# Manual for Sondovač 1.1

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Sondovač<sup>1</sup> is a script to create orthologous low-copy nuclear probes from transcriptome and genome skim data for target enrichment (Schmickl et al., 2016). See https://github.com/V-Z/sondovac/wiki.

#### Abstract

Phylogenetics benefits from using a large number of putatively independent nuclear loci in combination with other sources of information, such as the plastid and mitochondrial genome. Selecting such orthologous low-copy nuclear (LCN) loci is still a challenge for non-model organisms. In recently published phylogenies based on target enrichment of several hundred LCN genes, these loci were selected from transcriptomes, genomes, gene expression studies, the literature, or a combination of these sources. Automated bioinformatic pipelines for the selection of LCN genes are, however, largely absent. We created a user-friendly, automated and interactive script named Sondovač to design LCN loci by a comparison between transcriptome and genome skim data. The script is licensed under open-source license GPL v.3 allowing further modifications. It runs on major Linux distributions and Mac OS X. Strong bioinformatics skills and access to high-performance computer clusters are not required; Sondovač runs on a standard desktop computer equipped with modern CPU like Intel i5 or i7.

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<sup>&</sup>lt;sup>1</sup>English pronunciation is "Sondovach". The word is a Czech neologism meaning something like "The Prober" or "The Probe Maker".

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Sondovač is a script to create orthologous low-copy nuclear probes from transcriptome and genome skim data for target enrichment (Schmickl et al., 2016). For information and download see https://github.com/V-Z/sondovac/wiki.

# 1 Introduction

High-throughput sequencing (HTS) has the potential to greatly increase the amount of phylogenetically informative signal in molecular datasets (Parks et al., 2009, 2012) and overcome difficulties in phylogenetic reconstructions, such as polytomies and low support values, that are often the result of using only a small fraction of the genome. However, HTS also "opens the era of real incongruence" (Jeffroy et al., 2006), and even massive amounts of sequence data do not always result in strongly resolved phylogenies (Pyron, 2015).

Currently, target enrichment (sequence capture) of hundreds of loci is becoming increasingly popular in phylogenetics. In animal phylogenomics non-exonic or partly exonic ultraconserved elements and their more variable flanking regions are often utilized (e.g. Faircloth et al., 2012; Hedtke et al., 2013; Smith et al., 2014). For plant phylogenetics, low-copy nuclear (LCN) genes are targeted (Mandel et al., 2014; Weitemier et al., 2014; Grover et al., 2015; Heyduk et al., 2015; Mandel et al., 2015; Nicholls et al., 2015; Stephens et al., 2015a,b) due to the paucity of ultraconserved nuclear sequences (Reneker et al., 2012). Target sequencing strategies for plant nuclear genomes are largely lineage-specific, requiring the de novo design of target enrichment probes. Chamala et al. (2015) recently introduced a pipeline for phylogenetic marker development in angiosperms using transcriptomes, and they obtained several hundred putative LCN genes that can be utilized at three phylogenetic levels (genus, family, order); however empirical evidence for the phylogenetic utility of these loci was not demonstrated. Alternative phylogenetic marker developments, also utilizing transcriptomes (Pillon et al., 2014; Rothfels et al., 2013; Tonnabel et al., 2014), resulted in a much smaller number (up to 20) of mainly LCN loci, but these loci were evaluated with PCR in the empirical datasets, not target enrichment. In recently published phylogenies based on target enrichment of several hundred LCN genes, these loci were selected from transcriptomes, gene expression studies, the literature, or a combination of these sources (Mandel et al., 2014; Grover et al., 2015; Heyduk et al., 2015; Mandel et al., 2015; Nicholls et al., 2015; Stephens et al., 2015a,b). Weitemier et al. (2014) designed LCN probes for target enrichment based on a combination of transcriptome and genome data. The limitation of this probe design pipeline is that (draft) genomes are still infrequent, especially for non-model species, and are costly to generate. This limitation also applies to the approach of de Sousa et al. (2014), who selected 50 LCN loci from a genomic source and amplified them using target enrichment. Except for Chamala et al. (2015), who offer a user-friendly but empirically untested probe design pipeline, and Weitemier et al. (2014), whose Hyb-Seq pipeline is designed for more advanced users, no automated probe design pipeline for LCN genes is currently available.

In this study (Schmickl et al., 2016) we developed a novel probe design pipeline for targeting orthologous LCN loci for phylogenetic reconstruction by using genome skim and transcriptome data. In particular, genome skim data of one accession of the studied plant group were combined with a congeneric transcriptome from the 1000 Plants (1KP) initiative (http://onekp.com/). We implemented our software workflow in the user-friendly, automated and interactive BASH script Sondovač, which allows a straightforward design of LCN probes also catering for users with limited bioinformatics skills.

Sondovač workflow is divided into three parts (see details on page 5 and Figure 1):

- 1. Raw input data are analyzed by sondovac\_part\_a.sh.
- 2. Sequences obtained in part a are assembled by Geneious in a separate step by the user.
- 3. Final probes are produced by sondovac\_part\_b.sh.

### 1.1 Pipeline – how the data are processed

A transcriptome assembly and paired-end genome skim raw data are combined to get hundreds of orthologous LCN loci (Schmickl et al., 2016). Enrichment of multi-copy loci is minimized by using unique transcripts only, which are obtained by comparing all transcripts and removing those sharing >90\% sequence similarity using BLAT. Before matching the genome skim data against those unique transcripts, reads of plastid (and mitochondrial) origin are removed with Bowtie 2, SAMtools and bam2fastq, utilizing reference sequences. Paired-end reads are subsequently combined with FLASH. These processed reads are matched against the unique transcripts sharing >85\% sequence similarity with BLAT. Transcripts with >1000 BLAT hits (indicating repetitive elements) and BLAT hits containing masked nucleotides are removed before de novo assembly of the BLAT hits to larger contigs with Geneious, using the medium sensitivity / fast setting. After assembly, only those contigs that comprise exons of a minimum bait length (usually >120 bp in case of probe design for phylogenies) and have a certain minimum total locus length (multiple of the bait length, should not be too short in order to obtain sufficient phylogenetically informative signal; we recommend at least  $\geq 600$  bp) are retained. To ensure that probes do not target multiple similar loci, any probe sequences sharing >90% sequence similarity are removed using cd-hit-est, followed by a second filtering step for contigs containing exons of a minimum bait length and totaling minimum loci length (see comments above). To ensure that plastid sequences are absent from the probes, the probe sequences are matched against the plastome reference sharing  $\geq 90\%$  sequence similarity with BLAT and the hits removed from the probe set. The workflow of Sondovač is summarized in Figure 1. The direction of the workflow is indicated by arrows. An optional removal of reads of mitochondrial origin from the genome skim data is indicated by greyed text. The required input files of Sondovač are highlighted in bold.

The steps of Sondovač are consecutively numbered to aid comprehension. Sondovač has three parts: two script parts and an intermediate part using Geneious. The workflow is as follows:

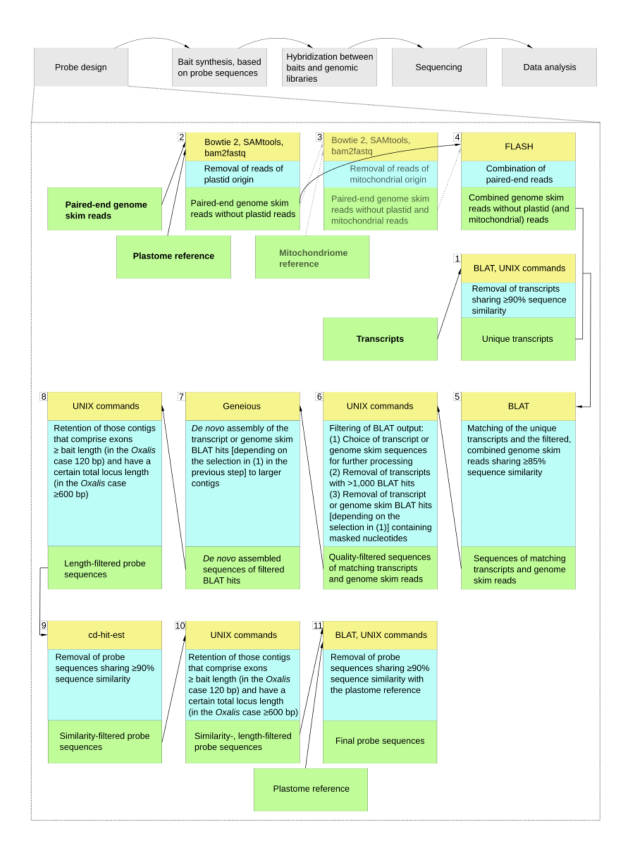
#### A. sondovac\_part\_a.sh: Covers steps 1 to 6.

- 1. Removal of transcripts sharing  $\geq 90\%$  sequence similarity.
- 2. Removal of reads of plastid origin.
- 3. Removal of reads of mitochondrial origin (optional).
- 4. Combination of paired-end reads.
- 5. Matching of the unique transcripts and the filtered, combined genome skim reads sharing  $\geq 85\%$  sequence similarity.
- **6.** Filtering of BLAT output:
  - **6.1.** Choice of transcript or genome skim sequences for further processing.
  - **6.2.** Removal of transcripts with >1000 BLAT hits.
  - **6.3.** Removal of transcript or genome skim BLAT hits [depending on the selection in (6.1)] containing masked nucleotides.

Input files for sondovac\_part\_a.sh are FASTA transcriptome data, FASTQ paired-end genome skim reads and a plastome (and possible also mitochondriome) reference. The input file for Geneious is the output of sondovac\_part\_a.sh.

- **B.** Geneious: Covers step 7 (see page 17).
  - 7. De novo assembly of the transcript or genome skim BLAT hits [depending on the selection in (6.1)] to larger contigs. Note that you need a copy of Geneious for this step.

The output files of Geneious are input files for sondovac\_part\_b.sh.



**Figure 1:** Workflow of the probe design script Sondovač. An overview of the main steps of Hyb-Seq are given in the top part of the figure; probe design is the first one. Each step of Sondovač is numbered and illustrated by three boxes: Software is highlighted in yellow, a summary of each step is given in light blue, and input/output of each step is depicted in light green. An optional removal of reads of mitochondrial origin from the genome skim data is marked by greyed text. The required input files of Sondovač are highlighted in bold. The direction of the workflow is indicated by arrows.

- C. sondovac part b.sh: Covers steps 8 to 11.
  - 8. Retention of those contigs that comprise exons  $\geq$  bait length and have a certain total locus length.
  - **9.** Removal of probe sequences sharing  $\geq 90\%$  sequence similarity.
  - 10. Retention of those contigs that comprise exons  $\geq$  bait length and have a certain total locus length.
  - 11. Removal of probe sequences sharing  $\geq 90\%$  sequence similarity with the plastome reference.

The output file of sondovac part b.sh is the final list of probes.

When Sondovač starts, a directory **bin** is created in the current working directory; Sondovač saves binaries of required software packages in this directory (if they are not available). The user can then add this directory to PATH, move or delete it afterwards.

### 1.2 General considerations before you start

The success of the probe design in terms of a high number of LCN genes of a sufficient minimum total length with Sondovač depends on various aspects of your transcriptome and genome skim input data:

- number of transcripts,
- read length of genome skim reads; longer reads and paired-end reads are preferable due to a higher quality de novo assembly of the reads to contigs (exons),
- number of nuclear genome skim reads,
- quality of nuclear genome skim reads,
- sequence divergence between transcriptome and genome skim data.

These aspects influence the number of probe sequences and the proportion of paralogous loci among the probe sequences. The usage of a transcriptome and genome skim data of **diploid** accessions is strongly recommended in order to account for orthology of the probe sequences. An example of how one aspect, the number of nuclear genome skim reads, can affect the probe design, is shown in Table 1 and Figure 2.

**Table 1:** Summary of two examples of a LCN probe design with Sondovač. The *Oxalis* example is from Schmickl et al. (2016), the *Curcuma* example is unpublished data from Tomáš Fér and Roswitha Schmickl. The respective Sondovač steps are listed; see Figure 1 for details regarding these steps. For both probe designs 250 bp paired-end reads were utilized. Input files are given in **typewriter** font. Quality control of the genome skim data, which is not part of Sondovač, is colored in grey.

Step of Sondovač	Substep of Sondovač	$Oxalis \ {f species}$	Curcuma species
Input file	Transcriptome taxon	$Oxalis$ $corniculata \ L.$	Curcuma longa L.
Input file	Genome skim taxon	$Oxalis\ obtusa$ Jacq.	Curcuma ecomata Craib
Input file	Plastome taxon	Ricinus $communis$ L.	Curcuma roscoeana Wall., Zingiber spectabile Griff.
Input file	Mitochondriome taxon	Ricinus $communis$ L.	Oryza sativa L. subsp. indica

... continued Table 1.

continued	Table 1.	_	
Step of Sondovač	Substep of Sondovač	$Oxalis \ { m species}$	Curcuma species
1	Number of transcripts	22,093	23,996
1	Number of unique transcripts	16,123	17,203
1	Total length of unique transcripts	11,799,393 bp	11,919,459 bp
	Number of genome skim raw reads		
2	(without quality-filtering and duplicate removal)	9,236,186	12,299,804
Quality control	Percentage of dropped quality-filtered genome skim reads	2%	2%
Quality control	Number of quality-filtered genome skim reads	8,525,040	11,340,170
Quality control	Percentage of duplicate quality-filtered genome skim reads	7%	3%
Quality control	Number of quality-filtered genome skim reads after duplicate removal	7,938,349	11,041,405
Quality control	Number of masked bases in quality-filtered genome skim reads after duplicate removal	3,775	7,725
	Number of nuclear genome skim raw		
3	reads (without quality-filtering and	8,240,470	11,636,852
	duplicate removal)	0,= -0, 0	,000,00_
4	Number of combined nuclear genome skim raw reads	2,619,197	3,834,278
	Combined nuclear genome skim raw		
4	reads as proportion of the total number of nuclear genome skim raw reads	64%	66%
4	Total length of combined nuclear genome skim raw reads	856,720,402 bp	1,218,798,300  bp
5	Mean sequence divergence between the unique transcripts and the combined nuclear genome raw skim reads	7%	6%
5	Mean sequence length of the match between the unique transcripts and the combined nuclear genome raw skim reads (genome skim data)	216 bp	204 bp
5	Mean sequence length of the match between the unique transcripts and the combined nuclear genome raw skim reads (transcripts)	194 bp	195 bp
7	Mean sequence depth of the contigs (exons) after the de novo assembly of the matching sequences	4	3
7	Mean sequence length of the contigs (exons) after the de novo assembly of the matching sequences	114 bp	169 bp
7	Mean pairwise identity between the assembled reads of the contigs (exons) after the de novo assembly of the matching sequences	99%	100%

... continued Table 1.

Step of Sondovač	Substep of Sondovač	$Oxalis \ { m species}$	Curcuma species
7	Minimum pairwise identity between the assembled reads of the contigs (exons) after the de novo assembly of the matching sequences	84%	94%
11	Number of exons $\geq 120$ bp	4,926	4,618
11	Number of genes	$1,164$ ( $\geq 600 \text{ bp}$ )	1,180 (≥960 bp)
11	Total length of probe sequences	1,127,2049 bp	1,571,800  bp

## 2 Installation of Sondovač

Sondovač is a simple BASH script, but it requires additional software to run successfully. The script will check for the presence of all required software and, if needed, will offer installation. The easiest way is just to launch the script (see chapter 2.4 on page 15) and let yourself to be guided through the whole process.

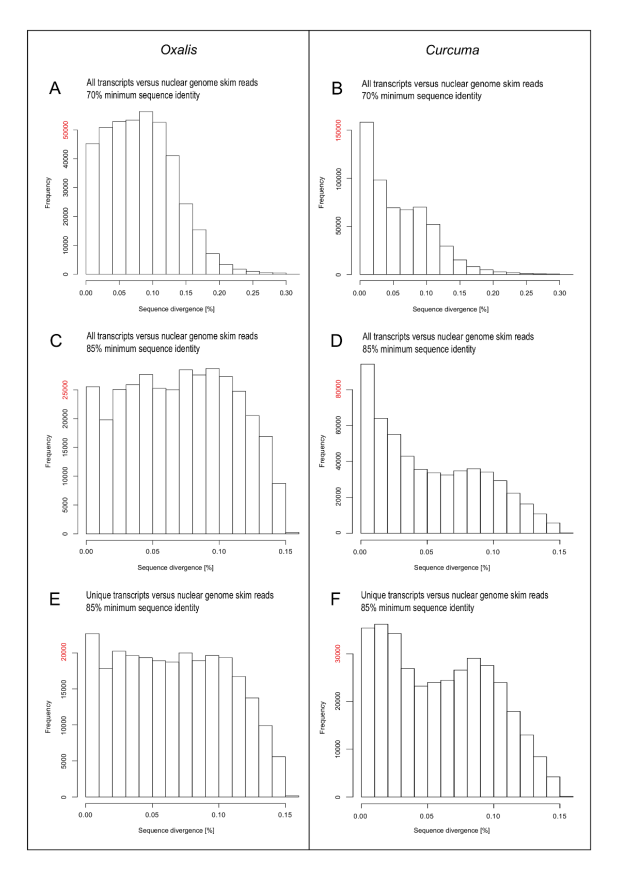
## 2.1 Requirements to run Sondovač

Sondovač is currently tested on major Linux distributions (in current versions) openSUSE, Debian, Ubuntu, Linux Mint, Fedora, Centos and Scientific Linux; and on Mac OS X (version 10.10 Yosemite).

In order to run Sondovač you need a UNIX-based operating system (preferably Linux, alternatively Mac OS X) equipped with BASH or a compatible shell interpreter (this should by default be available for any Linux distribution, Mac OS X and any other UNIX-based operating system like Solaris, BSD and its variants etc.). You should use the current operating system version supported by upstream, otherwise we will not be able to help you in case of problems. Older operating systems can have different versions of shell and system libraries, which can cause various problems and incompatibilities.

Sondovač uses several scientific software packages (namely bam2fastq, BLAT, Bowtie2, CD-HIT, FASTX toolkit, FLASH, Geneious, htsjdk, libgtextutils, and SAMtools – see required versions and links, Table 2), and basic UNIX tools (see below). Sondovač will check if those programs are installed – available in the PATH (i.e. if the shell application can locate and launch respective binaries, see also vocabulary at page 20). If you have those packages installed (in current versions, see Table 2), ensure that their binaries are in PATH. This should not be a problem for basic tools available in any UNIX-based operating system, as basic installation usually contains all needed tools. If you lack some of the required tools, the script will notify you, and you will have to install them manually. If this is needed, check the documentation for your operating system.

If required programs are not installed, Sondovač will offer you installation. You can use precompiled binaries available together with the script (this is the recommended option) or (sometimes) from the web. In case you would like to compile required software yourself, the script will guide you through this process. This is recommended only for advanced users, as compilation might sometimes be very tricky. Users of Mac OS X can install those applications also using Homebrew (see http://brew.sh/). For compilation you need Apache Ant, GNU G++, GNU GCC, GIT, Java/OpenJDK, libpng developmental files, and zlib developmental files. Ensure you have those tools available – they should be readily available for any UNIX-based operating system. Chapters 2.2 and 2.3 give details about requirements and their manual fulfilling. This is mainly a reference for more advanced users or users with special needs. For



**Figure 2:** Sequence divergence between all transcripts and nuclear genome skim data  $(\mathbf{A}-\mathbf{D})$  and unique transcripts and nuclear genome skim data  $(\mathbf{E},\mathbf{F})$  in the case of *Oxalis* and *Curcuma*. The generally larger number of *Curcuma* nuclear genome skim reads compared to the *Oxalis* nuclear genome skim reads is highlighted in red.

most users it should be fully sufficient to run the script and let it do this job (see chapter 2.4 on page 22).

The following UNIX tools are required to run Sondovač. They are usually readily available in UNIX systems (but see note for Mac OS X below), so there is usually no need to install them manually. The tools are awk, bc, bunzip2, cat, cp, curl or wget, cut, dirname, dos2unix, echo, egrep, cd, g++, gcc, grep, gunzip, join, less, lsb\_release, make, mkdir, paste, perl, pkg-config, pwd, python, sed, sort, tar, tr, uname, uniq, unzip, wc. Not all tools are required every time – some are used only during particular actions (e.g. when the user decides to compile the required software manually). And the user usually does not need to bother with them. See also details in the following subchapters for some common Linux distributions and Mac OS X.

See below for details about tools required by Sondovač and their manual installation. For most users it should be sufficient to be guided by the script to install needed tools automatically.

## 2.2 Installation of required software in Linux

Linux distributions have precise package management tools (similar, but with more functions, to various app stores known from Android, iOS or recent Mac OS X, MS Windows, etc.), but unfortunately Linux repositories commonly do not contain all needed scientific packages (or not enough recent versions). We recommend pre-compiled binaries of scientific applications available together with the script. If the user wishes to compile the software, for whatever reason, the script will guide through that process.

## 2.2.1 openSUSE and SUSE Linux Enterprise (SLE)

SUSE Linux Enterprise (https://www.suse.com/) and openSUSE (https://www.opensuse.org/) use for package management command zypper<sup>2</sup>. The script will check if all required software packages are installed, and if not, will install them. You can also install manually:

```
1 # Verify installation of basic tools (they are installed in 99.9%):
2 sudo zypper in bash gawk bc coreutils grep less lsb-release perl-base python \
    sed wget
4 # Install packages needed for compilation:
5 sudo zypper in gcc-c++ gcc make pkg-config bzip2 gzip tar unzip \
    patterns-openSUSE-devel_basis libpng12-devel zlib-devel gcc-java \
    java-1_7_0-openjdk java-1_7_0-openjdk-devel git-core ant
8 # Update installed packages:
9 sudo zypper up
10 # Remove package:
11 sudo zypper rm PACKAGE
12 # Search for package:
13 zypper se PACKAGE/KEYWORD
14 # More information about zypper usage:
15 zypper --help
16 man zypper
17 # Note backslash ("\") means that the code continues on the next line
```

Originally, those distributions used only rpm\* commands (see rpm --help and man rpm for basic usage).

<sup>&</sup>lt;sup>2</sup>See https://en.opensuse.org/Zypper and https://activedoc.opensuse.org/book/opensuse-start-up/chapter-9-managing-software-with-command-line-tools for details.

#### 2.2.2 Debian, Ubuntu, Linux Mint and derivatives

Debian (https://www.debian.org/), Linux Mint (http://linuxmint.com/), Ubuntu (http://www.ubuntu.com/) and all derived distributions<sup>3</sup> like Kubuntu (http://www.kubuntu.com/) use for package management commands apt-get (basic) and aptitude (text-based front-end for apt-get, recommended, not available by default in every DEB based distribution). There are more tools available<sup>4</sup>, we will describe only the basic usage needed for our purpose. The script will check if all required software packages are installed, and if not, will install them. You can also install manually:

```
# Verify installation of basic tools (they are installed in 99.9%):
19 sudo apt-get install bash gawk bc coreutils grep less lsb-release perl-base \
    python sed wget
21 # Install packages needed for compilation:
22 sudo apt-get install build-essential bzip2 gzip tar unzip gcc g++ cpp make \
    libpng12-dev zlib1g-dev openjdk-7-jre openjdk-7-jdk openjdk-7-source \
    git ant pkg-config realpath
25 # Update installed packages:
26 sudo apt-get update # Update list of available packages in repositories
27 sudo apt-get upgrade # Actually update installed packages
28 # Remove package:
29 sudo apt-get remove PACKAGE
30 sudo apt-get autoremove # Automatically remove orphaned unneeded packages
31 # Search for package:
32 apt-cache --help # Usage options
33 apt-cache show PACKAGE # Display information about PACKAGE
34 apt-cache search KEYWORD # Search for KEYWORD, including regular expressions
35 # More information about apt-get usage:
36 apt-get --help
37 man apt-get
38 # Interactive command-line package manager
39 sudo aptitude
40 # Help for aptitude
41 aptitude --help
42 man aptitude
43 # Note backslash ("\") means that the code continues on the next line
```

Note you can use **aptitude** in a similar way as **apt-\*** commands (e.g. **aptitude instal PACKAGE** etc.). For special package operations, there are plenty of **dpkg** commands for advanced management.

#### 2.2.3 RedHat, Fedora, Centos, Scientific Linux and derivatives

RedHat (https://www.redhat.com/), Fedora (https://getfedora.org/; until version 21), Centos (https://centos.org/) and Scientific Linux (https://www.scientificlinux.org/) and other related distributions<sup>5</sup> use for package management command yum<sup>6</sup>. The script will

 $<sup>^3</sup>$ For complete lists see http://distrowatch.com/search.php?basedon=Debian and http://distrowatch.com/search.php?basedon=Ubuntu.

<sup>&</sup>lt;sup>4</sup>See https://wiki.debian.org/PackageManagement for list of tools and https://www.debian.org/doc/manuals/debian-reference/ch02.en.html for exhaustive documentation. A shorter introduction is available at https://help.ubuntu.com/community/AptGet/Howto and http://ubuntuguide.org/wiki/Ubuntu\_Trusty\_Packages\_and\_Repositories. Ubuntu-specific information at https://help.ubuntu.com/stable/ubuntu-help/addremove.html.

<sup>&</sup>lt;sup>5</sup>See http://distrowatch.com/search.php?basedon=Fedora for complete list.

<sup>&</sup>lt;sup>6</sup>See http://yum.baseurl.org/ for details.

check if all required software packages are installed, and if not, will install them. You can also install manually:

```
# Verify installation of basic tools (they are installed in 99.9%):
sudo yum install bash coreutils gawk bc grep less lsb perl python sed wget
# Install packages needed for compilation:
sudo yum install bzip2 gzip pkgconfig unzip gcc gcc-c++ cpp libpng12-devel \
make zlib-devel java-1.8.0-openjdk java-1.8.0-openjdk-devel git ant tar
# Update installed packages:
sudo yum update
# Remove package:
sudo yum remove PACKAGE
# Search for package:
yum search PACKAGE/KEYWORD
# More information about yum usage:
yum --help
man yum
# Note backslash ("\") means that the code continues on the next line
```

Since version 22, Fedora uses the command dnf for package management. It replaces older yum, and yum commands are redirected to dnf. The basic usage is the same, so that one can just replace yum with dnf in the above examples. Originally, those distributions used only rpm\* commands (see rpm --help and man rpm for basic usage).

# 2.3 Installation of required software in Mac OS X

For Mac OS X users, Homebrew (see http://brew.sh/ and https://github.com/Homebrew/) will be installed by the script, and it will install (new software or newer versions) Apache Ant, BASH (the shell interpreter), GNU AWK, GNU coreutils, GNU GCC, git, GNU grep, GNU make, pkg-config, GNU sed, and wget. Mac OS X is missing some tools and contains outdated BSD versions for others (typically sed, grep or awk). The script will guide the user through the process, and the user can safely and easily remove these tools afterwards if necessary. Unfortunately, Mac OS X does not have usable build-in package management, and it has outdated versions of some required tools. Homebrew fills this gap. It is a simple command-line installer (similar to package managers known from Linux, BSD or Solaris) of various applications.

Homebrew requires Xcode<sup>7</sup> (set of tools required to compile software) to be installed. Unfortunately, it is not possible to easily and universally check if Xcode is installed, so that the script will ask if the user wishes to install it. If the user is unsure if Xcode is installed, it is safe to answer Yes and install it. The manual command to install Xcode is the following:

```
59 xcode-select --install
60 # Following error means Xcode has already been installed:
61 xcode-select: note: no developer tools were found at '/Applications/Xcode.app',
62 requesting install. Choose an option in the dialog to download the command
63 line developer tools.
64 # Verify Xcode installation by
65 xcode-select --print-path # Prints installation location of Xcode
66 xcode-select --version # Prints version of Xcode
```

If Xcode is not installed yet, the user will see windows similar to that on Figure 3, offering installation of Xcode. Select **Install** to continue. After installation, the script will exit, and the user must start it again.

The script will guide the user through all those steps and basic usage of Homebrew. Manual installation of Homebrew is also simple:

<sup>&</sup>lt;sup>7</sup>https://developer.apple.com/xcode/

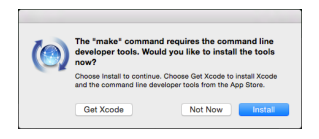


Figure 3: If the user uses the command requiring Xcode for the first time, the system will offer installation of Xcode.

```
67 # Install Homebrew
68 ruby -e "$(curl -fsSL https://raw.githubusercontent.com/Homebrew/install/ \
    master/install)"
70 # Basic help
71 brew help
72 # Install UNIX tools required by Sondovač
73 brew install coreutils gnu-sed gawk grep bash dos2unix pkg-config gcc make \
    git ant wget
75 # List of installed packages (brew formulae)
76 brew list
77 # Information about particular formula
78 brew info FORMULA
79 # Search for applications
80 brew search KEYWORD
81 # Update Homebrew
82 brew update
83 # Update all packages installed by Homebrew
84 brew upgrade
85 # Remove Homebrew package (formula)
86 brew uninstall FORMULA
87 # Cleaning after uninstallation
88 brew cleanup
89 # Completely remove Homebrew (after uninstallation of all formulae)
90 ruby -e "$(curl -fsSL https://raw.githubusercontent.com/Homebrew/install/ \
    master/uninstall)"
92 # Note backslash ("\") means that the code continues on the next line
```

Within the Homebrew project, there is also a scientific section (Homebrew Science, see http://brew.sh/homebrew-science/) containing plenty of software<sup>8</sup>. When the script checks for required scientific packages, it offers several ways to install missing software. Mac OS X users can also use Homebrew. It is the recommended way. If the user wishes to install the software manually, it is possible to use the following commands:

```
93 # Install required scientific packages using Homebrew Science
94 brew install homebrew/science/blat homebrew/science/bowtie2 \
95 homebrew/science/samtools homebrew/science/flash \
96 homebrew/science/fastx_toolkit
97 # Note backslash ("\") means that the code continues on the next line
```

Using Homebrew, software will be installed system-wide, and Homebrew easily allows checks for updates. The only software missing in Homebrew science is bam2fastq, but it is easily installed via another method offered by the script.

<sup>&</sup>lt;sup>8</sup>See https://github.com/Homebrew/homebrew-science/wiki/List-of-homebrew-science-formulae for complete list of available scientific packages.

### 2.4 First launch of Sondovač

Download the latest version from https://github.com/V-Z/sondovac/releases/ and unpack the archive. You can run Sondovač from any directory. In command line navigate to the directory with the unpacked Sondovač files (see Figure 4):

```
98 cd /path/to/directory_with_sondovac
and start it by
99 ./sondovac_part_a.sh -h
to see basic usage instructions. See chapter 3 on page 22 for more information.
```

#### 2.4.1 Examples

See page 22 for explanation of command line parameters. The basic and most simple usage (running in interactive mode, see chapter 3 on page 22):

```
100 ./sondovac_part_a.sh -i
      Specify some of the required input files, otherwise run interactively:
./sondovac_part_a.sh -i -f input.fa -t reads1.fastq -q reads2.fastq
      Running in non-interactive, automated mode (parameter "-n", see chapter 3 at page 22) with
   example data downloaded from https://github.com/V-Z/sondovac/wiki/Sample-data:
   ./sondovac_part_a.sh -f input1_JHCN_Oxalis_corniculata_transcriptome_data.fa \
102
      -c input2_Ricinus_communis_reference_plastid_genome.fsa -m \
     input5_Ricinus_communis_reference_mitochondrial_genome.fasta -t \
104
     input3_J12_Oxalis_obtusa_genome_skim_data_R1.fastq -q \
105
     input4_J12_Oxalis_obtusa_genome_skim_data_R2.fastq -n
106
   # Note backslash ("\") means that the code continues on the next line
      Modify parameter "-a", otherwise run interactively:
```

```
./sondovac_part_a.sh -i -a 300
```

Run in non-interactive mode (parameter "-n", see chapter 3 at page 22) – in such cases the user must specify all required input files (parameters "-f", "-c", "-m", "-t" and "-q"). Moreover, parameter "-y" is modified:

```
./sondovac_part_a.sh -n -f input.fa -c referencecp.fasta \
    -m referencemt.fsa -t reads1.fastq -q reads2.fastq -y 90
```

Modifying parameter "-s". Note that the interactive mode "-i" is implicit and does not need to be specified explicitly:

```
./sondovac_part_a.sh -s 950
```

We recommend launching Sondovač in interactive mode, at least for the first time, so that the script can verify all requirements and install missing tools where needed. We recommend using non-interactive mode for routine usage.

## 2.5 Help for usage of terminal

If you are not familiar with the use of command line, try some basic tutorials first. Some options include:

- https://doc.opensuse.org/documentation/leap/startup/html/book.opensuse.startup/part.bash.html
- https://help.ubuntu.com/community/UsingTheTerminal
- https://www.gnu.org/software/bash/manual/ (advanced full reference manual)
- https://www.debian.org/doc/manuals/debian-reference/ch01.en.html
- https://en.wikibooks.org/wiki/Guide\_to\_Unix

```
\square \square \times
 File Edit View Bookmarks Settings Help
vojta@veles:~> cd /home/vojta/dokumenty/botanak/oxalis/south_africa_target_enrich
ment_genome_skimming/script_probe_design_pipeline/script/
vojta@veles:~/dokumenty/botanak/oxalis/south_africa_target_enrichment_genome_skim
ming/script_probe_design_pipeline/script> ls -a
                                              .info
                                                                   README
                                                                   README~
                                              .info∼
bin
                                              INSTALL
                                                                   sondovac_functions
                                              INSTALL~
geneious_column_separator.pl
                                                                   sondovac_functions~
geneious_column_separator.pl~
                                              LICENSE
                                                                   sondovac_part_a.sh
                                                                   sondovac_part_a.sh~
                                              LICENSE~
.gitignore
                                              mac_aliases
                                                                   sondovac_part_b.sh
.gitignore∼
CHANGELOG
                                              mac_aliases~
                                                                   sondovac_part_b.sh~
                                              manual
                                                                   src
CHANGELOG~
                                              pkgs
vojta@veles:~/dokumenty/botanak/oxalis/south_africa_target_enrichment_genome_skim
ming/script_probe_design_pipeline/script> ./sondovac_part_a.sh -i
          Sondovač is a script to create orthologous low-copy nuclear probes
             Copyright (C) 2015 R. Schmickl, A. Liston, V. Zeisek and others
                  When using this script, please, cite Schmickl et al. 2016
            This is version 0.95 released 2
            For newest version check ht
            In case of problems not covered in README for user support see
           For basic usage see
            For detailed usage instructions see README or
This program is free software: you can redistribute it and/or modify it under
the terms of the GNU General Public License as published by the Free Software
Foundation, either version 3 of the License, or (at your option) any later
version. For more information see LICENSE, https://gnu.org/licenses/gpl.html
or "./sondovac_part_a.sh -l".
This program is distributed in the hope that it will be useful, but WITHOUT ANY
                           SE. See the GNU General Public License for more details.
This part is for filtering of raw data and their preparation for assembly in
Geneious. Results of Geneious assembly are processed in part B to get the final
list of low-copy nuclear probe sequences. See README and/or manual for details.
Running in interactive mode...
Press any key to continue... (or press Ctrl+C to exit the script).
                      script : sondovac part a
```

Figure 4: Starting terminal and navigating to Sondovač. First look at the terminal (command-line, shell) window, navigate to directory with Sondovač (using command cd), listing directory content (command ls) and preparing to launch Sondovač (./sondovac\_part\_a.sh).

- http://tldp.org/LDP/Bash-Beginners-Guide/html/Bash-Beginners-Guide.html
- https://trapa.cz/en/course-linux-command-line-2015
- http://linuxcourse.rutgers.edu/documents/Bash-Beginners-Guide/
- http://ryanstutorials.net/linuxtutorial/
- http://www.hypexr.org/bash\_tutorial.php
- http://mywiki.wooledge.org/BashGuide

#### 2.6 Geneious

For part **B** of the script the user must have Geneious (Kearse et al., 2012). Geneious is a DNA alignment, assembly, and analysis software and one of the most common software platforms used in genomics. It is utilized for de novo assembly in Sondovač. We plan to replace it with a free open-source command line tool in a future release of Sondovač. Visit <a href="http://www.geneious.com/">http://www.geneious.com/</a> for download, purchase, installation and usage of Geneious. After the input data are processed (interactively or not) by <a href="mailto:sondovac\_part\_a.sh">sondovac\_part\_a.sh</a>, the user must process its output manually with Geneious according to the instructions given below. The output of Geneious is then processed by <a href="mailto:sondovac\_part\_b.sh">sondovac\_part\_b.sh</a>, which produces the final probe set. Geneious versions 6, 7 and 8 have been tested and are compatible with this script.

## 2.7 Software used by Sondovač

Table 2 lists all software used by Sondovač, including minimal required versions and homepages. As long as you have a recently-updated version of your operating system and you use the automated installation of additional software offered by Sondovač, you do not need to worry about this. In case you installed some of the required scientific packages manually, ensure that you have the required minimal version. The following list refers to papers and web resources describing methods used by software utilized by Sondovač:

**Table 2:** Required software, its versions and homepages. "X" denotes any subversion of particular lineage and "v. >" denotes any version higher then noted. Generally, any current version should usually be fine.

Software	Version	Homepage
Apache Ant	1.9.X	https://ant.apache.org/
bam2fastq	1.1.0	http://gsl.hudsonalpha.org/information/software/bam2fastq
BASH	v. > 4	https://gnu.org/software/bash/bash.html
BLAT	v.36	http://genome.ucsc.edu/FAQ/FAQblat.html
Bowtie2	2.2.6	http://bowtie- bio.sourceforge.net/bowtie2/index.shtml
CD-HIT	4.6	http://weizhongli-lab.org/cd-hit/
FASTX, libgtextutils	0.0.13	http://hannonlab.cshl.edu/fastx_toolkit/
FLASH	1.2.11	http://sourceforge.net/projects/flashpage/
G++, $GCC$	v. > 4.2	http://gcc.gnu.org/
Geneious	v. > 6.1	http://www.geneious.com/
GIT	v. > 2.0	http://git-scm.com/
GNU core utils	8.X	https://gnu.org/software/coreutils/coreutils.html
grab_synglet- on_clusters.py	1.00	https://github.com/listonlab/Hyb-Seq_protocol/

#### ... continued Table 2.

Software	$\overline{ m Version}$	Homepage
Java/OpenJDK	v. > 7	https://www.java.com//http://openjdk.java.net/
libpng	1.6.X	http://www.libpng.org/
SAMtools,	1.2	http://www.htslib.org/
htsjdk Sondovač	0.9	https://github.com/V-Z/sondovac/wiki
zlib	1.2.8	http://zlib.net/

sondovac\_part\_a.sh requires (and will install) the following software packages:

- BLAT
- Bowtie2
- SAMtools
- bam2fastq (will be replaced by Picard in a future release)
- FLASH
- FASTX-toolkit

sondovac\_part\_b.sh requires (and will install) the following software packages:

- CD-HIT
- BLAT
- grab\_syngleton\_clusters.py (included with Sondovač)

Papers describing the software used by Sondovač:

**BLAT** Kent (2002): BLAT – the BLAST-like alignment tool.

Bowtie2 Langmead and Salzberg (2012): Fast gapped-read alignment with Bowtie 2.

**CD-HIT** There are several papers describing CD-HIT:

- Li et al. (2001): Clustering of highly homologous sequences to reduce the size of large protein databases.
- Li et al. (2002): Tolerating some redundancy significantly speeds up clustering of large protein databases.
- Li and Godzik (2006): Cd-hit: a fast program for clustering and comparing large sets of protein or nucleotide sequences.
- Fu et al. (2012): CD-HIT: accelerated for clustering the next generation sequencing data.
- Huang et al. (2010): CD-HIT Suite: a web server for clustering and comparing biological sequences.
- Niu et al. (2010): Artificial and natural duplicates in pyrosequencing reads of metagenomic data.
- Li et al. (2012): Ultrafast clustering algorithms for metagenomic sequence analysis.

**FASTX toolkit** Gordon and Hannon (2010): FASTX-Toolkit. FASTQ/A short-reads preprocessing tools.

- **FLASH** Magoč and Salzberg (2011): FLASH: fast length adjustment of short reads to improve genome assemblies.
- **Geneious** Kearse et al. (2012): Geneious Basic: An integrated and extendable desktop software platform for the organization and analysis of sequence data.
- grab\_syngleton\_clusters.py Weitemier et al. (2014): Hyb-Seq: Combining target enrichment and genome skimming for plant phylogenomics.

**SAMtools** There are several papers describing SAMtools:

- Li et al. (2009): The Sequence alignment/map (SAM) format and SAMtools.
- Li (2011a): A statistical framework for SNP calling, mutation discovery, association mapping and population genetical parameter estimation from sequencing data.
- Li (2011b): Improving SNP discovery by base alignment quality.

**Sondovač** Schmickl et al. (2016): Phylogenetic marker development for target enrichment from transcriptome and genome skim data: the pipeline and its application in southern African *Oxalis* (Oxalidaceae).

#### 2.8 The PATH variable

PATH (\$PATH) is a system variable used in every UNIX system. It lists directories (separated by colon ":") where the current shell (see also Chapter 2.9 Vocabulary at page 20) searches for binaries (commands), so that the user does not have to specify the full path to the software (e.g. just sed instead of /usr/bin/sed). If some software is installed outside standard locations, the user must specify the full path, even if the user is located in the same directory as the software (e.g. ./sondovac\_part\_a.sh - this is for security reasons). In the case of two commands with the same name (e.g. /bin/somecommand and /usr/bin/somecommand), the order of directories in \$PATH matters - the first occurrence is used, and any later commands are ignored (but this is usually a rare case). PATH can be managed using the following commands:

```
# See the $PATH variable
cho $PATH # Sample output is on the next line:
/home/$USER/bin:/usr/local/bin:/usr/bin:/opt/bin:/sbin:/usr/sbin
# Adding new directory to $PATH
export PATH=$PATH:/some/new/directory
# Do not do it in the following way - it would overwrite $PATH, and
# there would be only the new directory (not the original content)!
export PATH=/some/new/directory # Wrong! Old PATH is missing and will be lost!
# Removing possible duplicate entries in PATH with regular expressions and awk
export PATH="$(echo "$PATH" | awk 'BEGIN{RS=":";}{sub(sprintf("%c$",10),""); \
if(A[$0]){}else{A[$0]=1;printf(((NR==1)?"":":")$0)}}')"
echo $PATH # See it after modifications
# Note backslash ("\") means that the code continues on the next line
```

Sondovač requires certain software to be installed (see Table 2 on page 18), and if some software is missing, the script offers installation. By default, Sondovač creates a directory bin in the current working directory in which it installs the required software. It then temporarily modifies the content of the \$PATH variable to contain this new directory. Sondovač notifies the user about this, and the user can then – if it is wished to keep the newly installed software for later usage – (1) move the content of this directory to a preferred location or (2) add this directory to the \$PATH. This directory can also be safely removed. Permanent modification of the \$PATH variable is done by adding line export PATH=\$PATH:/some/new/directory (same as the above example) to file ~/.bashrc or ~/.bash\_profile (usage of those files varies slightly among UNIX systems, see manual for your operating system). On Mac OS X, installation

of Homebrew is required. For correct functioning of Sondovač, \$PATH is modified to contain directories /usr/local/opt/coreutils/libexec/gnubin and /usr/local/bin containing new and updated UNIX tools. The first directory must always be present, as it contains updated versions of basic command line utilities – replacing outdated versions provided with Mac OS X. All those modifications are temporary and used only within Sondovač scripts.

## 2.9 Vocabulary

- **Binary** An application in a form that is understandable by the computer, but usually not transferable among operating systems and/or hardware platforms. Binaries in Windows usually have the extension \*.exe, in UNIX there is usually no extension.
- **BASH** "The command line" fully featured programming scripting language accessible through the terminal of any UNIX-based operating system (any Linux, Mac OS X, Solaris, any variant of BSD and more). BASH scripts usually have the extension \*.sh.
- **BSD** Group of popular UNIX-based operating systems. See https://en.wikipedia.org/wiki/Berkeley\_Software\_Distribution.
- C Popular programming language. Source code must be compiled for each operating system. See https://en.wikipedia.org/wiki/C\_(programming\_language).
- C++ Popular programming language. Source code must be compiled for each operating system. See https://en.wikipedia.org/wiki/C++.
- Centos Popular Linux distribution. Community remake of RedHat Enterprise Linux. See <a href="https://centos.org/">https://centos.org/</a>.
- Compilation "Translation" of software application from the source code (text readable by human programmer) into binary form launchable by the computer. It requires special tools (compilers), and it usually must be done for every operating system and hardware platform.
- Console See "Shell".
- **Debian** One of the oldest and most popular Linux distributions. See https://www.debian.org/.
- **Fedora** Popular Linux distribution developed together with RedHat Linux as its free community testing platform. See https://getfedora.org/.
- GNU Major project providing free software widely used in many operating systems, see https://gnu.org/.
- **Homebrew** Tool primarily for Mac OS X (although there is also a Linux version available) replacing the virtually missing package manager for this system. Can be used to install plenty of various applications as well as updating tools already available in Mac OS X. See <a href="http://brew.sh/">http://brew.sh/</a>.
- Java Very popular programming language. It requires Java runtime environment to be installed, but the applications are very well transferable among operating systems. See <a href="https://www.java.com/">https://www.java.com/</a>.
- Library Pack of software tools and functions used by other applications.
- **License** Conditions under which software is distributed. Can be very restrictive (typically paid software) or permissive (typically free and open-source software).

- Linux One of the most common variants of UNIX-based operating systems. Linux kernel is used by many developers, so that there are plenty of Linux distributions ("flavors") from various sources (e.g. Ubuntu and derivatives, openSUSE, SLE, Debian, Linux Mint, Fedora, Centos, RedHat etc.). They share many features, although at first sight they can look different. See https://en.wikipedia.org/wiki/Linux.
- **Linux Mint** Popular Linux distribution based on Debian and Ubuntu, see http://linuxmint.com/.
- Mac OS X Popular operating system produced by Apple. The system kernel is based on UNIX, see https://www.apple.com/osx/.
- **Open-source** Generally, the source code of an application is available together with the application and can, under certain conditions, be defined in license modified, redistributed etc. See https://en.wikipedia.org/wiki/Free\_and\_open-source\_software.
- openSUSE Popular Linux distribution, see https://www.opensuse.org/.
- Operating system Basic system running on your computer typically MS Windows (not supported by Sondovač, although it might work), Mac OS X or some Linux distribution (Ubuntu and derivatives, openSUSE, SLE, Debian, Linux Mint, Fedora, Centos, RedHat etc.).
- Package Software or its part, group of tools, library etc. Basic unit of software management in most UNIX systems (mainly Linux, Solaris, BSD, practically missing in Mac OS X). Those systems usually have special applications (command line as well as graphical tools) to easily manage (install, remove, update) software.
- Parameter(s) Option(s) passed to any function/command line application to modify its usage. Some can be required, some are optional, and some can be used only in particular cases. In case of shell applications, parameters are usually given such as "application -X", "application -parameter", "application -Param SomeValue" and so on. See manual for particular application (e.g. "man application"), in case of Sondovač see page 22.
- **PATH** Directories in the computer where the system looks for installed software (in a UNIX-based system you can view it by the command "echo \$PATH"). If you need to modify it manually, see the documentation for your operating system.
- **Perl** Popular interpreted programming language excelling mainly in system tasks working with text. Perl scripts are easily transferable among operating systems. See <a href="https://www.perl.org/">https://www.perl.org/</a>.
- **Python** Very popular and powerful interpreted programming language used for wide variety of tasks. Python scripts are easily transferable among operating systems. See <a href="https://www.python.org/">https://www.python.org/</a>.
- **RedHat** Probably the biggest Linux company providing mainly solutions for big companies. See https://www.redhat.com/.
- **Repository** Internet folder (available through HTTP or FTP) containing software packages for UNIX systems.
- Scientific Linux Popular Linux distribution. Community remake of RedHat Enterprise Linux. See https://www.scientificlinux.org/.

- Script Software application. It requires an interpreter (application installed on the computer that is able to launch scripts written in a particular language), but the application itself is portable among operating systems and hardware architectures, and it is written in plain text, so that developers can easily modify it. Common examples are Python, Perl or BASH.
- **Shell** "The command line" the interface to interact with software using commands typed into the terminal window (See Figure 4).
- **Solaris** Popular (mainly on servers) UNIX-based operating system, now developed by Oracle and including several independent clones. See http://distrowatch.com/table.php?distribution=solaris.
- Source code Human-readable code written in any text editor used to develop any application. Applications written in interpreted languages (BASH, Perl, Python, ...) can be distributed just in form of a source code (nothing else is required). Other programming languages (C, C++, ...) require compilation to get a fully functional application.
- SUSE Linux Enterprise (SLE) Large Linux company providing mainly solutions for big companies. See https://www.suse.com/.

Terminal see "Shell".

- Ubuntu Popular Linux distribution, see http://www.ubuntu.com/. There are plenty of distributions based on Ubuntu. See http://distrowatch.com/search.php?basedon=Ubuntu.
- UNIX (UNIX-like, UN\*X, \*nix, ...) Family of operating systems sharing the same logic, software architecture and plenty of tools. See https://en.wikipedia.org/wiki/Unix-like for details.
- **Upstream** Developers usually support (e.g. by fixing of bugs) only newer versions of an application. If you use an older version and you encounter problems, no one can probably help you. Moreover, using old versions of software can be a security risk because of security issues fixed in newer versions.
- Variable Named value storing various information, one of the basic part of any programming language, application, operating system.

# 3 Usage of Sondovač

#### 3.1 Command line parameters

Sondovač has some parameters that are useful especially for advanced users, on remote servers, for repeated analyses and so on. We recommend starting with basic interactive usage – the script will ask for the input files and, if needed, also for installation of additional software.

```
# Go to directory with unpacked Sondovač (in terminal):

cd /path/to/directory_with_sondovac

# Run sondovac_part_a.sh in basic interactive mode:

./sondovac_part_a.sh -i

# Then run Geneious and continue with sondovac_part_b.sh:

./sondovac_part_b.sh -i
```

#### 3.1.1 General parameters

Shared by sondovac\_part\_a.sh as well as sondovac\_part\_b.sh.

- -h, -v Print help message and exit.
- -u Check for updates. If there is a newer version of Sondovač available on https://github.com/V-Z/sondovac/releases/, download of the newer version will be offered to the user.
- -1 Display LICENSE for license information (this script is licensed under GNU GPL v.3, other software under variable licenses). Exit viewing by pressing the ℚ key.
- -r Display README for detailed usage instructions. Exit viewing by pressing the Q key.
- Display INSTALL for detailed installation instructions. Exit viewing by pressing the Q key. See also page 9.
- -e Display detailed citation information and exit.
- Set name of output files. Output files will start with that name. Do not use spaces or special characters some software can't handle them correctly. The default value (if the user does not provide another name) is "output". See below for the list of produced output files.
- -i Running in interactive mode the script will on-demand ask for the required input files, installation of missing software etc.. This is the recommended default value (the script runs interactively without explicitly using option -n).
- -n Running in non-interactive mode. The user must provide at least the required input files (see below). You can use only one of the parameters -i or -n (not both of them). If the script fails to find some of the required software packages, it will exit. This is recommended for batch or repeated analysis, on remote servers and for more advanced users. The user must be sure that all required software is installed (see page 9).

## 3.1.2 Input files

Those parameters are required when running the script in non-interactive mode. The parameters are optional in default interactive mode. Please use file names without spaces and without special characters.

- **-f FILE** Transcriptome input file in FASTA format.
  - sondovac\_part\_a.sh
- -c FILE Plastome reference sequence input file in FASTA format.
  - sondovac\_part\_a.sh, sondovac\_part\_b.sh
  - Plastome reference sequences from taxa up to the same order of the studied plant group are suitable. See Straub et al. (2012).
- -m FILE Mitochondriome reference sequence input file in FASTA format (optional).
  - sondovac\_part\_a.sh
  - This step is optional, as plant mitochondrial genomes have largely variable sizes and high rearrangement rates.
- -t FILE Paired-end genome skim input file in FASTQ format (first file).
  - sondovac\_part\_a.sh

- -q FILE Paired-end genome skim input file in FASTQ format (second file).
  - sondovac\_part\_a.sh
- -x FILE Input file in TSV format (output of Geneious assembly).
  - sondovac part b.sh
- -z FILE Input file in FASTA format (output of Geneious assembly).
  - sondovac\_part\_b.sh

### 3.1.3 Optional parameters

See page 5 and Figure 1 for steps referred here. If those parameters are not provided, the default values are used, and it is not possible to change them any time later (not even in interactive mode).

- -a ### Maximum overlap length expected in approximately ≥90% of read pairs (parameter -M of FLASH, see its manual for details).
  - Step 4 of Sondovač, sondovac\_part\_a.sh.
  - FLASH can not combine paired-end reads that do not overlap by at least 10 bp (default minimum overlap length).
  - DEFAULT: 65
  - OPTIONS: Integer ranging from 10 to 300
- -y ## Sequence similarity between unique transcripts and the filtered, combined genome skim reads (parameter -minIdentity of BLAT, see its manual for details).
  - Step 5 of Sondovač, sondovac\_part\_a.sh.
  - Filtering for orthologs, using sequence similarity as criterion.
  - DEFAULT: 85 (highly recommended)
  - OPTIONS: Integer ranging from 70 to 100
- -g Choice of transcript or genome skim sequences for further processing.
  - Step 6.1 of Sondovač, sondovac\_part\_a.sh.
  - Depending on the phylogenetic depth that should be obtained, the probe sequences need to be designed from either the transcript or genome skim sequences, or it might not matter (if the taxa, from which the transcriptome and genome skim data were generated, are closely related).
  - DEFAULT: no usage of -g (probe design from genome skim sequences)
  - OPTIONS: usage of -g (probe design from transcript sequences)
- -s #### Number of BLAT hits per transcript when matching unique transcripts and the filtered, combined genome skim reads.
  - Step 6.2 of Sondovač, sondovac\_part\_a.sh.
  - Transcripts with a high number of BLAT hits, indicating repetitive elements, need to be removed from the putative probe sequences.
  - DEFAULT: 1000
  - OPTIONS: Integer ranging from 100 to 10000

- -b ### Minimum exon (bait) length.
  - Steps 8 and 10 of Sondovač, sondovac\_part\_b.sh.
  - The minimum exon length should not fall below the bait length in order to account for specific binding between genomic libraries and baits during hybridization.
  - DEFAULT: 120 (preferred length for phylogeny)
  - OPTIONS: 80, 100, 120
- -k ### Minimum total locus length.
  - Steps 8 and 10 of Sondovač, sondovac\_part\_b.sh.
  - When running the script in interactive mode, the user will be asked which value to use. A table summarizing the total number of LCN loci, which will be the result of the probe design for all minimum total locus lengths that the user can select (600 bp, 720 bp, 840 bp, 960 bp, 1080 bp, 1200 bp), will be displayed to facilitate this choice.
  - DEFAULT: 600
  - OPTIONS: 720, 840, 960, 1080, 1200
- -d 0.## Sequence similarity between probe sequences (parameter -c of cd-hit-est, see its manual for details).
  - Step 9 of Sondovač, sondovac\_part\_b.sh.
  - Probes that target multiple similar loci need to be removed.
  - DEFAULT: 0.9 (highly recommended)
  - OPTIONS: Decimal ranging from 0.85 to 0.95
- -y ## Sequence similarity between probe sequences and plastome reference (parameter -minIdentity of BLAT, see its manual for details).
  - Step 11 of Sondovač, sondovac part b.sh.
  - Some plastid reads might not have been removed in step 2; they should be removed in this step.
  - DEFAULT: 90 (highly recommended)
  - OPTIONS: Integer ranging from 70 to 100

### 3.2 Input and output files

All names of input files and paths to them must be without spaces and without special characters (some software has difficulties handling them). Important note: HTS data are big. The Sondovač pipeline is relatively long, and part A contains several format conversions and can (for some time) use dozens of GB of disk space. Temporary files not potentially useful to the user are deleted at the end of the pipeline – these files may be useful for debugging if something goes wrong. For example, input data of Schmickl et al. (2016) are approximately 4.5 GB, and the overall output of part A of the script is about 28 GB, of which less then half is kept by the pipeline. This analysis took less than an hour on an if 3.4 GHz CPU. Part B is very quick and does not consume a significant amount of disk space. All input files must have UNIX end of lines. The script checks for it and converts the files, if needed (using dos2unix; typically when user runs Geneious on Windows).

Script sondovac\_part\_a.sh requires as input files:

- 1. Transcriptome input file in FASTA format. **Note:** For technical reasons, the labels of FASTA sequences *must* be unique numbers (no other characters). Sondovač will check the labels, and if they are not in an appropriate form, a copy of this input file with correct labels will be created.
- 2. Plastome reference sequence input file in FASTA format.
- 3. Paired-end genome skim input file in FASTQ format (two files forward and reverse reads).
- 4. OPTIONAL: Mitochondriome reference sequence input file in FASTA format. This file is not required.

## Script sondovac\_part\_a.sh creates the following files:

- 1. \*\_renamed.fasta A copy of the transcriptome input file with the changed labels of the FASTA sequences (unique numbers corresponding to the line numbers in the original file). File \*\_old\_and\_new\_names.tsv then contains two columns: 1) the original sequence labels as in the user-provided transcriptome input file and 2) new sequence labels. This might be useful to trace back certain sequences/probes.
- 2. \*\_blat\_unique\_transcripts.psl − Output of BLAT (removal of transcripts sharing ≥90% sequence similarity).
- 3. \*\_unique\_transcripts.fasta Unique transcripts in FASTA format.
- 4. \*\_genome\_skim\_data\_no\_cp\_reads Genome skim data without cpDNA reads.
- 5. \*\_genome\_skim\_data\_no\_cp\_no\_mt\_reads Genome skim data without mtDNA reads only if mitochondriome reference sequence was used.
- 6. \*\_combined\_reads\_co\_cp\_no\_mt\_reads Combined paired-end genome skim reads.
- 7. \*\_blat\_unique\_transcripts\_versus\_genome\_skim\_data.pslx Output of BLAT (matching of the unique transcripts and the filtered, combined genome skim reads sharing ≥85% sequence similarity).
- 8. \*\_blat\_unique\_transcripts\_versus\_genome\_skim\_data.fasta Matching sequences in FASTA.
- 9. \*\_blat\_unique\_transcripts\_versus\_genome\_skim\_data-no\_missing\_fin.fsa Part A, final FASTA sequences for usage in Geneious (step 7, see chapter 3.3 at page 27, and page 5).

Files 1-8 are not necessary for further processing by this pipeline, but may be useful to the user. The last file (9) is used as input file for Geneious in the next step. An asterisk (\*) denotes the beginning of the output files' names specified by the user with parameter -o. If the user does not select a custom name, default value (output) will be used.

Geneious requires as input the last output file of sondovac\_part\_a.sh (file 9: \*\_blat\_uni que\_transcripts\_versus\_genome\_skim\_data-no\_missing\_fin.fsa). The output from Geneious consists of two files (see page 27):

- 1. Final assembled sequences exported as TSV.
- 2. Final assembled sequences exported as FASTA.

#### Script sondovac\_part\_b.sh requires as input files:

1. Plastome reference sequence input file in FASTA format.

- 2. Assembled sequences exported from Geneious as TSV.
- 3. Assembled sequences exported from Geneious as FASTA.

#### Script sondovac\_part\_b.sh creates the following files:

- 1. \*\_prelim\_probe\_seq.fasta Preliminary probe sequences.
- 2. \*\_prelim\_probe\_seq\_cluster\_100.fasta Unclustered exons and clustered exons with 100
- 3. \*\_prelim\_probe\_seq\_cluster\_90.clstr Unclustered exons and clustered exons with more than a certain sequence similarity (CLSTR file).
- 4. \*\_unique\_prelim\_probe\_seq.fasta Unclustered exons / exons with less than a certain sequence similarity.
- 5. \*\_similarity\_test.fasta Contigs that comprise exons ≥ bait length and have a certain total locus length.
- 6. \*\_target\_enrichment\_probe\_sequences.fasta Final probes in FASTA.
- 7. \*\_possible\_cp\_dna\_gene\_in\_probe\_set.pslx In case of any BLAT hits, the user needs to manually remove these plastid probe sequences from \*\_target\_enrichment\_probe\_se quences.fasta; the remaining ones are the final probe sequences in FASTA.

An asterisk (\*) denotes the beginning of the output files' names specified by the user with parameter -o. If the user does not select a custom name, the default value (output) will be used. By default, output files are created in the same directory from which Sondovač was launched. Output files can be saved in a custom directory by specifying an output directory with parameter -o:

```
# Find current directory (e.g. /home/user):

pwd

# Launching Sondovač located in directory /home/user/sondovac

# and save output to e.g. desktop (/home/user/Desktop):

//sondovac/sondovac_part_a.sh -o Desktop/MyFile

# Sondovač will save software (if needed) in "bin" directory

# located in directory from which it was launched, see it:

ls bin/*

# Output files are in desired directory, see them e.g. by:

ls -lh Desktop/MyFile*
```

#### 3.3 Geneious usage

Import the output file of part A of the script (sondovac\_part\_a.sh): go to menu File | Import | From File... This file is named as: \*\_blat\_unique\_transcripts\_versus\_genome\_skim\_data-no\_missing\_fin.fsa (see Figure 5).

Select the file and go to menu Tools | Align / Assemble | De Novo Assemble. In Data frame select Assemble by 1st (...) Underscore. In Method frame select Geneious Assembler (if you don't have other assemblers, this option might be missing) and Medium Sensitivity / Fast sensitivity (see Figure 6).

In Results frame check Save assembly report, Save list of unused reads, Save in subfolder, Save contigs (do not check Maximum) and Save consensus sequences (Click to Options – Save consensus used by assembler must be selected.). Do not trim. Otherwise keep defaults. Run it. Geneious may warn about possible hanging because of big file size. Do

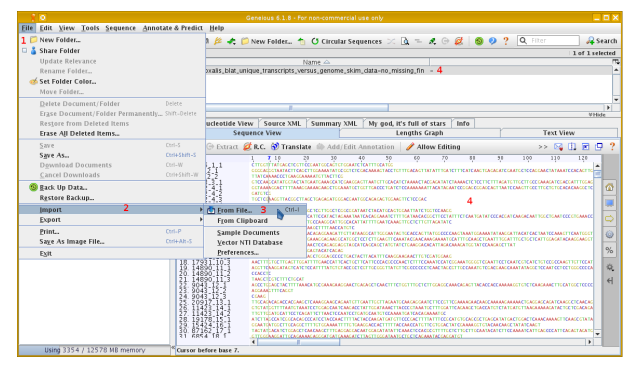


Figure 5: Import of output of sondovac\_part\_a.sh into Geneious for next processing (see page 5). Go to menu File (1) | Import (2) | From file...(3) and import output of sondovac\_part\_a.sh. You should see a similar result as (4).

not use Geneious for other tasks during the assembly. Running Geneious may take a long time (see Figure 6).

Select all resulting contigs (typically named \* Contig #) and export them (go to menu File | Export | Selected Documents...) as Tab-separated table values (\*.tsv). Save the following columns (Hold Ctrl key to mark more fields): # Sequences, % Pairwise Identity, Description, Mean Coverage, Name and Sequence Length. If this option is inaccessible to you, export all columns (see Figure 7). Warning! Do not select and export \* Consensus Sequences, \* Unused Reads or \* Report – only the individual \* contig # files (see Figure 7).

Select items **Consensus Sequences** and **Unused Reads** and export them as one **FASTA**. Go to menu **File** | **Export** | **Selected Documents...** and choose **FASTA** file type (see Figure 8).

Use the exported files from Geneious as input for part B of the script (sondovac\_part\_b.sh).

## 3.4 Record output of Sondovač

To record the whole output of Sondovač (regardless of used parameters), use utility tee. This will produce a plain text output with everything printed to the screen. It can be useful for reference or exploration if something went wrong. Use it as follows:

```
./sondovac_part_a.sh | tee records.log
man tee # See more options how tee can record the script's output
# "|" is a~pipe passing output of the 1st command as input for the 2nd command
less records.log # See the record. Quit viewing by "Q"
rm records.log # Delete the log file
```

You can use any command line arguments; the script will behave as usual. The plain text file records.log will then contain all its output. Unfortunately, tee usually wrongly records "invisible" characters – tabs and coloration used to highlight user messages in the script. If you see weird characters in textttrecords.log that disturb reading, use the following commands:

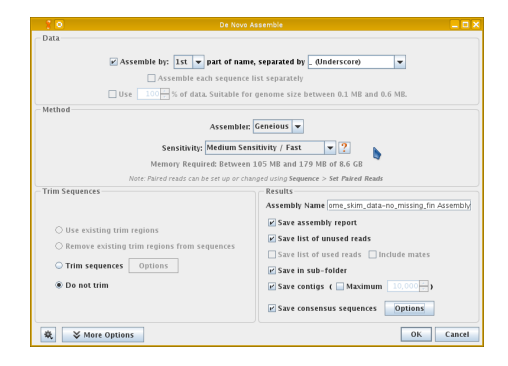


Figure 6: Settings of Geneious assembly as described in the main text. It can take a longer time to run it.

```
# Assume output of Sondovač is named "records.log"

sed -i 's/.\[[[:digit:]]\{1,2\}m//g' records.log

# Explanation of regular expression (find pattern and replace by nothing):

# any character, [, one/two number(s), m (sequence defining text formatting)

sed -i 's/.(B.\[m//g' records.log

# Explanation of regular expression (find pattern and replace by nothing):

# any character, (, B, [, m (sequence defining text formatting)

# Escaping \[ \] is required to search specifically for brackets []

# (NOT searching for any character within [...] - there is no escaping)

# but \{...\} define number of occurrences of previous character(s)
```

Note for Mac OS X users: the regular expressions above require GNU sed, not the version presented by default in Mac OS X. It is installed by Sondovač using Homebrew (see page 13), but to launch it you probably must use the command gsed instead of sed.

# 4 Sample data

Together with the script, we provide the ZIP archive (1.8 GB) that contains example input files for running the script: Oxalis genome skim data as well as the Ricinus cpDNA and mtDNA reference sequences. See https://github.com/V-Z/sondovac/wiki/Sample-data for download of sample data.

The package contains:

- 1. input2\_Ricinus\_communis\_reference\_plastid\_genome.fsa cpDNA reference (parameter -c), GenBank reference number NC\_016736.
- 2. input3\_J12\_0xalis\_obtusa\_J12\_genome\_skim\_data\_R1.fastq paired-end genome skim data, file 1 (forward reads, parameter -t).
- 3. input4\_J12\_0xalis\_obtusa\_J12\_genome\_skim\_data\_R2.fastq paired-end genome skim data, file 2 (backward reads, parameter -q).

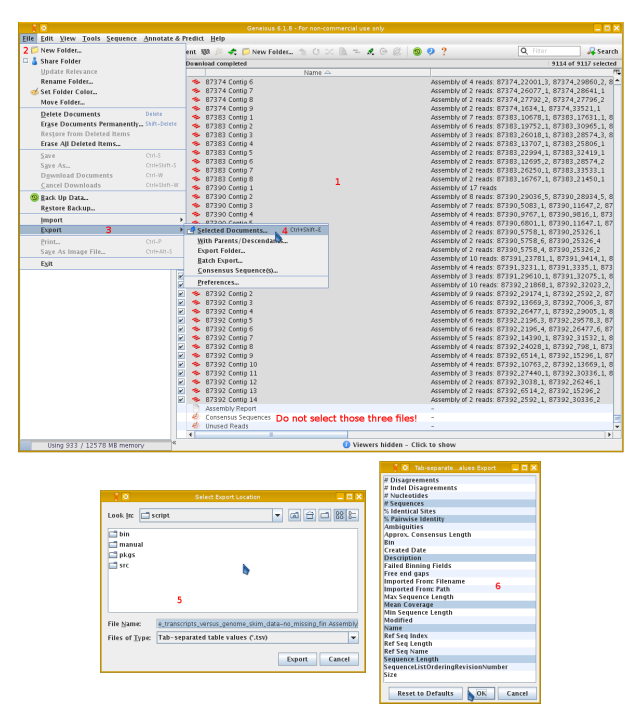


Figure 7: Select all (and only) \* Contig # files (1). Go to menu File (2) | Export (3) | Selected Documents...(4) and export them as Tab-separated table values (TSV) (5). Export only marked columns (6) (hold Ctrl to mark more fields).

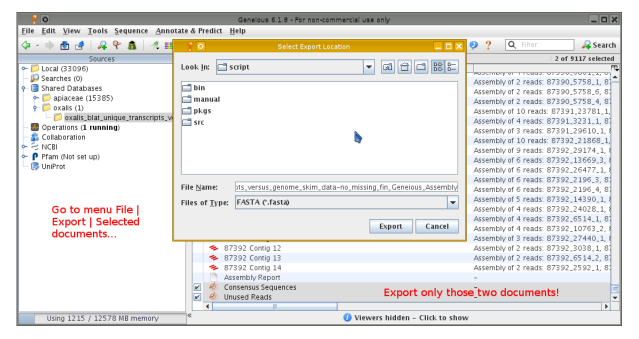


Figure 8: Select only documents Consensus Sequences and Unused Reads and export them as FASTA format (see also Figure 7).

4. input5\_Ricinus\_communis\_reference\_mitochondrial\_genome.fasta - mtDNA reference (parameter -m), GenBank reference number NC\_015141.

The transcriptome input file is unpublished data from G. K.-S. Wong et al.. As soon as the data are published, we will post them on GitHub. Data can now be found under

- http://www.onekp.com/
- http://www.onekp.com/samples/list.php
- http://www.onekp.com/samples/single.php?id=JHCN

The transcriptome FASTA file used for the probe design is named JHCN-SOAPdenovo-Trans-assembly.dnas.out and can be found under JHCN/Assembly/JHCN-SOAPdenovo-Transtranslated/. Information about how to get access to data download is given in Matasci et al. (2014).

Explanation of command line parameters is on page 22.

# 5 Questions not covered here, reporting bugs and wishes

If you have a question or encounter a problem, please see https://github.com/V-Z/sondovac/issues and feel free to ask any question and/or express any wish. The authors will do their best to help you.

# 6 Changelog

List of changes in released versions of Sondovač.

#### 6.1 Version 1.1 regular release released 2016-03-15

• Checking if input FASTA files are interleaved or not (required) and, if needed, FASTA files are converted not to be interleaved.

- Added requirement of Python.
- Removed BAM files (part A) they are not kept anymore.
- Language checking and enhancements of documentation.
- Modified CD-HIT-EST.
- Improved summary of the probe design in part B.
- More possibilities for minimum total locus length.
- Various smaller fixes.

### 6.2 Version 1.0 regular release released 2016-01-12

- Renaming of input FASTA sequences names is required it ensures correct working of part B.
- Added check if input files were created on Windows if so, they are converted into UNIX style EOL.
- Various smaller fixes.
- Better showing of the information in part B.
- Enhanced documentation.

#### 6.3 Version 0.99 release candidate released 2015-12-08

- Fixed error with some input files for part B.
- Finished coloration of command-line user interface.
- Added possibility to set minimal exon length of the loci.
- Various fixes and UI enhancements.
- Improved documentation.

## 6.4 Version 0.95 beta released 2015-11-27

- Offer the possibility to choose between transcripts or genome skim sequences for further processing in step 6.1, part A.
- Coloration of command-line user interface (incomplete).
- Added possibility to change -minIdentity parameter of BLAT in step 11, part B.
- Fixed problems with some transcriptome input files.
- Added possibility to set custom bait length.
- Added information about article in MER introducing Sondovač.

## 6.5 Version 0.9 beta released 2015-10-23

- Highly enhanced part B.
- Better handling of variable output from Geneious.
- Possibility to specify the name of the custom output file.
- Full support for Linux distributions using DEB Debian, Ubuntu, Linux Mint and derivatives.
- Enhanced documentation.
- Support for Mac OS X, package management using Homebrew.
- Support for RedHat based Linux distributions Fedora, Centos and Scientific Linux and derivatives.
- Better compilation and installation of required software.
- For downloading automatically select whether to use wget (preferred) or curl.
- · Various fixes.

# 6.6 Version 0.8 alpha released 2015-10-09

- Usage of mitochondrial reference sequence is optional.
- Better formatting of script messages.
- Various fixes and enhancements.

## 6.7 Version 0.7 alpha released 2015-10-06

- Fixed reported problems with sed differences among Linux and Mac OS X.
- Added more exhaustive documentation.
- Various fixes and enhancements.

## 6.8 Version 0.6 alpha released 2015-08-10

- Fixed problems with some versions of output of Geneious.
- Better compilation and installation of required additional software packages.
- Various fixes and enhancements.

#### 6.9 Version 0.5 alpha released 2015-07-24

• First public release, early alpha stage.

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a user through a computer network, with no transfer of a copy, is not conveying.

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