three parameters, FASTA_File, REGEX_SuffixForward and REGEX_SuffixReverse, that define the grouping rule to let sangeranalyseR automatically match correct reads in **FASTA** file and divide them into forward and reverse directions.

Note:



Fig. 57: Figure 2. SangerAlignment FASTA input file (4 out of 8 reads are showed).

- FASTA_File: this is the path to **FASTA** file that contains all sequences of reads, and it can be either an absolute or relative path. We suggest users to include only target reads inside this **FASTA** file and do not include any other unrelated reads.
- REGEX_SuffixForward: this is a regular expression that matches all read names in forward direction. grep1 function in R is used.
- REGEX_SuffixReverse: this is a regular expression that matches all read names in reverse direction. grep1 function in R is used.

If you don't know what regular expression is, don't panic - it's just a way of recognising text. Please refer to *What is a regular expression?* for more details. Here is an example of how it works in sangeranalseR:

So how sangeranalyseR works is that it first matches the forward and reverse reads by matching REGEX_SuffixForward and REGEX_SuffixReverse. Then, sangeranalyseR uses the str_split function to split and vectorize their read names into "contig name" and "direction-suffix" two parts. For those having the same "contig name" will be grouped into the same contig.

Therefore, it is important to have a consistent naming strategy. You need to make sure that reads in the **FASTA** file that are in the same contig group share the same contig name and carefully select your REGEX_SuffixForward and REGEX_SuffixReverse. The bad file-naming and wrong regex matching might accidentally include reverse reads into the forward read list or vice versa, which will make the program generate wrong results. So, how should we systematically name the reads? We suggest users to follow the file-naming regulation in *Figure_3*.

[Consensus Read Name] + _ + [index] + _ + [F,R]

Fig. 58: Figure 3. Suggested read naming regulation in FASTA file - SangerAlignment.

As you can see, the first part of the regulation is a consensus read name (or contig name), which helps sangeranalseR to identify which reads should be grouped into the same contig automatically. The second part of the regulation is an index; since there might be more than one read that is in the forward or reverse direction, we recommend you to number your reads in the same contig group. The Last part is a direction which is either 'F' (forward) or 'R' (reverse).

To make it more specific, let's go back to the true example. In *Figure_2*, there are eight reads in the FASTA file (fasta_FN; Sanger_all_reads.fa). First, we set REGEX_SuffixForward to "_[0-9]*_F\$" and REGEX_SuffixReverse to "_[0-9]*_R\$" to let sangeranalyseR match and group forward and reverse reads automatically. By the regular expression rule, $Achl_ACHLO006-09_1_F$, $Achl_ACHLO007-09_1_F$, $Achl_ACHLO040-09_1_F$, and $Achl_ACHLO041-09_1_F$, are categorized into forward reads, and $Achl_ACHLO006-09_1_R$, $Achl_ACHLO007-09_1_R$, $Achl_ACHLO007-09_1_R$, $Achl_ACHLO007-09_1_R$, and $Achl_ACHLO041-09_1_R$ are categorized into reverse reads. Then, str_split function is used to split each filename above into "contig name" and "direction-suffix". Four contig names are detected in this example which are $Achl_ACHLO006-09$, $Achl_ACHLO007-09$, $Achl_ACHLO040-09$, and $Achl_ACHLO041-09$. Last, a loop iterates through all contig names, and sangeranalseR creates each of them into a SangerContig instance. You can check *Advanced User Guide - SangerContig (FASTA)* to see how sangeranalyseR creates a *SangerContig* instance.

The reason why we strongly recommend you to follow this file-naming regulation is that by doing so, you can directly adopt the example regular expression matching values, "[0-9]*F" and "[0-9]*R", to group reads and reduce chances of error. Everything mentioned above will be done automatically.

After understanding how parameters work, please refer to *Creating SangerAlignment instance from FASTA* below to see how sangeranalseR creates *SangerAlignment* instance.

(2) "CSV file matching" SangerAlignment inputs (FASTA)

No doubt that read names in the original **FASTA** file do not follow the naming regulation, and you do not want to change the original **FASTA** file; thus, we provide a second grouping approach, CSV file matching method. sanger-

analyseR will group reads in the **FASTA** file based on the information in a CSV file automatically, and users do not need to alter the read names in the **FASTA** file. The note below shows the regulations:

Note: Here is an example CSV file (*Figure 4*)

"reads","direction","contig"
"Achl_ACHLO006-09_1_F","F","Achl_ACHLO006-09"
"Achl_ACHLO006-09_2_R","R","Achl_ACHLO006-09"
"Achl_ACHLO007-09_1_F", "F", "Achl_ACHLO007-09"
"Achl_ACHLO007-09_2_R","R","Achl_ACHLO007-09"
"Achl_ACHLO040-09_1_F","F","Achl_ACHLO040-09"
"Achl_ACHLO040-09_2_R","R","Achl_ACHLO040-09"
"Achl_ACHLO041-09_1_F","F","Achl_ACHLO041-09"
"Achl_ACHLO041-09_2_R","R","Achl_ACHLO041-09"

Fig. 59: Figure 4. Example CSV file for SangerAlignment instance creation.

- There must be three columns, "reads", "direction", and "contig", in the CSV file.
- The "reads" column stores the filename of AB1 files that are going to be included in the analysis.
- The "direction" column stores the direction of the reads. It must be "F" (forward) or "R" (reverse).
- The "contig" column stores the contig name that each read blongs. Reads in the same contig have to have the same contig name, and they will be grouped into the same contig.

There are two parameters, FASTA_File and CSV_NamesConversion, that define the grouping rule to help sangeranalseR to automatically match correct reads in the **FASTA** file and divide them into forward and reverse directions.

Note:

- FASTA_File: this is the path to **FASTA** file that contains all sequences of reads, and it can be either an absolute or relative path. We suggest users to include only target reads inside this **FASTA** file and do not include any other unrelated reads.
- CSV_NamesConversion: this is the path to the CSV file. It can be either an absolute or relative path.

The main difference between "CSV file matching" and "regular expression matching" is where the grouping rule is written. For "regular expression matching", rules are writtein in read names, and thus more naming requirements are required. In contrast, rules of "CSV file matching" are written in an additional CSV file so it is more flexible on naming reads.

So how sangeranalyseR works is that it first reads in the CSV file (with "*reads*", "*direction*", and "*contig*" columns), find the read names in the **FASTA** file that are listed in "*reads*", and assign directions to them based on "*direction*".

To make it more specific, let's go back to the true example. First, we prepare a CSV file (CSV_NamesConversion) and a fasta file (FASTA_File). In the CSV file, there are 8 rows and

4 distinct contig names. sangeranalyseR matches "*reads*" of these 8 rows to read names in the **FASTA** file. Then sangeranalyseR groups all matched reads, Achl_ACHL0006-09_1_F, Achl_ACHL0007-09_1_F, Achl_ACHL0040-09_1_F, Achl_ACHL0040-09_1_F, Achl_ACHL0007-09_1_R, Achl_ACHL0007-09_1_R, Achl_ACHL0040-09_1_R, and Achl_ACHL0041-09_1_R, into 4 distinct contigs which are Achl_ACHL0006-09, Achl_ACHL0007-09, Achl_ACHL0040-09, and Achl_ACHL0041-09, by the "contig" column. Last, the directions of reads in each contig are assigned by the "direction" column. Take Achl_ACHL0041-09 contig as an example. Its "forward read list" will include Achl_ACHL0041-09_1_R.

After understanding how parameters work, please refer to *Creating SangerAlignment instance from FASTA* below to see how sangeranalseR creates *SangerAlignment* instance.

7.9.2 Creating SangerAlignment instance from FASTA

After preparing the input directory, we can create a *SangerAlignment* instance by running SangerAlignment constructor function or new method. The constructor function is a wrapper for new method and it makes instance creation more intuitive. Their input parameters are same, and all of them have their default values. For more details about *SangerAlignment* inputs and slots definition, please refer to sangeranalyseR reference manual. We will explain two *SangerAlignment* instance creation methods, "regular expression matching" and "CSV file matching".

(1) "regular expression matching" SangerAlignment creation (FASTA)

The consturctor function and new method below contain three parameters, FASTA_File, REGEX_SuffixForward, and REGEX_SuffixReverse, that we mentioned in the previous section. In contrast to **AB1** input method, it does not include quality trimming and chromatogram visualization parameters. Run the following code and create my_sangerAlignmentFa instance.

```
# using `constructor` function to create SangerAlignment instance
my_sangerAlignmentFa <- SangerAlignment(inputSource = "FASTA",</pre>
                                        processMethod = "REGEX",
FASTA_File = fastaFN,
                                        REGEX_SuffixForward = "[0-9] \star F;",
                                        REGEX_SuffixReverse = "[0-9] * R;",
                                        refAminoAcidSeq
→ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNNI
\rightarrow ",
                                        minReadsNum
                                                              = 2.
                                        minReadLength
                                                              = 20,
                                                              = 0.5,
                                        minFractionCall
                                                             = 0.5,
                                        maxFractionLost
                                        geneticCode
                                                              = GENETIC_CODE,
                                        acceptStopCodons = TRUE,
                                        readingFrame
                                                              = 1,
                                                              = 1)
                                        processorsNum
my_sangerAlignmentFa <- new("SangerAlignment",</pre>
                            inputSource
                                                 = "FASTA",
                                                 = "REGEX",
                            processMethod
                            FASTA_File
                                                 = fastaFN,
                            REGEX_SuffixForward = "[0-9] \star F \$",
```

(continues on next page)

REGEX_SuffixReverse	= "_[0-9]*_R\$",
refAminoAcidSeq	=
	LIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLM <mark>I</mark> GAPDMAFPRMNN
minReadsNum	= 2,
minReadLength	= 20,
minFractionCall	= 0.5,
maxFractionLost	= 0.5,
geneticCode	= GENETIC_CODE,
acceptStopCodons	= TRUE,
readingFrame	= 1,
processorsNum	= 1)

In this example, 8 reads are detected and 4 distinct *SangerContig* instances are created. These *SangerContig* instances are stored in a "contig list" in my_sangerAlignmentFa, which will be used as the input for the following functions.

Inside the R shell, you can run my_sangerAlignmentFa to get basic information of the instance or run my_sangerAlignmentFa@objectResults@readResultTable to check the creation result of every Sanger read after my_sangerAlignmentFa is successfully created.

Here is the output of my_sangerAlignmentFa:

```
SangerAlignment S4 instance
Input Source : FASTA
Process Method : REGEX
Fasta File Name : /Library/Frameworks/R.framework/Versions/4.0/Resources/
→library/sangeranalyseR/extdata/fasta/SangerAlignment/Sanger_all_reads.fa
REGEX Suffix Forward : _[0-9]*_F$
REGEX Suffix Reverse : _[0-9]*_R$
Contigs Consensus : _
→TTATAYTTATTYTRGGCGTCTGAGCAGGAATGGTTGGAGCYGGTATAAGACTYCTAATTCGAATYGAGCTAAGACARCCRGGAGCRTTCCTRGGMAGI
SUCCESS [2021-14-07 04:33:57] 'SangerAlignment' is successfully created!
```

Here is the output of my_sangerAlignmentFa@objectResults@readResultTable:

readName creation	Result erro	rType error	Message inputS	ource	directi	on
1 Achl_ACHLO006-09_1_F	TRUE	None	None	FASTA	Forward	Read
2 Achl_ACHLO006-09_2_R	TRUE	None	None	FASTA	Reverse	Read
3 Achl_ACHLO007-09_1_F	TRUE	None	None	FASTA	Forward	Read
4 Achl_ACHLO007-09_2_R	TRUE	None	None	FASTA	Reverse	Read
5 Achl_ACHLO040-09_1_F	TRUE	None	None	FASTA	Forward	Read
6 Achl_ACHLO040-09_2_R	TRUE	None	None	FASTA	Reverse	Read
7 Achl_ACHLO041-09_1_F	TRUE	None	None	FASTA	Forward	Read
8 Achl_ACHLO041-09_2_R	TRUE	None	None	FASTA	Reverse	Read

(2) "CSV file matching" SangerAlignment creation (FASTA)

The consturctor function and new method below contain two parameters, FASTA_File, and CSV_NamesConversion, that we mentioned in the previous section. Run the following code and create my_sangerAlignmentFa instance.

```
csv_namesConversion <- file.path(rawDataDir, "fasta", "SangerAlignment", "names_
→conversion.csv")
```

(continues on next page)

```
(continued from previous page)
# using `constructor` function to create SangerAlignment instance
my_sangerAlignmentFa <- SangerAlignment(inputSource</pre>
                                                           = "FASTA",
                                                           = "CSV",
                                       processMethod
                                       FASTA_File
                                                           = fastaFN,
                                       CSV_NamesConversion = csv_namesConversion,
                                       refAminoAcidSeq
→ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNNI
∽",
                                                             = 2,
                                       minReadsNum
                                       minReadLength
                                                             = 20,
                                       minKeadLength
minFractionCall
                                                           = 0.5,
= 0.5,
                                       maxFractionLost
                                       geneticCode
                                                            = GENETIC_CODE,
                                       acceptStopCodons = TRUE,
                                       readingFrame
                                                            = 1,
                                       processorsNum
                                                             = 1)
my_sangerAlignmentFa <- new("SangerAlignment",</pre>
                            inputSource = "FASTA",
processMethod = "CSV",
                           FASTA_File
                                               = fastaFN,
                           CSV_NamesConversion = csv_namesConversion,
                            refAminoAcidSeq =
→ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNN
\rightarrow ",
                           minReadsNum
                                                = 2,
                           minReadLength
                                                = 20,
                           minFractionCall
                                                = 0.5,
                                                = 0.5,
                           maxFractionLost
                            geneticCode
                                                 = GENETIC_CODE,
                            acceptStopCodons
                                                 = TRUE,
                            readingFrame
                                                 = 1,
                            processorsNum
                                                 = 1)
```

First, you need to load the CSV file into the R environment. If you are still don't know how to prepare it, please check (2) "CSV file matching" SangerAlignment inputs (FASTA). Then, it will follow rules in the CSV file and create my_sangerAlignmentFa. After it's created, inside the R shell, you can run my_sangerAlignmentFa to get basic information of the instance or run my_sangerAlignmentFa@objectResults@readResultTable to check the creation result of every Sanger read after my_sangerAlignmentFa is successfully created.

Here is the output of my_sangerAlignmentFa:

Here is the output of my_sangerAlignmentFa@objectResults@readResultTable:

readName crea	ationResult errorT	ype erro	orMessage inputSo	urce	directi	Lon
1 Achl_ACHLO006-09_1_F	TRUE	None	None	FASTA	Forward	Read
2 Achl_ACHLO006-09_2_R	TRUE	None	None	FASTA	Reverse	Read
3 Achl_ACHLO007-09_1_F	TRUE	None	None	FASTA	Forward	Read
4 Achl_ACHLO007-09_2_R	TRUE	None	None	FASTA	Reverse	Read
5 Achl_ACHLO040-09_1_F	TRUE	None	None	FASTA	Forward	Read
6 Achl_ACHLO040-09_2_R	TRUE	None	None	FASTA	Reverse	Read
7 Achl_ACHLO041-09_1_F	TRUE	None	None	FASTA	Forward	Read
8 Achl_ACHLO041-09_2_R	TRUE	None	None	FASTA	Reverse	Read

7.9.3 Writing SangerAlignment FASTA files (FASTA)

Users can write the *SangerAlignment* instance, my_sangerAlignmentFa, to **FASTA** files. There are four options for users to choose from in selection parameter.

- reads_unalignment: Writing reads into a single FASTA file (only trimmed without alignment).
- reads_alignment: Writing reads alignment and contig read to a single FASTA file.
- contig: Writing the contig to a single FASTA file.
- all: Writing reads, reads alignment, and the contig into three different files.

Below is the oneliner for writing out FASTA files. This function mainly depends on writeXStringSet function in Biostrings R package. Users can set the compression level through writeFasta function.

Users can download the output FASTA file of this example through the following three links:

- (1) Sanger_contigs_unalignment.fa
- (2) Sanger_contigs_alignment.fa
- (3) Sanger_all_trimmed_reads.fa

7.9.4 Generating SangerAlignment report (FASTA)

Last but not least, users can save *SangerAlignment* instance, my_sangerAlignmentFa, into a report after the analysis. The report will be generated in **HTML** by knitting **Rmd** files.

Users can set includeSangerContig and includeSangerRead parameters to decide to which level the *SangerAlignment* report will go. Moreover, after the reports are generated, users can easily navigate through reports in different levels within the **HTML** file.

One thing to pay attention to is that if users have many reads, it will take quite a long time to write out all reports. If users only want to generate the contig result, remember to set includeSangerRead and includeSangerContig to FALSE in order to save time.

Here is the generated SangerAlignment html report of this example (FASTA). Users can access to 'Basic Information', 'Contigs Consensus', 'Contigs Alignment', 'Contigs Tree', and 'Contig Reports' sections inside it. Furthermore, users can also navigate through html reports of all contigs and forward and reverse SangerRead in this SangerAlignment report.

7.9.5 Code summary (SangerAlignment, FASTA)

(1) Preparing SangerAlignment FASTA inputs

```
rawDataDir <- system.file("extdata", package = "sangeranalyseR")
fastaFN <- file.path(rawDataDir, "fasta", "SangerAlignment", "Sanger_all_reads.fa")</pre>
```

(2) Creating SangerAlignment instance from FASTA

(2.1) "Regular Expression Method" SangerAlignment creation (FASTA)

```
# using `constructor` function to create SangerAlignment instance
my_sangerAlignmentFa <- SangerAlignment(inputSource</pre>
                                                          = "FASTA",
                                         processMethod
                                         processMethod
FASTA_File
                                                              = "REGEX",
                                                               = fastaFN,
                                         REGEX_SuffixForward = "[0-9] *_F$",
                                         REGEX_SuffixReverse = "[0-9] *_R$",
                                         refAminoAcidSeq
                                                               =
→ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNN
→")
my_sangerAlignmentFa <- new("SangerAlignment",</pre>
                             inputSource = "FASTA",
processMethod = "REGEX",
                             FASTA_File = fastaFN,
                             REGEX_SuffixForward = "[0-9] *_F \$",
                             REGEX_SuffixReverse = "[0-9] *_R$",
                             refAminoAcidSeq
→ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNNI
                                                                           (continues on next page)
```

Chapter 7. Documentation

Following is the R shell output that you will get.

(2.2) "CSV file matching" SangerAlignment creation (FASTA)

```
csv_namesConversion <- file.path(rawDataDir, "fasta", "SangerAlignment", "names_
↔ conversion.csv")
# using `constructor` function to create SangerAlignment instance
my_sangerAlignmentFa <- SangerAlignment(inputSource = "FASTA",</pre>
                                                   = "CSV",
                                  processMethod
                                  FASTA_File
                                                    = fastaFN,
                                  CSV_NamesConversion = csv_namesConversion,
                                  refAminoAcidSeq
                                                   =
→ " )
my_sangerAlignmentFa <- new("SangerAlignment",</pre>
                        inputSource
                                         = "FASTA",
                                        = "CSV",
                        processMethod
                        FASTA_File
                                        = fastaFN,
                        CSV_NamesConversion = csv_namesConversion,
`refAminoAcidSeq
                  =
↔ "SRQWLFSTNHKDIGTLYFIFGAWAGMVGTSLSILIRAELGHPGALIGDDQIYNVIVTAHAFIMIFFMVMPIMIGGFGNWLVPLMIGAPDMAFPRMNNI
→ " )
```

Following is the R shell output that you will get.

(3) Writing SangerAlignment FASTA files (FASTA)

writeFasta(my_sangerAlignmentFa)

Following is the R shell output that you will get.

You will get three FASTA files:

- (1) Sanger_contigs_unalignment.fa
- (2) Sanger_contigs_alignment.fa
- (3) Sanger_all_trimmed_reads.fa

(4) Generating SangerAlignment report (FASTA)

generateReport(my_sangerAlignmentFa)

You can check the html report of this SangerAlignment example (FASTA).

7.10 Q & A ...

7.10.1 What is a regular expression?

A regular expression (sometimes shortened as regex or regexp) is a sequence of characters that define a sequence pattern matching rule, mainly used for searching and replacing. It is used in all programming languages like C++, Python, Javascript, and in our case, R.

7.10.2 How to deal with secondary peaks

7.10.3 How to work with FASTA files for input

7.11 User Manual (functions)

Following are input parameters for **SangerRead**, **SangerContig**, and **SangerAlignment** constructors. For more detials about other functions, please refer to the sangeranalyseR user manual.

7.11.1 SangerRead Constructor Parameters

```
SangerRead(inputSource = "ABIF",
    readFeature = "",
    readFileName = "",
    fastaReadName = "",
    geneticCode = GENETIC_CODE,
    TrimmingMethod = "M1",
    M1TrimmingCutoff = 0.0001,
    M2CutoffQualityScore = NULL,
    M2SlidingWindowSize = NULL,
    baseNumPerRow = 100,
    heightPerRow = 200,
    signalRatioCutoff = 0.33,
    showTrimmed = TRUE)
```

- inputSource: The input source of the raw file. It must be "ABIF" or "FASTA". The default value is "ABIF".
- readFeature: The direction of the Sanger read. The value must be "Forward Read" or "Reverse Read".
- readFileName: The absolute filename of the target ABIF or FASTA file.
- **fastaReadName**: If *"inputSource"* is *"FASTA"*, then this value has to be the name of the read inside the FASTA file; if *"inputSource"* is *"ABIF"*, then this value is *"NULL"* by default.
- **geneticCode**: Named character vector in the same format as "*GENETIC_CODE*" (the default), which represents the standard genetic code. This is the code with which the function will attempt to translate your DNA sequences. You can get an appropriate vector with the "*getGeneticCode()*" function. The default is the standard code.
- **TrimmingMethod**: The read trimming method for the *SangerRead*. The value must be "*M1*" (the default) or "*M2*", which represents "*method 1*" or "*method 2*" respectively. M1 is the modified Mott's trimming algorithm that can also be found in Phred/Phrap and Biopython. M2 is like trimmomatic's sliding window method.
- **M1TrimmingCutoff**: The cutoff for the trimming method 1. If *TrimmingMethod* is "*M1*", then the default value is "0.0001". Otherwise, the value must be "*NULL*".
- **M2CutoffQualityScore**: The trimming cutoff quality score for the trimming method 2. If *TrimmingMethod* is "M2", then the default value is "20". Otherwise, the value must be "NULL". This parameter works with M2SlidingWindowSize.
- M2SlidingWindowSize: The trimming sliding window size for the trimming method 2. If *TrimmingMethod* is "M2", then the default value is "10". Otherwise, the value must be "NULL". This parameter works with M2CutoffQualityScore.
- **baseNumPerRow**: This parameter is related to chromatogram and defines maximum base pairs in each row. The default value is *"100"*.
- heightPerRow: This parameter is related to chromatogram and defines the height of each row in chromatogram. The default value is "200".
- **signalRatioCutoff**: The ratio of the height of a secondary peak to a primary peak. Secondary peaks higher than this ratio are annotated. Those below the ratio are excluded. The default value is "0.33". This parameter is related to chromatogram.
- **showTrimmed**: The logical value storing whether to show trimmed base pairs in chromatogram. The default value is *"TRUE"*.

7.11.2 SangerContig Constructor Parameters

SangerContig(inputSource	= "ABIF",	
fastaFileName	= "",	
namesConversionCSV	= NULL,	
parentDirectory	= "",	
contigName	= "",	
suffixForwardRegExp	= "_F.ab1",	
suffixReverseRegExp	= "_R.ab1",	
TrimmingMethod	= "M1",	
M1TrimmingCutoff	= 0.0001,	
M2CutoffQualityScore	= NULL,	
M2SlidingWindowSize	= NULL,	
baseNumPerRow	= 100,	
		(continues on next page)

heightPerRow	= 200,
signalRatioCutoff	= 0.33,
showTrimmed	= TRUE,
refAminoAcidSeq	= "",
minReadsNum	= 2,
minReadLength	= 20,
minFractionCall	= 0.5,
maxFractionLost	= 0.5,
geneticCode	= GENETIC_CODE,
acceptStopCodons	= TRUE,
readingFrame	= 1,
processorsNum	= NULL)

- inputSource: The input source of the raw file. It must be "ABIF" or "FASTA". The default value is "ABIF".
- fastaFileName: If "inputSource" is "FASTA", then this value has to be the name of the FASTA file; if "input-Source" is "ABIF", then this value is "NULL" by default.
- **namesConversionCSV**: The absolute filename of CSV file that provides read names following the naming regulation. If *"inputSource"* is *"FASTA"*, then users need to prepare the csv file or make sure the original names inside FASTA file are valid; if *"inputSource"* is *"ABIF"*, then this value is *"NULL"* by default.
- **parentDirectory**: The parent directory of all of the reads contained in ABIF format you wish to analyse. In SangerContig, all reads must be in the first layer in this directory.
- contigName: The contig name of all the reads in "parentDirectory".
- **suffixForwardRegExp**: The suffix of the filenames for forward reads in regular expression, i.e. reads that do not need to be reverse-complemented. For forward reads, it should be "_F.ab1".
- **suffixReverseRegExp**: The suffix of the filenames for reverse reads in regular expression, i.e. reads that need to be reverse-complemented. For reverse reads, it should be "_R.ab1".
- **TrimmingMethod**: The read trimming method for the *SangerRead*. The value must be "*M1*" (the default) or "*M2*", which represents "*method 1*" or "*method 2*" respectively. M1 is the modified Mott's trimming algorithm that can also be found in Phred/Phrap and Biopython. M2 is like trimmomatic's sliding window method.
- **M1TrimmingCutoff**: The cutoff for the trimming method 1. If *TrimmingMethod* is "*M1*", then the default value is "0.0001". Otherwise, the value must be "*NULL*".
- M2CutoffQualityScore: The trimming cutoff quality score for the trimming method 2. If *TrimmingMethod* is "M2", then the default value is "20". Otherwise, the value must be "NULL". This parameter works with M2SlidingWindowSize.
- M2SlidingWindowSize: The trimming sliding window size for the trimming method 2. If *TrimmingMethod* is "M2", then the default value is "10". Otherwise, the value must be "NULL". This parameter works with M2CutoffQualityScore.
- **baseNumPerRow**: This parameter is related to chromatogram and defines maximum base pairs in each row. The default value is "100".
- heightPerRow: This parameter is related to chromatogram and defines the height of each row in chromatogram. The default value is "200".
- **signalRatioCutoff**: The ratio of the height of a secondary peak to a primary peak. Secondary peaks higher than this ratio are annotated. Those below the ratio are excluded. The default value is "0.33". This parameter is related to chromatogram.
- **showTrimmed**: The logical value storing whether to show trimmed base pairs in chromatogram. The default value is *"TRUE"*.

- **refAminoAcidSeq**: An amino acid reference sequence supplied as a string or an AAString object. If your sequences are protein-coding DNA sequences, and you want to have frameshifts automatically detected and corrected, supply a reference amino acid sequence via this argument. If this argument is supplied, the sequences are then kept in frame for the alignment step. Fwd sequences are assumed to come from the sense (i.e. coding, or "+") strand. The default value is "".
- minReadsNum: The minimum number of reads required to make a consensus sequence, must be 2 or more. The default value is "2".
- **minReadLength**: Reads shorter than this will not be included in the readset. The default "20" means that all reads with length of 20 or more will be included. Note that this is the length of a read after it has been trimmed.
- **minFractionCall**: Minimum fraction of the sequences required to call a consensus sequence for SangerContig at any given position (see the ConsensusSequence() function from DECIPHER for more information). Defaults to 0.75 implying that 3/4 of all reads must be present in order to call a consensus.
- **maxFractionLost**: Numeric giving the maximum fraction of sequence information that can be lost in the consensus sequence for SangerContig (see the ConsensusSequence() function from DECIPHER for more information). Defaults to 0.5, implying that each consensus base can ignore at most 50 percent of the information at a given position.
- **geneticCode**: Named character vector in the same format as "*GENETIC_CODE*" (the default), which represents the standard genetic code. This is the code with which the function will attempt to translate your DNA sequences. You can get an appropriate vector with the getGeneticCode() function. The default is the standard code.
- acceptStopCodons: The logical value "*TRUE*" or "*FALSE*". "*TRUE*" (the defualt): keep all reads, regardless of whether they have stop codons; "*FALSE*": reject reads with stop codons. If "*FALSE*" is selected, then the number of stop codons is calculated after attempting to correct frameshift mutations (if applicable).
- readingFrame: "1", "2", or "3". Only used if "accept.stop.codons == FALSE". This specifies the reading frame that is used to determine stop codons. If you use a "refAminoAcidSeq", then the frame should always be "1", since all reads will be shifted to frame 1 during frameshift correction. Otherwise, you should select the appropriate reading frame.
- processorsNum: The number of processors to use, or NULL (the default) for all available processors.

7.11.3 SangerAlignment Constructor Parameters

SangerAlignment(inputSource	= "ABIF",
fastaFileName	= "",
namesConversionCSV	= NULL,
parentDirectory	= "",
suffixForwardRegExp	= "_F.ab1",
suffixReverseRegExp	= "_R.ab1",
TrimmingMethod	= "M1",
MlTrimmingCutoff	= 0.0001,
M2CutoffQualityScore	= NULL,
M2SlidingWindowSize	= NULL,
baseNumPerRow	= 100,
heightPerRow	= 200,
signalRatioCutoff	= 0.33,
showTrimmed	= TRUE,
refAminoAcidSeq	= "",

(continues on next page)

minReadsNum	= 2,
minReadLength	= 20,
minFractionCall	= 0.5,
maxFractionLost	<pre>= 0.5,</pre>
geneticCode	= GENETIC_CODE,
acceptStopCodons	= TRUE,
readingFrame	= 1,
minFractionCallSA	= 0.5,
maxFractionLostSA	= 0.5,
processorsNum	= NULL)

- inputSource: The input source of the raw file. It must be "ABIF" or "FASTA". The default value is "ABIF".
- fastaFileName: If "inputSource" is "FASTA", then this value has to be the name of the FASTA file; if "input-Source" is "ABIF", then this value is "NULL" by default.
- **namesConversionCSV**: The file path to the CSV file that provides read names that follow the naming regulation. If *"inputSource"* is *"FASTA"*, then users need to prepare the csv file or make sure the original names inside FASTA file are valid; if *"inputSource"* is *"ABIF"*, then this value is *"NULL"* by default.
- **parentDirectory**: The parent directory of all of the reads contained in ABIF format you wish to analyse. In SangerContig, all reads must be in the first layer in this directory.
- **suffixForwardRegExp**: The suffix of the filenames for forward reads in regular expression, i.e. reads that do not need to be reverse-complemented. For forward reads, it should be "_F.ab1".
- **suffixReverseRegExp**: The suffix of the filenames for reverse reads in regular expression, i.e. reads that need to be reverse-complemented. For reverse reads, it should be "_R.ab1".
- **TrimmingMethod**: The read trimming method for the *SangerRead*. The value must be "*M1*" (the default) or "*M2*", which represents "*method 1*" or "*method 2*" respectively. M1 is the modified Mott's trimming algorithm that can also be found in Phred/Phrap and Biopython. M2 is like trimmomatic's sliding window method.
- **M1TrimmingCutoff**: The cutoff for the trimming method 1. If *TrimmingMethod* is "*M1*", then the default value is "0.0001". Otherwise, the value must be "*NULL*".
- M2CutoffQualityScore: The trimming cutoff quality score for the trimming method 2. If *TrimmingMethod* is "M2", then the default value is "20". Otherwise, the value must be "NULL". This parameter works with M2SlidingWindowSize.
- M2SlidingWindowSize: The trimming sliding window size for the trimming method 2. If *TrimmingMethod* is "M2", then the default value is "10". Otherwise, the value must be "NULL". This parameter works with M2CutoffQualityScore.
- **baseNumPerRow**: This parameter is related to chromatogram and defines maximum base pairs in each row. The default value is *"100"*.
- heightPerRow: This parameter is related to chromatogram and defines the height of each row in chromatogram. The default value is "200".
- **signalRatioCutoff**: The ratio of the height of a secondary peak to a primary peak. Secondary peaks higher than this ratio are annotated. Those below the ratio are excluded. The default value is "0.33". This parameter is related to chromatogram.
- **showTrimmed**: The logical value storing whether to show trimmed base pairs in chromatogram. The default value is *"TRUE"*.
- **refAminoAcidSeq**: An amino acid reference sequence supplied as a string or an AAString object. If your sequences are protein-coding DNA seuqences, and you want to have frameshifts automatically detected and corrected, supply a reference amino acid sequence via this argument. If this argument is supplied, the sequences

are then kept in frame for the alignment step. Fwd sequences are assumed to come from the sense (i.e. coding, or "+") strand. The default value is "".

- minReadsNum: The minimum number of reads required to make a consensus sequence, must be 2 or more. The default value is "2".
- **minReadLength**: Reads shorter than this will not be included in the readset. The default "20" means that all reads with length of 20 or more will be included. Note that this is the length of a read after it has been trimmed.
- **minFractionCall**: Minimum fraction of the sequences required to call a consensus sequence for SangerContig at any given position (see the ConsensusSequence() function from DECIPHER for more information). Defaults to 0.75 implying that 3/4 of all reads must be present in order to call a consensus.
- **maxFractionLost**: Numeric giving the maximum fraction of sequence information that can be lost in the consensus sequence for SangerContig (see the ConsensusSequence() function from DECIPHER for more information). Defaults to 0.5, implying that each consensus base can ignore at most 50 percent of the information at a given position.
- **geneticCode**: Named character vector in the same format as "*GENETIC_CODE*" (the default), which represents the standard genetic code. This is the code with which the function will attempt to translate your DNA sequences. You can get an appropriate vector with the getGeneticCode() function. The default is the standard code.
- acceptStopCodons: The logical value "*TRUE*" or "*FALSE*". "*TRUE*" (the defualt): keep all reads, regardless of whether they have stop codons; "*FALSE*": reject reads with stop codons. If "*FALSE*" is selected, then the number of stop codons is calculated after attempting to correct frameshift mutations (if applicable).
- readingFrame: "1", "2", or "3". Only used if "accept.stop.codons == FALSE". This specifies the reading frame that is used to determine stop codons. If you use a "refAminoAcidSeq", then the frame should always be "1", since all reads will be shifted to frame 1 during frameshift correction. Otherwise, you should select the appropriate reading frame.
- **minFractionCallSA**: Minimum fraction of the sequences required to call a consensus sequence for Sanger-Alignment at any given position (see the ConsensusSequence() function from DECIPHER for more information). Defaults to 0.75 implying that 3/4 of all reads must be present in order to call a consensus.
- maxFractionLostSA: Numeric giving the maximum fraction of sequence information that can be lost in the consensus sequence for SangerAlignment (see the ConsensusSequence() function from DECIPHER for more information). Defaults to 0.5, implying that each consensus base can ignore at most 50 percent of the information at a given position.
- processorsNum: The number of processors to use, or NULL (the default) for all available processors.

7.12 Frequently Asked Questions

7.12.1 Q: What is the difference between two different trimming methods?

A: In sangeranalyseR, we provide two trimming methods, "M1" (the default) or "M2", which represents "method 1" or "method 2" respectively. M1 is the modified Mott's trimming algorithm that can also be found in Phred/Phrap and Biopython. M2 is like trimmomatic's sliding window method. If you want to set M1 as your trimming method, you need to assign "TrimmingMethod" to "M1" and "M1TrimmingCutoff" as the threshold that you want. Its default value is "0.0001". In contrast, you can assign "TrimmingMethod" to "M2" if you want to set M2 as your trimming method. "M2CutoffQualityScore" and "M2SlidingWindowSize" are two parameters that control M2 trimming and their default values are "20" and "10" respectively.

7.13 Conclusion

sangeranalyseR provides a new approach to do Sanger sequencing data analysis in R. The main features include wellstructured S4 classes, automated data analysis, interactive Shiny apps, exporting reads to FASTA and the generation thorough report. We hope it will be helpful for R users and the bioinformatics community!

7.14 License

MIT License

Copyright (c) 2019 Kuan-Hao Chao

Permission is hereby granted, free of charge, to any person obtaining a copy of this software and associated documentation files (the "Software"), to deal in the Software without restriction, including without limitation the rights to use, copy, modify, merge, publish, distribute, sublicense, and/or sell copies of the Software, and to permit persons to whom the Software is furnished to do so, subject to the following conditions:

The above copyright notice and this permission notice shall be included in all copies or substantial portions of the Software.

THE SOFTWARE IS PROVIDED "AS IS", WITHOUT WARRANTY OF ANY KIND, EXPRESS OR IMPLIED, INCLUDING BUT NOT LIMITED TO THE WARRANTIES OF MERCHANTABILITY, FITNESS FOR A PAR-TICULAR PURPOSE AND NONINFRINGEMENT. IN NO EVENT SHALL THE AUTHORS OR COPYRIGHT HOLDERS BE LIABLE FOR ANY CLAIM, DAMAGES OR OTHER LIABILITY, WHETHER IN AN ACTION OF CONTRACT, TORT OR OTHERWISE, ARISING FROM, OUT OF OR IN CONNECTION WITH THE SOFT-WARE OR THE USE OR OTHER DEALINGS IN THE SOFTWARE.

7.15 Contact

Contact here:

For now, please just use the issue tracker on GitHub for all contacts. That will help us keep up to date with things.

7.16 Help

If you need any help, feel free to contact me <kuanhao.chao@gmail.com>

7.16.1 Inside help test