User Manual

LRSDAY: Long-read Sequencing Data Analysis for Yeasts

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INTRODUCTION

Background

Twenty years ago, the genome sequence of the budding yeast Saccharomyces cerevisiae was published¹. As the first complete eukaryotic genome ever sequenced, this marked a major scientific milestone in biology. Since then, the genomes of many model and non-model organisms have been sequenced especially following the emergence of next-generation sequencing (NGS) technologies. Despite the notably improved throughputs, NGS technologies suffer from the limitation of short reads and usually result in highly fragmented genome assemblies containing numerous gaps and local mis-assemblies. The recently developed longread sequencing technologies represented by Pacific Biosciences (PacBio) and Oxford Nanopore offer compelling alternatives to overcome such hurdles, producing high-quality genome assemblies with significantly improved continuity and accuracy. Although initially tested in microbial genome sequencing, their recent applications in complex mammalian and plant genomes also achieved high-quality results²⁻⁶. With such new sequencing technologies, challenging genomic regions with enriched repetitive elements, strongly biased GC-content, or complex structural variants can often be correctly resolved. It is therefore anticipated that genome sequencing projects will routinely adopt long-read-based sequencing technologies in the coming years to gain insight in these complex genomic regions.

Yeast is a leading model organism with great importance in both basic biomedical research and biotechnological applications. Its small genome size makes it particularly suitable for long-read-based high-coverage genome sequencing. The resulting complete genome assembly with fully-resolved subtelomere structure can in turn illuminate the genetic basis of many complex phenotypic traits with unprecedented resolution. Recently, we used the long-read sequencing technologies to generate the first panel of population-level end-to-end reference genomes of 12 yeast strains representing major subpopulations of the partially domesticated *S. cerevisiae* and its sister species *Saccharomyces paradoxus*⁷. In addition, there have been a number of other studies carrying out long-read sequencing for many *S. cerevisiae* strains^{8–11}. Given the vast genomic and phenotypic diversity of *S. cerevisiae*, we expect the incoming collection of long-read-based high-quality genome assemblies of strains from widespread geographic locations and ecological niches will substantially deepen our understanding in the *S. cerevisiae* natural genetic variation and its associated biotechnological values.

Overview of the LRSDAY workflow

Here we present a highly organized and modular computational framework named Long-Read Sequencing Data Analysis for Yeasts (LRSDAY), which enables automated high-quality yeast genome assembly and annotation production from raw long-read sequencing data. The prototype of LRSDAY has been developed to generate the yeast population reference panel (YPRP) in our previous study⁷. Under the hood, LRSDAY contains a series of task-specific modules handling long-read-based de novo genome assembly, short-read-based assembly polishing, reference-quided assembly scaffolding, as well as comprehensive genomic feature annotations. These tasks can be run individually, selectively or coordinately depending on users' needs. LRSDAY supports both leading long-read sequencing technologies: PacBio and Oxford Nanopore. Running the full LRSDAY workflow, the final output is a chromosome-level genome assembly with high-quality annotations of centromeres, protein-coding genes, tRNAs, transposable elements (TEs; Ty1-Ty5 for S. cerevisiae and S. paradoxus), and telomere associated core X and Y' elements. LRSDAY is shipped with various auxiliary scripts, configuration files and supporting data that enable its semi-automatic installation, configuration, and execution with minimal manual intervention (Appendix 1). This design concept greatly alleviates the technical barrier for bench biologists with limited bioinformatics experiences. In addition, a real case example and its final outputs are also provided for users' test and comparison. All these task-specific modules, auxiliary files, installed tools, sample outputs, together with the user-created project directories for the testing example and users' own data are hosted under the same home directory (\$LRSDAY_HOME) in a self-contained fashion (Fig. 1). This design makes LRSDAY well-isolated from the rest of the system and therefore greatly improves its portability. To sum up, LRSDAY is a highly transparent, automated and powerful computational framework that handles both genome assembly and annotation, which suits the needs of the yeast community in performing long-read-based genome sequencing projects. In the PROCEDURE section of this article, we provide a step-by-step walkthrough on how to install, configure, and run LRSDAY with our prepared testing example.

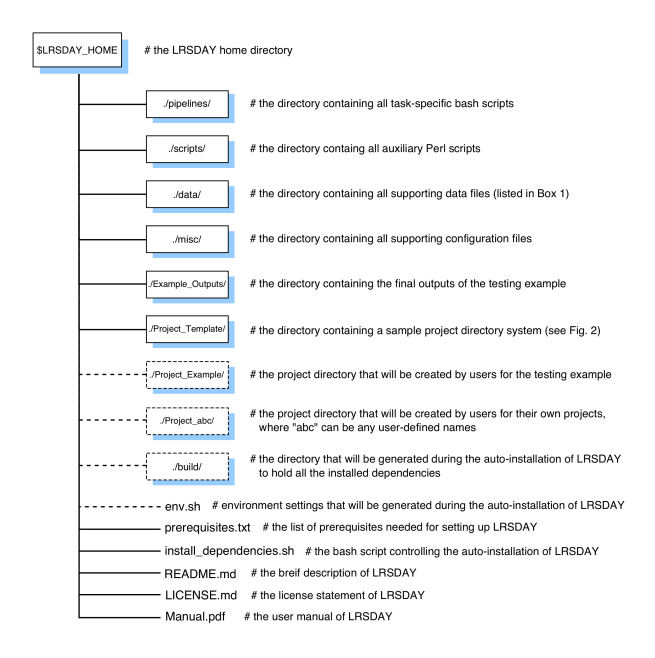


Figure 1. Overview of the LRSDAY directory system. All the top-level directories (boxes, solid lines) and individual files of LRSDAY are listed and briefly described. Additional directories and files will be generated during the installation or execution of LRSDAY (boxes, dashed lines).

Comparison with other methods

Genome assembly is a rapidly moving field, co-evolving with the fast-paced development of sequencing technologies. In recent years, both hybrid (i.e. using both long and short reads) and native (i.e. using long reads only) assemblers that support long-read sequencing data have

been developed and tested on many different organisms^{8,12–17}. As for gene annotation, there is also a wide range of choices that perform gene model prediction in an *ab initio* fashion or based on additional evidences (e.g. mRNA transcripts, protein-sequence alignment)^{18–20}. Specifically for yeasts, a web-based gene annotation tool has been developed that combines both approaches²¹. However, there is currently no integrated solution that handles both genome assembly and annotation in a seamless way. To fill this gap, we revamped our original workflow for deriving the yeast population-level reference genomes⁷ into a self-contained package to considerably streamline this process with modular design and automated implementation. Moreover, rather than simply combining the existing tools for genome assembly and gene annotation, LRSDAY assembled a well-integrated workflow with many other functionalities (e.g. reference-guided scaffolding, gene orthology identification, and additional genomic feature annotation) built in, which makes it a unique one-stop solution for high-quality genome assembly and annotation production from long-read sequencing data.

Experimental Design

Genome assembly and annotation are complex computational processes with many intermediate steps and inputs/outputs involved. With LRSDAY, we designed a highly structured project directory system to help users to run the whole workflow in an organized and modular way (Fig. 2). Within such project directory system, the three subdirectories holding the preshipped reference genome as well as the user-supplied long (PacBio or Oxford Nanopore) and short (Illumina) reads are numbered as "00" and the task-specific subdirectories are numbered sequentially from "01" to "13" according to their execution orders. For each subdirectory, a selfexplained name is attached after the number index to help users to navigate through the workflow. To run each task, users only need to edit (i.e. to specify the input and output file paths and certain task-specific parameters) and execute the task-specific pipeline scripts pre-placed in these subdirectories. These pipeline scripts will automatically set environment variables, process the data, and formulate the results. All computationally intensive tasks can be processed using multiple threads to substantially save computation time. Although LRSDAY is mainly designed for yeasts, most of these tasks can be further adapted for the analysis on any eukaryotic organisms (See Appendix 2 for details). Below we briefly describe the computational processes executed by each task-specific module in LRSDAY with the corresponding PROCEDURE Step labeled in parentheses.

- **01.** Long-read-based_Genome_Assembly (Step 13): Long reads generated from PacBio or Oxford Nanopore technologies are used to perform *de novo* genome assembly using an overlap-layout-consensus (OLC) algorithm.
- 02. Illumina-read-based_Assembly_Polishing (Step 15; optional): Illumina reads are first clipped and trimmed to remove potential sequencing adapters and low-quality regions. The cleaned reads are subsequently mapped to the raw long-read-based genome assembly. The resulting BAM file is further processed for alignment sorting, mate information and read group fixing, duplicates removal as well as local realignment. Finally, the processed bam file is used for correcting base-level errors of the long-read-based assembly.
- 03. Reference-guided_Assembly_Scaffolding (Step 16): The contigs from the polished genome assembly are first aligned to the reference genome to identify their shared sequence homology, based on which reference-guided assembly scaffolding is subsequently performed. The chromosomal identity of each scaffold is labeled accordingly. Structural rearrangements captured in the contigs will remain untouched during the scaffolding.
- **04. Centromere_Identity_Profiling** (Step 18): The pre-shipped *S. cerevisiae* centromere sequences are searched against the scaffolded assembly for chromosome-specific centromere identity profiling.
- **05. Mitochondrial_Genome_Assembly_Improvement** (Step 19): The polished contigs corresponding to the mitochondrial genome are re-collected from the scaffolded assembly. The mitochondrial contigs spanning over the designated starting point (the *ATP6* gene by default) are broken into subsegments to prevent assembly problems caused by the circular organization of the mitochondrial genome. The resulting contigs are then re-assembled into a single linear sequence, which is further circularized by the designated *ATP6* starting point. The nuclear scaffolds and the circularized mitochondrial sequence together form the improved genome assembly.
- 06. Supervised_Final_Assembly (Steps 20-22): A modification list containing the ordering, orientation and naming information of each sequence from the improved genome assembly is generated for users to review and to make manual adjustment when needed. The final genome assembly is further generated based on the user-edited modification list.
- **07. Centromere_Annotation** (Step 23): The pre-shipped *S. cerevisiae* centromere sequences are searched against the final genome assembly for centromere annotation.

- **08. Gene_Annotation** (Step 25): *De novo* protein-coding and tRNA gene annotations are performed for the final genome assembly, which are further leveraged by the mRNA transcripts and protein sequences alignments.
- **09. TE_Annotation** (Step 28): The pre-shipped curated TE library (containing the long terminal repeats (LTRs) and internal sequences of the *S. cerevisiae* and *S. paradoxus* Ty1-Ty5 by default) is searched against the final genome assembly to identify TEs. The identified TEs are further curated and classified into the full-length, truncated, and solo-LTRs of Ty1-Ty5.
- **10.** Core_X_Element_Annotation (Step 30): The pre-shipped curated hidden Markov model (HMM) of the *S. cerevisiae* core X elements is searched against the final genome assembly to annotate core X elements.
- **11. Y_Prime_Element_Annotation** (Step 32): The pre-shipped representative *S. cerevisiae* Y' element sequence is searched against the final genome assembly to annotate Y' elements. Note that Y' elements can have long, short or degenerated forms²², and we used a representative long-form Y' element as the query to maximize detection power.
- **12. Gene_Orthology_Identification** (Step 34): The annotated protein-coding genes are compared with the reference protein-coding genes based on both sequence homology and gene order conservation to identify gene orthology relationship between these two sets. Based on such gene orthology relationships, the *Saccharomyces* Genome Database (SGD; http://www.yeastgenome.org/) systematic names are assigned to the annotated protein-coding genes.
- **13. Annotation_Integration** (Step 35): The annotations of centromeres, TEs, protein-coding genes, tRNAs, as well as core X and Y' elements are combined and sorted to form a final integrated multi-feature annotation.

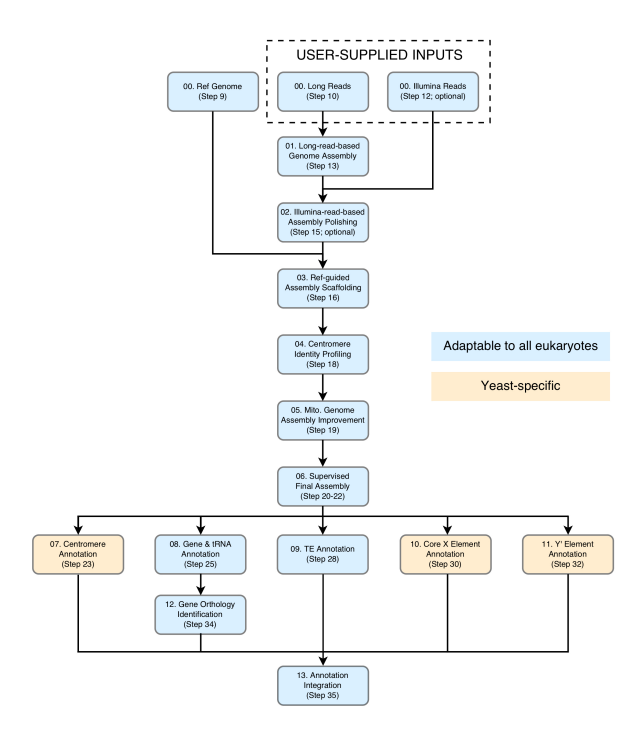


Figure 2. The workflow of LRSDAY. Each box represents an individual module. These modules are numbered according to the tasks described in Experimental Design, with the corresponding protocol step numbers also indicated. Modules that can be adapted for other eukaryotes are colored in light blue while those yeast-specific are colored in light orange.

Limitations and potential adaptation

In its distributed form, LRSDAY is tailored for the model budding yeast *S. cerevisiae* and its closely related sister species *S. paradoxus* with a number of pre-shipped auxiliary data files configured accordingly. However, given its modular design, the backbone of LRSDAY can be adapted for virtually any eukaryotic organisms to perform genome assembly, sequence polishing, reference-guided scaffolding, protein-coding genes and tRNA annotations, gene orthology identification, and annotation integration. In Appendix 2, we provide some tips with regard to such adaptation. Moreover, those assembly polishing, scaffolding and various annotation modules (See Experimental Design) can also be used to analyze existing genome assemblies derived from any or any combination of sequencing technologies. Such flexibility makes LRSDAY very useful for expanded use cases and therefore suits the needs of a broader audience.

Expected improvements

As thousands of yeast strains have been or are currently under sequencing^{23–25}, our knowledge of the overall genome content diversity²⁶ of this important model organism is expanding rapidly, revealing a whole new picture of the pan-genome diversity of *S. cerevisiae*. For example, our lab is currently working on characterizing the pan-genome of >1,000 *S. cerevisiae* isolates across the globe (The 1002 Yeast Genomes Project http://1002genomes.u-strasbg.fr/). Future developments of LRSDAY will incorporate such pan-genome dataset to provide additional annotation information for those non-reference genes, especially with regard to their evolutionary origin, population prevalence, and putative functions. Such information will greatly help users to dissect and interpret complex genotype-phenotype interactions in diverse ecological and biotechnological settings. Additional potential future direction include the direct integration with the downstream synteny analysis tools (e.g. CHROnicle²⁷, MCScanX²⁸, etc) to perform automatic large-scale structural variants discovery, which exploits one of the major benefits of having a high-quality genome assembly derived from long reads. Finally, we envision developing a dedicated web-based tool to implement such database and tool integration at a larger scale towards fully automated genomics analysis for the yeast community in the long run.

MATERIALS

Hardware, operating system and network

This protocol is designed for a desktop or computing server running an x86-64-bit Linux operating system. Multi-threaded processors are preferred to speed up the process since many steps can be configured to use multiple threads in parallel. For assembling and analyzing the budding yeast genomes (genome size = ~12 Mb), at least 16 Gb of RAM and 100 Gb of free disk space are recommended. When adapted for other eukaryotic organisms with larger genome sizes, the RAM and disk space consumption will scale up, majorly during de novo Canu¹⁷). assembly (performed bv Please refer genome to Canu's (http://canu.readthedocs.io/en/latest/) for suggested RAM and disk space consumption for assembling large genomes. Stable Internet connection is required for the installation and configuration of LRSDAY as well as for retrieving the testing data.

Software or library requirements

- Bash (https://www.gnu.org/software/bash/)
- Bzip2 (http://www.bzip.org/)
- Cmake (https://cmake.org/)
- GCC and G++ v4.7 or newer (https://gcc.gnu.org/)
- Ghostscript (https://www.ghostscript.com)
- Git (https://git-scm.com/)
- GNU make (https://www.gnu.org/software/make/)
- Gzip (https://www.gnu.org/software/gzip/)
- Java runtime environment (JRE) v1.8.0 or newer (https://www.java.com)
- Perl v5.12 or newer (https://www.perl.org/)
- Python v2.7.9 or newer (https://www.python.org/)
- Python v3.6 or newer (https://www.python.org/)
- Tar (https://www.gnu.org/software/tar/)
- Unzip (http://www.info-zip.org/UnZip.html)
- Virtualenv v15.1.0 or newer (https://virtualenv.pypa.io)
- Wget (https://www.gnu.org/software/wget/)
- Zlib (https://zlib.net/)

Input data

- Long reads: A single FASTQ file containing PacBio or Oxford Nanopore reads is needed, which will be used for long-read based *de novo* genome assembly (Task 01).
- Short reads: Short reads are optional for LRSDAY but LRSDAY could take advantage of short reads when such data is available to perform additional assembly polishing (Task 02). If paired-end Illumina sequencing is performed, two FASTQ files containing the forward and reverse Illumina reads respectively are needed. If only single-end Illumina sequencing data is available, one FASTQ file containing the single-end reads is needed.
- Reference genome: For the budding yeast *S. cerevisiae*, we pre-shipped two reference genome files (one original assembly and one with subtelomeres and chromosome-ends hard-masked based on our previous study⁷). The masked version is used for chromosomal scaffolding to minimize the confounding effect due to interchromosomal subtelomeric rearrangemnets. When working with organisms of which the subtelomeric regions are undefined, users can just use a single raw reference genome instead. The reference genome file(s) will be used for reference-guided scaffolding, mitochondrial genome assembly improvement and supervised final genome assembly (Task 03, 05 and 06 respectively).
- A number of S. cerevisiae-specific auxiliary data have been pre-shipped with LRSDAY for genomic feature annotation and gene orthology identification (Task 07-12).

Example data

• The S. cerevisiae reference genome pre-shipped with LRSDAY is taken from our previous study⁷ with the Genbank accession number GCA_002057635.1. The sequencing reads used for the testing example come from the same study, which consists of both PacBio and Illumina reads produced from the S. cerevisiae strain SK1. The PacBio reads can be retrieved with the ENA analysis accession number ERZ448251. The Illumina reads can be retrieved with the SRA sequencing run accession number SRR4074258. In LRSDAY, we have provided bash scripts to automatically download and setup these data for the testing example.

PROCEDURE

Download, install and configure LRSDAY • TIMING <1 h

- 1) Download the latest LRSDAY release (current version: v1.0.0) by entering the following commands in a terminal window:
 - \$ wget https://github.com/yjx1217/LRSDAY/releases/download/v1.0.0/LRSDAY-v1.0.0.tar.gz
 - \$ tar xvzf LRSDAY-v1.0.0.tar.gz
 - \$ cd LRSDAY-v1.0.0
 - \$ bash install_dependencies.sh

▲ CRITICAL STEP Make sure you have a fast and stable Internet connection when running this step since many tools will be downloaded here. Check that all the prerequisites (see *Software or library requirements* in EQUIPMENT or the prerequisite.txt file in the downloaded LRSDAY directory) have been installed on your system.

ACRITICAL STEP Upon the successful completion of executing the bash script, you should see a confirmation message prompted out: "LRSDAY message: This bash script has been successfully processed! :)". Otherwise, it means an error has occurred during the execution of the bash script, which interrupted the automatic installation process. This also applies to all the bash scripts used in Steps 8-35. Whenever an error is encountered, please check the error message and refer to the troubleshooting section when available. When the cause of the error is fixed, re-run this step to initiate a new installation. The installer will prompt for the confirmation of deleting the build directory generated by the old run, always answer "yes" to authorize such action so that the new installation can start.

▲ CRITICAL STEP Pay attention to the final message prompted by the installation script for a number of notes with regard to the additional manual setup steps as well as the licensing information for commercial users.

? TROUBLESHOOTING

- 2) Load the environment settings for LRSDAY by entering:
 - \$ source env.sh

After loading the pre-configured environment settings, the current directory should be assigned to the environment variable \$LRSDAY_HOME. You can check to see if the full path to your current directory is displayed after entering:

\$ echo \$LRSDAY_HOME

▲ CRITICAL STEP Although most required tools have been automatically installed and configured, manual downloading and/or configuration are needed for GATK²⁹ and RepeatMasker³⁰ due to license restriction of these tools or of their dependent libraries. Make sure to run this command to load the pre-configured environment settings before such manual setup (described in Steps 3-7). If you exited your terminal session before or in the middle of your manual setup, you need to re-load the environment settings before proceeding. These environment settings will be automatically loaded each time the task-specific bash pipelines of LRSDAY are executed.

? TROUBLESHOOTING

- 3) Download and set up GATK. Go to the official website of GATK (https://software.broadinstitute.org/gatk/download/archive) and download GATK v3.8 (the recently released GATK4 will not work). Registration might be needed for unregistered GATK users. Place the downloaded GATK package (file name: GenomeAnalysisTK-3.8-0-ge9d806836.tar.bz2) in the GATK installation directory and set up GATK by entering the following command in terminal:
 - \$ mv GenomeAnalysisTK-3.8-0-ge9d806836.tar.bz2 \$LRSDAY HOME/build/GATK
 - \$ cd \$LRSDAY HOME/build/GATK
 - \$ tar xjf GenomeAnalysisTK-3.8-0.tar.bz2
 - \$ mv ./GenomeAnalysisTK-3.8-0*/GenomeAnalysisTK.jar .
 - \$ chmod 755 GenomeAnalysisTK.jar
- 4) Set up TRF for RepeatMasker. Go to the TRF website (http://tandem.bu.edu/trf/trf.download.html) to download the TRF v4.09 executable built for 64-bit Linux. Place the downloaded file (file name: trf409.linux64) in the RepeatMasker installation directory and set up TRF by entering:
 - \$ mv trf409.linux64 \$LRSDAY HOME/build/RepeatMasker/
 - \$ cd \$LRSDAY_HOME/build/RepeatMasker/
 - \$ chmod 755 trf409.linux64
 - \$ In -s trf409.linux64 trf
- 5) Set up the RepBase library for RepeatMasker. Go to the RepBase website (http://www.girinst.org/repbase/) and register a user account (using a non-profit institutional email address, e.g. .com extension will not work) to download the RepBasker edition of the RepBase library (version: 20170127). Place the downloaded file (file name: RepBaseRepeatMaskerEdition-20170127.tar.gz) in the RepeatMasker installation directory and uncompress it by entering:

\$ mv RepBaseRepeatMaskerEdition-20170127.tar.gz \$LRSDAY_HOME/build/RepeatMasker/

\$ cd \$LRSDAY_HOME/build/RepeatMasker/

\$ tar xzf RepBaseRepeatMaskerEdition-20170127.tar.gz

? TROUBLESHOOTING

6) Get the installation path of TRF and rmblastn/makeblastdb by entering:

\$ echo \$trf_dir

\$ echo \$rmblast dir

Remember these two paths since they will be used for the RepeatMasker configuration in Step 7.

? TROUBLESHOOTING

7) Run the configuration script for RepeatMasker by entering:

\$ perl ./configure

This configuration script will prompt for several questions. Please do the following to answer these questions. Enter "env" for the question about the installation path of Perl. Just press enter for the question about the installation path of RepeatMasker. Enter the first path that you obtained in Step 6 for the question about the installation path of TRF. Enter "2" for the question about selecting a search engine. Then enter the second path that you obtained in Step 6 for the question about the installation path of rmblastn and makeblastdb. Just press enter for the question about the default search engine. And finally enter "5" to complete the configuration.

Run LRSDAY with the testing example • TIMING 42 h

8) Create the project directory. When running LRSDAY with your own data, it is recommended to make a copy of our Project_Template directory to create your own project directory such as Project_abc, where "abc" can be any string containing letters, numbers, or underscores. For this testing example, we make a copy of the Project_Template directory and name it as Project_Example by entering:

\$ cd \$LRSDAY HOME

\$ cp -r Project Template Project Example

▲ CRITICAL STEP Before proceeding to your own project, it is advised to first run our prepared testing example to check if LRSDAY is working properly as well as to get acquainted with the logic and workflow of LRSDAY.

9) Prepare the reference genome files. When running LRSDAY with your own data, you can directly put the reference genome (in FASTA format without compression) in the

00.Ref_Genomes subdirectory of your project directory (e.g. Project_abc). If your sequenced organism is *S. cerevisiae* or *S. paradoxus*, you can use the reference genome pre-shipped with LRSDAY. Here we prepare the pre-shipped reference genome for the testing example by entering:

- \$ cd ./Project_Example/00.Ref_Genome
- \$ bash LRSDAY.00.Prepare_Sc_Ref_Genome.sh
- 10) Prepare the long reads. When running LRSDAY with your own data, you can directly put the long reads in the 00.Long_Reads subdirectory of your project directory (e.g. Project_abc). The reads file should be in FASTQ or FASTA format. Compressed files with file extensions of ".gz", ".bz2", and ".xz" are also supported. For this testing example, you can download the sample long reads by entering:
 - \$ cd ./../00.Long Reads
 - \$ bash LRSDAY.00.Retrieve Sample PacBio Reads.sh
- 11) (Optional) If your long reads are generated from the PacBio Sequel platform, your reads are likely to be in BAM format. In this case, convert it to our supported format (FASTA or FASTQ) using the following commands:
 - \$ source ./../../env.sh
 - \$\$bedtools_dir/bedtools bamtofastq -i long_reads.bam -fq long_reads.fastq
 - \$ gzip long_reads.fastq
- 12) (Optional) Prepare the Illumina reads. Like for the long reads, directly put your Illumina reads in the 00.Illumina_Reads subdirectory of your project directory (e.g. Project_abc) when running LRSDAY for your own data. The reads file should be in FASTQ format with "gzip" comprehension (identified by the ".gz" extension). For this testing example, you can download the Illumina reads by entering:
 - \$ cd ./../00.Illumina Reads
 - \$ bash LRSDAY.00.Retrieve Sample Illumina Reads.sh

? TROUBLESHOOTING

- 13) Perform long-read-based *de novo* genome assembly by running the following commands:
 - \$ cd ./../01. Long-read-based Genome Assembly
 - \$ bash LRSDAY.01.Long-read-based Genome Assembly.sh
 - Upon the completion of this step, a summary file (SK1.canu.stats.txt for this testing example) will be generated to report some basic summary statistics (e.g. total assembly size, N50 (i.e. the contig length such that 50% of the total assembly size is contained in contigs of at least this size), L50 (i.e. the number of longest contigs such that 50% of the

total assembly size is contained), GC-content, etc) to assist gauging the genome assembly quality (Table 1). Two VCF files (SK1.canu.filter.mummer2vcf.SNP.vcf and SK1.canu.filter.mummer2vcf.INDEL.vcf for the testing example) will also be generated to report base-level differences between the raw genome assembly and the reference genome for their uniquely alignable regions, which could also help for assessing the genome assembly quality.

▲ CRITICAL STEP This step can be run with multiple threads to speed up the process. Depending on the CPU configuration of your Linux server/desktop, you can edit the "threads=" option in the bash script LRSDAY.01.Long-read-based_Genome_Assembly.sh to enable multi-threading. You can do the same for all the following tasks whenever the "threads=" option is provided in the corresponding task-specific bash script. Simple text editors such as emacs, vim, gedit or pico are highly recommended for such editing. Rich text editors might not work

▲ CRITICAL STEP Note this step will take long to finish (see TIMING), so we recommend running this step and all the other time-consuming steps with "nohup" (https://en.wikipedia.org/wiki/Nohup), which allows the process to continue running after you exit the terminal or logout from the server. As an example, you can run the bash script using nohup as follows:

\$ nohup bash LRSDAY.01.Long-read-based Genome Assembly.sh >run log.txt 2>&1 &

▲ CRITICAL STEP When running LRSDAY with your own data, modify the bash script to specify the input reads and reference genome, the input reads type (e.g. "pacbio_raw", "pacbio_corrected", "nanopore_raw" or "nanopore_corrected"), the estimated genome size for the assembled genome, as well as the prefix for the output data. Remember to do similar project-specific adjustment for all the following steps.

? TROUBLESHOOTING

14) (Optional) Polishing genome assembly with long-reads. When running LRSDAY for your own data, if you performed PacBio sequencing and also have access to a locally installed PacBio SMRT Analysis software package (http://www.pacb.com/products-and-services/analytical-software/smrt-analysis/), we recommend running the first-pass polishing for the assembly generated in Step 13 based on raw PacBio reads by using PacBio's own Quiver/Arrow pipeline 13. If you performed Oxford Nanopore sequencing, we recommend using nanopolish (https://github.com/jts/nanopolish) or equivalent tools to perform the assembly polishing based on raw Nanopore reads in this step.

15) (Optional) Polishing genome assembly with Illumina reads. When Illumina reads are available, we recommend running this additional polishing step either for the raw assembly generated in Step 13 (when Step 14 is skipped as in our testing example) or for the long-read-polished assembly generated in Step 14. Use the following commands to perform Illumina-read-based assembly polishing. This step can be run with multiple threads.

\$ cd ./../02.Illumina-read-based_Assembly_Polishing

\$ bash LRSDAY.02.Illumina-read-based_Assembly_Polishing.sh

TABLE 1 | Assembly statistics for the genome of *S. cerevisiae* strain SK1 assembled in the testing example.

Assembly statistics	Raw assembly	Scaffolded assembly	Final assembly
Total sequence count	22	20	20
Total sequence length (bp)	12282921	12292985	12261860
Min sequence length (bp)	24410	24410	24410
Max sequence length (bp)	1480192	1480198	1480198
Mean sequence length (bp)	558314.59	614649.25	613093.00
Median sequence length (bp)	558503.50	637681.50	637681.50
N50 (bp)	829956	923665	923665
L50	6	6	6
N90 (bp)	462498	462493	462493
L90	14	13	13
GC%	38.21	38.17	38.24
N%	0.00	0.08	0.04

Note

N50: the contig length such that 50% of the total assembly size is contained in contigs of at least this size.

L50: the number of longest contigs such that 50% of the total assembly size is contained.

N90: the contig length such that 90% of the total assembly size is contained in contigs of at least this size.

L90: the number of longest contigs such that 90% of the total assembly size is contained.

GC%: the percentage of guanine (G) and cytosine (C) bases in the nucleotide sequences.

N%: the percentage of the N bases in the nucleotide sequence. In genome assembly, the N bases are usually used to represent scaffolding gaps.

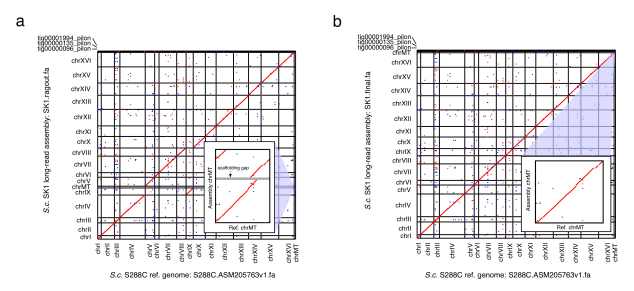


Figure 3. Genome-wide dotplots of the S. cerevisiae SK1 genome assembly generated in the LRSDAY testing example. Both the raw scaffolded assembly (panel a; generated in Step 16) and the final assembly (panel b, generated in Step 22) are analyzed. The forward and reverse sequence matches are depicted in red and blue respectively, while the zoomed-in views of the mitochondrial genome (chrMT) comparison are shown in insets. The scaffolding gap of the mitochondrial genome is indicated by the black arrow. In addition to the sixteen nuclear chromosomes and the mitochondrial genome, the scaffolded and final assemblies also contain three short contigs (tig00000096_pilon, tig00000135_pilon, and tig00001994_pilon) that are derived from highly repetitive regions.

16) Perform chromosome-level scaffolding for the long-read-based assembly. Run the following commands:

\$ cd ./../03.Reference-guided Assembly Scaffolding

\$ bash LRSDAY.03.Reference-guided Assembly Scaffolding.sh

This step can be run with multiple threads. Upon completion, a list of summary statistics (SK1.ragout.stats.txt for this testing example) will be generated for the scaffolded assembly (Table 1).

▲ CRITICAL STEP Please check the generated genome-wide dotplot (SK1.ragout.filter.pdf for the testing example) (Fig. 3a) to verify the correctness of chromosomal identity assignment performed by Ragout³¹ and apply manual adjustment in Step 21 when necessary. When running LRSDAY with your own data, you might see a single scaffold corresponds to more than one reference chromosomes, which could be

due to shared sequence homology between duplicated regions or interchromosomal rearrangements. Both types of events can be correctly interpreted based on the genome-wide dotplot generated in this Step. In either case, LRSDAY can correctly assign chromosomal identity of the corresponding scaffold based on its encompassed centromere identity as annotated in Step 18. Check the generated AGP file (SK1.ragout.agp for the testing example) for the details of reference-based scaffolding.

▲ CRITICAL STEP Due to the high AT and repeat contents and the circular conformation of the mitochondrial genome, multiple contigs corresponding to the mitochondrial genome are often obtained from the raw genome assembly, as shown in the generated mitochondrial genome dotplot (SK1.ragout.chrMT.filter.pdf for the testing example) (Fig. 3a, inset). A list of such mitochondrial contigs will also be generated (SK1.mt_contig.list for the testing example), which will be used in Step 19 for improving mitochondrial genome assembly.

17) (Optional) When running LRSDAY for your own data, if you have strong evidence for mis-scaffolding based on prior knowledge or other experimental data (e.g. mate-pair libraries or chromosomal contact data), break the corresponding ragout scaffolds back to contigs and re-joined them with corrected order using the pre-shipped Perl scripts break_scaffolds_by_N.pl, join_contigs_by_N.pl and extract_region_from_genome.pl in the \$LRSDAY_HOME/scripts directory by running the following commands:

perl \$LRSDAY_HOME/scripts/break_scaffolds_by_N.pl -i <the input FASTA file containing the scaffold sequence(s) to break> -o < the output FASTA file containing the scaffolds after the breaking> -g <the minimal length of runs of Ns in the input scaffold(s) for breaking, e.g. 100> perl \$LRSDAY_HOME/scripts/join_contigs_by_N.pl -i <the input FASTA file containing contigs for joining in a sequential order> -o <the output FASTA file containing scaffold sequence after the

contig joining> -g <gap size, i.e. the number of Ns to be inserted between two joined contigs> -t

<sequence name for the newly joined scaffold>

perl \$LRSDAY_HOME/scripts/extract_region_from_genome.pl -i <the input FASTA file containing the genome assembly> -o <the output FASTA file containing the extracted query sequence> -q <specially formatted query string (sequence:start-end:strand) containing the genomic coordinates for the region to be extracted, e.g. using the query string chrl:1000-4000:+ for extracting the sequence in the region 1000-4000 bp on the + strand of chrl in the input genome assembly> -f <the length of flanking sequences to be extracted as well, e.g. 100 for 100-bp flanking region>

A scenario for such use case is when the breakpoints of structural rearrangements are also the breakpoints of the genome assembly. In this case, the reference-based

- scaffolding will arrange contigs according to the reference genome configuration and therefore un-do the genome rearrangement.
- 18) Perform centromere profiling for the scaffolded genome assembly by running the following commands:
 - \$ cd ./../04.Centromere_Identity_Profiling
 - \$ bash LRSDAY.04.Centromere Identity Profiling.sh
 - ▲ CRITICAL STEP The chromosome-specific centromere identities profiled here will be used as another layer of information for the final chromosomal identity assignment in Step 21. The profiled centromere identities usually agree well with the chromosomal identities labeled in Step 16, so that chrl will have the CEN1 centromere and chrll will have the CEN2 centromere, etc. Exception can occur when interchromosomal rearrangements are involved in your sequenced genome. In such case, we recommend naming those rearranged chromosomes according to their encompassed centromeres in Step 21.
- 19) Perform mitochondrial genome assembly improvement by running the following commands:
 - \$ cd ./../05.Mitochondrial Genome Assembly Improvement
 - \$ bash LRSDAY.05.Mitochondrial_Genome_Assembly_Improvement.sh
 - ▲ CRITICAL STEP Check the generated final mitochondrial genome dotplot (SK1.mt_improved.chrMT.filter.pdf for the testing example) and compare it with the mitochondrial genome dotplot generated in Step 16 to see how the mitochondrial genome assembly has been improved when aligning with the reference mitochondrial genome (Fig. 3b, inset). When running this step for your own data, the degree of such improvement may vary because it depends on both the complexity of the assembled mitochondrial genome and the quality of library preparation and sequencing experiments.

? TROUBLESHOOTING

- 20) Generate the assembly modification list file for performing the final chromosome assignment by running the following commands:
 - \$ cd ./../06.Supervised Final Assembly
 - \$ bash LRSDAY.06.Supervised Final Assembly.1.sh
- 21) Edit the generated assembly modification list file (SK1.modification.list for the testing example) based on the genome-wide dotplot generated in Step 16 and the centromere profiles generated in Step 18. The modification list file consists of three commaseparated columns, which correspond to the original sequence name, sequence

orientation, and new sequence name respectively. With this file, you can do three types of editing:

If you need to change the current sequence order, you can move the corresponding rows upward or downward to reflect the correct order.

If you need to invert the orientation of a given sequence, you can change its orientation from "+" to "-" in column 2.

If you need to rename a given sequence, you can specify the new name in the third column.

For this testing example here, we need to move the row "chrIX,+,chrIX" downward to place it after the row "chrVIII,+,chrVIII", so that chrIX will be placed after chrVIII in the final assembly. Also, we need to change the row "chrMT_Contig1,+,chrMT_Contig1" to "chrMT_Contig1,+,chrMT" for renaming the assembled sequence corresponding to the mitochondrial genome.

- 22) Once all the modifications have been specified, run the following bash script to generate the final genome assembly as well as the associated genome-wide dotplot (Fig. 3b), assembly statistics (Table 1), and VCF files:
 - \$ bash LRSDAY.06.Supervised_Final_Assembly.2.sh
- 23) Re-run centromere annotation for the final genome assembly using the following commands:
 - \$ cd ./../07.Centromere Annotation
 - \$ bash LRSDAY.07.Centromere_Annotation.sh
- 24) (Optional) Customize the configuration file for gene annotation. When running LRSDAY with your own data, edit the configuration file \$LRSDAY_HOME/misc/maker_opts.customized.ctl if your sequenced organisms is neither *S. cerevisiae* nor *S. paradoxus* (See Appendix 2 for the details). If your sequenced organism is *S. cerevisiae* or *S. paradoxus*, no customization is needed unless you have native transcriptome or expressed sequence tag (EST) data for the strain that you sequenced. In this case, you can edit the line 16 of this file to provide the full path of the native transcriptome or EST assembly for your sequenced strain.
- 25) Annotate protein-coding genes and tRNAs for the final genome assembly, using the following commands. This step can be run with multiple threads.
 - \$ cd ./../08.Gene Annotation
 - \$ bash LRSDAY.08.Gene Annotation.sh

- 26) (Optional) A manual checklist file (SK1.EVM.manual_check.list for the testing example) containing a list of genes with suspicious annotations will be generated in Step 25. As labeled in this file, these annotated gene models could be fragmented, frameshifted or containing internal stop codons. Potentially, these genes could be good candidates for pseudogenes. Manually inspect the annotated gene models of these genes by loading the annotation result (SK1.EVM.gff3 for the testing example) together with the protein/EST-alignment evidence files generated during the annotation in Step 25 (SK1.protein_evidence.gff3 and SK1.est_evidence.gff3 for the testing example) into IGV³² to check how well these suspicious gene models are supported by the corresponding protein/EST-alignment evidence and to tag/remove those truly problematic ones in your downstream analysis.
- 27) (Optional) Perform dedicated mitochondrial protein-coding and RNA annotation. If you are interested in studying mitochondrial genomes, we highly recommend running dedicated mitochondrial feature annotation with specialized software such as MFannot (http://megasun.bch.umontreal.ca/cgi-bin/mfannot/mfannotInterface.pl). Although MFannot does not support local installation so far, you can run MFannot conveniently via its web portal. Be sure to select the correct genetic code table (e.g. "3 Yeast Mitochondrial" for annotating yeast mitochondrial genomes) for your analysis.
- 28) Annotate transposable elements (TEs) for the final genome assembly. This step can be run with multiple threads using the following commands:
 - \$ cd ./../09.TE_Annotation \$ bash LRSDAY.09.TE Annotation.sh
- 29) (Optional) TE activity can be highly dynamic in the genome with many complex cases such as fragmentation and nested insertion. In LRSDAY, we used REannotate³³ to automatically resolve these complex cases, which works well for most cases but it could occasionally misjoin two adjacent TEs when they are closely spaced. Further inspect and curate the LRSDAY TE annotation output (SK1.TE.gff3 for the testing example) by visualizing it in IGV³² together with the raw REannotate annotation (SK1.REannotate.gff for the testing example). For each TE found in the LRSDAY TE annotation output, examine its corresponding LTR and internal region structure based on the raw REannotate annotation to check for misjoinings. If needed, you can manually edit the corresponding TE annotation output file to decouple the misjoinings.
- 30) Annotate yeast telomere-associated core X elements for the final genome assembly, using the following commands:

```
$ cd ./../10.Core_X_Element_Annotation
$ bash LRSDAY.10.Core X Element Annotation.sh
```

- 31) (Optional) Inspect the alignment file for annotated core X element. In LRSDAY, we label the identified core X elements to be "partial" if they are shorter than 300 bp. This should work for most cases but we recommend inspecting the generated alignment file (SK1.X_element.aln.fa for the testing example) for further curation. Upon the curation, manually adjust the "partial" labeling in the annotation file (SK1.X_element.gff3 for the testing example) when needed.
- 32) Annotate yeast telomere-associated Y' elements for the final genome assembly by using the following commands:

```
$ cd ./../11.Y_Prime_Element_Annotation
$ bash LRSDAY.11.Y_Prime_Element_Annotation.sh
```

- 33) (Optional) Inspect the alignment file for annotated Y' element. Like for the core X element annotation, we used a hard length cutoff (3500 bp) to label if the identified Y' elements are "partial". We recommend manually inspecting the generated alignment file (SK1.Y_prime_element.aln.fa for the testing example) to check if the "partial" labeling is needed and edit the annotation file (SK1.Y_prime_element.gff3 for the testing example) accordingly.
- 34) Perform orthology identification for protein coding genes by using the following commands:

```
$ cd ./../12.Gene_Orthology_Identification
$ bash LRSDAY12.Gene_Orthology_Identification.sh
```

In this step, a gene orthology relationship list is created between the annotated proteome and the SGD *S. cerevisiae* reference proteome based on both sequence similarity and synteny conservation. Based on this list, we further attach SGD systematic names to our gene annotation as shown in the "Name=" field of the generated GFF3 file (SK1.updated.gff3 for the testing example). For a given annotated gene, when more than one orthologous gene can be found in the SGD reference proteome, we will label all of its co-orthologs in the "Name=" filed with "/" in between, whereas when no orthologous gene can be found, we will label its gene name as "Name=NA". This step can be run with multiple threads.

35) Integrate the annotation of different genomic features into a unified GFF3 file by using the following commands:

```
$ cd ./../13.Annotation_Integration
$ bash LRSDAY.13.Annotation Integration.sh
```

? TROUBLESHOOTING

Troubleshooting advice can be found in Table 2.

TABLE 2 | Troubleshooting table.

Step	Problem	Possible reason	Solution
1	Downloading errors or unresponsive remote servers	Unstable internet connection or temporary problems of the remote servers where the tools for downloading are hosted.	Stabilizing the internet connection and give another try.
1	Compilation or installation error for specific tools.	Prerequisites not satisfied or corner cases due to your specific system settings.	If missing prerequisites are found (see <i>software and library requirements</i> in EQUIPMENT), install the missing prerequisites and give another try as described above. Otherwise, record the error message and email it to the developers of the problematic tools or the authors of this protocol for further problem diagnosis.
2	Cannot find the file "env.sh".	Step 1 has failed.	Check the error message that you got when running Step 1 and refer to the troubleshooting for Step 1.
5	Rejected by RepBase for the license registration.	A personal or commercial email address (e.g. with a .com extension) was used for the registration.	Use a non-commercial institutional email address for the registration or purchase the commercial license if you work in a commercial entity. Also, the installation of RepBase can be potentially skipped although not recommended by RepeatMasker. Users can still follow the remaining steps of this protocol even if they skip the RepBase installation here.

6	"echo \$rmblast_dir" returns nothing.	Step 2 has failed or your terminal session got interrupted before this step.	Re-load environment settings in Step 2 and then re-try this step.
12	Downloading warnings/errors encountered.	Temporary SRA server problems.	Directly download the sample Illumina reads for the testing example by running the following commands: \$ wget ftp://ftp.sra.ebi.ac.uk/vol1/fastq/SR R407/008/SRR4074258/SRR4074 258_1.fastq.gz \$ In -s SRR4074258_1.fastq.gz SRR4074258_pass_1.fastq.gz \$ wget ftp://ftp.sra.ebi.ac.uk/vol1/fastq/SR R407/008/SRR4074258/SRR4074 258_2.fastq.gz \$ In -s SRR4074258_2.fastq.gz \$ In -s SRR4074258_2.fastq.gz SRR4074258_pass_2.fastq.gz
13	The total assembly size is much smaller than the expected value or the assembly is too fragmented.	Insufficient sequencing depth of coverage.	Obtain more reads. We recommend a minimal sequencing depth of 50X.
19	Only partial assembly is obtained for the mitochondrial genome.	High AT and repeat content of the mitochondrial genome is challenging for <i>de novo</i> genome assembly.	If the sequencing is done with PacBio, obtaining more reads should solve the problem. If sequencing is done with Oxford Nanopore data, currently there is no satisfying solution. Future improvement of the Oxford Nanopore sequencing technology will likely solve this problem given the rapid development of this

TIMING

The following timing information was measured on a Linux computing server with an Intel Xeon CPU E5-2630L v3 (1.80GHz) using 4 threads. Enabling multithreading can substantially decrease the processing time. All optional steps except Steps 12 and 15 were not processed when measuring the computation time.

- Steps 1-7, set up LRSDAY: < 1h.
- Step 8, prepare the project directory for the testing example: 5 s.
- Step 9, prepare the reference genomes for the testing example: 1 s.
- Step 10, prepare the long reads for the testing example: 22 min.
- Step 12 (optional), prepare the Illumina reads for the testing example: 20 min.
- Step 13, de novo assembly using long-reads: 18 h.
- Step 15 (optional), assembly polishing using Illumina reads: 1h.
- Step 16, reference-guided scaffolding for the raw assembly: 8 min.
- Step 18, centromere identity profiling for the scaffolded assembly: 1 s.
- Step 19, mitochondrial genome assembly improvement: 4 min.
- Step 20, generate the assembly modification list file: 1s.
- Step 21, manual editing the assembly modification list file: 30 s.
- Step 22, finalize genome assembly based on the assembly modification list file: 20 s.
- Step 23, centromere annotation for the final assembly: 1s.
- Step 25, protein-coding gene and tRNA annotation for the final assembly: 19 h.
- Step 28, TE annotation for the final assembly: 5 min.
- Step 30, core X element annotation for the final assembly: 2 min.
- Step 32, Y' element annotation for the final assembly: 2.5 min.
- Step 34, gene orthology identification for the annotated protein-coding genes: 20 min.
- Step 35, final annotation integration: 1 min.

ANTICIPATED RESULTS

Upon the completion of the LRSDAY workflow described above, users can expect to obtain a chromosome-level genome assembly with comprehensive genomic feature annotation, which will lay a solid foundation for all kinds of downstream genomic and functional analyses. As demonstrated in Fig. 3b, the final genome assembly is highly continuous, with each chromosome assembled in an end-to-end fashion. Both genome-wide dotplots (like those shown in Fig. 3) and summary statistics (like those listed in Table 1) will be generated to help users to evaluate the genome assembly quality both graphically and quantitatively. As for the annotation, LRSDAY profiles a full spectrum of genomic features for the assembled yeast genome, which include centromeres, protein-coding genes, tRNAs, Ty1-Ty5 transposable elements, as well as the telomere-associated core X and Y' elements. The availability of such rich information can be very valuable for users working on diverse biological questions. In Table 3, we further summarized the major outputs of the testing example. The final genome assembly and annotation outputs generated with this testing example are also provided in the directory Example_Outputs for users to make direct comparison with their own results.

TABLE 3 | Major LRSDAY outputs from the testing example.

Task	Step	Output file or directory	Explanation for the output
01	13	SK1.canu.fa	The long-read-based de novo genome assembly containing all the contigs assembled by Canu ¹⁷ .
		SK1.canu.stats.txt	The summary table reporting basic assembly statistics, such as the number of the assembled sequences, the total length of the assembled sequences, the minimal, maximal, mean and median lengths of the assembled sequences, the N50, L50, N90, and L90 of the assembled sequences, as well as the base composition (A%, T%, G%, C%, AT%, GC% and N%) of the assembled sequences. Users can compare this file with the similar file generated in Step 16 and Step 22. We also summarized such comparison in Table 1.
		SK1.canu.filter.pdf	The genome-wide dotplot for the comparison between the raw genome assembly and the reference genome.
		SK1.canu.filter.mummer2vcf.SNP .vcf	The VCF files to show SNP differences between the raw genome assembly and the reference genome. These two files can be used for assessing the raw assembly quality.

		SK1.canu.filter.mummer2vcf.IND EL.vcf	The VCF files to show INDEL differences between the raw genome assembly and the reference genome. These two files can be used for assessing the raw
			assembly quality.
		SK1_canu_out	the directory containing all the output files of Canu
02	15	SK1.pilon.fa	The polished genome assembly generated by Pilon ³⁴ .
		SK1.pilon.vcf	The VCF file reporting the variants identified by Pilon based on short reads
		SK1.pilon.changes	mapping against the input genome assembly. The space-delimited record of all the changes that Pilon made during the assembly polishing. The four columns
			are: the original sequence coordinate, the new sequence coordinate after the
			correction, the original base, the new
		SK1.realn.bam	base after the correction. The BAM file of short reads mapping
03	16	SK1.ragout.fa	against the input genome assembly. The scaffolded genome assembly based
00	10	Olt i.iagout.ia	on the reference genome.
		SK1.ragout.stats.txt	The summary table reporting basic
		· ·	assembly statistics of the scaffolded
			genome assembly. Users can compare
			this file with the similar file generated in
			Step 13 and Step 22. We also
		SK1.ragout.agp	summarized such comparison in Table 1. the AGP file reporting the order and
		Civilagoutagp	orientation of each input contig used
			during scaffolding.
		SK1_ragout_out	The directory containing all the output files of Ragout ³¹ .
		SK1.ragout.filter.pdf	The genome-wide dotplot for the
		-	comparison between the scaffolded
			assembly and the reference genome.
		SK1.mt_contig.list	The list of assembled contigs
			corresponding to the mitochondrial genome. This file will be used for Step 19.
		SK1.mt_contig.fa	The assembled contig sequences
		OKT.IIIL_Confug.la	corresponding to the mitochondrial
		SK1.ragout.chrMT.filter.pdf	genome. The dotplot for the comparison between the scaffolded mitochondrial genome assembly and the reference mitochondrial
			genome.
04	18	SK1.centromere.gff3	The profiled centromere identities for the scaffolded genome assembly.
05	19	SK1.mt_improved.fa	The improved genome assembly with better processing (re-assembling and
			circularization) of the mitochondrial
		SK1.mt_improved.chrMT.filter.pdf	genome. The dotplot for the comparison between
		Str.mc_mproved.onnvrr.mter.pur	the improved mitochondrial genome

			assembly and the reference mitochondrial
			genome. You should see improved
			collinearity in this plot than the similar plot
			generated in Step 16.
06	20-22	SK1.modification.list	The assembly modification list file for
			manual editing to guide the final genome
		01/4 ("1 (-	assembly.
		SK1.final.fa	The final genome assembly generated by
		CK1 final filter adf	LRSDAY.
		SK1.final.filter.pdf	The genome-wide dotplot for the comparison between the final genome
			assembly and the reference genome.
		SK1.final.stats.txt	The summary table reporting basic
		OKT.IIIIdi.Stato.txt	assembly statistics of the final genome
			assembly. Users can compare this file
			with the similar file generated in Step 13
			and Step 16. We also summarized such
			comparison in Table 1.
		SK1.final.filter.mummer2vcf.SNP.	The VCF file to show SNP differences
		vcf	between the final genome assembly and
			the reference genome. This file can be
			used for assessing the final assembly
			quality.
		SK1.final.filter.mummer2vcf.INDE	The VCF file to show INDEL differences
		L.vcf	between the final genome assembly and
			the reference genome. This file can be
			used for assessing the final assembly quality.
07	23	SK1.centromere.gff3	The centromere annotation for the final
01	20	erri.com.cro.gno	
80	25	SK1.maker.raw.gff3	genome assembly. The raw MAKER ³⁵ annotation for protein-
		· ·	coding genes and tRNA genes.
		SK1.EVM.gff3	The final EVM ³⁶ annotation for protein-
			coding genes and tRNA genes with
			systematically assigned gene IDs.
		SK1.protein_evidence.gff3	The protein-to-genome alignment
			evidences generated by MAKER. This file
			can be used for manual curation of those
		01/41 14 170	suspicious annotations.
		SK1.est_evidence.gff3	The EST-to-genome alignment evidences
			generated by MAKER. This file can be used for manual curation of those
			suspicious annotations.
		SK1.EVM.trimmed_cds.fa	The CDSs of the final EVM protein-coding
		OKT.EVW.ammed_odo.id	annotation with the out-of-frame parts
			trimmed.
		SK1.EVM.trimmed_cds.log	The log file of the CDS trimming for the
			final EVM protein-coding gene annotation.
		SK1.EVM.pep.fa	The translated protein sequences of the
			trimmed CDSs derived from the final EVM
			protein-coding gene annotation.
		SK1.EVM.manual_check.list	The list of suspicious gene annotations for
		0144 = 0.045 =	manual curation.
		SK1.EVM.PoFF.gff	The gene synteny information derived
			from SK1.EVM.gff3, which will be used for
			Task 12 (Step 34).

SK1.EVM.PoFF.ffn SAMe as SK1.EVM.trimmed_cds.fa but with simpler sequence IDs, which could be used for Task 12 (Step 34). Same as SK1.EVM.popf ab tur with simpler sequence IDs, which will be used for Task 12 (Step 34). SAME as SK1.EVM.popf ab tur with simpler sequence IDs, which will be used for Task 12 (Step 34). The firal TE annotation from Reannotate. This file can be used for further curating TE annotation. SK1.E.gff3 The final TE annotation from LRSDAY. The final TE annotation from LRSDAY. The final Core-X element annotation from LRSDAY. The sequence of all the annotated core X elements. The sequence alignment of all the annotated core X elements. The sequence alignment of all the annotated feature is complete or partial as well as whether this is consistent with the labeling in the annotated Y elements. SK1.Y_prime_element.aln.fa SK1.Y_prime_element.aln.fa SK1.Y_prime_element.aln.fa SK1.Y_prime_element.aln.fa The sequence of all the annotated Y elements. The sequence alignment of all the annotated Y elements. The sequence of all the annotated Y elements. The sequence alignment of all the annotated Y elements. The sequence alignment of all the annotated Y elements. The sequence alignment of all the annotated Y elements. The sequence alignment of all the annotated Y elements. The sequence of all the annotated Y elements. The sequence so fall the annotated Y elements. The sequence of all the annotated Y elements of further checking whether the annotated Y elements. The sequence of all the annotated Y elements of the annotated Y elements of the annotated Y elements of the annotated Y elements. The sequence of all the annotated Y elements of the annotated Y elements of the sequence of the annotated Y elements of				
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APPENDIX

Appendix 1: Pre-shipped supporting data for LRSDAY

With LRSDAY, we pre-ship the following supporting datasets for the automatic execution of LRSDAY. Unless labeled otherwise, most of these pre-shipped datasets were either described or generated in our previous study⁷.

- 1) ATP6.cds.fa # The coding sequence (CDS) of the S. cerevisiae S288C ATP6 gene.
- 2) fuzzy_defragmentation.txt # the fuzzy defragmentation file for REannotate³³.
- 3) Proteome_DB_for_annotation.CDhit_I95.fa # our curated proteome dataset for *S. cerevisiae* and other closely related yeast *sensu stricto* species.
- 4) query.Y_prime_element.long.fa # the sequence of a representative *S. cerevisiae* Y' element.
- 5) S288C.ASM205763v1.fa.gz # the S. cerevisiae S288C genome assembly.
- 6) S288C.ASM205763v1.noncore_masked.fa.gz # the *S. cerevisiae* S288C genome assembly with subtelomeres and chromosome-ends hard-masked.
- 7) S288C.centromere.fa # the centromere sequence of S. cerevisiae S288C.
- 8) S288C.gene.hmm # the hidden Markov model (HMM) for *de novo* gene annotation based on *S. cerevisiae* S288C.
- 9) S288C.X_element.hmm # the hidden Markov model (HMM) for the core X element annotation based on *S. cerevisiae* S288C.
- 10) Sc-meth.sites # the S. cerevisiae methylation sites (shipped with snoScan³⁷).
- 11) Sc-rRNA.fa # the S. cerevisiae rRNA sequences (shipped with snoScan³⁷).
- 12) SGDref.PoFF.faa # the proteinortho proteome file generated for the SGD reference genome.
- 13) SGDref.PoFF.ffn # the proteinortho CDS file generated for the SGD reference genome.
- 14) SGDref.PoFF.gff # the proteinortho gene order gff file generated for the SGD reference genome.
- 15) te_proteins.fasta # protein sequences for genes encoded within TEs (shipped with MAKER³⁵).
- 16) TY2_specific_region.fa # the sequence of a Ty2 specific regions for differentiating Ty1 and Ty2.
- 17) TY_lib.Yue_et_al_2017_NG.fa # a custom RepeatMasker library for Ty annotation in *S. cerevisiae* and *S. paradoxus*.

18) TY_lib.Yue_et_al_2017_NG.LTRonly.fa # representative Ty LTR sequences of S. cerevisiae and S. paradoxus.

Appendix 2: Tips for adapting LRSDAY to other eukaryotic organisms

The backbone modules of LRSDAY can be easily adapted for other eukaryotic organisms. Here are some tips with regard to this:

- For Task 01 (long-read-based genome assembly; Step 13), be sure to adjust the genome size parameter in line 13 of the bash script LRSDAY.01.Long-readbased_Genome_Assembly.sh based on the estimated genome size of the organism that you sequenced.
- 2) For Task 03 (reference-guided assembly scaffolding; Step 16), be sure to modify the bash script LRSDAY.03.Reference-guided_Assembly_Scaffolding.sh to provide the reference genome file of your sequenced organisms to guide the scaffolding and chromosome assignment. It is very likely that the chromosomal cores and subtelomeres of your reference genome have not been clearly defined. In such case, you can provide the same reference genome file for both the "ref_genome_raw=" and "ref_genome_noncore_masked=" parameters. The "chrMT_tag=" and "gap_size=" parameters should also be adjusted for your own project.
- 3) By default, Task 03 (reference-guided assembly scaffolding; Step 16) performs reference-guided scaffolding using the whole genome alignment constructed by Sibelia³⁸. Sibelia is designed for processing small genomes (<100 Mb). For organisms with large should install Progressive genomes, you Cactus (https://github.com/glennhickey/progressiveCactus) and use it to build the whole genome alignment in HAL format and feed it directly to Ragout³¹. Please refer to Ragout's user manual (http://fenderglass.github.io/Ragout/usage.html) for this advanced usage. We have pre-shipped the HAL tools (https://github.com/ComparativeGenomicsToolkit/hal) to enable Ragout to process the HAL file generated by Progressive Cactus.
- 4) For Task 05 (mitochondrial genome assembly improvement; Step 19), be sure to modify the "gene_start=", "ref_genome_raw=", and "chrMT_tag=" parameters in the bash script LRSDAY.05.Mitochondrial_Genome_Assembly_Improvement.sh based on your own project.
- 5) While many of the genomic feature annotation tasks are *S. cerevisiae* and *S. paradoxus* specific, Task 08 (protein-coding genes and tRNA annotation; Step 25) can be adapted for any eukaryotic organism. In general, you will need to edit the lines 16, 22, 34, 36, 44, 45, and 68-71 in the \$LRSDAY_HOME/misc/maker_opts.customized.ctl file to feed organism-specific parameters into MAKER. Also, please refer to MAKER's own Wiki page (http://weatherby.genetics.utah.edu/MAKER/wiki/index.php/Main_Page) and

- protocols (https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4286374/) for technical details and advanced usage. Similarly, you can learn more about EVM from its website (http://evidencemodeler.github.io/).
- 6) While Task 09 (TE annotation; Step 28) is heavily tuned for yeasts, the same tools that we used here (RepeatMasker³⁰ and REannotate³³) can be used for any eukaryotic organism. We recommend reading their respective manuals (distributed with the corresponding software) for adapting these tools in your own study.
- 7) For Task 12 (gene orthology identification; Step 34), you need to edit the "ref_PoFF_faa=" and "ref_PoFF_gff=" parameters based on the reference gene annotation that you used. Please check ProteinOrtho's manual (https://www.bioinf.uni-leipzig.de/Software/proteinortho/manual.html) for more details on required file formats. The pre-shipped Perl script prepare_PoFFfaa_simple.pl and prepare_PoFFgff_simple.pl in the \$LRSDAY_HOME/scripts directory should help for this. You can run these two scripts as follows:

\$ source ./../../env.sh \$ perl \$LRSDAY_HOME/scripts/prepare_PoFFffn_simple.pl -i prefix.pep.fa -o prefix.PoFF.faa \$ perl \$LRSDAY_HOME/scripts/prepare_PoFFgff_simple.pl -i prefix.raw.gff -o prefix.PoFF.gff